

An algorithm for automatic fitting and formula assignment in atmospheric mass spectra

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Abstract. Mass spectrometry is an established method for studying the chemical composition of gases and particles in the atmosphere. Using this technique, signals corresponding to thousands or even tens of thousands of compounds may be detected from ambient air. The process of identifying all the peaks in the mass spectra is often arduous and time-consuming, in particular when multiple overlapping peaks are present. This manual peak fitting and identification may take even experienced analysts anywhere from weeks to months to complete, depending on the desired accuracy and completeness.

In this work, we attempted to automate the fitting and formula assignment workflow and evaluate how far the process can get using a "one-button" algorithm. The algorithm constructed in this work takes in commonly known parameters specific to the instrument type, and by pressing one button it runs and ultimately provides a list of likely peaks for the mass spectrum. The algorithm utilizes weighted-leastsquares fitting and a modified version of the Bayesian information criterion along with an iterative formula assignment process. We applied it to synthetic mass spectra and both a gas-phase chemical-ionization mass spectrometer (CIMS) dataset and an aerosol mass spectrometer (AMS) dataset. The results were largely comparable with manual peak fitting and identification done previously but were achieved in a fraction of the time. Erroneous assignments mainly appeared at low-intensity signals, with interference from nearby higherintensity signals, a case that is challenging also for manual peak fitting. This algorithm provides an excellent starting point for a peak list, which, if needed, can be manually revised.

The main result of this study is the algorithm itself. While further improvements and tweaks are possible, the algorithm presented here is currently being implemented into the commonly used Tofware analysis software package to allow easy utilization by the broader community. We hope this can save valuable time of researchers for data interpretation rather than data processing and curation.

1 Introduction

Volatile organic compounds (VOCs), emitted into the atmosphere from a multitude of activities and processes, both anthropogenic and biogenic (Fowler et al., 2009; Goldstein and Galbally, 2007), are key components of atmospheric chemistry. These compounds are oxidized in the atmosphere, forming a vast number of different species, some with lowenough volatility to contribute to aerosol formation (Jokinen et al., 2015; Ehn et al., 2014; Zhang et al., 2012; Riipinen et al., 2011; Kulmala et al., 1998). Understanding the dynamics and impacts of these trace gases and particles on both health and climate requires knowledge about their chemical composition and the processes that form and transform them (Masson-Delmotte et al., 2021; Shrivastava et al., 2017; Heal et al., 2012).

To study these compounds and their chemistry, whether it is in the gas or particle phase, mass spectrometers are commonly used (Zhang et al., 2023; Huey, 2007). There is a vast array of variations of different instruments, targeted at different classes of compounds. These instruments utilize various ionization methods (Riva et al., 2019b; Rissanen et al., 2019; Lopez-Hilfiker et al., 2019; Canagaratna et al., 2007), inlets (Häkkinen et al., 2023; Eichler et al., 2015; Lopez-Hilfiker et al., 2014), and mass analyzers (Boesl, 2017; Batey, 2014; Hu et al., 2005). In all cases peak fitting to the mass spectra is required to be able to identify all the compounds of interest. The Orbitrap mass analyzer exhibits around an order of magnitude better resolving power than time-of-flight (ToF) mass analyzers and can in most cases unambiguously separate all ions encountered in the mass spectrum (Riva et al., 2019a; Zuth et al., 2018), but ToF mass analyzers are far more common in the field of atmospheric science. ToF mass analyzers often have a large number of partially overlapping signals that require careful peak fitting to separate (Stark et al., 2015; Junninen et al., 2010).

Peak fitting and identification are therefore necessary steps in the analysis of data pertaining to the chemistry of the atmosphere. This process can be arduous for analysts, potentially requiring experts with understanding and intuition of the chemical properties of the studied system going through each signal in the spectrum individually. Depending on the desired accuracy of the analysis, this process may take researchers from several days up to months to complete for a newly acquired mass spectral dataset. Thus, there have been several attempts and discussion about ways to facilitate the analysis process (Sandström et al., 2024; Alton et al., 2023; Zhang et al., 2019; Stark et al., 2015). However, most of these studies have not focused on automating the peak list generation process but rather on improving complementary techniques. Stark et al. (2015) did present an algorithm for automated peak list generation, but the work was focused on obtaining bulk chemical properties of the dataset rather than accurate individual fits. In this work we present an attempt at a fully automatic peak assignment process to establish a baseline for how accurate such an approach can be made. The ultimate aim is to be able to provide an algorithm that can dramatically decrease the time analysts need to spend on data processing in the form of peak fitting and formula assignment.

We will here describe our "one-button" algorithm, which, given a number of inputs (mass-calibrated spectrum, resolution and peak shape functions, and restrictions on the type of ions to be expected from the instrument), provides the user with a list of chemical formulas that are likely to be present in the sample. We describe the working principles of the algorithm in detail and apply it to both real and synthetic datasets to understand and evaluate its usefulness.

2 Methods

This section outlines the methods used to design the algorithm and also includes brief descriptions of the testing datasets. Note that when discussing these methods, a single charge is assumed for the signals, and therefore "mass" and "mass-to-charge ratio" are used interchangeably. When discussing peaks, the term "position" is used to describe the mass-to-charge ratio where the signal distribution is centered. This is not necessarily the same as the mean or the peak of that distribution but depends entirely on the definition of the peak shape function. The peak shape function is a function that describes the shape of the expected signal from a single type of ion. The algorithm requires four inputs: a mass-calibrated mass spectrum, a resolution function, a peak shape function, and a list of potential formulas. The former three are standard concepts in high-resolution peak fitting and will not be discussed in more detail here (Stark et al., 2015; Junninen et al., 2010). The list of potential formulas was generated specifically for this work, and the method is described in Sect. 2.2.4. In addition to the four necessary inputs, an optional input of baseline may be provided. A number of algorithm parameters mentioned in later sections may also be tweaked but default values are used for all datasets described here, unless mentioned otherwise.

2.1 Algorithm structure

The algorithm can roughly be divided into two parts. The first, or the free-fitting part, provides data and initial guesses for where peaks may be for the following part. The second part, or the peak assignment part, iteratively assigns formulas to the fits from the free-fitting part and updates the free fit after every formula assigned.

In more detail, the free-fitting section of the algorithm simply fits between zero and n_{max} peaks at each unit mass. At this point there is still no decision made about the number of peaks, so the algorithm starts by fitting zero peaks, then moves on to one peak, and so on. The previous fit is used to initialize the next fit. The data obtained from these fits are later used in determining the number of peaks to fit at each unit mass and to initialize the peak assignment part of the algorithm. n_{max} is chosen to be higher than the highest number of peaks that could realistically be identified at any unit mass. Picking a higher number than necessary only costs additional computational resources. For the tests and gas phase data in this paper, the value $n_{\text{max}} = 12$ was used, while $n_{\text{max}} = 10$ was used for the particle phase dataset. It is important to note that the free-fitting part does not utilize chemical information in any way. The fitting method is described in the following section (Sect. 2.1.1). The assignment part of the algorithm is described in detail in Sect. 2.1.3.

2.1.1 Peak fitting

Peak fitting is a common process for analyzing mass spectra. It is an attempt to describe the mass spectral signal as a superposition of signals from individual ions and a background signal. These peaks have a known shape, which is often empirically determined, and the position of the peak is determined by the mass-to-charge ratio of the ion. The position of the peak is commonly defined as the maximum of the fitted peak shape function. It is important that the definition is consistent between mass calibration and the fitted ions to accurately determine the mass-to-charge ratio of the detected ions. In practice these peaks are often located by a fitting algorithm that minimizes the residual or unexplained signal. In this work the function used in the fitting process used for the algorithm is minimizing the χ^2 value given by

$$\chi_n^2 = \sum_{i}^{k} \frac{(y_i - \hat{y}_{i,n})^2}{\hat{y}_{i,n}},$$
(1)

where k is the number of data points fit to, y_i is data point *i* in the spectrum, and \hat{y}_i is the fit value to this data point. Note that \hat{y}_i includes both the fit signals and the baseline estimate. Index *n* denotes the number of peaks included in the fit resulting in this particular value of χ_n^2 . The algorithm is also adapted to be able to fit several spectra simultaneously, for example, those obtained from factorization techniques (Zhang et al., 2019). In this case $\chi_{n,tot}^2$ is calculated as the sum of the χ^2 values for each individual spectrum. In following sections, whenever the peaks are fit, it refers to finding the positions and heights of the peaks that minimize the value of χ^2 . Later the χ^2 is also utilized for evaluating the number of peaks that provide the best fit.

Except for the slightly different definition of the minimized function, the algorithm uses the same approach for fitting peaks as Junninen et al. (2010), and therefore the method will not be described here in greater detail. The code used to conduct the peak fitting is also based on the code in the tofTools software developed by Junninen et al. (2010).

2.1.2 Determining number of peaks

One of the most difficult problems to solve for the algorithm is to decide what number of peaks to fit for a given unit mass. An example of some results from free fitting is presented in Fig. 1, and the data used for the fit are presented later in Sect. 2.2.2. As can be seen from the figure, the inclusion or exclusion of relatively small peaks may result in a significant shift in the positions of the peaks contributing the majority of the signal at a unit mass. This highlights the importance of choosing the correct number of peaks to fit. To determine the number of peaks fit, the following function is used:

score(n) = A
$$\cdot \frac{\chi_n^2}{\chi_{n_{\max}}^2} + n \cdot \ln(k).$$
 (2)

Here, like before, k and n are the number of data points and the number of peaks, respectively; χ^2 comes from Eq. (1); and A is an internal parameter to the algorithm, which will be discussed shortly. Further theoretical motivation of this expression for the score, which is based on the Bayesian information criterion (Neath and Cavanaugh, 2012), is given in Appendix Sect. A1. However, the main justification for this function is not from theory but from the results of testing, described in Sects. 2.3 and 3.1. The number of peaks to fit is determined by the n value that results in the lowest score. However, the final number of peaks may increase or decrease later, when chemical information is incorporated, as discussed in the following section.

A somewhat intuitive description of the *A* parameter is that it determines how much the algorithm values goodness of fit relative to the sparsity of fit peaks. If we select *A* such that two fits with different number of peaks n_1 and n_2 have an equal score, we see that *A* is proportional to the ratio between the difference in peak number (n) and the difference in goodness of fit (χ^2) :

$$A = \chi_{n_{\max}}^2 \cdot \ln(k) \frac{n_2 - n_1}{\chi_{n_1}^2 - \chi_{n_2}^2}.$$
(3)

This results in a direct relationship between A and the total number of peaks fit by the algorithm N_{fit} (Fig. 2a), meaning that determining an optimal value of A is of great importance. Note that A is a single parameter that is used for all unit masses in the entire dataset; however, the score function, where A is applied, results in a different number of peaks at each unit mass. The optimal value of this parameter may vary between datasets, so to make the algorithm as general as possible it is automatically determined before the peak identification portion of the algorithm.

After the free-fitting portion of the algorithm is complete, a value of *A* is determined, which is then used for deciding the number of peaks using the score function (Eq. 2). By examining both the synthetic and real datasets analyzed for this paper, the following method for determining a suitable value of *A* was arrived at. After some number of peaks (roughly five peaks per unit mass on average in Fig. 2b), the average relative χ^2 value (i.e., mean $(\chi_n^2/\chi_{n_{max}}^2)$) over all unit masses was roughly proportional to $\exp(-b(\bar{n}_{max} - \bar{n}(A))^2)$, where \bar{n} is the average number of peaks fit per unit mass. A fit was performed to the higher range of \bar{n} , and *A* was defined as a point where this fit starts to deviate from the data (Fig. 2b). A more precise description of the procedure is described in Appendix A3.

2.1.3 Assigning formulas

The algorithm iteratively assigns peaks to one integer mass at a time, starting from the lowest mass specified and proceeding to the following integer mass after completing the assignment process at a given mass. The flowchart in Fig. 3 outlines the general approach of the peak assignment. Equations (1) and (2) are both central to this part of the algorithm as well. The general idea behind the structure of this part of the algorithm is to find formulas that match the peaks as well as possible, starting from the most clearly distinguishable peaks. To determine which peak is the easiest to find, the concept of peak significance is used, explained below in Step 2 of the process. The algorithm does, however, have the option to change a previously assigned peak in cases where the presence of a formula becomes much less clear at some



Figure 1. (**a**–**d**) Free fits of 2, 3, 4, and 5 peaks to the unit mass at 326 Th in the gas phase dataset, illustrating the difficulty of the problem. The fitted peaks in the legends are sorted from highest to lowest intensity (top to bottom). It is quite clear that there are at least three peaks present at the mass. However, the addition of the fourth peak is not as obvious, despite it being assigned far higher signal than the third one included. This illustrates how much the overlap of peaks complicates this problem. It is even less clear if the inclusion of the fifth peak is necessary, and despite its size it changes the distribution of signal at two of the higher peaks significantly. Both the manual fit and the algorithm fit to this unit mass are presented in Fig. 5.

later stage in the process (see Step 4 below). In this sense this part of the algorithm mimics the process a human analyst may use when evaluating which formulas are present at an integer mass.

Step 1 of the assignment process is to update the preliminary results using the expected isotopic signals from lower masses. Since the number of peaks may be re-evaluated many times during the assignment process, the algorithm sets the maximum number of peaks that can be fit, n_{max} , at one more than the number of peaks that was determined optimal by the score function, based on the free fits. This saves significant time, and it is also rare that the number of peaks to fit would increase after this point. Often times the number decreases since isotopes from lower masses explain part of the signal. The isotopic signal is calculated from the expected isotopic ratios of each formula at lower integer masses. For the common elements detected in the ambient considered here, the rarer isotopes all have higher mass than the most common one, so isotopes from ions detected at the current integer mass or higher do not need to be considered. Then the algorithm subtracts the expected isotopic signal from the spectrum and refits the spectrum at the currently analyzed integer mass (this follows the process described in Step 5).

Step 2 evaluates the significance of each peak. This is done by removing the peak and refitting the heights of the other peaks while keeping their positions locked. The increase in χ^2 from the fit with all peaks determines the significance of the peak. This difference in χ^2 is then used to ensure that formulas are first assigned to the peaks most important for the overall fit. In this paper, mentions of the significance of a peak refer to this difference in χ^2 . The use of the word "significance" should not be confused with a strict statistical significance, even though it does serve a similar purpose.

Step 3 assigns a formula to a selected peak. All the potential formulas are provided to the algorithm as a list; the lists used for this paper are discussed in Sect. 2.2.4. The selected peak is the most significant peak (according to Step 2) for which a nearby potential formula can be found. This is done by first locating all the potential formulas within the allowed assignment interval, defined as one-fifth of the full width at half maximum (FWHM) of the peak by default. The option resulting in the lowest χ^2 is then assigned to the selected peak. When computing the χ^2 values of the different options, the positions of other peaks are locked in place. Locking the other peaks is to prevent them from moving into the position that the selected peak is adjusted around but conveniently also saves computing time.

Step 4 is introduced to account for cases where the most recent formula assignment greatly decreases the significance of a peak with a previously assigned formula. A previously assigned formula may be removed if the corresponding peak fulfills two conditions. First, the peak with a previously assigned formula must have lower significance than the peak with the most recently assigned formula (recall that the for-



Figure 2. (a) How the number of peaks increases with the value of parameter *A*. As *A* increases, the number of peaks per unit mass approaches the selected value of $n_{\text{max}} = 12$. (b) How the mean relative goodness-of-fit score improves with increasing average number of peaks. The fit curve is as described in Sect. 2.1.2. The vertical black line in both figures denotes the algorithm selected value of *A*.

mulas are assigned in order of descending peak significance, meaning that the formula assignments must have caused the order to change from the initial situation). Second, the significance of the peak with a previously assigned formula must be below 10% (default value) of what it was before the most recent assignment. If these conditions are met for any peak with an assigned formula, this formula assignment is removed, and the peak itself is appended to the list of free peaks. The list of free peaks is then again sorted based on significance. The algorithm is limited to remove peaks at most 10 times for each integer mass. This is to prevent rare situations where the algorithm ends up in an infinite loop of assigning and removing peaks. Although removing peak assignments is a relatively common occurrence, reaching the cap of 10 removals was very rare and unlikely in situations where the algorithm was not stuck.

Step 5 re-evaluates the number of free peaks and their positions every time a new formula is assigned. This follows a similar process to the preliminary fits. The lowest number of peaks that may be fit, n_{\min} , is the number of peaks



Figure 3. Flowchart describing the peak assignment process on a general level for each integer mass in the spectrum. A more detailed description of how each step marked with a number works is provided in Sect. 2.1.3. (*) The significance of a peak refers to how much the omission of that peak would increase the χ^2 value of the fit. (**) The assignment interval can be adjusted but is set to the default value of 0.2. FWHM for all of the runs presented in this paper.

that have been assigned a formula, while n_{max} is adjusted in Step 1. The algorithm then performs fits with numbers of peaks ranging from n_{min} to n_{max} peaks, starting from n_{min} . The peaks that have an assigned formula are locked in place, while free peaks are free to change their positions. Each time the number of peaks increases, the position of the added peak is initialized by the most significant remaining free peak in the list of free peaks from the previous step. When the number of peaks to fit exceeds the number of free peaks in the list, the positions are instead initialized using the residual of the fit with one fewer peaks. The fit used for the next iteration of the assignment process is the one resulting in the lowest value of the score function (Eq. 2).

At the end of the assignment process there may still be free peaks left that have not been assigned any formula due to there not being any available options within the assignment interval. These are allowed and are simply labeled "unknown". Although this may be useful for locating some peaks, it should not be relied upon, since isotopes of unknown formulas cannot be accounted for and may lead to problems at other masses. The default assignment interval of $0.2 \cdot$ FWHM represents the minimum distance between a peak and any potential formula for the peak to be labeled "unknown". Since all the potential formulas that are within the interval are tested by fitting, the value is primarily selected to be wide enough not to exclude the correct formula. Tweaking this value may be relevant if the list of potential compounds is more or less restrictive.

If the list of potential elements includes an element whose most common isotope is not the one with lowest mass, the handling of isotopes mentioned in Step 1 warrants reconsideration. However, atmospheric mass spectra with large numbers of peaks generally consist mostly of formulas made up from the elements C, H, O, and N. The algorithm does not currently check if the isotopes for an assigned formula are present. This may be a useful future improvement, but testing showed it would very rarely be useful in the datasets tested here.

2.2 Data sources and description

2.2.1 Synthetic data

Synthetic data were used frequently during development of the algorithm and for sensitivity test presented in this article (Sect. 3.1). The synthetic spectra were generated as Poisson distributed signals to match the noise that is expected in real mass spectra. This method of data generation is commonly used when attempting to replicate mass spectral signals (Cubison and Jimenez, 2015; Lee and Marshall, 2000). First, a noiseless spectrum, $\lambda(m/z)$, was generated as a sum of a constant background level and signals from individual ions.

$$\lambda(m/z) = \mathrm{BL} + \sum_{i}^{n} I_{i} \cdot f(m/z, \mu_{i})$$
(4)

Here BL is the baseline signal; I_i is the intensity of the peak indexed i; and $f(m/z, \mu_i)$ is the distribution function describing the peak shape, centered at the mass μ_i , accounting for resolution. The peak shape used was empirically estimated from the gas phase dataset (Sect. 2.2.2), and the peak locations correspond to real formulas randomly selected from the list of potential formulas for the gas phase data (Sect. 2.2.4). The numbers of peaks per unit mass were uniformly distributed integers between 0 and 8. The generated datasets span the unit mass range from 200-400 Th. The total intensity allocated to all peaks at an integer mass was sampled from a lognormal distribution. That signal was then distributed between peaks at the same unit mass by weights sampled from a uniform distribution. This results in a high variety in intensity between integer masses and a high variety of signal-to-background ratios, while restricting the number of unit masses where a single peak contributes nearly all of the signal.

The noise was included by randomly sampling the signal, y(m/z), at each point from a Poisson distribution with expected value determined by the noiseless $\lambda(m/z)$ (i.e., $y(m/z) \sim \text{Pois}(\lambda(m/z))$, Fig. A1). This results in a signal-dependent noise level that mimics the expected signal distribution in ToF mass analyzers (Cubison and Jimenez, 2015; Lee and Marshall, 2000). Datasets corresponding to mass spectral resolutions of 4000 and 13 000 were generated to mimic the performances of commonly used highresolution ToF (H-ToF) and long-ToF (L-ToF) mass analyzers (Peräkylä et al., 2020).

2.2.2 Gas phase data

The gas phase data used in this work have been previously analyzed and published by Peräkylä et al. (2020). This dataset was collected using a nitrate chemical-ionization atmospheric-pressure-interface time-of-flight mass spectrometer (Jokinen et al., 2012) during chamber experiments of α -pinene ozonolysis. The instrument contained a long time-of-flight mass analyzer with a resolving power around 13 000. During the experiment, clean air; α -pinene; ozone; and sometimes NO₂, water vapor, and inorganic aerosol particles were added to the chamber. The data are only used as a reference of what peaks may be identified during thorough manual analysis of resolution-limited datasets, and therefore the spectra were simply interpolated to a common mass axis and averaged over the entire measurement period.

2.2.3 Particle phase data

The particle phase data were collected using an aerosol mass spectrometer (AMS) (Canagaratna et al., 2007) with an L-ToF mass analyzer at the SMEAR II station in Hyytiälä, Finland, during the spring of 2016 and have previously been analyzed and published by Graeffe et al. (2023). The resolution of the instrument is approximately 5000 at 100 m/z. The site is surrounded by boreal forest, and the main anthropogenic influence comes from a sawmill about 7 km away. Similarly to the gas phase data, this dataset was averaged over the measurement period to obtain a single average spectrum, to which the algorithm was applied.

2.2.4 Formula lists

This work utilized two different lists of potential formulas: one for the gas phase nitrate CIMS dataset and the other for the particle phase AMS dataset. These lists were generated on broad expectations of what types of ions we expect to detect with the different instruments.

The lists of potential formulas provided to the algorithm for the gas phase analysis were generated by providing some combinations of atoms that were sequentially added to form complete molecules. Additional constraints were placed on the constructed molecules by providing limits for the number of atoms of each element, as well as O-to-C and H-to-C ratios. For a complete description of these constraints, see Sect. A2. The gas phase list was also used for both generating and analyzing the synthetic dataset. The particle phase list was far less constrained, since there are a lot fewer potential formulas at the lower mass range, and the AMS utilizes electron impact ionization, fragmenting the compounds, which means much fewer constraints for ion compositions (Canagaratna et al., 2007). The list is generated mostly by combining atoms of common elements and does not include rules motivated by chemistry. For simplicity, only the elements C, H, O, N, and S, were included.

2.3 Evaluating algorithm performance

Evaluating the performance of a peak identification algorithm for atmospheric mass spectra is not without its own challenges. Even an experienced analyst cannot be certain about the accuracy of all their fits, and when analyzing mass spectra there is not always a need to attempt to identify all the peaks that may be in the data, since they might not be of relevance or be too uncertain for further analysis. This is the reason why synthetically generated data for which all the correct peak positions and signal intensities are known were used for most of the development and testing of the algorithm.

For the real data, it is common that there are peaks clearly present in the data that have not been fitted during manual analysis, either because the peak is not relevant to further analysis or because it is difficult to find a formula corresponding to the signal. In other instances the manual analysis has included peaks that are more or less clearly not present in the list as a part of a series of formulas. Therefore, it is important to remember that the fits used to evaluate the algorithm results are in no way perfect. However, they do represent the information that a typical analyst wants to obtain from the dataset. With this in mind, we use the term "match" rather than "correct" when referring to a formula that was identified by both the manual analyst and the algorithm. This is to remain conscious of the incompleteness and fallibility of manual identification of peaks.

The quantities used to compare the list provided by the algorithm with either lists of generated peaks or lists provided by manual analysis are presented in Table 1. Here, and in the rest of this article, the letters N and S refer to total number of peaks and total area of signal, while n refers to the number of peaks at one unit mass and s refers to the area of the signal of one peak. The subscripts "fit", "gen", and "corr" refer to "fit by algorithm", "generated", and "correct", respectively. When discussing the real data, the subscript "gen" is replaced by "man" to denote "manually fit" instead of "generated", and the subscript "corr" is replaced by "match" to denote "a match between manual and algorithm fits". A peak is considered correct, or matching, if it has the same formula as a peak in the list of generated peaks or manually fit peaks, depending on the dataset. s_{corr} (or s_{match}) is defined as the smaller one of $s_{\rm fit}$ and $s_{\rm gen}$ (or $s_{\rm man}$) for a correctly fit (or matching) peak. For an incorrectly fit (or not matching) peak $s_{\rm corr} = 0.$

2.3.1 Synthetic data tests

Four different tests were conducted using synthetic data. The first one investigated how the algorithm performed with different selections of the value for parameter A, evaluated using the metrics from Table 1. The results of this test were later used to inform the selection of A explained in Sect. 2.1.2. For this test, and this test only, all the isotopes were removed from the dataset since the inclusion of isotopes makes it difficult to clearly define whether a free peak is in a correct location or not, and this test mostly relied on algorithm results before formula assignment.

The other three tests evaluated how sensitive the algorithm performance was to imprecise inputs from a potential user. The inputs tested were the list of potential formulas, resolution function, and mass calibration. The effect of the potential formula list was tested by adding up to an additional zero to four molecules of an imaginary element X to each formula in the existing list. Element X had a single isotope with a mass of exactly 1 atomic mass unit, and this addition resulted in many more overlapping formulas for the algorithm to choose from when assigning compositions.

The sensitivity to the resolution function used was tested by applying a resolution scaling factor between 0.9-1.1 to the resolution used in the fit. For example, a factor of 0.9 results in the resolution of the fit peaks being 10% lower than the resolution of the peaks that were generated for the dataset; i.e., the algorithm fits peaks that are too wide.

Calibration error refers to an error in the definition of the mass axis of the mass spectrum. This results in the signals from all ions being offset from their actual mass in the spectrum. The test for calibration error was done by shifting the generated spectrum by 0-16 ppm before running the algorithm, resulting in a corresponding offset between the peaks in the spectrum and their correct formulas. This results in all the correct formulas being offset by some amount from their actual signal.

3 Results

The results of all testing and evaluation of the algorithm are presented in this section. The results are divided into one subsection for each type of data: synthetic, gas phase, and particle phase. The synthetic results focus on the sensitivity tests mentioned in the previous section, as well as some motivations for the methods the algorithm uses. The results of the application of the algorithm to real data are focused on comparing the algorithm list with the manual list to evaluate how useful the algorithm is as a tool to facilitate that analysis.

3.1 Synthetic test results

Synthetic datasets were widely used when developing this algorithm, and many of the methods mentioned earlier, such as the score function (Eq. 2), were derived from tests using syn-

Synthetic data	Number	Signal	Real data	Number	Signal
algorithm fit	N _{fit}	S _{fit}	algorithm fit	N _{fit}	S _{fit}
generated data	N _{gen}	S _{gen}	manual fit	N _{man}	S _{man}
correctly peaks	N _{corr}	S _{corr}	matching peaks	N _{match}	S _{match}

Table 1. Quantities used when evaluating algorithm performance.

thetic data. The method used for selecting the value of the parameter A was mentioned earlier in Sect. 2.1.2. Here we briefly discuss more about the importance of this parameter and how it impacts the algorithm results. Figure 4a shows how the accuracy of algorithm results changes with this parameter. There is a critical value of A, at around A = 100in this case, beyond which there is a steep increase in erroneously fit peaks, seen by the sharp increase in the total number of peaks and simultaneous decrease in the correctly fit fraction of peaks. Although this point becomes less clear with the inclusion of isotopes and the limited knowledge of fit parameters for real data, the main objective in selecting a value of A is to stay below this critical point. To the left of this point there is an interval of decent options, depending on whether a more conservative (fewer peaks) or exploratory (more peaks) approach is desired. The chosen method for selecting an appropriate value for A focuses on landing somewhere fairly close to but below the critical value. The mean relative χ^2 value from the critical value of A onward follows the fit curve outlined in Sect. 2.1.2 and displayed in Fig. 2b.

The sensitivity tests for input parameters (Fig. 4b–d) all show expected behavior, with more poorly defined input parameters resulting in worse performance. The test with the expanded list of potential formulas shows the importance of limiting the number of formulas considered. This test was very challenging, with an up to 5-fold increase in the number of potential formulas. The challenge with more potential formulas is not only that there are more options to choose from, but that each erroneously assigned formula leads to the subtraction of expected isotopic signal that is not actually present, while the isotopic signal of the correct formula remains in the data. This problem can be minimized by limiting the number of formulas that are considered or by having higher-resolution data (Fig. A2b).

Regarding resolution sensitivity, the best results are achieved at the correct resolution of fits, i.e., when the resolution scaling factor is 1. However, there is some asymmetry between lower and higher scaling factors (Fig. 4c), with fits using slightly too high a resolution resulting in worse results than fits using slightly too low a resolution. This may be due to peaks that are too narrow compensating with additional peaks, which results in erroneous assignments, whereas peaks that are too wide only limit assignment of other peaks in instances where there are two neighboring peaks of similar magnitude. However, both of these errors will lead to increased difficulty in assigning smaller peaks nearby.

The final sensitivity test addresses mass calibration and shows that poor mass calibration has a fairly strong impact on the results. Unsurprisingly, a better mass calibration results in better fits. Even small improvements in the mass calibration leads to significantly better fit results, which makes this one of the most important parameters when utilizing the algorithm.

Overall these tests show that optimizing all of these inputs will improve the results of the algorithm. However, realistic accuracy that can be achieved with currently widely used analysis tools will not lead to the algorithm being useless. This is further supported by the results of the application of the algorithm to real datasets, where the peak shape and resolution functions were determined empirically and therefore are as precise as one could expect for a real dataset. These results are presented in the following sections and show what kind of results or accuracy can be expected from using the algorithm with real data.

3.2 Gas phase results

Some example comparisons between the algorithm and manual assignments are presented in Fig. 5. In general, the algorithm is able to adapt the number of peaks required, and there is good agreement between the algorithm and the manual fits, at least for the most dominant peaks. These examples also show the challenge between a direct comparison with manual analysis. The two formulas, $C_{13}H_{26}O_7N_1^-$ and $C_{12}H_{24}O_9N_1^-$, that contribute close to no signal have probably been included as a part of a series of formulas during manual analysis. Meanwhile, other peaks that were not relevant for the manual analysis due to their negative mass defect were not included in the manual analysis at all. This makes a one-to-one comparison between algorithm and manual fits misleading, and the following analysis will focus on how much agreement there was between algorithm and manual fits in terms of signal, as well as the characteristics of the peaks that the algorithm was able, or failed, to find. Formulas, like $C_{13}H_{26}O_7N_1^-$, in the manually compiled list that contributed no signal at all were also omitted from the summary statistics.

Table 2 presents summary statistics of both gas and particle phase results. Overall the algorithm found 76 % of all the peaks included in the manual dataset; the found peaks result



Figure 4. Results of sensitivity tests conducted for synthetic data with a resolution around 4000. (a) How results vary depending on parameter *A*. The solid lines depict how the scores vary for the free fit, while crosses depict the results after the formula assignment. For the free fits a fit is considered correct if within 0.2 FWHM (50 ppm at 300 Th) of the generated location; after assignment, a fit is considered correct only if the precise formula assigned is the same as was generated. Both before and after assignment a sharp drop in the correctly fit fraction of peaks can be seen at around A = 100 due to a large number of incorrect peaks being added. (b) Influence of the number of compositions in the list of potential formulas. (c) Influence of errors in mass calibration. (d) Influence of incorrect resolution of fit peaks on fit results. Corresponding results for resolution of 13 000 in Fig. A2.

in a 97 % match in assigned signal between the algorithm and the manual analysis.

A more detailed overview of individual peaks and masses is presented in Fig. 6. Each marker in Fig. 6a corresponds to a formula that was identified in the manual analysis but was not identified by the algorithm. The x-axis value shows how close the closest algorithm fit was to that formula. The red markers represent formulas that were also included in the list of potential formulas provided to the algorithm, and the black markers represent formulas that were not included in this list. If a formula is included in the list of potential formulas, there are three factors influencing how likely it is that the algorithm does not identify a peak in the manual list. The peak may have very low signal compared to other peaks, making it hard to discern; the peak may be located very close to another peak, also making it hard to discern; or there is another formula in the list of potential formulas with a very similar mass, resulting in the algorithm misidentifying the peak.

From right to left in the plot, the distance to the nearest fit decreases, and red markers with higher signal start to show up, as the shorter distance between peaks makes higher relative signals harder to identify or the algorithm finds signal in the right spot but assigns it to the wrong formula. The vertical black line shows the assignment interval. This is the maximum distance allowed between a formula and a given peak, where the formula may be assigned to that peak (see Sect. 2.1.3). For red markers to the left of the vertical black line, there was a free peak in roughly the correct location, but the algorithm decided on another formula in the list of potential formulas. However, the distance to other nearby peaks and relative signal intensity continue to influence how likely misidentification of a peak is. To summarize the meaning of the red markers, the algorithm is most likely to miss peaks that have low signal, that are located close to another peak, or if there are multiple potential formulas within the assignment interval of the peak.

There are also a number of black crosses in Fig. 6a. These represent formulas that were not included in the list of potential formulas given to the algorithm and therefore could not have been identified by the algorithm. The position of many of these markers in the plot, to the left of the vertical black line, shows that it is in several cases very likely that the algorithm would have located these formulas had they been on the list and a peak in their close proximity was instead labeled as "unknown". This shows that even when some unexpected formulas are present, the algorithm may be useful for finding them.

The distribution of the results for individual unit masses is shown in Fig. 6b. Each unit mass is assigned a bin based on



Figure 5. Comparisons between manual (**a**, **c**, **e**) and algorithm (**b**, **d**, **f**) fits to three example unit masses of different complexities. The fitted peaks in the legends are sorted from highest to lowest intensity (top to bottom). The free fits at 326 m/z were presented earlier in Fig. 1. In addition to the examples here, figures of fits to all of the analyzed integer masses are provided in the Supplement for a better overview of the results.

Table 2. Summary statistics for the gas phase and particle phase datasets.

Dataset	N _{fit}	N _{man}	N _{match}	$\begin{vmatrix} N_{\rm match} / \\ N_{\rm fit} \end{vmatrix}$	N _{match} / N _{man}	S _{match} / S _{man}
Gas	1557	844	644	41.4 %	76.3 %	97.1 %
Particle	330	349	241	73.0 %	69.1 %	96.8 %

the fraction of signal that matched between the manual and algorithm fits (S_{match}/S_{man}). The y axis shows the fraction of the total signal area in the spectrum in each bin (blue bars) or the fraction of unit masses in the spectrum in each bin (red bars). First, the algorithm performs exceptionally well for the vast majority of the signal area. Figure 6b shows that over 97% of the total signal is located at unit masses in the highest accuracy bin, i.e., unit masses where over 94% of the signal matches the manual result. Second, even when looking

at just the number of unit masses, the algorithm matches over 80% of manually fitted signal at 80% of unit masses.

Looking at the numbers of peaks fit by the two methods (Fig. 6c), it is clear that the algorithm often fits many more peaks than the manual analysis has. However, even when the algorithm does fit many more peaks, it often does not result in poor attribution of signal. Often the higher number of peaks in algorithm fits is due to the manual analysis not attempting to identify every peak. However, there are also a



Figure 6. (a) Summary of all the formulas that were located in the manual analysis but not by the algorithm. Red crosses are formulas that were in the list of potential formulas provided to the algorithm. Black markers show formulas that were not on that list. The x axis is the relative difference in mass between the formula and the closest peak provided by the algorithm (absolute units of mass in Fig. A5). The y axis shows what fraction of the total signal area the formula contributes at its integer mass. The vertical black line marks the assignment interval. (b) The distribution of the matching fraction of signal at each unit mass. (c) Summary of matching fraction of signal according to the number of peaks fit by each method. Marker area is proportional to the total manually fitted signal.

few instances where the algorithm seems to add an excessive number of peaks.

3.3 Particle phase results

In general, the algorithm was more restrained in adding peaks when analyzing the particle phase data compared to the gas phase data and fitted a similar number of peaks to the manual analysis. The reason for this may be related to greater inaccuracies in peak shape and resolution functions. Compared to



Figure 7. Same plots as Fig. 6 but for the particle phase data. For this dataset the manual list contained more formulas than the algorithm list. However, the general patterns and overview of the results are very similar.

the gas phase data, the resolution and peak shape used were not as precisely defined, which can be seen from the particle phase fits and residuals (Fig. A9), which may have led to a more conservative fit in general. This resulted in a higher rate of agreement between the manual analysis and the algorithm. In terms of signal, the algorithm achieved a 97 % match with the manual analysis also for this dataset. One of the main factors lowering this fraction was the exclusion of trace elements from the list of potential formulas in the interest of simplicity (Sect. 2.2.4).

A more detailed summary of the fits to particle phase data is presented in Fig. 7. In general, the results are very similar to those of the gas phase dataset, which reinforces the points made about the algorithm in the previous section. The one difference is that for this dataset the manual list includes slightly more formulas than the algorithm list. This may be in part due to the more conservative fitting to this dataset mentioned earlier and in part due to the manual fitting process. For AMS data it is common to use a long list of compounds and exclude formulas from that list rather than include them. This may lead to more formulas being used than if starting from an empty list and adding formulas one by one.

3.4 Discussion

The results of testing with real datasets show that, despite being a first proof of concept, this algorithm can provide accurate results. The vast majority of the signal is correctly allocated, and a clear majority of the manually identified peaks are also located by the algorithm. These results are very promising, especially considering the strict definition placed on the correctly assigned signal used (Sect. 2.3) and the uncertainty present even among manually identified formulas. Although the algorithm does fit a lot more peaks than the manual analysis for the gas phase dataset, this is often due to the incompleteness of the manual analysis. Despite there being some occurrences of over-fitting by the algorithm, this has not affected the peaks with more significant signal. There is also a natural tool already included in the algorithm to avoid this issue, as a user could adjust parameter A to a lower value to decrease the willingness of the algorithm to fit additional peaks.

Recently, utilizing factorization methods to facilitate peak separation and identification has also been suggested (Zhang et al., 2019). Since the algorithm can fit several spectra at once, this is another area of potential improvement. However, whether the score function can be successfully applied to spectra derived from factor analysis is not certain, since the details of how noise in the data is transferred to the factors are unclear. Therefore, further work would have to be conducted to effectively utilize these two methods together.

The total runtime of the algorithm for the gas and particle phase datasets was around 40 and 5 min, respectively, on a standard laptop. This difference in speed is a combination of the difference in the number of integer masses analyzed, the number of peaks present per integer mass on average, and the set value of n_{max} . The algorithm was not optimized for speed, as we believe accuracy should be prioritized over expediency in this case, as the peak list generation would typically be performed only once for a given dataset. As such, we believe that tens of minutes of additional runtime are worth the investment if it results in far less time ultimately spent to reach the final results.

The sensitivity tests show that fitting parameters also impact algorithm performance. Although the algorithm already achieves good results with fitting parameters of standard accuracy, further improvements in determining these parameters may also result in better algorithm fits. As evidenced by the results, the most challenging problem with peak identification is the peaks with lower signal overlapped by other peaks. These peaks are more significantly impacted by errors in mass calibration, peak shape, or assignments since the errors of overlapping peaks add up. The difference in signal also means that a small inaccuracy in the peak shape for a large peak can lead to a large impact on a smaller peak in the vicinity. Therefore, we believe the primary means of improving the results in general, but most significantly for the lower-intensity signals, is to improve the methods to accurately determine these fitting parameters.

The algorithm results themselves may also be used to better define these parameters, and future works may utilize this for better definitions of peak shape, resolution, and mass calibration functions. As more ions are identified in a dataset, there is more information available for determining these parameters. Potentially the definition of these can even be improved within the automated process for even better fits.

However, since peak fitting is a statistical tool, there will always be an inherent level of uncertainty. Whether the identification is done manually or by an algorithm, there will be some peaks that cannot be identified with the desired confidence. Where this limit is encountered depends on the input data. The goal of the algorithm is to save time, so we do not think it is necessary to demand it to be able to identify more peaks than manual fitting.

Another future improvement would be for the algorithm to reconsider formulas whose isotopic signals do not match the data. As mentioned previously, this was found to be relevant very rarely during testing – in part due to most organics having fairly similar isotopic patterns and in part because the algorithm mostly misidentified peaks with comparatively low signals. Even for datasets containing halogens or other elements with isotopic patterns that deviate from organics, the different mass defect should result in accurate identification of these formulas in a majority of cases. However, this may be an improvement for future consideration.

This new algorithm is completely automated, and the inputs required are parameters commonly used in analysis of mass spectra. Therefore, the threshold of adopting this method among users should be low, especially as a part of existing analysis software. Work is currently in progress to incorporate this process in Tofware (https://www.tofwerk. com/software/tofware/, last access: 18 March 2025), which is commonly used for analyzing atmospheric mass spectra. All internal parameters for the algorithm have been determined in a way that a potential user does not need to worry about adjusting them for their specific dataset, although it is possible for expert users to do so if desired. The potentially biggest obstacle for adoption of this algorithm is the need for a potential formula list. However, there are already tools within established analysis software providing lists of formulas containing select elements that can be readapted for generating such lists. There have also been calls for better data infrastructure and databases for a more-data-driven approach to the analysis of mass spectra (Sandström et al., 2024). We believe the method described here can complement this development well, both benefiting from easily accessible lists

of formulas and facilitating the establishment of such lists as well as reference spectra.

4 Conclusions

Mass spectrometers have been a driving force behind recent advances in atmospheric science and remain a widely utilized method in the field. However, due to the large number of unique compounds in the atmosphere, peak fitting and identification from complex mass spectra are a challenging and extremely time-consuming task. Thus, methods that facilitate this task can save researchers a lot of time.

We have here presented an algorithm for complete automation of the peak-fitting and assignment process. The main result of this work is this algorithm itself. The algorithm was tested on real data from two different mass spectrometers, and the results show that this algorithm can be a very useful tool for the peak identification process, with a 97 % match between manually identified and algorithmically identified signal intensity for both datasets. The goal of the algorithm is to save time during the process of analyzing atmospheric mass spectra, and given run times of 40 and 5 min for the gas and particle phase datasets, respectively, it is clearly much faster for a user to revise the algorithm-generated peak list than to start from scratch.

Sensitivity tests using synthetic data show the importance of correctly defined fitting parameters. Reasonably well defined parameters do yield good results, as the tests with real data indicate. However, the algorithm results may be significantly improved by more accurately defined parameters, particularly a good mass calibration. Future work could focus on improving these parameters within the algorithm itself.

As a proof of concept we believe this work has shown that automated peak fitting and identification can achieve excellent results. The algorithm described here can already save users a lot of time during peak identification. With further improvements and more users providing feedback, the automated fits can likely be improved even further. Work is currently in progress to include the methods described here in established analysis software, such as Tofware, which would allow easy utilization of these methods by a wider community.

Appendix A

A1 Score function

The Bayesian information criterion (BIC) is defined as

$$BIC = -2\ln\left(L\left(\Theta_n, \mathbf{y}\right)\right) + n\ln(k), \tag{A1}$$

where Θ_n is the parameter vector with *n* parameters, and *k* is the number of data points in the data matrix **y**. Assuming every data point y_i is normally distributed around a fit $\hat{y}_{n,i}$,



Figure A1. Example of synthetic data generation at integer mass 306 with a mass resolution of 4000. Three ions, with exact positions displayed in the legend, along with the baseline contribute to the expected signal λ . Each point in the generated spectrum, *y*, is sampled from a Poisson distribution with expected value λ , resulting in noise with a standard deviation of $\sqrt{\lambda}$.

with variance σ^2 , the likelihood function becomes

$$L(\mathbf{\Theta}_n, \mathbf{y}) = \prod_{i=1}^k \frac{1}{\sigma(x_i)\sqrt{2\pi}} \exp\left(-\frac{1}{2}\left(\frac{y_i - \hat{y}_{n,i}}{\sigma(x_i)}\right)^2\right), \quad (A2)$$

and the log-likelihood function is

$$\ln\left(L\left(\mathbf{\Theta}_{n},\mathbf{y}\right)\right) = k \ln\left(\frac{1}{\sqrt{2\pi}}\right) + \sum_{i=1}^{k} \\ \ln\left(\frac{1}{\sigma(x_{i})}\right) - \frac{1}{2} \sum_{i=1}^{k} \left(\frac{y_{i} - \hat{y}_{n,i}}{\sigma(x_{i})}\right)^{2}.$$
(A3)

Since we are looking for the number of parameters *n* that minimizes BIC, the first term is irrelevant, as it is independent of *n*. The second term is also omitted since the change in estimated error should not vary significantly between fits. Counting statistics suggest $\sigma(x_i) \propto \sqrt{\hat{y}_{n,i}}$. Thus, including a proportionality constant, *A*, yields

BIC
$$\approx A \sum_{i=1}^{n} \frac{(y_i - \hat{y}_{n,i})^2}{\hat{y}_{n,i}} + n \ln(k)$$
 (A4)

or

BIC
$$\approx A\chi_n^2 + n\ln(k)$$
. (A5)

Additional testing of the algorithm showed that the normalization of χ^2 improved results by making the value of *A* where the numbers of peaks starts to sharply increase be more consistent over a wide range of masses. This finally results in the used score function (Eq. 2).



Figure A2. Results of sensitivity tests conducted for synthetic data with a resolution around 13 000. (a) How results vary depending on parameter *A*. The solid lines depict how the scores vary for the free fit, while crosses depict the results after the formula assignment. For the free fits a fit is considered correct if within $0.2 \cdot \text{FWHM}$ (14.7 ppm at 300 Th) of the generated location; after assignment, a fit is considered correct only if the precise formula assigned is the same as was generated. (b) Influence of the number of compositions in the list of potential formulas. (c) Influence of errors in mass calibration. (d) Influence of incorrect resolution of fit peaks on fit results.

A2 Generation of potential formula lists

Potential compositions are generated by defining the following parameters:

- 1. *seeds*, which are the initial parts that carbon chains are built from, for which each composition must contain one and only one seed;
- 2. *ions*, which are parts that lead to charging the molecule, for which each composition must contain one and only one ion; and
- 3. *parts*, which are the parts that may be added to the carbon chain, for which each composition may contain between 0 and some maximum number of each part; below, when parts are specified the maximum number is before the formula for the part for example, 5(CH₂) means formulas may include between zero and five CH₂ units.

Some parts may contain negative numbers of certain elements, such as $-(H^+)$ (deprotonation) or NO₂(-H) (adds NO₂ and removes one H).

Gas phase formulas

Additional constraints for gas phase formulas are the following (all upper/lower limits are inclusive):

- the O-to-C ratio must be between 0.1 and 2.8 (for each nitrogen in the formula, three oxygen are subtracted from the number used to calculate this ratio);
- the H-to-C ratio must be between 0.6 and 2 (fluorine is counted as H for this ratio);
- the upper limits for atoms of each element are 20C, 36H, 24O, 3N, and 20F;
- the lower limits for atoms of each element are 4C, 0H, 2O, 0N, and 0F;
- the formulas without fluorine with fewer than four oxygen must be removed; and
- the formulas with exactly three nitrogen and an odd number of hydrogen must be removed.

The last criterion on the list is to make sure the formulas with three nitrogen follow the nitrogen rule. The nitrogen rule is not considered for other numbers of nitrogen atoms since formulas with zero to two nitrogen are more abundant, and even radicals can be observed.

The fluorine-containing compounds are commonly observed in datasets from negative ion CIMS instruments and originate from the Teflon tubing, chamber, and fan used in the experimental setup (Mattila et al., 2024; Zhang et al., 2020; Ehn et al., 2012).



Figure A3. Manually identified peaks also identified by the algorithm for the CIMS dataset. The x axis shows the relative distance to the nearest other peak fit by the algorithm, and the y axis shows the contribution of that peak to the total signal area at the same unit mass. Note that the closest fit for all of these formulas is actually zero, since the algorithm found the formula in the precise correct location; therefore, "other fit" is used for the x-axis label to denote that it is the closest fit not corresponding to the correct formula.

In the complete list there are three groups of formulas with individually defined parameters.

- Group 1 consists of fluorinated carboxylic acids and dicarboxylic acids:
 - seeds: CH₂O₂, CHO₂F, and CO₂F₂;
 - ions: NO_3^- and $-(H^+)$;
 - parts: 20(CF₂), 1(CO₂), 1(O), and 1(CHF).
- Group 2 consists of closed-shell carbon chains/rings:
 - seeds: CH_4 , CH_2O_2 , and $C_2H_2O_4$;
 - ions: NO_3^- and $-(H^+)$;
 - parts: 20(CH₂), 10(CO), 3(C), 4(O), 20(CH₂O), 2(NO₂(-H)), 2(NH), and 1(HNO₃).
- Group 3 consists of radicals similar to Group 2 molecules:
 - seeds: CH₃, CHO₂, and C₂HO₄;
 - ions: NO_3^- ;
 - parts: 20(CH₂), 5(CO), 3(C), 4(O), and 20(CH₂O).

Particle phase formulas

Additional constraints for particle phase formulas are the following (all upper/lower limits are inclusive):

- the O-to-C ratio must be between 0 and 2;
- the H-to-C ratio must be between 0 and 4, and additionally the number of H atoms must not be less than the number of C atoms minus 4;



Figure A4. Manually identified peaks also identified by the algorithm for the AMS dataset. The x axis shows the relative distance to the nearest other peak fit by the algorithm, and the y axis shows the contribution of that peak to the total signal area at the same unit mass. Note that the closest fit for all of these formulas is actually zero, since the algorithm found the formula in the precise correct location; therefore, "other fit" is used for the x-axis label to denote that it is the closest fit not corresponding to the correct formula.

- the upper limits for atoms of each element are 12C, 26H, 8O, 1N, and 1S;
- the lower limit for atoms of each element is 0.

Both the constraints and the parts used to build the formulas in this list are less general than for the gas phase dataset. This is because the AMS mostly detects fragments of molecules so valency rules cannot be used to constrain the list.

In the list there are two groups of formulas with individually defined parameters.

- Group 1 consists of compounds without sulfur:
 - seeds: C; - ions: $-e^-$;
 - parts: C, H, O, and N.
- Group 2 consists of compounds with sulfur:
 - seeds: S;
 - ions: $-e^-$;
 - parts: C, H, and O.

For Group 2 the upper limit for each element was lowered to 4C, 6H, 6O, and 1S.

A3 Fitting to select A value

This section describes the fitting used to select a good value of parameter A, such as the fit shown in Fig. 2b. The fitted function is $\exp(-b(\overline{n}_{\max} - \overline{n}(A))^2)$ as mentioned in Sect. 2.1.2. The fitting is done only within set intervals of



Figure A5. Similar to Figs. 6a and A3, showing all the manually fit formulas to the gas phase dataset, but with absolute units of mass difference. Note that for the blue markers the x-axis distance is to the nearest other fit, similarly to Fig. A3. Certain common differences in mass defect are marked with lines in the figure. The lines are labeled according to the two collections of atoms that the difference in mass corresponds to; for example, CH₄ - O refers to the difference in mass between CH₄ and O (all elemental symbols refer to the most common isotopes of that element). As a further example, if both the manual fit and the algorithm fit identified the two formulas $C_{10}H_{16}O_{13}N_3^-$ and $C_{11}H_{16}O_{14}N_1^-$, then this would result in one blue and one red marker on the dashed vertical line labeled $N_2 - CO$, given that there were no other closer peaks. Note that the fact that some mass defects only contain red markers does not suggest that the algorithm always selects the wrong option between the defects but rather that the algorithm never attempts to fit both of these options. The absence of blue markers is due to the fact that if the algorithm was to select the correct option, there would not be any other nearby fit, which would move that blue marker somewhere further to the right. Here the black markers are also more clearly separated depending on whether they were labeled as "unknown" or some other formula. If a black marker did not lie on a common mass defect line, it was very likely labeled as "unknown", since the position of the closest fit does not precisely correspond to a common mass difference. Had it been misidentified, the marker would instead likely lie on one of the common mass difference lines, since both the algorithm and manual fits would have assigned it a different chemical formula.

 $[\overline{n}_{\text{start}}, \overline{n}_{\text{max}}]$, where $\overline{n}_{\text{start}}$ is varied from 0 to $\overline{n}_{\text{max}}$. For the final fit the lowest $\overline{n}_{\text{start}}$ first local minimum of the mean squared error of the fit is selected. If there is no local minimum, the point with the slowest change in mean squared error is selected instead. The purpose of this selection of $\overline{n}_{\text{start}}$ is to get a fit where the two curves in Fig. 2b match well



Figure A6. Same as Fig. A5 but for particle phase dataset.

on the right-hand side, without being biased by the left-hand side. The point where the mean relative χ^2 value is more than 5 % higher than the fit value is then selected for parameter A.



Figure A7. Summary of fits to gas phase data in a similar way to Fig. 6c but this time instead plotting the number of matching peaks versus the number of manually fit peaks and the number of algorithmically fit peaks.



Figure A8. Summary of fits to particle phase data in a similar way to Fig. 7c but this time instead plotting the number of matching peaks versus the number of manually fit peaks and the number of algorithmically fit peaks.



Figure A9. Comparisons between manual (a, c, e) and algorithm (b, d, f) fits to three example unit masses of different complexities for the AMS data. The fitted peaks in the legends are sorted from highest to lowest intensity (top to bottom).

-	200	-	226	-	201
Integer mass:	308	Integer mass:	326	Integer mass:	386
$C_5H_2O_5N_1F_8^-$	307.9811	$C_5H_1O_5N_1F_9^-$	325.9716	$C_6H_1O_8N_1F_9^-$	385.9564
$C_8H_6O_{12}N_1^{-1}$	307.9895	$C_6H_4O_{13}N_3^{-1}$	325.975	$C_9H_8O_{16}N_1^{-1}$	385.9849
$C_7H_6O_{11}N_3^{-}$	308.0008	$C_8H_8O_{13}N_1^{-1}$	326.0001	$C_8H_8O_{15}N_3^{-1}$	385.9961
$C_9H_{10}O_{11}N_1^-$	308.0259	$C_7H_8O_{12}N_3^{-1}$	326.0113	$C_{13}H_8O_{13}N_1^-$	386.0001
$C_8H_{10}O_{10}N_3^{-}$	308.0372	$C_{12}H_8O_{10}N_1^-$	326.0154	$C_{12}H_8O_{12}N_3^{-1}$	386.0113
$C_{13}H_{10}O_8N_1^{-}$	308.0412	$C_{11}H_8O_9N_3^{-1}$	326.0266	$C_{10}H_{12}O_{15}N_1^-$	386.0212
$C_{10}H_{14}O_{10}N_1^{-}$	308.0623	$C_9H_{12}O_{12}N_1^-$	326.0365	$C_9H_{12}O_{14}N_3^{-1}$	386.0325
$C_9H_{14}O_9N_3^{-1}$	308.0736	$C_8H_{12}O_{11}N_3^{-1}$	326.0477	$C_{14}H_{12}O_{12}N_1^-$	386.0365
$C_{14}H_{14}O_7N_1^-$	308.0776	$C_{13}H_{12}O_9N_1^{-1}$	326.0518	$C_{13}H_{12}O_{11}N_3^{-}$	386.0477
$C_{11}H_{18}O_9N_1^{-}$	308.0987	$C_{12}H_{12}O_8N_3^{-1}$	326.063	$C_{11}H_{16}O_{14}N_1^{-1}$	386.0576
$C_{10}H_{18}O_8N_3^{-}$	308.1099	$C_{10}H_{16}O_{11}N_1^-$	326.0729	$C_{10}H_{16}O_{13}N_3^{-}$	386.0689
$C_{15}H_{18}O_6N_1^{-}$	308.114	$C_9H_{16}O_{10}N_3^{-1}$	326.0841	$C_{15}H_{16}O_{11}N_1^{-1}$	386.0729
$C_{12}H_{22}O_8N_1^{-}$	308.1351	$C_{14}H_{16}O_8N_1^{-1}$	326.0881	$C_{14}H_{16}O_{10}N_3^{-}$	386.0841
$C_{16}H_{22}O_5N_1^{-1}$	308.1503	$C_{11}H_{20}O_{10}N_1^-$	326.1093	$C_{12}H_{20}O_{13}N_1^{-}$	386.094
$C_{13}H_{26}O_7N_1^{-1}$	308.1715	$C_{10}H_{20}O_9N_3^{-1}$	326.1205	$C_{11}H_{20}O_{12}N_3^{-1}$	386.1052
$C_{17}H_{26}O_4N_1^{-}$	308.1867	$C_{15}H_{20}O_7N_1^{-}$	326.1245	$C_{16}H_{20}O_{10}N_1^{-}$	386.1093
		$C_{12}H_{24}O_9N_1^{-}$	326.1457	$C_{15}H_{20}O_9N_3^{-1}$	386.1205
		$C_{16}H_{24}O_6N_1^{-1}$	326.1609	$C_{13}H_{24}O_{12}N_1^-$	386.1304
		$C_{17}H_{28}O_5N_1^{-1}$	326.1973	$C_{12}H_{24}O_{11}N_3^{-1}$	386.1416
		$C_{18}H_{32}O_4N_1^{-}$	326.2337	$C_{17}H_{24}O_9N_1^{-3}$	386.1457
		1		$C_{16}H_{24}O_8N_3^{-1}$	386.1569
				$C_{14}H_{28}O_{11}N_1^-$	386.1668
				$C_{18}H_{28}O_8N_1^{-1}$	386.182
				$C_{19}H_{32}O_7N_1^{-1}$	386.2184
				$C_{20}H_{36}O_6N_1^{-1}$	386.2548
				1	

Table A1. Examples of all the potential compounds for the gas phase dataset at the integer masses presented in Fig. 5.

Integer mass:	66	Integer mass:	100	Integer mass:	137
$H_2S_2^+$	65.9592	$C_{3}S_{2}^{+}$	99.9436	$C_2H_1O_3S_2^+$	136.9362
$H_2O_2S_1^+$	65.977	$C_{3}O_{2}S_{1}^{+}$	99.9614	$C_3H_5O_2S_2^{\mp}$	136.9725
$C_{3}O_{1}N_{1}^{+}$	65.9974	$H_4O_2S_2^+$	99.9647	$C_3H_5O_4S_1^{\mp}$	136.9903
$C_4H_2O_1^+$	66.01	$C_{3}O_{4}^{+}$	99.9791	$C_3H_5O_6^+$	137.0081
$C_4H_4N_1^{\uparrow}$	66.0338	$H_4O_4S_1^+$	99.9825	$C_{6}H_{3}O_{3}N_{1}^{+}$	137.0107
$C_5H_6^+$	66.0464	$C_4H_4O_1S_1^+$	99.9977	$C_7H_5O_3^+$	137.0233
		$C_3H_2O_3N_1^+$	100.0029	$C_3H_7O_5N_1^+$	137.0319
		$C_4H_4O_3^+$	100.0155	$C_4H_9O_5^+$	137.0444
		$C_8H_4^+$	100.0308	$C_7H_7O_2N_1^+$	137.0471
		$C_4H_6O_2N_1^+$	100.0393	$C_8H_9O_2^+$	137.0597
		$C_5H_8O_2^+$	100.0519	$C_4H_{11}O_4N_1^+$	137.0683
		$C_5H_{10}\bar{O}_1N_1^+$	100.0757	$C_5H_{13}O_4^+$	137.0808
		$C_6H_{12}O_1^+$	100.0883	$C_8H_{11}O_1N_1^+$	137.0835
		$C_{6}H_{14}N_{1}^{+}$	100.1121	$C_9H_{13}O_1^+$	137.0961
		$C_7 H_{16}^+$	100.1247	$C_5H_{15}O_3N_1^+$	137.1046
				$C_6H_{17}O_3^+$	137.1172
				$C_9H_{15}N_1^+$	137.1199
				$C_{10}H_{17}^+$	137.1325
				$C_6H_{19}O_2N_1^+$	137.141
				$C_7H_{21}O_2^+$	137.1536
				$C_7H_{23}O_1N_1^+$	137.1774
				$C_8H_{25}O_1^+$	137.19

Table A2. Examples of all the potential compounds for the particle phase dataset at the integer masses presented in Fig. A9.

Code and data availability. The algorithm is available in its entirety on GitHub (https://github.com/VMickwitz/ PeakIdentificationAlgorithm, last access: 19 March 2025) or through Zenodo (https://doi.org/10.5281/zenodo.14753810, Mickwitz, 2025) along with the lists of potential formulas and fit results for the analyzed datasets.

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Author contributions. VM designed the algorithm, analyzed the results, and prepared the manuscript. ME, OP, FG, and DW provided constructive comments on methods and result interpretation. OP and FG provided the manual analyses for the gas and particle phase datasets, respectively. ME assisted in the preparation of the manuscript.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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References

- Alton, M. W., Stark, H. J., Canagaratna, M. R., and Browne, E. C.: Generalized Kendrick analysis for improved visualization of atmospheric mass spectral data, Atmos. Meas. Tech., 16, 3273– 3282, https://doi.org/10.5194/amt-16-3273-2023, 2023.
- Batey, J. H.: The Physics and Technology of Quadrupole Mass Spectrometers, Vacuum, 101, 410–415, https://doi.org/10.1016/j.vacuum.2013.05.005, 2014.

- Boesl, U.: Time-of-Flight Mass Spectrometry: Introduction to the Basics, Mass Spectrom. Rev., 36, 86–109, https://doi.org/10.1002/mas.21520, 2017.
- Canagaratna, M., Jayne, J., Jimenez, J., Allan, J., Alfarra, M., Zhang, Q., Onasch, T., Drewnick, F., Coe, H., Middlebrook, A., Delia, A., Williams, L., Trimborn, A., Northway, M., DeCarlo, P., Kolb, C., Davidovits, P., and Worsnop, D.: Chemical and Microphysical Characterization of Ambient Aerosols with the Aerodyne Aerosol Mass Spectrometer, Mass Spectrom. Rev., 26, 185– 222, https://doi.org/10.1002/mas.20115, 2007.
- Cubison, M. J. and Jimenez, J. L.: Statistical precision of the intensities retrieved from constrained fitting of overlapping peaks in high-resolution mass spectra, Atmos. Meas. Tech., 8, 2333– 2345, https://doi.org/10.5194/amt-8-2333-2015, 2015.
- Ehn, M., Kleist, E., Junninen, H., Petäjä, T., Lönn, G., Schobesberger, S., Dal Maso, M., Trimborn, A., Kulmala, M., Worsnop, D. R., Wahner, A., Wildt, J., and Mentel, Th. F.: Gas phase formation of extremely oxidized pinene reaction products in chamber and ambient air, Atmos. Chem. Phys., 12, 5113–5127, https://doi.org/10.5194/acp-12-5113-2012, 2012.
- Ehn, M., Thornton, J. A., Kleist, E., Sipilä, M., Junninen, H., Pullinen, I., Springer, M., Rubach, F., Tillmann, R., Lee, B., Lopez-Hilfiker, F., Andres, S., Acir, I.-H., Rissanen, M., Jokinen, T., Schobesberger, S., Kangasluoma, J., Kontkanen, J., Nieminen, T., Kurtén, T., Nielsen, L. B., Jørgensen, S., Kjaergaard, H. G., Canagaratna, M., Maso, M. D., Berndt, T., Petäjä, T., Wahner, A., Kerminen, V.-M., Kulmala, M., Worsnop, D. R., Wildt, J., and Mentel, T. F.: A Large Source of Low-Volatility Secondary Organic Aerosol, Nature, 506, 476–479, https://doi.org/10.1038/nature13032, 2014.
- Eichler, P., Müller, M., D'Anna, B., and Wisthaler, A.: A novel inlet system for online chemical analysis of semi-volatile submicron particulate matter, Atmos. Meas. Tech., 8, 1353–1360, https://doi.org/10.5194/amt-8-1353-2015, 2015.
- Fowler, D., Pilegaard, K., Sutton, M. A., Ambus, P., Raivonen, M., Duyzer, J., Simpson, D., Fagerli, H., Fuzzi, S., Schjoerring, J. K., Granier, C., Neftel, A., Isaksen, I. S. A., Laj, P., Maione, M., Monks, P. S., Burkhardt, J., Daemmgen, U., Neirynck, J., Personne, E., Wichink-Kruit, R., Butterbach-Bahl, K., Flechard, C., Tuovinen, J. P., Coyle, M., Gerosa, G., Loubet, B., Altimir, N., Gruenhage, L., Ammann, C., Cieslik, S., Paoletti, E., Mikkelsen, T. N., Ro-Poulsen, H., Cellier, P., Cape, J. N., Horváth, L., Loreto, F., Niinemets, Ü., Palmer, P. I., Rinne, J., Misztal, P., Nemitz, E., Nilsson, D., Pryor, S., Gallagher, M. W., Vesala, T., Skiba, U., Brüggemann, N., Zechmeister-Boltenstern, S., Williams, J., O'Dowd, C., Facchini, M. C., de Leeuw, G., Flossman, A., Chaumerliac, N., and Erisman, J. W.: Atmospheric Composition Change: Ecosystems-Atmosphere Interactions, Atmos. Environ., 43, 5193-5267, https://doi.org/10.1016/j.atmosenv.2009.07.068, 2009.
- Goldstein, A. H. and Galbally, I. E.: Known and Unexplored Organic Constituents in the Earth's Atmosphere, Environ. Sci. Technol., 41, 1514–1521, 2007.
- Graeffe, F., Heikkinen, L., Garmash, O., Äijälä, M., Allan, J., Feron, A., Cirtog, M., Petit, J.-E., Bonnaire, N., Lambe, A., Favez, O., Albinet, A., Williams, L. R., and Ehn, M.: Detecting and Characterizing Particulate Organic Nitrates with an Aerodyne Long-ToF Aerosol Mass Spectrometer, ACS Earth Space Chem., 7, 230– 242, https://doi.org/10.1021/acsearthspacechem.2c00314, 2023.

- Häkkinen, E., Zhao, J., Graeffe, F., Fauré, N., Krechmer, J. E., Worsnop, D., Timonen, H., Ehn, M., and Kangasluoma, J.: Online measurement of highly oxygenated compounds from organic aerosol, Atmos. Meas. Tech., 16, 1705–1721, https://doi.org/10.5194/amt-16-1705-2023, 2023.
- Heal, M., Kumar, P., and M. Harrison, R.: Particles, Air Quality, Policy and Health, Chem. Soc. Rev., 41, 6606–6630, https://doi.org/10.1039/C2CS35076A, 2012.
- Hu, Q., Noll, R. J., Li, H., Makarov, A., Hardman, M., and Graham Cooks, R.: The Orbitrap: A New Mass Spectrometer, J. Mass Spectro., 40, 430–443, https://doi.org/10.1002/jms.856, 2005.
- Huey, L. G.: Measurement of Trace Atmospheric Species by Chemical Ionization Mass Spectrometry: Speciation of Reactive Nitrogen and Future Directions, Mass Spectrom. Rev., 26, 166–184, https://doi.org/10.1002/mas.20118, 2007.
- Jokinen, T., Sipilä, M., Junninen, H., Ehn, M., Lönn, G., Hakala, J., Petäjä, T., Mauldin III, R. L., Kulmala, M., and Worsnop, D. R.: Atmospheric sulphuric acid and neutral cluster measurements using CI-APi-TOF, Atmos. Chem. Phys., 12, 4117–4125, https://doi.org/10.5194/acp-12-4117-2012, 2012.
- Jokinen, T., Berndt, T., Makkonen, R., Kerminen, V.-M., Junninen, H., Paasonen, P., Stratmann, F., Herrmann, H., Guenther, A. B., Worsnop, D. R., Kulmala, M., Ehn, M., and Sipilä, M.: Production of Extremely Low Volatile Organic Compounds from Biogenic Emissions: Measured Yields and Atmospheric Implications, P. Natl. Acad. Sci. USA, 112, 7123–7128, https://doi.org/10.1073/pnas.1423977112, 2015.
- Junninen, H., Ehn, M., Petäjä, T., Luosujärvi, L., Kotiaho, T., Kostiainen, R., Rohner, U., Gonin, M., Fuhrer, K., Kulmala, M., and Worsnop, D. R.: A high-resolution mass spectrometer to measure atmospheric ion composition, Atmos. Meas. Tech., 3, 1039– 1053, https://doi.org/10.5194/amt-3-1039-2010, 2010.
- Kulmala, M., Toivonen, A., Mäkelä, J. M., and Laaksonen, A.: Analysis of the Growth of Nucleation Mode Particles Observed in Boreal Forest, Tellus B, 50, 449–462, https://doi.org/10.1034/j.1600-0889.1998.t01-4-00004.x, 1998.
- Lee, H.-N. and Marshall, A. G.: Theoretical Maximal Precision for Mass-to-Charge Ratio, Amplitude, and Width Measurements in Ion-Counting Mass Analyzers, Anal. Chem., 72, 2256–2260, https://doi.org/10.1021/ac991256u, 2000.
- Lopez-Hilfiker, F. D., Mohr, C., Ehn, M., Rubach, F., Kleist, E., Wildt, J., Mentel, Th. 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), Atmos. Meas. Tech., 7, 983–1001, https://doi.org/10.5194/amt-7-983-2014, 2014.
- Lopez-Hilfiker, F. D., Pospisilova, V., Huang, W., Kalberer, M., Mohr, C., Stefenelli, G., Thornton, J. A., Baltensperger, U., Prevot, A. S. H., and Slowik, J. G.: An extractive electrospray ionization time-of-flight mass spectrometer (EESI-TOF) for online measurement of atmospheric aerosol particles, Atmos. Meas. Tech., 12, 4867–4886, https://doi.org/10.5194/amt-12-4867-2019, 2019.
- Masson-Delmotte, V., Zhai, P., Pirani, A., Connors, S. L., Pean, C., Berger, S., Caud, N., Chen, Y., Goldfarb, L., Gomis, M. I., Huang, M., Leitzell, K., Lonnoy, E., Matthews, J. B. R., Maycock, T. K., Waterfield, T., Yelekci, O., Yu, R., and Zhou, B.: IPCC, 2021: Climate Change 2021: The Physical Science

Basis. Contribution of Working Group I to the Sixth Assessment Report of the Intergovernmental Panel on Climate Change, Report, Cambridge University Press, Cambridge, UK, ISBN 9781009157896, 2021.

- Mattila, J. M., Krug, J. D., Roberson, W. R., Burnette, R. P., Mc-Donald, S., Virtaranta, L., Offenberg, J. H., and Linak, W. P.: Characterizing Volatile Emissions and Combustion Byproducts from Aqueous Film-Forming Foams Using Online Chemical Ionization Mass Spectrometry, Environ. Sci. Technol., 58, 3942– 3952, 2024.
- Mickwitz, V.: Code and analyzed datasets, Zenodo [code and datasets], https://doi.org/10.5281/zenodo.14753810, 2025.
- Neath, A. A. and Cavanaugh, J. E.: The Bayesian Information Criterion: Background, Derivation, and Applications, WIREs Comput. Stat., 4, 199–203, https://doi.org/10.1002/wics.199, 2012.
- Peräkylä, O., Riva, M., Heikkinen, L., Quéléver, L., Roldin, P., and Ehn, M.: Experimental investigation into the volatilities of highly oxygenated organic molecules (HOMs), Atmos. Chem. Phys., 20, 649–669, https://doi.org/10.5194/acp-20-649-2020, 2020.
- Riipinen, I., Pierce, J. R., Yli-Juuti, T., Nieminen, T., Häkkinen, S., Ehn, M., Junninen, H., Lehtipalo, K., Petäjä, T., Slowik, J., Chang, R., Shantz, N. C., Abbatt, J., Leaitch, W. R., Kerminen, V.-M., Worsnop, D. R., Pandis, S. N., Donahue, N. M., and Kulmala, M.: Organic condensation: a vital link connecting aerosol formation to cloud condensation nuclei (CCN) concentrations, Atmos. Chem. Phys., 11, 3865–3878, https://doi.org/10.5194/acp-11-3865-2011, 2011.
- Rissanen, M. P., Mikkilä, J., Iyer, S., and Hakala, J.: Multi-scheme chemical ionization inlet (MION) for fast switching of reagent ion chemistry in atmospheric pressure chemical ionization mass spectrometry (CIMS) applications, Atmos. Meas. Tech., 12, 6635–6646, https://doi.org/10.5194/amt-12-6635-2019, 2019.
- Riva, M., Ehn, M., Li, D., Tomaz, S., Bourgain, F., Perrier, S., and George, C.: CI-Orbitrap: An Analytical Instrument To Study Atmospheric Reactive Organic Species, Anal. Chem., 91, 9419– 9423, https://doi.org/10.1021/acs.analchem.9b02093, 2019a.
- Riva, M., Rantala, P., Krechmer, J. E., Peräkylä, O., Zhang, Y., Heikkinen, L., Garmash, O., Yan, C., Kulmala, M., Worsnop, D., and Ehn, M.: Evaluating the performance of five different chemical ionization techniques for detecting gaseous oxygenated organic species, Atmos. Meas. Tech., 12, 2403–2421, https://doi.org/10.5194/amt-12-2403-2019, 2019b.
- Sandström, H., Rissanen, M., Rousu, J., and Rinke, P.: Data-Driven Compound Identification in Atmospheric Mass Spectrometry, Adv. Sci., 11, 2306235, https://doi.org/10.1002/advs.202306235, 2024.
- Shrivastava, M., Cappa, C. D., Fan, J., Goldstein, A. H., Guenther, A. B., Jimenez, J. L., Kuang, C., Laskin, A., Martin, S. T., Ng, N. L., Petaja, T., Pierce, J. R., Rasch, P. J., Roldin, P., Seinfeld, J. H., Shilling, J., Smith, J. N., Thornton, J. A., Volkamer, R., Wang, J., Worsnop, D. R., Zaveri, R. A., Zelenyuk, A., and Zhang, Q.: Recent Advances in Understanding Secondary Organic Aerosol: Implications for Global Climate Forcing, Rev. Geophys., 55, 509–559, https://doi.org/10.1002/2016RG000540, 2017.
- Stark, H., Yatavelli, R. L. N., Thompson, S. L., Kimmel, J. R., Cubison, M. J., Chhabra, P. S., Canagaratna, M. R., Jayne, J. T., Worsnop, D. R., and Jimenez, J. L.: Methods to Extract Molecular and Bulk Chemical Information from Series of Complex Mass

Spectra with Limited Mass Resolution, Int. J. Mass Spectrom., 389, 26–38, https://doi.org/10.1016/j.ijms.2015.08.011, 2015.

- VMickwitz: PeakIdentificationAlgorithm, GitHub [data set], https: //github.com/VMickwitz/PeakIdentificationAlgorithm (last access: 19 March 2025), 2025.
- Zhang, R., Khalizov, A., Wang, L., Hu, M., and Xu, W.: Nucleation and Growth of Nanoparticles in the Atmosphere, Chem. Rev., 112, 1957–2011, https://doi.org/10.1021/cr2001756, 2012.
- Zhang, W., Xu, L., and Zhang, H.: Recent Advances in Mass Spectrometry Techniques for Atmospheric Chemistry Research on Molecular-Level, Mass Spectrom. Rev., 43, 1091–1134, https://doi.org/10.1002/mas.21857, 2023.
- Zhang, Y., Peräkylä, O., Yan, C., Heikkinen, L., Äijälä, M., Daellenbach, K. R., Zha, Q., Riva, M., Garmash, O., Junninen, H., Paatero, P., Worsnop, D., and Ehn, M.: A novel approach for simple statistical analysis of high-resolution mass spectra, Atmos. Meas. Tech., 12, 3761–3776, https://doi.org/10.5194/amt-12-3761-2019, 2019.
- Zhang, Y., Peräkylä, O., Yan, C., Heikkinen, L., Äijälä, M., Daellenbach, K. R., Zha, Q., Riva, M., Garmash, O., Junninen, H., Paatero, P., Worsnop, D., and Ehn, M.: Insights into atmospheric oxidation processes by performing factor analyses on subranges of mass spectra, Atmos. Chem. Phys., 20, 5945–5961, https://doi.org/10.5194/acp-20-5945-2020, 2020.
- Zuth, C., Vogel, A. L., Ockenfeld, S., Huesmann, R., and Hoffmann, T.: Ultrahigh-Resolution Mass Spectrometry in Real Time: Atmospheric Pressure Chemical Ionization Orbitrap Mass Spectrometry of Atmospheric Organic Aerosol, Anal. Chem., 90, 8816–8823, https://doi.org/10.1021/acs.analchem.8b00671, 2018.