Supplement of

Resolving anthropogenic aerosol pollution types – deconvolution and exploratory classification of pollution events

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S.1 PMF – on robust mode and rotational ambiguity

Since the amount of weight given in PMF to an observation (here: the value of a variable \(i\) at a certain time \(j\)) by the iterative process is proportional to the square of \(E_{i,j}/\sigma_{i,j}\), outliers with abnormally high squared signal or low variance may end up dominating the model solution. This phenomenon is especially relevant in environmental observations, as there are several types of outliers that would conceivably cause this behaviour, such as errors in the functioning of the measurement instrument or extreme, rare events that are considered contamination from the point of view of the analysis.

Therefore a robust mode for PMF was introduced (Paatero, 1997). The approach in short is to introduce a limit \(\alpha\), for the weight given to a point \((E_{i,j}/\sigma_{i,j})\) beyond which the point is considered an outlier, and dynamically down-weighted to negate its disproportional effect on the objective function \(Q\). For a complete explanation on outliers and the robust mode, we refer the reader to the original work (Paatero, 1997).

The main issues of PMF and indeed most factor analytic or linear algebraic methods are: 1) The rotational ambiguity of the solutions, i.e. the existence of multiple, sometimes very different, mathematical solutions with equally high rate of explanation.
of the observed (weighted) variance (Paatero, 1997; Paatero et al., 2002; Paatero et al., 2014). Exploring the rotations and selecting the best solution from the “solution space” needs to be done by the analyst, often based mainly on interpretability of the results in the context of the particular research topic at hand. 2) The selection of number of factors, \( f \). While exploring the rate of decrease of \( Q \) when increasing \( f \) can be considered an indicator of the amount of factors present in the set of data (Paatero and Tapper, 1993; Ulbrich et al., 2009; Reff et al., 2007), it rarely gives unambiguous answers. In the end it is up to the analyst to decide \( f \) based on both the diagnostics offered by \( Q \) and the interpretability of the result. These two subjective selections are often considered the most debatable part of a factor analysis (Ulbrich et al., 2009; Reff et al., 2007; Kim and Mueller, 1978). An additional constraint of the method is that it regards the chemical composition of a factor invariable, and as such is less than ideal for atmospheric conditions where physicochemical processes constantly alter the aerosol composition (Canonaco et al., 2015).

In this work we utilise PMF in a non-standard way, to resolve the time series and mass spectral profiles explaining “anomalous” observations often discarded from a PMF analysis: the periods with air pollution spikes and plumes. PMF analysis is done for each air pollution event individually, altering the time window of the analysis around the event to include both the pollution episode and some background before and after the event. The advantage of studying this type of short relatively short term phenomena is, that we can easily evaluate fulfilment of the criteria outlined in Sect. 2.2.2, and we can additionally discriminate between mathematically equal solutions, mostly evading the issue of rotational ambiguity. Essentially knowing beforehand what the (qualitative) temporal behaviour of a pollution and background factors should be like, (i.e. the time series of the factors should be uncorrelated), we explore the number of factors and the solution space to select the solution best fulfilling our criteria for a physically correct solution. Adhering to these criteria, we strive to minimise the ambiguity related to our selection of solutions, as well as considerably reduce the effect of subjectivity with regard to selection of solutions.

We note the inclusion of the robust mode, hard coded in our user interface of choice (SoFi 4.8; Canonaco et al., 2013) is a potential issue for events with temporally very short plumes of only a few time points, but our testing confirmed it did not noticeably hinder the algorithm finding the expected, physically realistic solutions. It was noted if the solution returned by the algorithm was not driven by the pollution plume-like temporal behaviour, the time window of the analysis was often too wide, and applying a narrower window reproduced the plume factor in most such cases. In the few cases, when despite using a narrow window an acceptable solution was not reached, the event was discarded from further analysis to avoid any errors in extraction of pollution features.

In extracting pollution spectra, in cases where an acceptable solution was not found despite a large number (100) of seed runs, we additionally explored the rotational solution space using non-zero Fpeak values (Paatero and Hopke, 2009). In 7 cases (included in the total of 81 spectra) an acceptable rotation was found this way. The criteria for PMF solution acceptance remained the same, and when non-zero Fpeak was applied, extra care was taken not to allow factors with profiles that exhibit “unrealistic/unphysical behaviour” such as only containing noise-like spectra or e.g. no contribution to m/z 43 or 44 Th.
The iterative method to achieve this operates as follows:

1. (Initialisation) In the very first step, called initialisation, a pre-set amount of $k$ starting cluster centres are defined: this can be done in a number of ways, but often involves random selection. Distances between each object $x$ and every cluster centre $c$ are calculated.

2. (Step 1: Assignation) All sample objects are assigned to their nearest cluster centre, based on the selected metric of distance (dissimilarity). Every object now belongs to exactly one cluster.

3. (Step 2: Updating) Cluster centres $c$ are re-calculated as a mean $\mu$ of the now updated cluster members $x_i$ of each cluster. After updating the cluster centres the assignation is performed again (move to step 1).

4. (Convergence) This cycle is repeated until no more changes are made in the assignments in consecutive iterations. The algorithm has reached a convergence.

The input parameters required for the operation of k-means include:

1. Number of clusters, $k$. This value must be selected by the user. It can be based on $a$ priori, external information revealing the number of clusters to be expected present, or in an exploratory analysis it can be set $a$ posteriori, based on diagnostics values indicating the quality of solution for various $k$ values and/or based on analyst expert opinion (reasonability and interpretability of the result). This requires calculating clustering solutions for a range of $k$ values.

2. A metric for distance used in calculation of $J(C_n)$. Typically Euclidean squared “distance” $\|x_i - \mu_n\|^2$ is used, (as in Eq. (5), main text), but other option also exist. In lieu of an actual distance between two objects, a metric describing the similarity or (conversely dissimilarity) of the said objects may be used instead (Anderberg, 1973).

3. Initialisation cluster centres. They can either be selected by the user or randomly chosen. Random selection can be obtained sampling from among all the objects at random (or uniformly at random), or by performing pre-clustering with a subset of data ($e.g.$ 10% of the objects selected randomly) and using these as initialisation for the final clustering. The selection of initialisation, (along with the number of random repetitions of the clustering) may influence the likelihood of finding the global minimum of $J(C)$ instead of a local one.

Regarding initialisation, it has been shown by Arthur and Vassilvitskii (2007) that selecting the initialisation points not uniformly at random but spreading them out via stepwise selection from a weighted distribution improves the performance of the k-means algorithm. We also adopt this k-means initialisation method, resulting in an algorithm commonly known as “k-means++”.

S.2 k-means clustering, parameter selection
As there are no general rules for the selection of metric or number of clusters, but they rather depend on the type of data and application at hand, we will experimentally study their effects and try to select the settings fitting the classification of AMS mass spectra.

S.3 On spectral (dis)similarity

When applying clustering methodologies, the analyst’s choice of an appropriate distance or dissimilarity metric, constructed from the variables studied in the analysis, is inherently dependent on the class of data variables (i.e. nominal, ordinal, interval, ratio scale) on hand. When dealing with aerosol mass spectrometric results, the data consists of a set of ratio scale variables, i.e. fractions of signals at various $m/z$ ratios that together form a mass spectra (an observation or object). Ratio scale measurement fulfil the strictest requirements of quantitative measurements (i.e. the unit difference between two values is meaningful as is the zero point of the scale), and contain the most information of these metric categories (Kaufman and Rousseeuw, 2009). They also impose the least restrictions on the choice of a distance metric.

A significant benefit of the AMS as an instrument is, that the mass spectrometric signal is well quantified and linear. All the variables in our data adhere the same units and scale, so they can be considered homogenous and harmonised relative to each other. While the variables are not completely independent, when dealing with fractions of total signal, the dependence is generally not dominant over the actual variations in the signal. Typically for the AMS the dynamic range of individual normalised signals is quite moderate compared to many other mass spectrometers, which partly avoids the problem of large signals completely dominating the clustering outcome – and also with the aforementioned issue with the signal fractions being co-dependent. Still, we are far from immune to these effects, and will discuss and test the effect signal intensity and possible scaling options.

Authors of classic textbooks such as Anderberg (1973), Kaufman and Rousseeuw (2009) and Spath (1980) also discuss the dissimilarity metrics’ theoretical background. It is noted e.g. by Anderberg (1973) that the correlation and cosine methods are very closely related, and their advantage is they are invariant to uniform multiplicative scaling, unlike the Euclidean and city block distances. For mass spectra this means if a spectra B is a scalar multiplicative of spectra A (as a thought experiment we disregard breaking of normalisation here), they are rightly considered identical by the cosine and correlation methods while the city block and Euclidean distances would find them different. With proper normalisation, this is less of a problem in practice, although, there are cases when this could conceivably cause issues. As an example, consider a case when we have two otherwise identical spectra, but in the other we let us say double $m/z$ 44 Th, diminishing the other signals respectively via the normalisation. The single difference in aerosol chemistry would now be considered the change of oxidation level, but the Euclidean distance method would now find all the $m/z$ signals to be dissimilar, the correlation and cosine methods would (perhaps more correctly) consider the dissimilarity increased only with regard $m/z$ 44 Th. The feature is further exacerbated by Euclidean distances’ proneness to giving excessive weight to outlying values of single variable (Cormack, 1971). We therefore hypothesise the Euclidean distance will discriminate the AMS spectra more with regard to their highest signals, such
as 44 and 43 Th ions often linked to aerosol oxidation level (Aiken et al., 2007; Aiken et al., 2008), and possible outliers. Conversely, cosine and correlation would rather focus on the (dis)similarity of the higher end of the spectrum. This hypothesis is supported in light of the results in Sect. (3.3), where it is shown the squared Euclidean algorithm does best in sorting aerosols types by their oxidation state and finds outlier groups, but struggles to separate classes with minor variation in higher up spectral structures.

On the more subtle differences between the closely related methods of cosine and correlation, Anderberg (1973) states:

“[...]the distinction [between cosine and correlation] is precisely the difference between ratio and interval scale variables, respectively. Thus, the cosine makes use of ratio scale information, while the correlation coefficient only uses interval scale information.”,

and recommends cosine to be used when the origin is meaningful and well established, which is the case with our AMS mass spectra (and derivatively the PMF results). This can be considered as a good argument for selecting the cosine metric. Anderberg (1973) additionally notes the correlation metric is invariant to any linear transformations, such as uniformly adding a constant to all the elements (here: spectrum m/z’s) and therefore less discriminating, which in the case of clustering isn’t a favourable quality, than the cosine metric. Previous AMS clustering studies (e.g. Marcolli et al., 2006) have also utilised a “dot-product” similarity metric

\[
d(u, v) = 1 - u \cdot v,
\]

(S.1)

which when normalized by the vector lengths, as is done for our mass spectra from PMF results, becomes exactly equal to the cosine metric (Eq. 7).

There also exist experimental evaluations for the performance of different metrics, the most comprehensive known to us being the aforementioned work of Stein and Scott (1994), focusing on evaluating the (dis)similarity metrics used to automatically identify mass spectra. The compared metrics include the dot product (cosine) metric and the Euclidean distance, and finds the dot product metric to perform highest in matching the primary NIST library spectrum and an alternative spectrum from the same compound, with 75% identification accuracy. Euclidean distance is the runner-up among the five metrics included, with 72% accuracy. (The three others being absolute value distance (68%), probability based matching (68%) and the Hertz et al. method (64%); Stein and Scott, 1994). Stein and Scott therefore conclude dot product (cosine) to be the best algorithm for mass spectra matching.

Additionally, in connection to the hypothesis about oxidation level indicator signals presented above, and in the spirit of Stein & Scott’s mass weighting rationale (“[mass weighting] deemphasizes the more variable and less characteristic lower mass range in a spectrum and emphasizes the more informative higher mass ions near the molecular ion”) we also tested completely omitting the oxidation-sensitive signal range, below 45 Th, with proper re-normalisation, of course, to see if we can alter the basis of discrimination in the clustering. The results proved intriguing, as can be found in Figure S.1, but were considered similar to the ones derived with the more elegant mass weighting processing, which we consider preferential over the omission method. The silhouette maximum seen at \(k = 4\) (Figure S.1, left panel) for \(m/\ell < 45\) Th omission derives from discrimination
against outlier groups, discussed in main text (Sect. 3.4.3) and was therefore seen as not incorporating new information over the selected method of mass scaling.

Figure S.1. Silhouettes for clustering solutions over \( k \) values from 2 to 20. Left: dataset processed with mass scaling. Right: \( m/z < 45 \) Th omitted. These curves can be further compared to the case of the original set of data (Figure 2), which can be considered a baseline for any pre-processing tests.

S.4 Additional methods for evaluating clustering solutions

The alternative methods of solution quality evaluation were briefly examined. Evaluation results and references are presented in Figures S.2 to S.4.

Figure S.2. Calinski-Harabasz criterion; (suggests max value \( k = 3 \); Caliński and Harabasz, 1974).
S.5 Notes on silhouette value

From the main defining Eq. (11) it follows that

\[-1 \leq s(i) \leq 1. \]  \hspace{1cm} (S.2)

The definition of $s(i)$ in this fashion is invariant to multiplicative matrix operations, e.g. multiplying the distances by a positive number, but not additive operations such as adding a positive constant to the distances. The silhouette method not only provides a metric for the robustness of a clustering solution but also introduces a graphical display for the description of n-dimensional results, an immensely helpful feature for an analyst when having to judge the “goodness” of a particular solution offered up by the clustering algorithm. A typical way to display silhouette values is a horizontal bar plot, such as the graph presented for the solution “corr” $k = 8$ of this study in Figure S.5.
Figure S.5. Silhouette plot for correlation, $k = 8$ ($s_m = 1.21$) solution. Numeric clusters 1 – 8 correspond (from top down) to clusters S-I, S-II, W-I, W-II, W-III, O-I, O-II and O-III, interpreted further in Sect. 3.4.

S.6 Posteriori weighting by silhouette value, effect on mass spectra

Posteriori weighting by silhouette values had minimal effect on cluster centroids, but did affect calculated intra-cluster variability values somewhat. The difference between unweighted and weighted cases is shown in Figure S.6.
Figure S.6. Effect of cluster centroid weighting by silhouette for the S-X and W-X clusters’ mass spectra. Above (5 uppermost spectra): unweighted cluster centroid spectra (mean of objects in cluster). Below (lowest 5 spectra), similar spectra weighted by the silhouettes of the objects. Correlation between the respective clusters is effectively unity ($r_s^2 > 0.994$), so the main difference is seen for the intra-cluster variabilities, depicted by the error bars.
S.7 Mass and intensity weighting, effect on solution quality

Scaling effect on mass spectra is further illustrated in Figures S.7 to S.9 below. Histograms of mass scaled silhouette values are presented in Figure S.10.

Figure S.7. Improvement in (solution) absolute silhouette values for mass scaling.
Figure S.8. Solution absolute silhouette values with intensity scaling.

Figure S.9 Silhouette improvement / degradation for intensity scaling.
Figure S.10. Histograms of total solution silhouette value distributions for mass scaled data, corresponding to results presented in Figure 3. Value of 0.45 (red line) was chosen as a lower limit for solution to be included in a more detailed examination (Sect. 3.3; Table 2). Note: for city block metric the x-axis scaling differs – the minimum x-axis value of other panels, 0.25, is marked with a dashed line.

S.8. Clusters’ cross correlations and diagnostics values

Some of the main diagnostics values and correlations for the clustering result are presented in Tables S.1 to S.3.

Table S.1. Similarity matrix for clustering results, correlation, $k = 8$. Scaled similarity values ($r_s^2; s_m=1.36; p < 0.05$) are used.

<table>
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<tr>
<th></th>
<th>sawmill-SOA</th>
<th>A-LV-OOA</th>
<th>A-SV-OOA</th>
<th>HOA</th>
<th>COA</th>
<th>&quot;amine-58&quot;</th>
<th>&quot;amine-100&quot;</th>
<th>&quot;amine-86&quot;</th>
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<td>-</td>
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<td>0.16</td>
<td>0.15</td>
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<td>0.54</td>
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<td>f44</td>
<td>f55</td>
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Table S.2. Cluster diagnostics values: population, silhouette, f-values, estimated O:C ratio

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<th>spectrum name</th>
<th>reference</th>
<th>r_s^2 vs</th>
<th>SV(HOA)</th>
<th>SV(COA)</th>
<th>SV(BBOA)</th>
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<td>0.51</td>
<td>0.35</td>
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<td>COA cluster (W-V)</td>
<td>(this study)</td>
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S.9 Wind direction dependences and diurnality of pollution events:

Average wind directions at 0-64 m height and local time at peak concentration was recorded for all events, and is shown in Figure S.11. Sawmill-SOA seems connected to directions with lumber mills (see Liao et al., 2011) and the A-SV-OOA similarly to the Juupajoki/Korkeakoski direction. HOA seems to originate from the direction with the nearest road (western sector), and the A-LV-OOA that we find connected to wood burning, is mostly seen with incoming air passing over the Hyytiälä forestry station buildings, saunas and the nearby cottages and houses. COA directionality is already more evenly distributed, but includes the forestry station as one major direction of origin. For the outlier spectra there are too few data points available, but we note there is a cattle farm to the east direction, and agricultural fields to the south, which would be possible sources of the amine compounds. Also the forest clear-cut area is to the south.

As for the diurnality analysis presented in a histogram in Figure S.12., the results are inconclusive at best. For sawmill-SOA the most peaks arrive after midnight, but include observations at all times of a day. For A-LV-OOA and A-SV-OOA the observations are more frequent during night time. In the case of A-LV-OOA domestic heating and cooking in the evening may play a part. HOA plumes occur mostly daytime, while the COA (potentially mixed with fresh BBOA) is evenly distributed. The few amine plumes occur night-time or in the early morning, potentially suggesting the compounds could be semi-volatile and therefore sensitive to diurnal temperature changes. Again, all of the above is speculative due to statistically too low sample sizes. Also it should be noted these are times when the plume is observed at the receptor location (SMEAR II), and therefore delayed by an unknown amount of time from the actual time of the emission.

Figure S.11. Wind rose “histogram” plot for (correlation k= 8) clusters: sawmill-SOA, A-LV-OOA, A-SV-OOA, HOA, COA, “amine-58”, “amine-100” and “amine-86”. Bar length depicts number of pollution cases with wind direction from the sector in question.
Figure S.12. Diurnal plots for the correlation, $k=8$ clusters. x-axis: local time when the plume is observed at SMEAR II, y-axis: plume count. Due to very low sample sizes not many conclusions can be drawn.

S.10 Local and regional sources identification

To support source identification, wind direction analysis (Figure S.11) was combined with geographical information (Figures S.13 to S.16) of the area nearby SMEAR II (Hyytiälä) measurement station.
Figure S.13. Potential close-proximity aerosol sources – the main close range sources are related to the activities at the Hyytiälä forestry station, and the local road “Hyytiäläntie”.

Figure S.14. Potential local aerosol sources – despite the remote location of SMEAR II there is scattered housing and small scale agriculture nearby.
Figure S.15. The two sawmills and adjoined pellet factory are situated some seven kilometres south-east of SMEAR II, at the village of Korkeakoski.

Figure S.16. Likely regional sources of anthropogenic aerosols. The town of Orivesi and the city of Tampere are highlighted. Sawmill locations taken from Liao et al. (2011) are marked by the arrows.
S.11 On additional important dimensions driving the clustering results

The source-indicative $f_{55}:f_{57}$ dimension is what we interpret separates the sawmill aerosol type from the other semi-oxidised aerosol types with similar O:C ratios and $f_{44}$ contributions (WI-III, A-SV-OOA types). While we should not draw stretched conclusions based on the separation of this one instance of a biogenic source, we find the sawmill aerosol type is additionally characterised by the high 53 to 57 Th ratio (or “$f_{53}:f_{57}$”); the biogenic sawmill aerosol has very low $m/z$ 57 Th contribution, while the relatively high 53 Th signal clearly sets it apart from e.g. COA with sometimes similarly low $f_{57}$ but high $f_{55}$. We find a ratio of $f_{53}:f_{57} > 2$ seems indicative of aerosol originating from the SOA conversion of sawmill monoterpane emissions (Figure S.17).

As for the HOA vs COA separation, we feel equally conclusive indicators as $f_{55}:f_{57}$ of the same division are found higher up in the mass spectrum $m/z$ scale; the odd $m/z$ value structures from 65 to 71 Th and 77 to 85 Th seem to offer equally good markers. Specifically we suggest a low ratio of $f_{65}$ to $f_{71}$ and $f_{85}$ to $f_{77}$ can be used as an alternative a marker for HOA as opposed to COA (possibly mixed with fresh BBOA) (Figure S.18).

Figure S.17. Correlation, k = 8 clustering solution projected onto 2-d axes corresponding to $f_{44}$ derived oxidation level (estimated O:C; Aiken et al., 2008) and $f_{53}:f_{57}$. Marker size corresponds to silhouette value of the point, ranging from zero to one. Cluster centroid locations are marked separately with darker colours. Outlier clusters are shown in grey, without centroids.
Figure S.18. The ratios $f_{77}:f_{85}$ (upper panel) and $f_{65}:f_{71}$ (lower panel) seem to offer good additional indicators for separating the COA (W-III) and HOA (W-II) clusters. Marker size corresponds to silhouette value of the point, ranging from zero to one. Cluster centroid locations are marked separately with darker colours.
S.12 Outlier clusters chemical interpretation and source hypotheses

Within AMS measurements, amines have been postulated from unit resolution mass spectra (Aiken et al., 2009) and confirmed via high mass resolution analysis (Huffman et al., 2009; Sun et al., 2011) in studies conducted in the heavily populated and industrialised Megacities of New York, and Mexico City. Also Allan and co-workers (2006), performing the first quadrupole AMS measurements at the SMEAR II site in 2003 already speculated on the possibility of amines explaining the extra nitrate signal at 30 Th, not explicable by ammonium nitrate alone. The amines’ contribution at 30 Th peak corresponds to the \( \text{NH}_2\text{CH}_2^+ \) ion, but is often obscured in unit-resolution data by other organic fragments and the \( \text{NO}^+ \) fragment from common \( \text{NH}_4\text{NO}_3 \). Of the aforementioned studies only the amine containing aerosol from New York was available, submitted to the AMS high resolution database (Ulbrich et al., 2009; http://cires1.colorado.edu/jimenez-group/HRAMSsd/) with reference to after a later similar analysis by Docherty et al. (2011). This high-resolution AMS spectra was translated to unit mass resolution and compared with our samples. A moderate similarity \( (r_s^2 = 0.67) \) was found between the library specimen and our cluster O-III, lending some confidence to the assertion of similarity. Although the other amine spectra were unavailable for mathematical correlation check, we note the general spectra of the aerosols reported by Huffman and others. (2009) for Mexico City and Sun et al. (2011) for New York exhibit some similar features to the spectra of O-I and O-III clusters, namely increased \( m/z \) 58, 72 and 86 Th signals, but in different relative fractions.

The aerosol described by Sun et al. has the major nitrate containing peaks at \( m/z \) (56), 58, 59 and 72 Th, but shows only a small peak at 86 Th and no significant contribution at 100 Th. The two Mexico City spectra reported by Huffman et al. (2009) include major peaks at 58 and 86 Th, but little to no contribution at 100 Th. By visual inspection the Aiken et al. (2009) amine aerosol specimen doesn’t seem to contain any of the peaks discussed here, so we consider it not to be a relevant reference in this particular case.

Additionally, some very similar spectra we encountered in literature were from those laboratory SOA formation study by Murphy et al. (2007), who measured using an AMS, secondary aerosol generated from various aliphatic amines. The spectra they report for trimethylamine photo-oxidation product aerosol has multiple similar features at 58, 86 and 100 Th, albeit in different signal fractions \( (m/z \ 86 \text{ Th signal} > m/z \ 100 \text{ Th signal}) \) to ours. Looking up the 70 eV electron impact ionisation spectrum of trimethylamine, available from the U.S. National Institute of Standards and Technology (NIST), we find a specimen with the same peaks as our but again with different ratios for \( m/z \) 58, 72, 86 and 100 Th.

Closest match for the O-I and O-II clusters within the AMS spectral database was the marine aerosol \( (r_s^2 = 0.50 \text{ O-I}; 0.68 \text{ O-II}) \) reported by Chang and co-authors (2011) for Arctic Ocean marine biogenic aerosol. It does contain a small peak at 58 Th, but only low signals at 86 or 100 Th.

As for the sources, in addition to marine and industrial emissions, amines have been reported from biodegradation of forest organic material such as leaves and needles In spring they could be released from their trapping below the layer of snow and subsequently be evaporated from the solar-heated ground surface at and after snowmelt. Amines are known to be common products of biomaterial degradation processes and to be volatile (Kieloaho et al., 2013; Kuhn et al., 2011). Alkylamines are
also known to be produced at the forest floor and their concentrations are found to be temperature dependent and peaking during autumn forest litterfall (Kieloaho et al., 2013). Our AMS measurements coincided with the snow melt period at the site, but to our knowledge no data amine data from snowmelt period at SMEAR II exists, regrettably.

Also manure application in crop fields has been shown to be a potential source of amines (Schade and Crutzen, 1995; Ge et al., 2011). It would be plausible some nearby field that has been fertilised with manure, would release a considerable amount of volatile amine compounds upon drying. This hypothesis has been questioned, however, by the findings of Kuhn et al. (2011) who instead conclude the amine emissions are more likely from the animals’ exhalation and feeding. Also more specific studies on cattle amine emission can be found; for example, emission of trimethylamine from cattle was reported by Sintermann and co-authors (2014). There exists a cattle farm some two kilometres to the east of the field station and some agricultural fields closer by (supporting information Fig. S.14).

Finally, a nearby patch of forest of 0.8 hectares of area, at a distance of 300 to 500 meters from the site, was cut clear a month prior to the start of our measurements (Virkkula et al., 2014). Amines have been found to be emitted from tree trunks and needles of live coniferous trees of the boreal zone (Kieloaho et al., 2013), and it would therefore be imaginable the process of clear-cutting a forest stand using a harvester vehicle and the subsequent transportation of the trunks away from the site would cause considerable emissions of volatile organic compounds, including monoterpenes and amines for several weeks after the process.

Besides amines, other possible, but in our opinion unlikely explanations for the signals observed in our outlier clusters may exist. A laboratory study by Rollins and co-workers (2010) also reported increased signals for m/z 58 and 86 Th when synthesised hydroxynitrates were measured with an AMS. However, their spectra seem to be dominated by peaks not observed in our outlier spectra, and the said series (CnH2n+2N+) only forms a minor part of the whole signal reported by Rollins and others. Additionally, Wolf et al. (2015) reported some atmospheric bacteria related to ice-nucleation producing a 70 eV EI MS signal at 86 Th (C5H12N+), but this sample lacks the other signals present in our series.

We additionally note that similar homologous series exist for some aliphatic ketones (CR2=C(OH)R+) following McLafferty rearrangement, producing a unit mass series at 58, 72, 86 and 100 Th (McLafferty, 1959). Conclusively differentiating between these two organic groups would benefit from high-resolution ToF data, which we unfortunately do not have available at this time. While we cannot definitely rule out the possibility of high concentrations of ketone compounds, we do not find references to this type of observations in the aerosol mass spectrometry related literature.

References


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