A robust clustering algorithm for analysis of composition-dependent

2 organic aerosol thermal desorption measurements

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17 Abstract

18 One of the challenges of understanding atmospheric organic aerosol (OA) particles stems from 19 its complex composition. Mass spectrometry is commonly used to characterize the compositional 20 variability of OA. Clustering of a mass spectral data set helps identify components that exhibit 21 similar behavior or have similar properties, facilitating understanding of sources and processes 22 that govern compositional variability. Here, we developed an algorithm for clustering mass 23 spectra, Noise-Sorted Scanning Clustering (NSSC), appropriate for application to thermal 24 desorption measurements of collected OA particles from the Filter Inlet for Gases and AEROsols 25 coupled to a chemical ionization mass spectrometer (FIGAERO-CIMS). NSSC, which extends the 26 common DBSCAN algorithm, provides a robust, reproducible analysis of the FIGAERO 27 temperature-dependent mass spectral data. The NSSC allows for determination of thermal 28 profiles for compositionally distinct clusters of mass spectra, increasing the accessibility and 29 enhancing the interpretation of FIGAERO data. Applications of NSSC to several laboratory 30 biogenic secondary organic aerosol (BSOA) systems demonstrate the ability of NSSC to 31 distinguish different types of thermal behaviors for the components comprising the particles 32 along with the relative mass contributions and chemical properties (e.g. average molecular 33 formula) of each mass spectral cluster. For each of the systems examined, more than 80% of the 34 total mass is clustered into 9-13 mass spectral clusters. Comparison of the average thermograms 35 of the mass spectral clusters between systems indicate some commonalty in terms of the thermal 36 properties of different BSOA, although with some system-specific behavior. Application of NSSC 37 to sets of experiments in which one experimental parameter, such as the concentration of NO, is 38 varied demonstrates the potential for mass spectral clustering to elucidate the chemical factors 39 that drive changes in the thermal properties of OA particles. Further quantitative interpretation

40 of the thermograms of the mass spectral clusters will allow for more comprehensive41 understanding of the thermochemical properties of OA particles.

42 **1. Introduction**

43 Atmospheric particles are composed of hundreds to thousands of individual compounds (e.g., Hamilton et al., 2004; Goldstein and Galbally, 2007), reflecting the many different sources 44 45 and the variety of chemical pathways that lead to their formation and growth. Various mass 46 spectrometry (MS) methods provide for characterization of this compositional variability, among 47 other techniques. Individual MS methods yield different insights into particle composition, 48 dependent upon the chemical selectivity of the method. Application of various data reduction 49 methods, such as clustering or matrix factorization, helps to reduce the inherent compositional 50 complexity and develop understanding of the sources and chemical transformations that 51 determine particle composition. Clustering and matrix factorization are complementary methods. 52 In this work, we develop and apply a new clustering method to measurements of the evolved gas 53 composition derived from thermal desorption of organic aerosol, specifically to mass spectral 54 measurements from the Filter Inlet for Gases and AEROsols (Lopez-Hilfiker et al., 2014) coupled 55 with chemical ionization mass spectrometry (Lee et al., 2014) (FIGAERO-CIMS). The mass spectral 56 clustering method developed here facilitates interpretation of variability in organic aerosol 57 composition and volatility, and how these depend on formation conditions.

58 Clustering methods applied across many research fields have aided in the interpretation 59 and understanding of large data sets. Clustering methods work by classifying data into several 60 groups according to the similarity between one or more properties. In the field of atmospheric 61 chemistry, clustering methods have been applied to a variety of data types. Examples include: 62 back trajectories of trace gases (Cape et al., 2000) or particles (Abdalmogith and Harrison, 2005; 63 Pinero-Garcia et al., 2015), helping to elucidate the origin and transport of pollutants; particle 64 size distributions, providing information on aerosol emission and formation (Beddows et al., 2009; 65 Wegner et al., 2012); and, the morphology of and organic functional groups comprising individual 66 particles, allowing for classification of the types of organic carbon (Takahama et al., 2007).

67 Beyond the above examples, clustering methods have been extensively applied to the 68 interpretation of single particle mass spectra, serving to characterize variability in their chemical 69 composition and identify the sources and extent of chemical processing (e.g., Gaston et al., 2013; 70 Lee et al., 2015). While clustering is a general method, a variety of specific algorithms have been 71 developed for application to a given particle mass spectral dataset. The algorithms applied to 72 analysis of single particle mass spectra include: K-means (Giorio et al., 2012; Liu et al., 2013; Lee 73 et al., 2015); fuzzy c-means (Kirchner et al., 2003; Roth et al., 2016); density-based special clustering of applications with noise (DBSCAN) (Zhou et al., 2006); neural network-based 74 75 methods, such as an algorithm derived from Adaptive Resonance Theory (ART-2a) (Song et al., 76 1999; Zhao et al., 2008; Giorio et al., 2012); hierarchical clustering (Murphy et al., 2003; Rebotier 77 and Prather, 2007); and, some combined algorithms (Zhao et al., 2008; Reitz et al., 2016). Each 78 clustering algorithm has strengths and weaknesses. In some cases, different algorithms are 79 equally effective and lead to similar categorization of the same data set, while in other cases 80 quite different results are obtained (Zhao et al., 2008). For example, K-means and ART-2a gave 81 broadly similar results on a regional particle data set (Giorio et al., 2012), and K-means performed 82 as well as a variant of hierarchical clustering method on four particle data sets (Rebotier and 83 Prather, 2007).

84 Here, we describe and apply a clustering method, an extension of DBSCAN appropriate for 85 analysis of combined thermal desorption-mass spectral measurements of organic particle 86 composition, specifically applied to data from the FIGAERO-CIMS. FIGAERO-CIMS has been 87 increasingly used in field (e.g. Gaston et al., 2016; Lee et al., 2016; Lopez-Hilfiker et al., 2016; 88 Mohr et al., 2017; Huang et al., 2018; Le Breton et al., 2019) and laboratory studies (e.g.Lopez-89 Hilfiker et al., 2015; D'Ambro et al., 2017; Wang and Ruiz, 2018) to develop understanding of the 90 molecular composition of organic aerosols. A key feature of FIGAERO-CIMS is the ability to 91 characterize the thermal behavior of organic compounds in particles on a near molecular level 92 (Lopez-Hilfiker et al., 2014). The use of chemical ionization, a relatively soft ionization method, 93 facilitates detection and characterization of both monomeric and oligomeric parent compounds 94 in organic aerosols. In FIGAERO-CIMS, particles are collected and then thermally desorbed, with 95 mass spectra of the evolved gases measured as a function of temperature. This can also be 96 displayed as a thermogram: the concentration of an ion or sum of ions as a function of desorption 97 temperature. The temperature at which a thermogram reaches maximum signal, or T_{max} , provide 98 information on the volatility, while particularly broad desorption shapes can indicate thermal
99 decomposition, suggesting the presence of lower volatility, possibly oligomeric, material (Lopez100 Hilfiker et al., 2014). A typical FIGAERO-CIMS mass spectrum of either ambient or
101 laboratory-generated organic aerosol consists of hundreds of individual ions and thermograms,
102 (D'Ambro et al., 2018; Lee et al., 2018).

103 Previous studies using FIGAERO-CIMS provided insights into particle composition, including 104 the presence of lower volatility material, based on analysis of the thermograms of several major 105 ions (Lopez-Hilfiker et al., 2014; D'Ambro et al., 2017; D'Ambro et al., 2018; Lee et al., 2018). We 106 expand on this previous work through the application of cluster analysis to FIGAERO-CIMS 107 thermograms. Clustering of FIGAERO-CIMS data provides a means to expand the understanding 108 developed from single-ion thermograms and establish the contributions of different types of 109 thermograms to the bulk particles. One previous study clustered FIGAERO-CIMS data using the 110 K-means algorithm using two parameters: the ion molecular weight and the maximum 111 desorption temperature (Faxon et al., 2018). What distinguishes our work is that we cluster the 112 thermogram across the entire desorption period for each ion, with ions grouped according to the 113 similarity of their overall volatility distribution. We have considered the performance of various 114 clustering algorithms (including K-means), ultimately concluding that a variant of the DBSCAN 115 algorithm, which we develop here and name noise-sorted scanning clustering (NSSC), provides 116 robust performance and has several advantages over other existing algorithms for FIGAERO-CIMS 117 data. The NSSC algorithm is applied to several laboratory data sets of secondary organic aerosol 118 (SOA) formed from various precursors and under various conditions, some are previously 119 described (D'Ambro et al., 2018). In this work we do not aim to provide comprehensive 120 interpretation of the resulting clustered thermograms in terms of their thermo-chemical 121 properties (Schobesberger et al., 2018), only to illustrate the potential of clustering to enhance 122 interpretation of FIGAERO-CIMS and other similar data.

123

2. Clustering Method Description

124 Application of a given clustering algorithm to a particular data type involves a number of 125 steps. Below, we discuss the specific steps for clustering of FIGAERO-CIMS data, including a

description of our noise-sorted scanning clustering algorithm. A brief discussion of otheralgorithms is also provided.

128 **2.1. Data Preprocessing**

129 **2.1.1.** Exclusion of anomalous thermograms

The quality of the data set should be examined prior to clustering. A typical thermogram exhibits a continuous evolution to a peak, peaking during a temperature ramping period, after which there is a steady decrease in signal-to-background over time during a constanttemperature soaking period; the background-corrected signal at all temperatures remains above zero or around zero within the uncertainties. See section 3.1 for further details of the FIGAERO-CIMS. An anomalous thermogram, however, contains negative signal with large magnitude.

136 Anomalous thermograms should be excluded from the clustering to assure the quality of 137 the results, although most such thermograms do not end up clustered with other ions. 138 Anomalous thermograms are identified as follows. (i) Estimate a reference noise level (σ_{ref}) for 139 each thermogram as the standard deviation of the last 100 points (corresponding to 500 seconds) 140 of the thermogram at the end of the constant-temperature soaking period, during which the 141 signals are usually relatively constant. Use of more points incorporates times when the signals 142 were still decreasing, while use of fewer points provides a less robust estimate of the noise level. 143 (ii) Find the minimum in the thermogram and calculate the average of this and the 50 points 144 (corresponding to 250 seconds, or 100 points) before and after the minimum, Amin. This provides 145 for consistency with the determination of σ_{ref} (iii) Identify thermograms for which $A_{min} < -3^* |\sigma_{ref}|$ 146 as anomalous and exclude these associated ions from further analysis. In other words, when a 147 thermogram has a valley with averaged negative values exceeding the magnitude of three times 148 of the reference noise level, then it is considered anomalous. The specific criteria specified above 149 were determined based on consideration of thermograms from 10 distinct SOA experiments. 150 While these criteria should be robustly applicable to other FIGAERO-CIMS datasets, they can be 151 adjusted depending on the specific application, data quality, and needs.

152 Ideally, when anomalous ions are identified the original data would be inspected to identify153 the likely origin of the anomalous behavior. Possible origins include problems with background

subtraction when the blank has substantially higher signal levels than the particle samples, which can happen when there is residual contamination or incomplete separation of ions having the same nominal mass. It is also possible that the components detected for the same ion are different for the particle and blank measurements. In the example systems considered here, we identified up to five anomalous ions out of what is typically a few hundred total ions.

In some cases, it is desirable to compare thermograms between related experiments, for example the experiments discussed here that investigated the influence of NO concentration on SOA formation (Section 4.3) and the impact of isothermal dilution on SOA composition and volatility (Section 4.4). In such cases, ions identified as anomalous for one experiment are excluded from analysis for all related experiments to ensure consistency.

164

2.1.2. Euclidean Distance

Any clustering algorithm requires a metric to determine the similarity between two members in the data set. Here, we use the commonly used Euclidean Distance (ED) as the metric. A smaller *ED* indicates greater similarity. A FIGAERO thermogram has *n* points, with all thermograms having an equal number of points in a data set. A data set here is defined as the collection of thermograms for all individual ions measured for a single desorption event. The *ED* between two thermograms *a* and *b* is calculated as:

171

$$ED_{a,b} = \sum_{n} \sqrt{(a_n - b_n)^2} \tag{1}$$

173

174 An individual ED value is obtained for every pair of ions in the mass spectrum, resulting in an n x 175 *n* matrix of *ED* values with the diagonal elements all zero. The signal levels between individual 176 ions differ substantially, reflecting their relative abundances. Therefore, the ED calculation uses 177 normalized thermograms, allowing for comparison between thermogram profiles irrespective of 178 signal magnitude. Normalization is achieved by dividing each point of the original thermogram 179 by the thermogram maximum, where the maximum is determined after smoothing using a 180 35-point boxcar moving average with the end points excluded from the smoothed thermogram. 181 Use of the smoothed maximum instead of the unsmoothed maximum reduces the influence of 182 noise on normalization. In the FIGAERO datasets used in this study, a typical thermogram has a 183 temperature resolution of $\Delta T \simeq 0.7$ °C during the ramping period, and a 35-point smooth 184 corresponds to smoothing over ~24.5 °C. Typical FIGAERO thermograms exhibit peaks ca. 40 °C 185 wide, and thus a 35-point smoothing retains the main peak shape while reducing the influence 186 of noise. In the constant temperature part of the thermogram (soaking period), signal levels 187 change slowly with time, on average less than 5 % for a 35 points (~3 minutes) period, so a 188 35-point smoothing is also appropriate. We note that the unsmoothed profiles are those that are 189 normalized; smoothing relates only to determining the maximum signal values used for 190 normalization.

191 The ED calculation from Eqn. 1 gives equal weight to all points in the thermogram. However, 192 in a FIGAERO thermogram, equal weighting may not be appropriate. The desorption process has 193 two stages, ramping and soaking, with the soaking period comprising approximately 70% of the 194 time points in thermograms. However, most thermograms are featureless in the soaking period. 195 In contrast, many thermograms exhibit a peak, or some otherwise characteristic behavior, in the 196 ramping period. Since the behavior in the ramping period provides greater information as to the 197 overall similarity between individual thermograms, we recommend down-weighting the soaking 198 period such that the ramping and soaking periods ultimately carry approximately 4:1 weight in 199 the calculation of the ED. We have tested weighting of 1:1, 2:1 and 10:1. Weighting of 4:1 200 provides for the most robust clustering results for the example datasets. We do not recommend 201 completely excluding the soaking period as this period still carries informational content 202 (Schobesberger et al., 2018). Specifically, in calculating ED we use all data from the ramping 203 period while down-weighting the data in the soaking period by calculating and using ten-point 204 averages.

In summary, we calculate the *ED* based on the following steps: (i) smooth the original thermogram (with absolute signal) to find the maximum value; (ii) normalize the original thermogram to the smoothed maximum; (iii) average every 10 points in the soaking period; and (iv) calculate the *ED* between every two normalized, down-weighted thermograms.

209 **2.1.3.** Dealing with noise

210 Noise is an inherent property of any measurement. Noise in the FIGAERO thermograms 211 results from various sources, including detector noise, background subtraction, and imperfect 212 fitting of mass spectra. Noise influences the ED calculated between two thermograms, typically 213 increasing the ED. Here, the level of noise, ξ , is characterized for each thermogram by calculating 214 the average difference between the smoothed and unsmoothed normalized thermograms for 215 the ramping period. The use of only the ramping period in assessing the noise level is consistent 216 with the generally more characteristic behavior compared to the soaking period. The use of the 217 normalized thermograms, rather than absolute, allows for comparison of noise between 218 thermograms.

The noise level generally varies inversely with the fractional mass contribution of the ions, illustrated for a case study of the α -pinene + OH SOA (Experiment 1 in **Table 1** and **Figure 1**). This indicates that ions contributing more to the total signal generally have a lower noise level. Detector noise is nominally independent of ion identity, and thus the low-signal ions have enhanced ξ after normalization.

Discussed further in section 2.3, clustering algorithms often perform poorly when overly 224 225 noisy data are included in the clustering. This is especially the case for algorithms such as k-means 226 and partitioning around medoids, which assign all the members to a cluster. Clustering methods 227 that do not require assignment of all members, such as DBSCAN or our NSSC, are generally less 228 sensitive to the influence of overly noisy members. However, we have found that the explicit exclusion of noisy thermograms up front serves to provide for more robust behavior and also 229 230 removes the need to consider each noisy thermogram as a possible single-member cluster. The 231 inclusion of overly noisy peaks might obscure the underlying structure of clustered thermograms. 232 Noisy thermograms are identified as follows. First, the 5% of ions having the lowest noise are identified. The $\boldsymbol{\xi}$ value of the noisiest ion from this subset of low-noise ions is defined as the 233 234 reference noise level, ξ_{ref} . Small differences in the choice of this threshold (e.g. using the lowest 235 7% of ions) do not materially influence the results. Ions for which $\xi_n > 3 \xi_{ref}$ are considered noisy 236 and excluded from the initial clustering. For the experiments we examined, there are 88-120 out 237 of ~300 ions left after noise screening, contributing 83.5% - 92.5% to the total particle mass.

238 2.2. Noise-sorted Scanning Clustering (NSSC)

239 2.2.1. Algorithm description

240 The noise-sorted scanning clustering (NSSC) algorithm developed here is a variant of the 241 commonly used DBSCAN. In NSSC, identification and clustering of thermograms occurs based on 242 their similarity to seed thermograms. When the ED between a given thermogram and the seed is 243 less than a specified ED criterion (ε) the two members belong to the same cluster. Importantly, 244 in NSSC the selection of the seed thermograms occurs based on their respective noise levels. The 245 least noisy thermogram is selected as the initial seed, the next noisiest is selected as the second 246 seed (assuming it is not already clustered), and so on. We have found that low-noise 247 thermograms typically have more well-defined and characteristic shapes and comprise a 248 substantial fraction of the total mass. The choice to select seeds based on the noise level leads 249 to overall more robust and reproducible clustering compared to random selection of seeds.

The optimal value of the distance criterion, ε , is not known *a priori*, but must be determined by the user, discussed in Section 2.2.3. A valid cluster must contain at least N_{min} members, inclusive of the seed. We use $N_{min} = 2$. Consideration and inspection of individual unclustered thermograms exhibiting unique behavior occurs as a post-clustering process (Section 2.2.2).

254 The flow of the noise-sorted scanning clustering algorithm is shown in Figure 2 and 255 summarized here. Clustering proceeds in two rounds. For the initial round, the thermograms are 256 sorted by the noise (ξ), and the ED values between all pairs of thermograms are calculated 257 accordingly. All of the thermograms are identified according to whether they have been already 258 used as seeds (SEED = 0 or 1, with 1 for thermograms used as seeds) and whether they have been 259 already included in a cluster (CLUSTER = 0 or 1, with 1 for already clustered thermograms). At the 260 start, SEED = 0 and CLUSTER = 0 for all thermograms. Clustering begins using the least noisy 261 thermogram having SEED = 0 and CLUSTER = 0 as the initial seed. The state of that seed is then 262 changed to SEED = 1. All thermograms having ED < ε for that seed and with CLUSTER = 0 are 263 identified from the ED matrix; these thermograms are considered neighbors of the seed 264 thermogram. The seed does not evolve as neighbors are added to the cluster during this step. If the number of neighbors plus the seed is greater than or equals N_{min} , the cluster is valid and 265

266 stored, with the states of all the thermograms in the cluster changed to CLUSTER = 1. Otherwise, 267 the cluster is dismissed, and CLUSTER = 0 for all the members. In this case, the current seed (with 268 SEED = 1 and CLUSTER = 0) will no longer be used as a seed in the future steps but can still end 269 up clustered as a neighbor in the other clusters. The above steps are repeated until all the 270 thermograms have either SEED = 1 or CLUSTER = 1.

271 Because a cluster must have at least N_{min} elements, not all the thermograms may end up 272 clustered. Some of these unclustered thermograms may nonetheless have very similar shapes to 273 the clustered thermograms. Here, an iterative, second round of clustering potentially adds these 274 initially unclustered thermograms to the initial clusters, using the signal-weighted average 275 thermograms for the clusters from the first round as the initial seeds. A matrix of ED values is 276 calculated between the individual unclustered thermograms and the new seeds. For each 277 unclustered thermogram, the minimum *ED*, corresponding to only one of the seeds, is identified. 278 When this minimum ED is less than ε , the unclustered thermogram is added into that cluster. A 279 new signal-weighted average thermogram for the cluster is calculated and this process repeats 280 until no additional unclustered thermograms can be added to existing clusters. The mass 281 contribution of the remaining unique unclustered thermograms after this second round can be 282 substantial or negligible, ranging from <0.05% to 2.6% in the experiments presented here, and 283 depends largely on the choice of ε . Some of these unclustered thermograms are defined as 284 additional one-member clusters, discussed in the following section.

285

2.2.2. Post-clustering Processes

286 After thermograms are clustered, we perform two post-clustering analyses to better 287 understand the whole data set: 1) identifying additional one-member clusters and 2) sorting of 288 the clusters.

289 Some of the remaining unclustered thermograms have significant individual mass 290 contributions and should be considered as one-member clusters. The criterion of "significant" 291 mass contribution is user-defined. We recommend determining the significance criterion as follows: (i) sorting all the ions (before the noise-filtering process) from largest to smallest 292 293 individual mass concentration; (ii) calculating the cumulative mass fraction for this sorted list;

and (iii) defining as "significant" all those ions contributing to a cumulative mass contribution upto 80%.

296 The number of significant ions in a data set depends on the specific chemical system, 297 varying from only a few to tens of ions. Significant unclustered ions are identified as additional 298 one-member clusters. In some cases, the thermograms for these one-member clusters are 299 unique compared to the previously identified clusters. In others, their shapes are visually similar 300 to the previously identified clusters but where the one-member clusters are sufficiently distinct 301 that they were not clustered. For the purpose of automation, these one-member clusters are all 302 included in the final clustering results and the number of one-member clusters serves as one of 303 the parameters to determine the optimal ε . User can also choose to exclude them or some of 304 them manually from the final clustering results based on their judgement. For the example 305 systems considered in Section 4, there are only a few one-member clusters (ranging from 0 to 4) 306 for the optimal ε used.

307 Sorting of clustered thermograms facilitates visual presentation and identification of the 308 similarities and dissimilarities among the clusters. The specific method of sorting can be varied 309 depending on the application and system under consideration. Here, we use the temperature 310 where 50% of the mass is desorbed (T_{m50}) for the weighted-average cluster thermogram as a first 311 criterion. The T_{m50} is typically similar to, but slightly larger than the temperature at which the 312 signal reaches a maximum. As such, the T_{m50} is approximately related to the saturation vapor 313 pressure of the desorbing compound, at least for compounds that desorb directly (e.g., Lopez-314 Hilfiker et al., 2014). When two or more clustered average thermograms have identical T_{m50} , a 315 rare but occasional occurrence, they are further sorted by T_{m75} , the temperature where 75% of 316 the mass is desorbed. The temperature difference between T_{m50} and T_{m75} indicates the slope of 317 the thermogram between these two temperatures, with larger values indicating slower decay. 318 Therefore, these two parameters generally illustrate the shape of a thermogram. The T_{m50} and 319 T_{m75} are determined by calculating the cumulative desorbed mass and finding the temperatures 320 where 50% and 75% are reached.

The sorting process tends to organize the cluster-specific thermograms such that clusters having lower peak temperatures (lower T_{m50}) and steeper downslopes after the peak (lower T_{m75})

323 come first. Thermograms of this type are indicative of major contributions from higher-volatility 324 monomers (Schobesberger et al., 2018). Thermograms having higher T_{m50} generally have broader 325 peaks, and shallower downslopes, indicative of substantial contributions from low-volatility 326 compounds or decomposition of oligomers. Further discussion of the interpretation of 327 thermogram shapes is provided in Section 3.2.

328

2.2.3. Choosing the optimal $\boldsymbol{\epsilon}$

329 NSSC is a distance-based clustering method, so the choice of the distance criterion, ε , is a 330 crucial step. For small ε , members within a cluster have high similarity, but few thermograms end 331 up clustered. In contrast, for large ε the majority of the thermograms are clustered into only a 332 few clusters having comparably low intra-cluster similarity. The choice of the optimal ε value is 333 guided here by consideration of several parameters that vary with ε . The overall aim is to simultaneously (i) minimize the unclustered mass fraction ($f_{m,unclustered}$) while (ii) maximizing the 334 335 number of clusters (N_c) having two or more members and (iii) minimizing the number of one-336 member clusters ($N_{c,one}$) yet (iv) maintain inter-cluster separation ($R_{interClst}$).

337 In general, N_c increases with ε for small ε because more thermograms of different shapes 338 get clustered and fewer thermograms remain unclustered. As ε further increases, some clusters 339 are combined and a greater number of thermograms are assigned to a single cluster. 340 Consequently, as ε increases the N_c generally increases, reaches a maximum level, and then 341 decreases. The maximum N_c and the ε at which the maximum occurs depends on the exact size 342 and the properties of dataset being examined. We have found that a typical SOA system usually 343 has 9-13 distinct thermogram clusters. We recommend selecting an ε that provides for N_c at or 344 near the maximum as this captures the greatest number of thermogram types.

The mass fraction of unclustered thermograms, $f_{m,unclustered}$, includes only the unclustered thermograms that were not excluded based on the noise filtering. In general, a smaller $f_{m,unclustered}$ is preferable as this indicates a greater amount of the OA mass is included in a cluster (including one-member clusters). The $f_{m,unclustered}$ generally decreases with ε , then plateaus above a certain value of ε ; ideally this plateau occurs at $f_{m,unclustered} = 0$. The ε where the plateau starts is indicated as ε_{MF} , where MF stands for mass fraction. Given that significant one-member clusters are

allowed, the unclustered thermograms that remain above ε_{MF} have individually small mass contributions and are either truly unique in their shapes or have a sufficiently high noise level that they cannot be clustered, even after the noise-screening process. We generally recommend selecting $\varepsilon \ge \varepsilon_{MF}$ to minimize the unclustered mass.

The number of one-member clusters, $N_{c,one}$, generally decreases with ε , as these ions are incorporated into multi-member clusters. Ideally, these one-member clusters would exhibit clear, visually distinct behavior compared to other one-member clusters and to multi-member clusters. However, we find this is often not the case, especially at smaller ε . Thus, the number of onemember clusters should generally be minimized; we suggest $N_{c,one}$ be held to five or fewer in general.

361 The inter-cluster separation parameter, $R_{interClst}$, characterizes the dissimilarity between 362 clusters, and is the ratio between the average inter-cluster distance ($ED_{seed,avg}$) and ε , where: 363

$$364 R_{interClst} = \frac{ED_{seed,avg}}{\varepsilon} = \frac{\sum_{i=1}^{N_{c,total}} \sum_{j=1}^{N_{c,total}} ED_{seed,i,j}}{N_{c,total} \cdot (N_{c,total} - 1) \cdot \varepsilon} (2)$$

365

and $ED_{\text{seed},i,j}$ is the distance between the seeds for the different clusters *i* and *j* and $N_{c,\text{total}} = N_c + N_{c,\text{one}}$. For a 2D data set, the seed can be visualized as the center of a circle and ε the radius of the circle. Thus, when $ED_{\text{seed},i,j}/\varepsilon < 2$, the two circles defining the boundaries of these two clusters have overlapping areas. Good separation (i.e. cluster dissimilarity) is indicated when $ED_{\text{seed},i,j}/\varepsilon > 2$. Although our data set is more than two dimensions, this illustrates the idea of establishing the level of similarity (or dissimilarity) between clusters, i.e., the extent to which they are unique. We recommend selecting an ε that results in $R_{\text{interClst}} \ge 2$, when possible.

All four parameters should be considered when determining the optimal ε . Consideration of the parameters individually may not result in the same optimal ε . Ultimately, the user must consider each parameter and aim to select an optimal ε that balances the different information provided in each parameter. This can be achieved by plotting the above parameters as a function of ε , and then selecting as the optimal value the ε that results in (i) a small $f_{m,unclustered}$ with (ii) N_c near the maximum and (iii) a small $N_{c,one}$ and (iv) $R_{interClst}$ near or above two. In addition, visual comparison of the clustering results, illustrated as the average thermogram of each cluster, can
be helpful. For the example data considered below, we find that the optimal ε tends to fall within
a relatively narrow range of values.

382 **2.2.4.** Summary

383 The NSSC allows for clustering of ion peaks in temperature-dependent mass spectra 384 measured by the FIGAERO-CIMS, from which mass thermograms of the different clusters are 385 determined. The NSSC emphasizes contributions of ions having high signal-to-noise by selecting 386 seeds for the mass spectral clusters according to decreasing signal-to-noise. The NSSC also 387 accounts for the full temperature-dependent behavior of each ion, weighted towards the 388 temperature ramping period during which the ions generally exhibit more characteristic 389 desorption profiles. However, the NSSC requires as user input a distance criterion, ε , which 390 characterizes the minimum similarity required between a selected seed ion thermogram and all 391 other (non-clustered) ion-specific thermograms for the non-seed ion to be considered part of the 392 mass spectral cluster. The appropriate ε value must be uniquely determined for a given 393 experiment or set of experiments, but we recommend should be selected to provide both the 394 greatest amount of clustered mass and number of mass spectral clusters having two or more 395 members while also maintaining the greatest average separation between the mass spectral 396 clusters. Altogether, these steps facilitate robust, reproducible determination of mass spectral 397 clusters from a given data set.

398

2.3. Alternative Clustering Methods

399 We have alternatively considered the performance of some of the most commonly used 400 clustering algorithms (k-means, k-medoids, mean-shift, DBSCAN) and a less-commonly used one 401 (FPClustering (Gonzalez, 1985)) for interpreting FIGAERO-CIMS observations. The clustering 402 methods considered are summarized in Table 2, with some of their pros and cons listed, and 403 described in further detail in Appendix A. We discuss them briefly here in the context of FIGAERO-404 CIMS data. All the methods considered require input of at least one key user-specified parameter. 405 These parameters and the associated clustering algorithms can be generally classified into two 406 categories: number-based and distance-based. Number-based clustering algorithms require

407 specifying the desired number of retrieved clusters; this includes k-means and k-medoids. 408 Number-based algorithms usually assign all members to clusters. The extent of similarity among 409 members of a cluster can vary greatly since there is no strict distance criterion for each cluster. 410 When applied to FIGAERO-CIMS thermograms, we have found these number-based algorithms 411 are particularly sensitive to the presence of noisy members and the initialization method. In 412 contrast, some clustering algorithms require specification of distance (similarity) criterion. This 413 includes the mean-shift, DBSCAN, and our NSSC algorithms. These distance-based algorithms 414 need not cluster all members of the initial population and generally emphasize intra-cluster 415 similarity or the density of the points. The methods differ in terms of the method used for 416 selection of the initial seed or center and the extent to which they emphasize point density versus 417 cluster similarity. Noisy members tend to naturally be excluded from any clusters. NSSC is a 418 variant of DBSCAN. It does, however, differ from the standard DBSCAN algorithm because NSSC 419 only searches for neighbors of the seed, while DBSCAN also searches for neighbors of the 420 neighbors. In doing so, NSSC emphasizes cluster similarity rather than point density. This is 421 particularly useful when clustering thermograms, as the behavior of the entire thermogram is 422 considered; inclusion of neighbors of neighbors may cluster together thermograms that exhibit 423 especially similar behavior in one region (e.g., the soaking period) but not another, an undesirable 424 result. Accordingly, the sorting of seeds by noise levels is a key aspect of the NSSC algorithm, 425 which we have found provides for more robust clustering results.

426 Most of these clustering algorithms, including k-means, k-medoids, and mean-shift, are 427 initialized with a random choice of the initial cluster centers (or seeds). For large data sets, this 428 randomness usually leads to different results of clustering with different runs. The extent to 429 which this impacts analysis and clustering of FIGAERO-CIMS data is considered using SOA from 430 the α -pinene + OH SOA system as the case study (Section 4.1). For the FIGAERO-CIMS data we 431 find that the various clustering results exhibit a moderate sensitivity to how the initial seeds are 432 selected for all of these algorithms, although the final clusters are generally similar between 433 different runs for the same input parameter. This may reflect either the relatively small size of 434 the data set (~300 members originally and ~100 members after noise screening) or that there are 435 generally characteristic peak shapes with overall good separation. However, some differences

436 between independent clustering runs result, which is undesirable. For FIGAERO-CIMS data we 437 know that not all thermograms are of equal quality, i.e. they have different noise levels reflecting 438 in part their different overall contributions to the total mass. The standard clustering methods 439 do not account for this information. The NSSC algorithm developed here takes into account this 440 measure of data quality and uses it to identify the seeds for clustering. This provides for an 441 entirely reproducible clustering and generally emphasizes the behavior of the ions that 442 contribute most to the FIGAERO-CIMS signal while still allowing for consideration of contributions 443 of low-signal ions.

444 We find that different clustering algorithms can result in similar numbers of clusters with 445 the cluster-averaged thermograms having visually similar shapes when each is run with 446 appropriate user-selected parameters, although the details and robustness of each cluster vary 447 method by method. The "appropriate" parameters however are different from the "optimal" 448 parameters. There is usually different guidance for different algorithms on how to find the 449 optimal parameters that result in the greatest similarity within clusters and dissimilarity among 450 clusters. In the case of k-medoids, for example, the average silhouette indicates an optimal 451 number of clusters of two for the case study system. Yet, this is certainly too few clusters based 452 on the other methods.

In summary, we propose NSSC as the preferred algorithm in dealing with the FIGAERO data set based on: (i) the ability to generate similar results as the other commonly used clustering algorithms; (ii) good reproducibility and stability of results due to accounting for the noise of individual thermograms; (iii) good control over the similarity within the clusters by using a user-definable distance criterion; and (iv) a capability to identify unique thermograms as one-member clusters.

459 **3. FIGAERO Measurements and Experiments**

460

3.1. Instrument and experiment description

The FIGAERO-CIMS instrument has been described previously in detail (Lee et al., 2014; Lopez-Hilfiker et al., 2014). A brief description is provided here, with some additional details in the Supplemental Material. The FIGAERO-CIMS measures the evolved gases from filter-collected

464 particles during temperature programmed thermal desorption. Thermal desorption of particles 465 occurs in two-stages: a "ramping" and "soaking" period. During ramping, the temperature 466 increases from room temperature to 200 °C, typically at 10 °C min⁻¹. Most OA mass desorbs 467 during the ramping stage. The temperature is held at 200 °C for ca. 30–40 mins during the soaking 468 period to facilitate evaporation of the remaining, low-volatility organic mass from the filter. The 469 evolved gas-phase compounds are measured using CIMS with the iodide (I⁻) reagent ion, 470 appropriate for characterization of generally highly oxygenated components comprising most 471 secondary organic aerosol (Lopez-Hilfiker et al., 2016; Isaacman-VanWertz et al., 2017; Lee et al., 472 2018). The resulting signal or mass concentration versus temperature (or equivalently time) 473 curves for each ion constitute a thermogram. All individual thermograms are background 474 corrected by subtracting the observed thermograms from appropriate blank experiments. The 475 overall bulk thermogram is obtained by summing together the individual thermograms.

476 Several example applications of the clustering on FIGAERO-CIMS data are discussed in 477 Section 4. These cover laboratory experiments on SOA derived from: (1) OH + α -pinene and (2) 478 OH + Δ -3-carene, both at low-NO_x conditions; (3) OH + α -pinene as a function of [NO]; and (4) 479 $O_3 + \alpha$ -pinene, but where the SOA is allowed to isothermally evaporate at 80% RH for varying 480 amounts of time prior to thermal desorption. These experiments are summarized in **Table 1**, with 481 further details in the Supplemental Material and associated publications (D'Ambro et al., 2018; 482 D'Ambro et al., 2019); all data are publicly available (Cappa et al., 2019). All the experiments were 483 done in a 10.6 m³ Teflon environmental chamber at Pacific Northwest National Laboratory (PNNL) 484 (Liu et al., 2012; Liu et al., 2016).

485

3.2. General interpretation of FIGAERO-CIMS thermograms

This work focuses on development of the clustering method, rather than on interpretation of the FIGAERO-CIMS thermograms; an illustrative thermogram is shown in **Figure 3**b. However, discussion of the clustering results is aided by a general understanding of how FIGAERO-CIMS thermograms have been previously interpreted. Ions contributed by semi- and low-volatility compounds that desorb directly tend to exhibit strongly peaked, Gaussian-like thermograms with single-mode peaks between around 50 °C to 120 °C; the lower the peak desorption temperature (T_{peak}) the higher the volatility of the desorbing compound (Lopez-Hilfiker et al., 2014; 2015). We

493 therefore refer to thermograms, or portions of thermograms, having this general shape as the 494 "monomeric" content of the ion hereafter; direct evaporation of thermally stable dimers or other 495 oligomers is possible, although will typically occur at higher temperatures due to the comparably 496 lower volatility of these compounds. When multiple monomeric compounds having different 497 vapor pressures contribute to the same ion, the resulting thermogram exhibits a broader peak 498 and shallower slopes or, in particular cases, multiple, distinct peaks (Lopez-Hilfiker et al., 2015). 499 However, very broad thermograms, especially those that peak at higher temperatures (> 120 °C 500 or so), can also indicate contributions from thermal decomposition of very low-volatility 501 monomers, dimers, and oligomers (Lopez-Hilfiker et al., 2015; Gaston et al., 2016; Schobesberger 502 et al., 2018). Dimers and oligomers can evaporate directly, without thermal decomposition, as 503 observed for isoprene-derived SOA (D'Ambro et al., 2017) and ambient monoterpene oxidation 504 products (Mohr et al., 2017). However, fragments of dimers or oligomers are generally more 505 abundant, indicating the importance of thermal decomposition for desorption of these low-506 volatility compounds. Both direct evaporation of extremely low-volatility compounds and 507 decomposition of large molecules or oligomers can lead to high signal levels above ~120 °C. We 508 refer to both peaks and the slowly varying signal above ~120 °C as the "oligomeric" content of 509 the ion hereafter. We use the terms monomer and oligomer in a qualitative manner. A more 510 quantitative analysis of the thermograms can help distinguish between direct evaporation, 511 thermal decomposition, and the contributions of monomers versus oligomers (Schobesberger et 512 al., 2018), yet is beyond the scope of the current work.

513

4. Example Applications

To illustrate the broad utility of NSSC for interpretation and analysis of FIGAERO-CIMS data, we apply NSSC to the laboratory-generated SOA systems described above. The systems include: SOA formed from a single precursor under NO_x-free conditions; SOA formed from a single precursor as a function of input [NO]; and, SOA formed from a single precursor with thermal desorption following isothermal evaporation.

519 **4.1.** α-pinene + OH SOA

520 A total of 298 ions were characterized by FIGAERO-CIMS for SOA generated from the α -pinene + OH reaction (**Table 1**). Four ions were characterized as anomalous and excluded from 521 522 further analysis (see Section 2.1.1). The mass concentration of each ion was calculated by 523 integrating the signal across the entire desorption period and assuming an equal sensitivity of 524 CIMS for all the compounds. The total mass concentration is the sum of all the non-anomalous 525 ions. The mass spectrum and bulk thermogram of the remaining 294 ions are shown in Figure 3, 526 with the bulk thermogram shown versus both temperature (Figure 3b) and time (Figure 3c) to 527 illustrate the difference between the ramping and soaking periods. The individual thermograms 528 exhibited a variety of shapes. The noise threshold for this data set was ξ_{ref} = 0.020893. A total of 529 188 ions were screened out via noise filtering. The remaining 106 ions contribute 92.5% to the 530 total mass detected by FIGAERO-CIMS. The optimal ε was established through consideration of 531 the co-dependencies of N_c , $N_{c,total}$, $f_{m,unclustered}$ and $R_{interClst}$ on ε (Figure 4; Table 3). For this data 532 set, we determine the optimal ε = 2.6. Choice of a much smaller ε , around 1.5, gives a maximum 533 in N_c, but leaves a large fraction of the mass unclustered. Choice of ε = 2.1 or 2.2 yields larger N_c 534 and $R_{\text{interClst}}$ than ε = 2.6, with a reasonably small $f_{\text{m,unclustered}}$. However, there is one type of 535 thermogram (Clst#11 in Figure 5) that is only captured with $\varepsilon \ge 2.6$ and this yields $f_{m,unclustered} = 0$. Using $\varepsilon \ge 2.7$ also yields $f_{m,unclustered} = 0$ and $N_{c,one} = 0$, but N_c and $R_{interClst}$ decrease from $\varepsilon = 2.6$, 536 537 indicating increasing similarity between clusters with fewer types of shapes captured. The choice 538 of ε = 2.6 provides a compromise between maximizing N_c, minimizing $f_{m,unclustered}$, and keeping 539 R_{interClst} above two. The parameters and thresholds used for this data set are summarized in **Table** 540 3.

A total of 11 clusters are identified with no one-member clusters. The unweighted and mass-weighted average thermograms for each cluster are shown along with the thermograms of individual members in **Figure 5**a. The differences between weighted and unweighted average clusters are negligible, in general. Clusters are organized and numbered (as Clst#*N*) from low to high T_{m50} , with deeper to shallower downslope. Clst#1 through Clst#6 all have a clear peak below 120 °C, but with different peak widths and downslopes. Clst#7 and Clst#8 are a bit noisier with

only a few members each, exhibiting a sharp upslope and shallow downslope. Clst#9 has a very
broad peak. Clst#10 peaks at around 150 °C after an initial rise and temporary plateau. Clst#11
exhibits behavior somewhat like Clst#10, but with a peak that occurs just into the soaking period,
evident if viewed in time space, at 200 °C with a rapid drop afterwards.

551 The total mass concentration of a given cluster ($M_{c,N}$) is the sum across all cluster members, 552 calculated by integrating the summed mass concentration across the entire desorption period. 553 The percentage mass contribution of each cluster, and of the unclustered and the noise-filtered 554 ions, as well as the number of members for each cluster are shown in Figure 5b and Error! Reference source not found.. Clst#2 and Clst#3 contain the majority of the mass (20.1% and 555 556 44.3%, respectively) and consist of nearly half of the clustered ions (11 and 42, respectively). 557 Clst#4 and Clst#9 also contain a notable percentage of the total mass (8.2% and 9.8%, 558 respectively) and include a notable number of ions (13 and 17, respectively). Other clusters 559 contribute relatively little to the total mass and contain a small fraction of ions.

560 The mass-weighted average molecular formulas ($C_xH_yO_zN_m$) differ between clusters, as do 561 the O:C and H:C atomic ratios (Error! Reference source not found.). There is no clear relationship between T_{m50} (or cluster number) and the number of carbon atoms, MW, or O:C. There is, 562 however, a reasonable, inverse correlation between T_{m50} and H:C ($r^2 = 0.78$). The number of 563 564 carbon atoms is notably larger for Cluster 6 (x = 11.1) and Cluster 7 (x = 15.3); if those two clusters 565 are excluded there is an inverse relationship between T_{m50} and the number of carbon atoms (r^2 566 = 0.79) and with MW (r^2 = 0.59). While the reason for these two clusters having comparably large 567 numbers of carbon atoms is unknown, this nonetheless suggests that the contribution of 568 oligomer decomposition might increase for clusters having higher T_{m50} values.

Interpretation of previous FIGAERO-CIMS studies have largely focused on the behavior of the bulk thermogram or of several major ions or sums of ions based on common factors such as the number of carbon atoms (Lopez-Hilfiker et al., 2016; D'Ambro et al., 2017; D'Ambro et al., 2018; Stolzenburg et al., 2018; Wang and Ruiz, 2018; Joo et al., 2019). The normalized thermograms of the top five ions contributing most to the total mass for the experiments here are shown in **Figure 5**c, along with the bulk thermogram. Together these five ions make up nearly 30% of the total mass, and exhibit very similar thermogram shapes to each other and to the bulk

thermogram and belong solely to either Clst#2 or Clst#3. Thus, examining these ions only would
capture only a fraction of the overall diversity in thermal behaviors. The clustering method
developed here provides a means to investigate more comprehensively the variability in volatility
between aerosol components.

580

4.2. Δ -3-carene + OH SOA

581 A total of 298 ions were characterized by FIGAERO-CIMS for SOA generated from the 582 reaction of Δ -3-carene + OH (**Table 1**). Five were identified as having anomalous thermograms 583 and excluded from further analysis. The mass spectrum and bulk thermograms of Δ -3-carene + 584 OH SOA are shown in **Figure 6**. Compared to the α -pinene +OH SOA described above, the mass 585 spectrum of Δ -3-carene SOA is quite different, with one ion (C₈H₁₂O₅) dominant. The bulk 586 thermograms of the two SOA systems both look bell-like, but with the Δ -3-carene SOA 587 thermogram having a peak temperature ca. 9 °C higher. After noise-filtering, 110 ions remained 588 for clustering, contributing 90.7% to the total mass. The optimal ε = 2.1, established again by 589 considering the system-specific dependence of N_c , $N_{c.one}$, $f_{m.unclustered}$ and $R_{interClst}$ on ε (Error! 590 Reference source not found.), with the parameters and thresholds summarized in Table 3.

591 Ten clusters are identified, including one one-member cluster, with thermograms shown in 592 Figure 7a and the mass contribution and number of ions in a cluster in Figure 7b. Chemical 593 properties of each cluster are summarized in Error! Reference source not found.. The general 594 characteristics of thermograms identified in the Δ -3-carene + OH SOA are similar to those of low-595 $NO_x \alpha$ -pinene + OH SOA described above, but with different mass contributions. For example, 596 Clst#4 has nearly identical shape of the thermogram as Clst#3 in the α -pinene SOA but 597 contributes less to the total mass, 28.0% compared to 44.3%. Clst#6 in the Δ -3-carene SOA 598 contributes 14.8% to the total mass and resembles Clst#5 in the α -pinene SOA, which contributes 599 only 4.0% to the total mass.

600 In general, Clst#1 – 6 in the Δ -3-carene SOA all exhibit a peak below 120 °C, with clear peaks 601 of varying width and downslopes of varying steepness, but nominally in order of narrow to wide 602 and steep to shallow, respectively. These clusters carry the majority of the desorbed mass. Clst#7 603 and Clst#8 both exhibit relatively flat thermograms in the ramping period after their initial rise,

and contribute 9% to the total mass. Clst#9 has a peak temperature above 150 °C and Clst#10
reaches a maximum during the soaking period. These last two clusters contribute little to the
total mass (0.6% and 0.3%, respectively).

The thermograms of the five largest ions are shown in **Figure 7**c. These five ions together carry ~35% of the SOA mass. A wider variety of thermogram shapes are captured by the top five ions compared to the α -pinene SOA system. However, thermograms characteristic of Clst#7–10 are not represented by these top five ions; this remains true even if the top 10 ions are considered (not shown).

612 There are ultimately three major differences between the two SOA systems. For one, there 613 is a different relationship between fractional contribution and cluster number (and thus $T_{m,50}$) 614 between the two. Secondly, the α -pinene SOA contains ions with especially narrow peaks at ca. 615 100 °C (i.e., Clst#7 & 8), that are not observed with Δ -3-carene SOA (compare Figure 5 with Figure 616 7). Lastly, the thermograms of the top five ions for Δ -3-carene SOA differ to a greater extent than 617 for α -pinene SOA. Although we are unable to determine the reasons for these differences here, 618 this illustrates the potential for clustering to help identify and understand differences between 619 different SOA systems.

620

$\Delta = 4.3. \quad \alpha$ -pinene + OH + NO SOA

621 Thermograms from SOA generated from the reaction of α -pinene + OH at varying NO 622 concentrations (5 ppb, 10 ppb and 25 ppb; Table 1) are considered as a set of experiments. 623 Together, differences between them illustrate the impact of changes to the fate of RO₂ peroxy 624 radical intermediates on the SOA composition and thermal properties (Praske et al., 2018; Zhao 625 et al., 2018). Clustering proceeds here using two complementary approaches. In the single 626 clustering method, clustering is performed for one reference experiment (i.e., at one NO 627 concentration, 5 ppb, Expt#3a). Then, average thermograms are calculated for the other 628 experiments in the set using the same cluster members as identified in the reference experiment. 629 In the multiple clustering method, clusters are independently determined for each experiment in 630 the set, and the shapes, relative abundances, and contributing ions are compared between

experiments. For all three experiments, the same initial set of 298 ions were characterized byFIGAERO-CIMS.

633 4.3.1. Single Clustering

634 The ions identified as anomalous in each experiment differed. This most likely results from 635 shifts in the background signal levels between experiments. To maintain consistency between 636 the three experiments, ions identified as anomalous in any of the experiments were excluded 637 from all the experiments, with four ions excluded in total. A total of 88 ions were kept for 638 clustering after noise-filtering using the 5 ppb NO reference experiment, contributing 84.5% to 639 the total mass. The optimal ε = 2.2 (Error! Reference source not found. and **Table 3**), resulting in 640 ten clusters with one one-member cluster. The same sets of ions were then used to calculate the 641 cluster-average thermograms for the 10 ppb and 25 ppb NO experiments. Chemical 642 characteristics of the clusters are summarized in Error! Reference source not found..

643 Mass spectra for the three experiments are compared in Figure 8a and the bulk thermograms shown in Figure 8b and c. The 5 ppb NO and 10 ppb NO SOA mass spectra are 644 645 nearly identical. The mass spectrum for the 25 ppb NO experiment, however, exhibits a notable 646 shift of the most abundant ions towards lower m/z. The bulk thermograms for the 5 ppb and 10 647 ppb NO experiments are nearly identical, peaking near 80 °C. The 25 ppb NO bulk thermogram 648 similarly peaks near 80 °C, but exhibits a much slower decay as temperature increases further. 649 Additionally, the change in slope at the transition from the ramping to soaking period is more pronounced in the 25 ppb NO experiment. Overall, a greater fraction of the mass desorbs above 650 651 100 °C and during the soaking period for the 25 ppb NO experiment compared to lower-NO 652 experiments.

Despite the differences in the bulk thermograms, the shapes of the weighted-average thermograms of clusters for all the NO experiments are generally similar, with the exception of Clst#6 (**Figure 9**a). In particular, the 25 ppb thermogram shape of Clst#6 differs substantially from those of low-NO conditions, with a much reduced initial peak (around 80 °C) and an more pronounced second peak at high temperature (around 200 °C). However, this cluster contributes negligibly to the overall mass. There is some suggestion of similar behavior for Clst#10, although

to a lesser extent. For the three most abundant clusters, Clst#1, 2 and 4, there is a slightly
increased relative contribution of the 100-200 °C tail for 25 ppb NO, consistent with differences
in the bulk thermograms.

662 The most notable NO-dependent change is in the relative abundances of the clusters 663 between the 5 and 10 ppb NO experiments and the 25 ppb NO experiment (Figure 9b). The 664 cluster mass fractions are nearly identical between the 5 and 10 ppb NO experiments. The 665 relative contributions of higher-number clusters (which have been ordered according to 666 increasing $T_{m,50}$ increase for the 25 ppb NO experiment. This is consistent with the increased 667 persistence of the 25 ppb NO bulk thermogram to higher temperatures and the nearly identical 668 nature of the 5 ppb and 10 ppb NO bulk thermograms (Figure 8b). The clustering analysis suggests 669 that differences in the bulk thermogram arise from shifts in the relative contributions of the 670 various SOA components that result from the altered photochemical environment. These 671 observations generally suggest an increasing fraction of oligomeric content, or less-volatile 672 compounds, formed in the particle phase—or potentially the gas phase—when the SOA was 673 generated under higher chamber NO conditions (Schobesberger et al., 2018).

674

4.3.2. Multiple Clustering

675 With multiple clustering, each experiment was processed and clustered independently, with experiment-specific ξ_{ref} , N_c , and ε , among other parameters (Error! Reference source not 676 677 found. and **Table 3**). The clustered thermograms from the three experiments are compared in 678 Figure 10a-c. The number of clusters identified increases with NO concentration. Comparison 679 between the shapes of the clusters from the 5 ppb NO (Figure 10a) and 10 ppb NO (Figure 10b) 680 experiments indicates generally similar types of thermograms, consistent with the single 681 clustering method. Ten of the 11 total 10 ppb clusters match with a 5 ppb cluster. The one 682 additional, unique cluster at 10 ppb NO (Clst#9), is a one-member cluster with a sharp, narrow 683 peak at low temperatures and a broader, shallow second peak at high temperatures. This ion was 684 filtered out due to high noise level in the 5 ppb NO experiment.

The 25 ppb NO experiment (**Figure 10**c) results in more clusters compared to the lower NO experiments; 13 for the 25 ppb NO experiment versus 10 and 11 for the 5 and 10 ppb experiments, respectively. Some of the 25 ppb NO clusters have shapes similar to the lower NO experiments,

but many differ substantially. For example, two of the unique 25 ppb NO clusters (Clst#12 and
#13) have thermograms for which the signal increases continuously through the ramping period
and even into the soaking period. These clusters were not found in the single clustering analysis
because the 5 ppb NO experiment was used as the reference.

692 The new types of thermograms observed in the 25 ppb NO experiment indicates either 693 formation of new compounds or a change in the relative contributions of different components 694 to the same ions. Either could result from a change in the fate of the peroxy radical intermediates 695 as the NO concentration increases, leading to notably different products. There were numerous 696 nitrogen-containing ions observed for the three experiments. These N-containing ions belong to 697 Clst#1 – 7 for all the three [NO] conditions (Error! Reference source not found.). The higher-698 number clusters did not include N-containing ions, also indicating a limited influence of the 699 N-containing products on these lower-volatility thermograms, although fragmentation 700 complicates the interpretation. Overall, the formation of new N-containing compounds at the 701 high NO condition does not seem to explain the unique thermograms in the 25 ppb NO 702 experiments.

703 The percent contribution of different clusters to total mass, along with the noise-filtered 704 and unclustered ions, differ between experiments (Figure 10d). Note that for the multiple 705 clustering method, clusters having the same index number are not necessarily directly 706 comparable between experiments because different sets of ions are included. For example, while 707 Clst#1 in the 5 ppb and 10 ppb NO experiments are comparable, the most similar cluster in the 708 25 ppb experiment is Clst#2. Nonetheless, there are some common features shared by the same, 709 or closely indexed, clusters. For example, Clst#1 - 4 in all three experiments exhibit a narrow, 710 single peak with the peak temperature below 120 °C. The mass contribution of Clst#1 – 4 is similar 711 between the 5 and 10 ppb NO experiment, but ~15% lower in the 25 ppb NO experiment. Clusters 712 that reach their maximum signal at or above 150 °C (Clst#9, 10 for 5 ppb, Clst#10, 11 for 10 ppb 713 and Clst#10 – 13 for 25 ppb) together contribute ~6% in the low NO experiments and ~13% in 714 the high NO experiments. Thus, there is some evidence that at higher NO there is an increased 715 contribution of oligomeric compounds, indicated by the increased contribution of clusters that 716 peak at higher temperatures and exhibit broader overall thermograms. However, overall these

observations suggest complex shifts in the distribution of products, both monomeric and
oligomeric, with sufficient increases in NO to change the fate of the peroxy radical intermediates.

719

4.4. α -pinene + O₃ SOA

SOA formed from dark ozonolysis of α -pinene was collected and then allowed to isothermally evaporate for varying amounts of time (0 h, 1 h, 3 h, 6 h and 24 h) before thermal desorption (**Table 1**, Expt#4). As above for the SOA formed at varying NO concentrations, these experiments are considered as a set and interpreted using both the single-clustering and multiple-clustering approaches. The single-clustering approach uses the 0 h (no-wait) experiment as the reference for initial clustering. In this set of experiments, 312 ions were characterized by FIGAERO-CIMS for each experiment.

727

4.4.1. Single Clustering

728 Only a few ions, if any, were identified as anomalous in each experiment; a total of ten ions 729 were removed from all the experiments to maintain consistency between experiments. The mass 730 spectra and bulk thermograms of the remaining 302 ions for the five experiments are shown in 731 **Figure 11**. As the isothermal evaporation time increases, the mass spectrum changes significantly, 732 as previously reported by D'Ambro et al. (2018). In the no-wait experiment, the mass spectrum 733 is dominated by one ion, C₁₀H₁₄O₆. Upon isothermal evaporation, the relative abundance of this 734 ion notably decreases, with the extent of decrease increasing with wait time; over time, a greater 735 number of ions contribute to the total mass, both at lower and higher m/z. With isothermal 736 evaporation, the bulk thermograms also exhibit a shift from a more peaked shape, reminiscent 737 of that from a single compound (Lopez-Hilfiker et al., 2014), to a more flattened peak with a 738 shallower rise (Figure 11). In other words, with increasing isothermal evaporation the majority 739 of the mass desorbed during thermal desorption shifts from a lower to higher temperature region. 740 This behavior largely reflects the loss of comparably more volatile compounds during isothermal 741 evaporation, leaving behind SOA that is overall less volatile (Error! Reference source not found.a). 742 It can also in part be due to higher molecular weight, lower volatility compounds being produced 743 with time via accretion reactions in the condensed phase.

744 There are 12 clusters determined from the no-wait experiment, exhibiting a wide variety of 745 the shapes (Figure 12a), with the parameters used for data pre-processing and clustering 746 reported in **Table 3** and shown in Error! Reference source not found.. Focusing first on the no-747 wait experiment, the cluster thermogram shapes include those having clear peaks at relatively 748 low temperatures (~60 °C) and with a sharp rise and fall (e.g., Clst#1-3), those having sharp peaks 749 at relatively low temperatures but with a shallow downward slope (e.g., Clst#6), those with a broad peak at somewhat higher temperatures (~100 °C) and long tails (e.g., Clst#7), and those 750 751 having a wide peak at even higher temperatures ~120 °C with a very broad rise and fall (e.g., 752 Clst#10).

753 Changes to the shapes of the thermograms that occur upon isothermal evaporation differ 754 between the clusters. Some of the clusters exhibit almost step changes from the no-wait to the 755 longer time experiments (e.g., Clst#2 and 6), while others exhibit more continuous changes (e.g., 756 Clst#3 and 5). However, in all cases the clusters shift to have peaks that occur at higher 757 temperatures with generally broader thermograms. In other words, the T_{m50} of all the clusters 758 increase as a function of evaporation time, but with larger increases observed for the clusters 759 having initially lower $T_{m,50}$ (Figure 12b). For some of the clusters with a clear peak below 100 °C, 760 such as Clst#1–6, the peaks broaden to become less obvious and shift to higher temperatures 761 with longer isothermal evaporation. For clusters that originally have very wide peaks, such as 762 Clst#8–10 and 12, isothermal evaporation engenders a general shift in the thermograms towards 763 higher temperatures. Different from the clusters described above, thermograms for two clusters, 764 Clst#7 and Clst#11, exhibit only minor shift of peak temperature and shapes. Thermograms of 765 these two clusters share the common features of a moderate-width peak that reaches a 766 maximum between 100 – 120 °C. The T_{m50} of these two clusters correspondingly exhibit small 767 changes compared to other clusters.

Isothermal evaporation generally leads to a reduction of the monomeric character of clusters, leaving behind components that exhibit increased oligomeric content. Differences in how the individual cluster thermograms evolve with isothermal evaporation are therefore likely indicative of differing relative contributions of monomeric versus oligomeric components. For example, Clst#1 and Clst#10 have distinctly different shapes in the 0-h wait experiment, but very

773 similar shapes in the 24-h wait experiment. This indicates that ions in Clst#1 are not contributed 774 from a single component, as might be inferred from the single-mode peak in the 0-h wait 775 experiment. Instead, they are contributed by multiple components, though initially dominated 776 by monomeric compounds, so the shift in peak temperature and broadness is substantial. On the 777 other hand, ions in Clst#10 must also derive from multiple components, but with only a small 778 fraction of monomeric compounds that evaporate in the 24 hours. Consequently, the loss of 779 low-temperature mass is apparent yet small. In contrast, ions in clusters such as Clst#7 and 11 780 must be composed of only low-volatility components because they exhibit minimal changes in 781 the thermograms shapes.

782 The extent of mass loss with isothermal evaporation differs between clusters. In general, 783 clusters that exhibit larger changes in shape have greater total mass loss, although with variability 784 (Error! Reference source not found.c). Consequently, the mass contributions of the clusters 785 evolve with isothermal evaporation (Figure 12b). The contribution of Clst#1 decreases 786 significantly and most notably as wait time increases. The most prominent ion in the no-wait 787 experiment, $C_{10}H_{14}O_6$, is grouped in Clst#1. The continuous mass loss of Clst#1 indicates the rapid 788 evaporation of its members. The mass contributions of the other clusters that exhibited similar 789 changes in shape as Clst#1 (Clst#3, 5, and 6) remain comparably constant, although with Clst#3 790 decreasing slightly. The relative abundances of the clusters for which the thermograms shapes 791 changed negligibly (Clst#7 and 11) increase continually, implying of the slowest evaporation of 792 the ions in these two clusters in the 24-hr evaporation period.

793 For comparison, D'Ambro et al. (2018) reported changes in the shapes of the thermograms 794 for the five most abundant individual ions from the no-wait to 24-hr experiment, together 795 carrying ~15% of the particle mass. They observed the individual ion thermograms generally all 796 evolved in a manner similar to our Clst#1, 3 and 5, shifting from narrower, more peaked profiles 797 towards broader profiles with a shallower rise, less evident peak, and increased evaporation at higher temperatures. Here, with the clustering of data, we are able to track the change of thermal 798 799 behaviors of ions carrying ~87% of the initial mass. We are able to confirm that ~70 % of the mass 800 exhibit similar thermal behaviors and responses to isothermal evaporation as the top five ions. 801 However, we are also able to identify another ~17% of the mass having initial thermograms not

characterized by the top five ions, including 12% of the mass (Clst#7 and 11) that behavesdistinctly different upon evaporation at room temperature.

804 4.4.2. Multiple Clustering

805 The number of clusters identified with the multiple-clustering method, using experiment-806 specific optimal ε values (**Table 3** and Error! Reference source not found.), decreases with 807 isothermal evaporation time, from 13 (no-wait) to 12 (1 h) to 11 (3 h) and then to 9 (6 h and 24 808 h) (Figure 13b-f). The noise levels of the thermograms increase with evaporation time due to 809 decreasing absolute particle mass. Nonetheless, the typical shapes of the cluster-specific 810 thermograms clearly evolve with increasing isothermal evaporation. For short isothermal 811 evaporation times, many cluster-specific thermogram profiles are relatively narrow, peaking at 812 lower temperatures (70-120 °C) and with rapid rises and evident downslopes. For longer 813 isothermal evaporation times, the cluster-specific profiles instead have broad peaks with slow 814 rises and most of the mass desorbing at higher temperatures.

815 To aid further general interpretation, the cluster-specific thermograms with T_{m50} < 120 °C 816 are grouped together as higher-volatility clusters. The number of higher-volatility clusters 817 decreases with isothermal evaporation, from ten for the no-wait experiment, to five in the 1-h 818 experiment, two in the 3-h and 6-h experiment, to none in the 24-h experiment (Figure 14). The 819 mass contributions of the higher-volatility clusters decrease from 81.9% to 60.4%, 17.2%, 9.4% 820 and to 0.0%, with increasing isothermal evaporation time. This overall behavior is consistent with 821 results from the single-clustering method and indicates the compounds with a wide range of 822 volatilities make up much of the mass in the initial particles, while the SOA after isothermal 823 evaporation is composed of compounds having lower volatilities.

After isothermal evaporation, some cluster-specific thermograms have signals that increase continuously during the ramping period, for example Clst#11 and 12 in the 1-h experiment; such clusters were not observed in the no-wait experiment. The relative abundance of these very lowvolatility clusters increases with isothermal evaporation, from 1.7% in the 1-h experiment (Clst#11 and 12) to 13.4% in the 24-hr experiment (Clst#7 and 9). The absence of these clusters for the no-wait experiment suggests that they are formed over time through condensed-phase

reactions. Their increasing contribution over time may reflect both evaporation of higher
volatility components and continued formation. Clusters having thermograms with very broad
peaks, such as Clst#11 and 13 in the 0-h experiment are also observed in all the other experiments,
with increasing contribution to the total mass.

834 The multiple-clustering method reveals the disappearance of certain types of thermograms, 835 (e.g., the no-wait Clst#3) and the emergence of other types of thermograms (e.g., the 1-h Clst#11) 836 as evaporation time increases. This complements the single-clustering method, which illustrates 837 gradual changes in the shapes of cluster-specific thermograms, by allowing for identification of 838 completely new thermogram shapes and divergent behavior between ions within initial clusters. 839 The multiple-clustering method also confirms the decrease of the diversity of the desorption 840 profiles, as suggested by the single-clustering method. The two methods complement each other 841 and together provide a detailed look into (i) how the desorption profiles of sets of ions evolve 842 with isothermal evaporation and (ii) how the fraction of different types of thermograms change 843 with evaporation time.

5. Conclusions

845 We developed a new clustering algorithm, the noise-sorted scanning clustering (NSSC) 846 algorithm, for application to FIGAERO-CIMS data sets. The NSSC algorithm provides a robust 847 method for clustering of FIGAERO-CIMS thermograms having distinct thermal desorption profiles 848 and of determining the mass contribution of each cluster. Each of the ions contributing to a 849 cluster results from one or more molecules sharing similar thermochemical properties. These 850 molecules either evaporate directly or decompose and then evaporate. Compared to other 851 existing clustering algorithms, NSSC is strictly similarity-based, reproducible, and takes into 852 consideration differences in noise levels between individual ions. The application of NSSC has the 853 potential to make FIGAERO data more accessible to the atmospheric chemistry community.

For the four different SOA systems we examined, more than 80% of the total mass is clustered, with the number of clusters ranging from 9 to 13. The shapes of the cluster-specific average thermograms exhibit substantial variation for a given system. Some have relatively sharp peaks, others broad peaks with slowly decreasing signal as heating continues, and others still

having signals that continually increase up to very high temperatures or long desorption times.
The mass contribution of a cluster varies from 0.2% to 44.3%. A few (2-3) clusters usually contain
more than 50% of the total mass in all the chemical systems examined. Comparison of the clusterspecific thermogram shapes between different SOA systems allows for qualitative assessment of
the similarity or uniqueness.

We also demonstrated the potential of the NSSC for guiding interpretation of sets of 863 864 experiments where one experimental condition varies (e.g., NO concentration and evaporation 865 time). For such experiments, two complementary methods are suggested: (i) the single clustering 866 method, where one experiment is used to determine the ions belonging to individual clusters 867 and then clusters comprising the same ions are calculated for the other experiments, and (ii) the 868 multiple clustering method, where each experiment is clustered independently and then 869 compared. The first approach helps establish how the properties of individual clusters evolve as 870 a set, while the second approach helps identify changes in the diversity of cluster-specific 871 thermogram shapes, properties, and mass contributions. The two approaches complement each 872 other and provide guidance for future efforts to cluster ambient observations having long time-873 series.

874 This paper focuses only on the description of the clustering algorithm and its potential as a 875 tool to characterize the thermal properties of organic aerosol in further detail. The application of 876 NSSC can be potentially expanded to any other composition-resolved data sets, such as diurnal 877 changes of different compounds measured in ambient air, temporal changes of different 878 generations of species in a smog chamber, and composition-dependent size distributions. All of 879 the above data sets share a common property that the noise of the curve/spectrum is related to 880 the composition. Therefore, NSSC would facilitate the analysis by taking noise into consideration. 881 Interpretation of the cluster-specific thermograms using frameworks such as that of 882 Schobesberger et al. (2018) will allow for more comprehensive understanding of the 883 thermochemical properties of the organic aerosol, the subject of future work. This will provide 884 insights into the thermal behavior of organic aerosol and the relative contributions of thermally 885 stable (e.g., monomer) versus thermally unstable (e.g., dimers or oligomers) compounds, the

volatility distribution of the thermally stable compounds, and the T-dependent rate coefficientsfor oligomer dissociation and formation.

888 **6. Data Availability**

All data and the NSSC algorithm used in this publication are archived in the UC DASH data repository (Cappa et al., 2019). The NSSC algorithm is also available at GitHub (<u>https://github.com/chriscappa/NSSC</u>), with the version used for this publication available as Li and Cappa (2019).

893

7. Author Contributions

ZL developed the NSSC algorithm. ELD, SS, CJG, FDL-H, JL, JES, and ZL performed measurements. ELD and SS performed detailed data processing. ZL and CDC analyzed data and wrote the manuscript, with contributions from all co-authors.

897 8. Acknowledgements

898 This work was supported by the National Science Foundation under Grant No. ATM-899 1151062. The experimental work described here was supported by the U.S. Department of 900 Energy ASR grants DE-SC0011791 and DE-SC0018221. E.L.D. was supported by the National 901 Science Foundation Graduate Research Fellowship (grant no. DGE-1256082) and S.S. was supported by the Academy of Finland (grant nos. 272041 and 310682). The SOAFFEE campaign 902 903 was done at Pacific Northwest National Laboratory, supported by the U.S. Department of Energy 904 (DOE) Office of Science, Office of Biological and Environmental Research, as part of the 905 Atmospheric Systems Research (ASR) program. PNNL is operated for DOE by Battelle Memorial 906 Institute under contract DE-AC05-76RL01830.

907 **9. References**

Abdalmogith, S. S., and Harrison, R. M.: The use of trajectory cluster analysis to examine the long range transport of secondary inorganic aerosol in the UK, Atmos Environ, 39, 6686-6695,
 https://doi.org/10.1016/j.atmosenv.2005.07.059, 2005.

Beddows, D. C. S., Dall'Osto, M., and Harrison, R. M.: Cluster Analysis of Rural, Urban, and
Curbside Atmospheric Particle Size Data, Environ Sci Technol, 43, 4694-4700,
https://doi.org/10.1021/es803121t, 2009.

Cape, J. N., Methven, J., and Hudson, L. E.: The use of trajectory cluster analysis to interpret trace
gas measurements at Mace Head, Ireland, Atmos Environ, 34, 3651-3663,
https://doi.org/10.1016/S1352-2310(00)00098-4, 2000.

- 917 Cappa, C. D., Li, Z., D'Ambro, E. L., Schobesberger, S., Shilling, J. E., Lopez-Hilfiker, F., Liu, J., Gaston,
- 918 C. J., and Thornton, J. A.: Initial application of the noise-sorted scanning clustering algorithm to
- 919 the analysis of composition-dependent organic aerosol thermal desorption measurements, UC
 920 Davis Dash, Dataset, <u>https://doi.org/10.25338/B87S43</u>, 2019
- 921 D'Ambro, E. L., Lee, B. H., Liu, J. M., Shilling, J. E., Gaston, C. J., Lopez-Hilfiker, F. D., Schobesberger,
- 922 S., Zaveri, R. A., Mohr, C., Lutz, A., Zhang, Z. F., Gold, A., Surratt, J. D., Rivera-Rios, J. C., Keutsch,
- 923 F. N., and Thornton, J. A.: Molecular composition and volatility of isoprene photochemical
- 924 oxidation secondary organic aerosol under low- and high-NOx conditions, Atmospheric Chemistry
- 925 and Physics, 17, 159-174, <u>https://doi.org/10.5194/acp-17-159-2017</u>, 2017.
- 926 D'Ambro, E. L., Schobesberger, S., Zaveri, R. A., Shilling, J. E., Lee, B. H., Lopez-Hilfiker, F. D., Mohr,
- C., and Thornton, J. A.: Isothermal Evaporation of alpha-Pinene Ozonolysis SOA: Volatility, Phase
 State, and Oligomeric Composition, Acs Earth Space Chem, 2, 1058-1067,
 https://doi.org/10.1021/acsearthspacechem.8b00084, 2018.
- 930 D'Ambro, E. L., Schobesberger, S., Gaston, C. J., Lopez-Hilfiker, F. D., Lee, B. H., Liu, J., Zelenyuk,
- A., Bell, D., Cappa, C. D., Helgestad, T., Li, Z., Guenther, A., Wang, J., Wise, M., Caylor, R., Surratt,
- 932 J. D., Riedel, T., Hyttinen, N., Salo, V. T., Hasan, G., Kurtén, T., Shilling, J. E., and Thornton, J. A.:
- 933 Chamber-based insights into the factors controlling IEPOX SOA yield, composition, and volatility,
- 934 Atmos. Chem. Phys. Discuss., 2019, 1-20, <u>https://doi.org/10.5194/acp-2019-271</u>, 2019.
- Faxon, C., Hammes, J., Le Breton, M., Pathak, R. K., and Hallquist, M.: Characterization of organic
 nitrate constituents of secondary organic aerosol (SOA) from nitrate-radical-initiated oxidation
 of limonene using high-resolution chemical ionization mass spectrometry, Atmospheric
 Chemistry and Physics, 18, 5467-5481, https://doi.org/10.5194/acp-18-5467-2018, 2018.
- 939 Gaston, C. J., Quinn, P. K., Bates, T. S., Gilman, J. B., Bon, D. M., Kuster, W. C., and Prather, K. A.:
- 940 The impact of shipping, agricultural, and urban emissions on single particle chemistry observed 941 aboard the R/V Atlantis during CalNex, J Geophys Res-Atmos, 118, 5003-5017, 942 https://doi.org/10.1002/jord.50427.2012
- 942 <u>https://doi.org/10.1002/jgrd.50427</u>, 2013.
- Gaston, C. J., Lopez-Hilfiker, F. D., Whybrew, L. E., Hadley, O., McNair, F., Gao, H. L., Jaffe, D. A.,
- and Thornton, J. A.: Online molecular characterization of fine particulate matter in Port Angeles,
- WA: Evidence for a major impact from residential wood smoke, Atmos Environ, 138, 99-107,
 https://doi.org/10.1016/j.atmosenv.2016.05.013, 2016.
- 947 Giorio, C., Tapparo, A., Dall'Osto, M., Harrison, R. M., Beddows, D. C. S., Di Marco, C., and Nemitz,
- 948 E.: Comparison of three techniques for analysis of data from an Aerosol Time-of-Flight Mass
- 949 Spectrometer, Atmos Environ, 61, 316-326, <u>https://doi.org/10.1016/j.atmosenv.2012.07.054</u>, 950 2012.
- 951 Goldstein, A. H., and Galbally, I. E.: Known and unexplored organic constituents in the earth's
- 952 atmosphere, Environ Sci Technol, 41, 1514-1521, <u>https://doi.org/10.1021/es072476p</u>, 2007.
- 953 Gonzalez, T. F.: Clustering to Minimize the Maximum Intercluster Distance, Theor Comput Sci, 38,
- 954 293-306, <u>https://doi.org/10.1016/0304-3975(85)90224-5</u>, 1985.

- Hamilton, J. F., Webb, P. J., Lewis, A. C., Hopkins, J. R., Smith, S., and Davy, P.: Partially oxidised
 organic components in urban aerosol using GCXGC-TOF/MS, Atmospheric Chemistry and Physics,
 4, 1279-1290, <u>https://doi.org/10.5194/acp-4-1279-2004</u>, 2004.
- 958 Huang, W., Saathoff, H., Pajunoja, A., Shen, X. L., Naumann, K. H., Wagner, R., Virtanen, A., Leisner,
- T., and Mohr, C.: alpha-Pinene secondary organic aerosol at low temperature: chemical
 composition and implications for particle viscosity, Atmospheric Chemistry and Physics, 18, 28832898, <u>https://doi.org/10.5194/acp-18-2883-2018</u>, 2018.
- 962 Isaacman-VanWertz, G., Massoli, P., O'Brien, R. E., Nowak, J. B., Canagaratna, M. R., Jayne, J. T., 963 Worsnop, D. R., Su, L., Knopf, D. A., Misztal, P. K., Arata, C., Goldstein, A. H., and Kroll, J. H.: Using 964 advanced mass spectrometry techniques to fully characterize atmospheric organic carbon: 965 capabilities and remaining gaps, Faraday Discussions, 200, 579-598, current 966 https://doi.org/10.1039/c7fd00021a, 2017.
- Joo, T., Rivera-Rios, J. C., Takeuchi, M., Alvarado, M. J., and Ng, N. L.: Secondary Organic Aerosol
 Formation from Reaction of 3-Methylfuran with Nitrate Radicals, Acs Earth Space Chem,
 https://doi.org/10.1021/acsearthspacechem.9b00068, 2019.
- Kirchner, U., Vogt, R., Natzeck, C., and Goschnick, J.: Single particle MS, SNMS, SIMS, XPS, and
 FTIR spectroscopic analysis of soot particles during the AIDA campaign, Journal of Aerosol Science,
 34, 1323-1346, https://doi.org/10.1016/S0021-8502(03)00362-8, 2003.
- Le Breton, M., Psichoudaki, M., Hallquist, M., Watne, A. K., Lutz, A., and Hallquist, A. M.:
 Application of a FIGAERO ToF CIMS for on-line characterization of real-world fresh and aged
 particle emissions from buses, Aerosol Science and Technology, 53, 244-259,
 <u>https://doi.org/10.1080/02786826.2019.1566592</u>, 2019.
- Lee, A. K. Y., Willis, M. D., Healy, R. M., Onasch, T. B., and Abbatt, J. P. D.: Mixing state of
 carbonaceous aerosol in an urban environment: single particle characterization using the soot
 particle aerosol mass spectrometer (SP-AMS), Atmospheric Chemistry and Physics, 15, 18231841, <u>https://doi.org/10.5194/acp-15-1823-2015</u>, 2015.
- Lee, B., Lopez-Hilfiker, F. D., D'Ambro, E. L., Zhou, P. T., Boy, M., Petaja, T., Hao, L. Q., Virtanen,
 A., and Thornton, J. A.: Semi-volatile and highly oxygenated gaseous and particulate organic
 compounds observed above a boreal forest canopy, Atmospheric Chemistry and Physics, 18,
- 984 11547-11562, <u>https://doi.org/10.5194/acp-18-11547-2018</u>, 2018.
- Lee, B. H., Lopez-Hilfiker, F. D., Mohr, C., Kurten, T., Worsnop, D. R., and Thornton, J. A.: An Iodide-
- Adduct High-Resolution Time-of-Flight Chemical-Ionization Mass Spectrometer: Application to
 Atmospheric Inorganic and Organic Compounds, Environ Sci Technol, 48, 6309-6317,
 <u>https://doi.org/10.1021/es500362a</u>, 2014.
- Lee, B. H., Mohr, C., Lopez-Hilfiker, F. D., Lutz, A., Hallquist, M., Lee, L., Romer, P., Cohen, R. C., Iyer, S., Kurten, T., Hu, W. W., Day, D. A., Campuzano-Jost, P., Jimenez, J. L., Xu, L., Ng, N. L., Guo,
- H. Y., Weber, R. J., Wild, R. J., Brown, S. S., Koss, A., de Gouw, J., Olson, K., Goldstein, A. H., Seco,
- 92 R., Kim, S., McAvey, K., Shepson, P. B., Starn, T., Baumann, K., Edgerton, E. S., Liu, J. M., Shilling,
- 93 J. E., Miller, D. O., Brune, W., Schobesberger, S., D'Ambro, E. L., and Thornton, J. A.: Highly
- 994 functionalized organic nitrates in the southeast United States: Contribution to secondary organic
 995 aerosol and reactive nitrogen budgets, P Natl Acad Sci USA, 113, 1516-1521,
 996 https://doi.org/10.1073/pnas.1508108113, 2016.
- Li, Z., and Cappa, C. D.: Noise Sorted Scanning Clustering Algorithm (Version v1.0.3), Zenodo,
 https://doi.org/10.5281/zenodo.3361797, 2019

- 999 Liu, J. M., D'Ambro, E. L., Lee, B. H., Lopez-Hilfiker, F. D., Zaveri, R. A., Rivera-Rios, J. C., Keutsch,
- F. N., Iyer, S., Kurten, T., Zhang, Z. F., Gold, A., Surratt, J. D., Shilling, J. E., and Thornton, J. A.:
 Efficient Isoprene Secondary Organic Aerosol Formation from a Non-IEPDX Pathway, Environ Sci
- 1002 Technol, 50, 9872-9880, https://doi.org/10.1021/acs.est.6b01872, 2016.
- Liu, S., Shilling, J. E., Song, C., Hiranuma, N., Zaveri, R. A., and Russell, L. M.: Hydrolysis of Organonitrate Functional Groups in Aerosol Particles, Aerosol Science and Technology, 46, 1359-1369, <u>https://doi.org/10.1080/02786826.2012.716175</u>, 2012.
- Liu, S., Russell, L. M., Sueper, D. T., and Onasch, T. B.: Organic particle types by single-particle measurements using a time-of-flight aerosol mass spectrometer coupled with a light scattering module, Atmospheric Measurement Techniques, 6, 187-197, <u>https://doi.org/10.5194/amt-6-</u> 1009 <u>187-2013</u>, 2013.
- Lopez-Hilfiker, F. D., Mohr, C., Ehn, M., Rubach, F., Kleist, E., Wildt, J., Mentel, T. F., Lutz, A.,
 Hallquist, M., Worsnop, D., and Thornton, J. A.: A novel method for online analysis of gas and
 particle composition: description and evaluation of a Filter Inlet for Gases and AEROsols
 (FIGAERO), Atmospheric Measurement Techniques, 7, 983-1001, https://doi.org/10.5194/amt-
- 1014 <u>7-983-2014</u>, 2014.
- Lopez-Hilfiker, F. D., Mohr, C., Ehn, M., Rubach, F., Kleist, E., Wildt, J., Mentel, T. F., Carrasquillo,
 A. J., Daumit, K. E., Hunter, J. F., Kroll, J. H., Worsnop, D. R., and Thornton, J. A.: Phase partitioning
 and volatility of secondary organic aerosol components formed from α-pinene ozonolysis and OH
 oxidation: the importance of accretion products and other low volatility compounds,
 Atmospheric Chemistry and Physics, 15, 7765-7776, https://doi.org/10.5194/acp-15-7765-2015,
 2015.
- 1021 Lopez-Hilfiker, F. D., Mohr, C., D'Ambro, E. L., Lutz, A., Riedel, T. P., Gaston, C. J., Iyer, S., Zhang,
- Z., Gold, A., Surratt, J. D., Lee, B. H., Kurten, T., Hu, W. W., Jimenez, J., Hallquist, M., and Thornton,
 J. A.: Molecular Composition and Volatility of Organic Aerosol in the Southeastern U.S.:
 Implications for IEPOX Derived SOA, Environ Sci Technol, 50, 2200-2209,
- 1025 <u>https://doi.org/10.1021/acs.est.5b04769</u>, 2016.
- 1026 Mohr, C., Lopez-Hilfiker, F. D., Yli-Juuti, T., Heitto, A., Lutz, A., Hallquist, M., D'Ambro, E. L.,
- 1027 Rissanen, M. P., Hao, L. Q., Schobesberger, S., Kulmala, M., Mauldin, R. L., Makkonen, U., Sipila,
- M., Petaja, T., and Thornton, J. A.: Ambient observations of dimers from terpene oxidation in the
 gas phase: Implications for new particle formation and growth, Geophysical Research Letters, 44,
- 1030 2958-2966, <u>https://doi.org/10.1002/2017gl072718</u>, 2017.
- 1031 Murphy, D. M., Middlebrook, A. M., and Warshawsky, M.: Cluster analysis of data from the 1032 Particle Analysis by Laser Mass Spectrometry (PALMS) instrument, Aerosol Science and 1033 Technology, 37, 382-391, <u>https://doi.org/10.1080/02786820300971</u>, 2003.
- Pinero-Garcia, F., Ferro-Garcia, M. A., Chham, E., Cobos-Diaz, M., and Gonzalez-Rodelas, P.: A
 cluster analysis of back trajectories to study the behaviour of radioactive aerosols in the southeast of Spain, J Environ Radioactiv, 147, 142-152, <u>https://doi.org/10.1016/j.jenvrad.2015.05.029</u>,
 2015.
- 1038 Praske, E., Otkjaer, R. V., Crounse, J. D., Hethcox, J. C., Stoltz, B. M., Kjaergaard, H. G., and
- 1039 Wennberg, P. O.: Atmospheric autoxidation is increasingly important in urban and suburban
- 1040 North America, P Natl Acad Sci USA, 115, 64-69, https://doi.org/10.1073/pnas.1715540115, 2018.

- 1041 Rebotier, T. P., and Prather, K. A.: Aerosol time-of-flight mass spectrometry data analysis: A 1042 benchmark of clustering algorithms, Anal Chim Acta, 585, 38-54, 1043 https://doi.org/10.1016/j.aca.2006.12.009, 2007.
- Reitz, P., Zorn, S. R., Trimborn, S. H., and Trimborn, A. M.: A new, powerful technique to analyze single particle aerosol mass spectra using a combination of OPTICS and the fuzzy c-means algorithm, Journal of Aerosol Science, 98, 1-14, <u>https://doi.org/10.1016/j.jaerosci.2016.04.003</u>, 2016.
- 1048 Roth, A., Schneider, J., Klimach, T., Mertes, S., van Pinxteren, D., Herrmann, H., and Borrmann, S.:
- Aerosol properties, source identification, and cloud processing in orographic clouds measured by
 single particle mass spectrometry on a central European mountain site during HCCT-2010,
 Atmospheric Chemistry and Physics, 16, 505-524, https://doi.org/10.5194/acp-16-505-2016,
- 1052 2016.
- Schobesberger, S., D'Ambro, E. L., Lopez-Hilfiker, F. D., Mohr, C., and Thornton, J. A.: A model framework to retrieve thermodynamic and kinetic properties of organic aerosol from composition-resolved thermal desorption measurements, Atmospheric Chemistry and Physics, 18, 14757-14785, https://doi.org/10.5194/acp-18-14757-2018, 2018.
- 1057 Song, X. H., Hopke, P. K., Fergenson, D. P., and Prather, K. A.: Classification of single particles 1058 analyzed by ATOFMS using an artificial neural network, ART-2A, Anal Chem, 71, 860-865, 1059 https://doi.org/10.1021/ac9809682, 1999.
- 1060 Stolzenburg, D., Fischer, L., Vogel, A. L., Heinritzi, M., Schervish, M., Simon, M., Wagner, A. C., 1061 Dada, L., Ahonen, L. R., Amorim, A., Baccarini, A., Bauer, P. S., Baumgartner, B., Bergen, A., Bianchi,
- 1062 F., Breitenlechner, M., Brilke, S., Mazon, S. B., Chen, D. X., Dias, A., Draper, D. C., Duplissy, J.,
- 1063 Haddad, I., Finkenzeller, H., Frege, C., Fuchs, C., Garmash, O., Gordon, H., He, X., Helm, J.,
- 1064 Hofbauer, V., Hoyle, C. R., Kim, C., Kirkby, J., Kontkanen, J., Kuerten, A., Lampilahti, J., Lawler, M.,
- 1065 Lehtipalo, K., Leiminger, M., Mai, H., Mathot, S., Mentler, B., Molteni, U., Nie, W., Nieminen, T.,
- 1066 Nowak, J. B., Ojdanic, A., Onnela, A., Passananti, M., Petaja, T., Quelever, L. L. J., Rissanen, M. P.,
- 1067 Sarnela, N., Schallhart, S., Tauber, C., Tome, A., Wagner, R., Wang, M., Weitz, L., Wimmer, D.,
- 1068 Xiao, M., Yan, C., Ye, P., Zha, Q., Baltensperger, U., Curtius, J., Dommen, J., Flagan, R. C., Kulmala,
- 1069 M., Smith, J. N., Worsnop, D. R., Hansel, A., Donahue, N. M., and Winkler, P. M.: Rapid growth of 1070 organic aerosol nanoparticles over a wide tropospheric temperature range, P Natl Acad Sci USA, 115, 0122, 0127, https://doi.org/10.1072/page.1007501115, 2018
- 1071 115, 9122-9127, <u>https://doi.org/10.1073/pnas.1807604115</u>, 2018.
- 1072Takahama, S., Gilardoni, S., Russell, L. M., and Kilcoyne, A. L. D.: Classification of multiple types1073of organic carbon composition in atmospheric particles by scanning transmission X-ray1074microscopy1075https://doi.org/10.1016/j.atmosenv.2007.08.051, 2007.
- 1076 Wang, D. S., and Ruiz, L. H.: Chlorine-initiated oxidation of n-alkanes under high-NOx conditions:
 1077 insights into secondary organic aerosol composition and volatility using a FIGAERO-CIMS,
 1078 Atmospheric Chemistry and Physics, 18, 15535-15553, https://doi.org/10.5194/acp-18-15535-
- 1079 2018, 2018.
- 1080 Wegner, T., Hussein, T., Hameri, K., Vesala, T., Kulmala, M., and Weber, S.: Properties of aerosol
- 1081 signature size distributions in the urban environment as derived by cluster analysis, Atmos
- 1082 Environ, 61, 350-360, https://doi.org/10.1016/j.atmosenv.2012.07.048, 2012.

1083Zhao, W. X., Hopke, P. K., and Prather, K. A.: Comparison of two cluster analysis methods using1084singleparticlemassspectra,AtmosEnviron,42,881-892,1085https://doi.org/10.1016/j.atmosenv.2007.10.024, 2008.

1086 Zhao, Y., Thornton, J. A., and Pye, H. O. T.: Quantitative constraints on autoxidation and dimer

- formation from direct probing of monoterpene-derived peroxy radical chemistry, P Natl Acad Sci
 USA, 115, 12142-12147, <u>https://doi.org/10.1073/pnas.1812147115</u>, 2018.
- 1089 Zhou, L. M., Hopke, P. K., and Venkatachari, P.: Cluster analysis of single particle mass spectra
- 1090 measured at Flushing, NY, Anal Chim Acta, 555, 47-56, https://doi.org/10.1016/j.aca.2005.08.061,
- 1091 2006.
- 1092

10. Tables

Exp #	Precursor		Oxidant		Seeds			т	RH	NO ^{#\$}	<i>M</i> _p ^{#&}	FIGAERO
	Туре	Conc. [#] (ppb)	Туре	Conc. ^{##} (ppm)	Туре	<i>D_p</i> #* (nm)	UV	(°C)	(%)	(ppb)	(μg/m ³)	Operation \$
1*	lpha-pinene	10	OH (H ₂ O ₂)	1.0	AS ^{&}	50	On	25	50	-	5.1	Normal
2	Δ -3-carene	10	OH (H ₂ O ₂)	0.25	AS	50	On	25	50	-	5.2	Normal
3a		10	OH (H ₂ O ₂)	1.0	AS	50	On	25	50	5	8.3	Normal
3b	α -pinene									10	9.2	
3c										25	9.1	
4a	_	e 10	O ₃	0.1	PS ^{&&}	50	Off	25	80	-	4.0	Normal
4b	_											1 h wait
4c	α -pinene											3 h wait
4d												6 h wait
4e												24 h wait

Table 1. Details of SOA formation and chamber conditions for all the example SOA systems.

* Experiment #1 is a case study used to test the performances of different clustering algorithms

[#] Conc. of precursors are the concentrations expected in the chamber with the absence of any chemistry

^{##} For OH, conc. refers to concentration of H_2O_2 injected into the chamber; for O_3 , conc. refers to steady-state concentration of O_3 in the chamber during SOA formation

** Seed particles are size-selected in all the experiments

^{#\$} NO concentration refers to the targeted NO concentration when NO is injected into the chamber. The actual steady-state concentration of NO is lower than targeted. "-" indicates that no external NO is added to the chamber

^{#&} M_p is the estimated mass concentration of particles including SOA and seeds measured by SMPS when the chamber is at steady-state, except for experiment 4 where M_p is the mass concentration of SOA only

^{\$} Normal operation mode means the desorption process starts immediately after collection period. X h wait means that particles are isothermally diluted for X hours before the desorption process is initiated

[&] AS = ammonium sulfate

^{&&} PS = potassium sulfate

•		0 0				
Clustering Algorithms	k-means	k-medoids	Mean-shift	DBSCAN	FPClustering	NSSC
Assign all the members?	Yes	Yes	No	No	Yes	No
Identify single-member clusters?	No	No	Yes	No	No	Yes
Robust solution?	No	No	No	Yes	No	Yes
Controlled distance from the center of clusters?	No	No	Yes	No	No	Yes
Influence of noise?	large	large	small	small	large	Small
Key preset parameters	Nc	Nc	ε, N _{min}	3	Initial seed	ε, N _{min}
Software used in this study	lgor	R	Python	lgor	lgor	lgor

Table 2. Comparison of different clustering algorithms

Expt #			Pre-processing						Clustering				
	SOA type		N _{total}	Nanomalous	N filtered	$f_{\sf m, filtered}$	ξref	$\mathbf{f}_{m,ref}$	3	Nc	N _{c,one}	$f_{\sf m, unclustered}$	RinterClst
1	α-pinene + OH		298	4	188	7.5	0.021	0.67	2.6	11	0	0.00	2.01
2	Δ -3-carene + OH		298	5	183	9.3	0.019	0.57	2.1	9	1	0.27	2.36
3a				6	204	15.3	0.025	0.55	2.2	9	1	1.52	2.06
3b		Single	298	6	204	17.5	-	-	-	9	1	1.72	-
3c	α -pinene +			6	204	21.0	-	-	-	9	1	2.27	-
3a	OH + NO)		2	208	15.5	0.025	0.55	2.2	9	1	1.52	2.06
3b		Multi	298	3	195	12.6	0.027	0.54	2.3	10	1	1.29	2.10
3c				6	200	12.8	0.028	0.43	2.5	12	1	1.21	1.96
4a				10	185	11.5	0.025	0.42	2.2	10	2	0.67	2.28
4b				10	185	14.0	-	-	-	10	2	0.79	-
4c		Single	312	10	185	14.0	-	-	-	10	2	0.84	-
4d				10	185	13.8	-	-	-	10	2	0.83	-
4e	lpha-pinene +			10	185	17.6	-	-	-	10	2	0.82	-
4a	O ₃			1	191	11.4	0.025	0.41	2.2	11	2	1.04	2.22
4b				0	210	16.5	0.044	0.41	3.3	8	4	0.00	2.02
4c		Multi	312	5	205	14.3	0.048	0.42	3.1	9	2	1.06	1.66
4d				3	203	12.8	0.055	0.39	3.3	8	1	2.50	1.80
4e				3	213	16.1	0.053	0.41	3.4	7	2	0.98	1.97

Table 3. Parameters and thresholds used for the data processing and noise-sorted scanning clustering for all the example experiments.

 N_{total} – Total number of ions characterized by CIMS

 $N_{\text{anomalous}}$ – Number of anomalous ions

N_{filtered} – Number of ions filtered out from the following clustering due to high levels of noises

 $f_{m,filtered}$ – Mass fraction of the ions filtered out due to high levels of noises, expressed in %

 ξ_{ref} – Noise threshold. Ions with noise levels above this threshold are excluded from clustering

 $f_{m,ref}$ – The threshold of mass contribution (%) to identify an ion as significant

 ϵ – distance criterion

 $N_{\rm c}$ – Number of clusters determined with two or more members

 $N_{c,one}$ – Number of clusters determined with only one member

 $f_{\rm m,unclustered}$ – Mass fraction of unclustered ions, expressed in %

 $R_{\text{interClst}}$ – The ratio of the average inter-cluster distance over the distance criterion ϵ

11. Figures

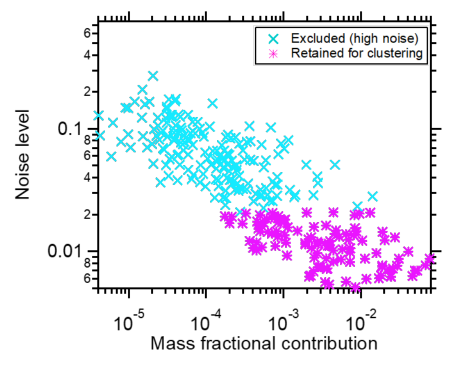


Figure 1: The relationship between thermogram noise levels and the fractional contributions of the corresponding ions to total mass, for α -pinene + OH SOA. The noise threshold, $\xi_{ref} = 0.021$ and is used to distinguish high-noise thermograms (cyan markers) from thermograms having acceptable noise levels (pink markers).

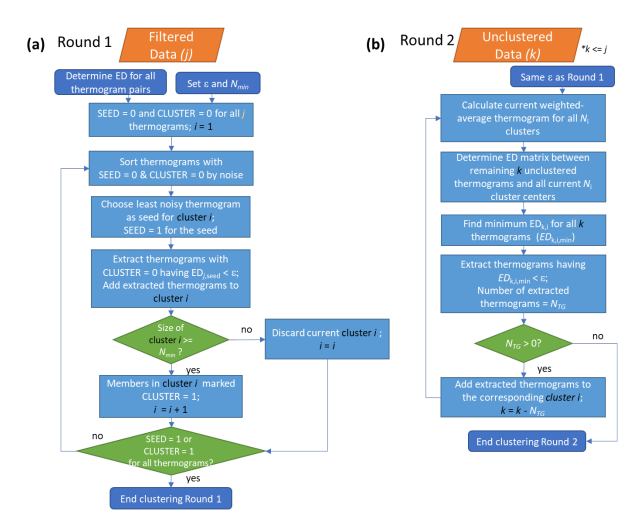


Figure 2: Flow of the noise-sorted scanning clustering. There are two rounds of clustering. (a) Round 1: The ED between all thermogram pairs are calculated and two parameters, ε and N_{min}, are set. Each thermogram is initialized with state SEED = 0 and CLUSTER = 0. Only thermograms with SEED = 0 and CLUSTER = 0 can be added to new clusters. The procedure terminates when all the thermograms are marked either SEED = 1 or CLUSTER = 1. (b) Round 2: Seeds are specified as the weighted-average thermogram for each cluster, and any remaining unclustered thermograms from Round 1 are potentially added to these clusters. With the indexing, *j* refers to the total number of thermograms, *i* to the number of clusters, and *k* to the number of unclustered thermograms after Round 1.

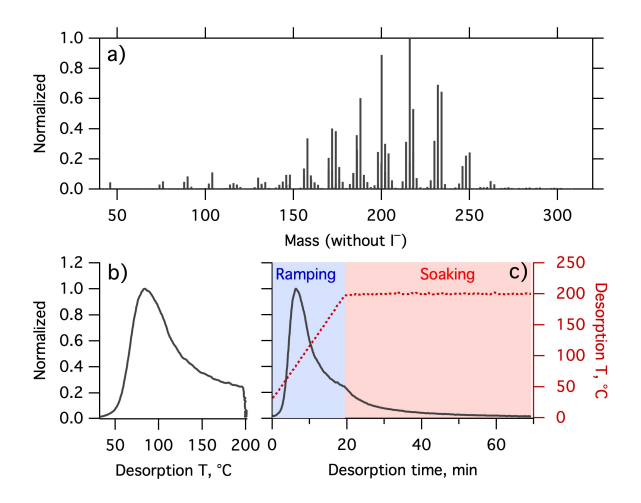


Figure 3. (a) Mass spectrum of α -pinene + OH SOA measured by FIGAERO-CIMS. The mass excludes iodine. (b) Normalized thermogram of the bulk SOA versus temperature. (c) Normalized thermogram of the bulk SOA versus time (black line) and the variation in desorption temperature with time (dark red dashed line). The long tail during the soaking period is evident when the thermogram is considered in time space. The light blue shaded area denotes the ramping period and the pink shaded area the soaking period.

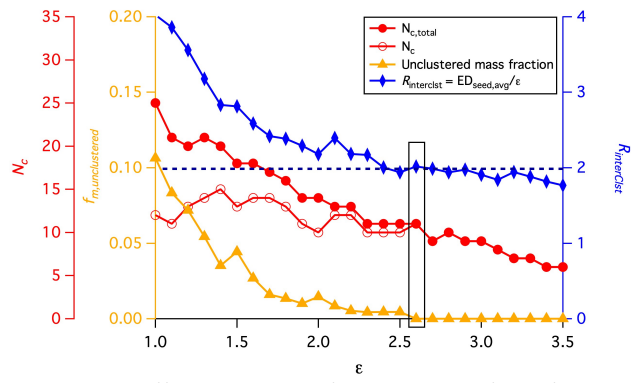


Figure 4. The variation of four parameters, N_c, N_{c,total}, f_{m,unclustered} and R_{interClst} as a function of the distance criterion ε . The black horizontal dashed line guides the judgement for R_{interClst} ≥ 2 . The values highlighted by a rectangle are the values corresponding to the optimal ε used for the clustering analysis.

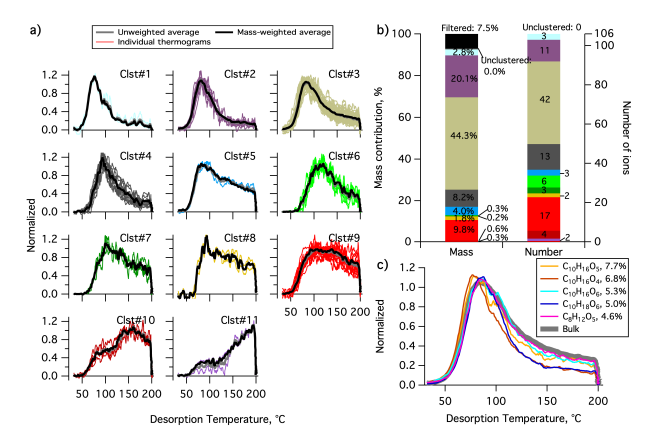


Figure 5. Clustering results for α -pinene + OH SOA. (a) Unweighted average thermograms (bold grey lines), mass-weighted average thermograms (bold black lines) and individual members (colored lines) of the 11 clusters identified. (b) Percentage contribution of each cluster to the total mass, as well as the filtered out and unclustered mass percentage (left bar), and the number of ions in each cluster and the unclustered number of ions (right bar). (c) Thermograms of the top 5 ions in terms of mass contribution. The cluster colors are consistent between (a) and (b).

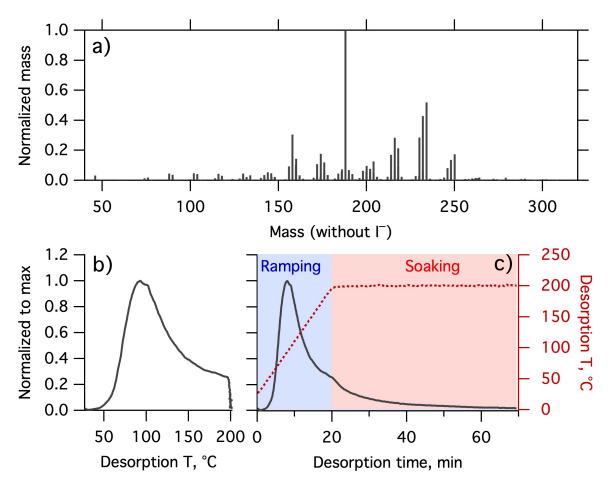


Figure 6. Same as Figure 3, but for Δ -3-carene + OH SOA. (a) SOA mass spectrum measured by FIGAERO-CIMS. The mass excludes iodine. The normalized thermogram of the bulk SOA versus (b) temperature and (c) time. In (c) the light blue shaded area denotes the ramping period and the pink shaded area the soaking period.

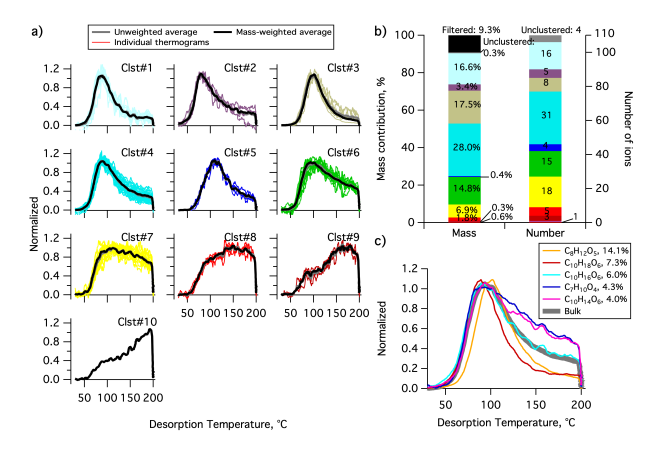


Figure 7. Same as Figure 5, but for Δ -3-carene + OH SOA. (a) Unweighted average thermograms (bold grey lines), mass-weighted average thermograms (bold black lines) and individual members (colored lines) of the ten clusters identified. (b) Percentage contribution of each cluster to the total mass, as well as the filtered out and unclustered mass percentage (left bar) and number of ions in each cluster and the unclustered number of ions (right bar). (c) Thermograms of the top 5 ions in terms of mass contribution. The cluster colors are consistent between (a) and (b).

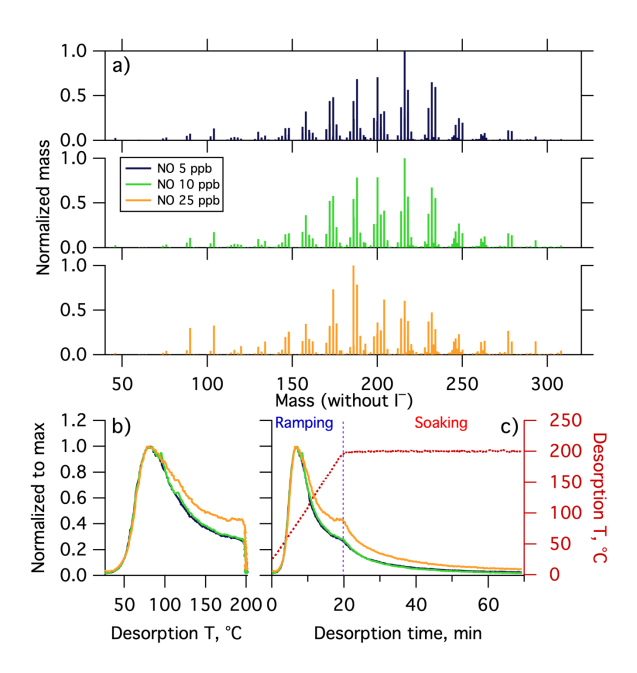


Figure 8. (a) Mass spectra of α -pinene + OH SOA formed with different NO concentrations, normalized to the most abundant ions mass concentration. The mass excludes iodine. Normalized thermograms of the bulk SOA versus (b) temperature and (c) desorption time, with the desorption temperature shown in dark red dashed line. The vertical purple dashed line delineates between ramping and soaking. In all the panels, colors correspond to the NO concentration (see legend).

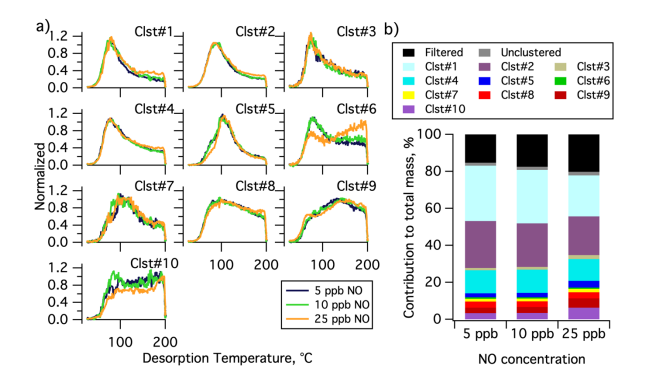


Figure 9. Single clustering results for α -pinene + OH SOA as a function of NO concentration. (a) Comparison of the normalized, weighted average thermograms of the ten clusters for the 5 ppb NO (navy), 10 ppb NO (green) and 25 ppb NO (orange) experiments. (b) Contribution of each cluster to the total mass, including the contribution from filtered out ions (black) and unclustered ions (gray). The total mass is calculated independently for each experiment.

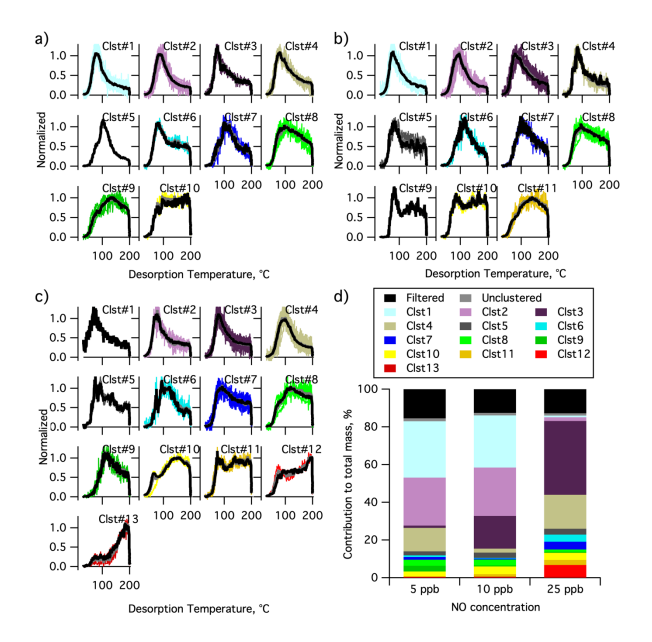


Figure 10. Multiple clustering results for α -pinene + OH SOA as a function of NO concentration. Clustering results are separately shown for the (a) 5 ppb NO, (b) 10 ppb NO, and (c) 25 ppb NO experiments. Each panel includes unweighted average thermograms (grey lines), mass-weighted average thermograms (black lines) and individual cluster members (colored lines). (d) Contribution of each cluster to the total mass for each experiment. The mass contribution of filtered-out ions (black bar) and unclustered ions (gray bar) are also shown.

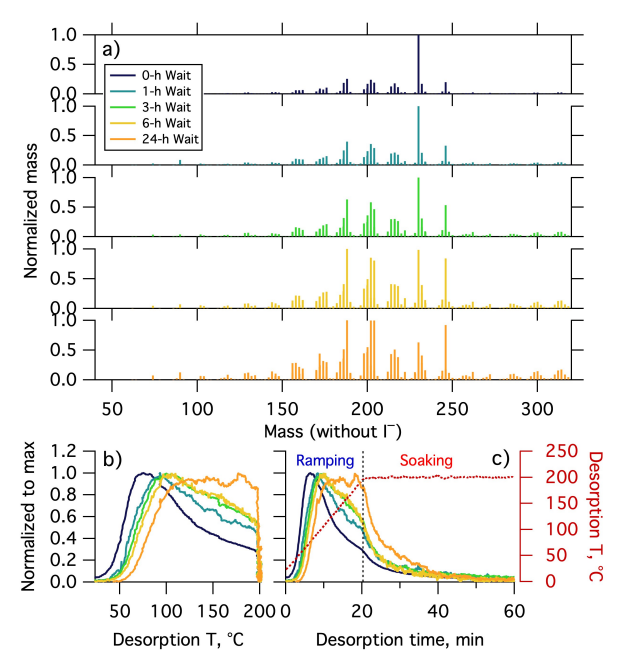


Figure 11. (a) Normalized mass spectra of α -pinene + O₃ SOA measured after different extents of isothermal evaporation at room temperature. The mass excludes iodine. The normalized thermograms of bulk SOA versus (b) temperature and (c) time, with the desorption temperature shown as a red dashed line. The vertical black dashed line in (c) delineates between ramping and soaking. The mass spectrum or thermogram colors indicate the isothermal evaporation time (see legend), with darker colors indicating shorter times.

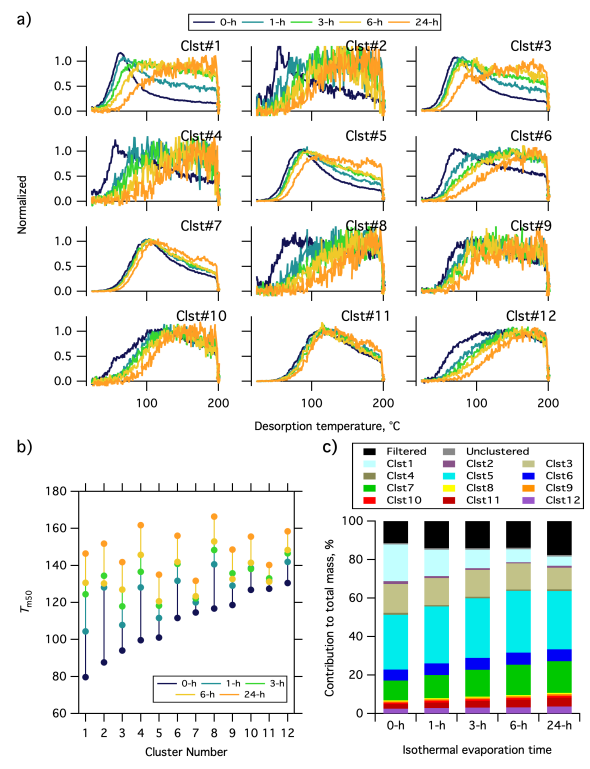


Figure 12. Single clustering results for α -pinene + O₃ SOA for different isothermal evaporation times. (a) Comparison of the normalized, weighted-average thermograms of the 12 clusters of 0-h wait (navy), 1-h wait (blue), 3-h wait (green), 6-h wait (yellow) and 24-h wait (orange) experiments. Note that the absolute signals of all of the clusters decrease with evaporation, but to varying extents (**Error! Reference source not found.**).

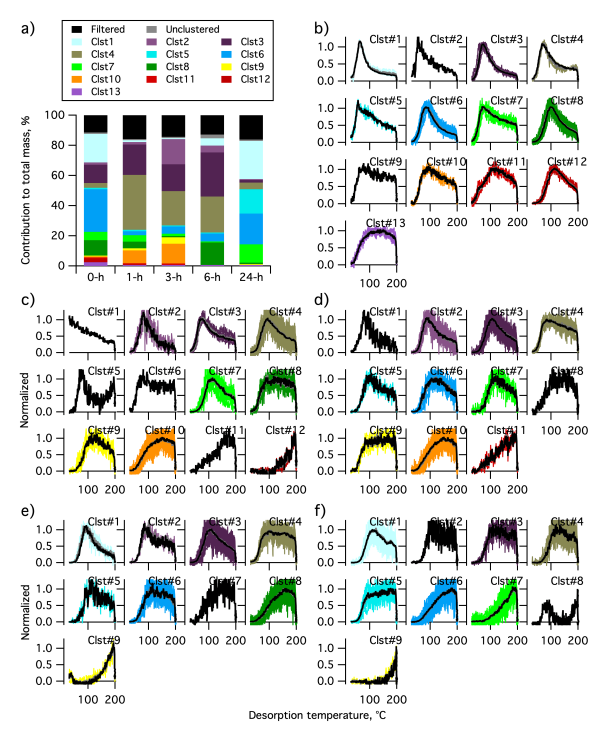


Figure 13. Multiple clustering results for α -pinene + O₃ SOA as a function of isothermal evaporation time. (a) Contribution of each cluster to the total mass for each experiment, along with the contributions of filtered-out ions (black bar) and unclustered ions (gray bar). The number of clusters obtained generally decreases with isothermal evaporation time. (b-f) The unweighted average (gray) and mass-weighted average (black) thermograms, along with the thermograms of individual members of clusters for the (b) 0-h, (c) 1-h, (d) 3-h, (e) 6-h, and (f) 24-h wait experiments. The cluster colors are consistent between panels.

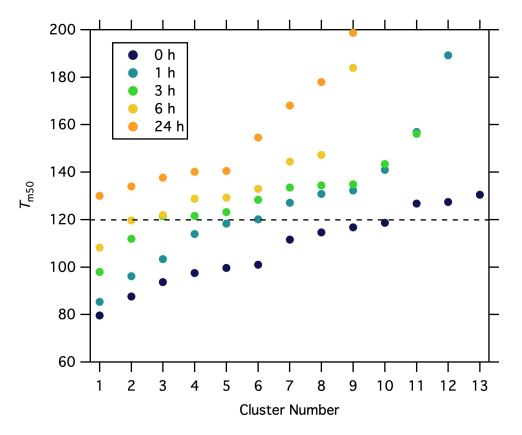


Figure 14. The T_{m50} values of the cluster-specific thermograms from multiple clustering for the five isothermal evaporation experiments.