

1 **Argon offline-AMS source apportionment of organic**  
2 **aerosol over yearly cycles for an urban, rural and marine**  
3 **site in Northern Europe**

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18  
19 **Abstract**

20 The widespread use of Aerodyne aerosol mass spectrometers (AMS) has greatly improved  
21 real-time organic aerosol (OA) monitoring, providing mass spectra that contain sufficient  
22 information for source apportionment. However, AMS field deployments remain expensive  
23 and demanding, limiting the acquisition of long-term datasets at many sampling sites. The  
24 offline application of aerosol mass spectrometry entailing the analysis of nebulized water  
25 extracted filter samples (offline-AMS) increases the spatial coverage accessible to AMS  
26 measurements, being filters routinely collected at many stations worldwide.

27 PM<sub>1</sub> (particulate matter with an aerodynamic diameter <1 μm) filter samples were collected  
28 during an entire year in Lithuania at three different locations representative of three typical

1 environments of the South-East Baltic region: Vilnius (urban background), Rūgšteliškis (rural  
2 terrestrial), and Preila (rural coastal). Aqueous filter extracts were nebulized in Ar, yielding  
3 the first AMS measurements of water-soluble atmospheric organic aerosol (WSOA) without  
4 interference from air fragments. This enables direct measurement of the  $\text{CO}^+$  fragment  
5 contribution, whose intensity is typically assumed to be equal to that of  $\text{CO}_2^+$ . Offline-AMS  
6 spectra reveal that the water soluble  $\text{CO}_2^+:\text{CO}^+$  ratio not only shows values systematically  $>1$   
7 but is also dependent on season, with lower values in winter than in summer.

8 AMS WSOA spectra were analyzed using positive matrix factorization (PMF), which yielded  
9 4 factors. These factors included biomass burning OA (BBOA), local OA (LOA) contributing  
10 significantly only in Vilnius, and two oxygenated OA (OOA) factors, summer OOA (S-OOA)  
11 and background OOA (B-OOA) distinguished by their seasonal variability. The contribution  
12 of traffic exhaust OA (TEOA) was not resolved by PMF due to both low concentrations and  
13 low water solubility. Therefore, the TEOA concentration was estimated using a chemical  
14 mass balance approach, based on the concentrations of hopanes, specific markers of traffic  
15 emissions. AMS-PMF source apportionment results were consistent with those obtained from  
16 PMF applied to marker concentrations (i.e. major inorganic ions, OC/EC, and organic  
17 markers including polycyclic aromatic hydrocarbons and their derivatives, hopanes, long-  
18 chain alkanes, monosaccharides, anhydrous sugars, and lignin fragmentation products). OA  
19 was the largest fraction of  $\text{PM}_{10}$  and was dominated by BBOA during winter with an average  
20 concentration of  $2 \mu\text{g m}^{-3}$  (53% of OA), while summer-OOA (S-OOA), probably related to  
21 biogenic emissions was the prevalent OA source during summer with an average  
22 concentration of  $1.2 \mu\text{g m}^{-3}$  (45% of OA).

23 PMF ascribed a large part of the  $\text{CO}^+$  explained variability (97%) to the OOA and BBOA  
24 factors. Accordingly we discuss a new  $\text{CO}^+$  parameterization as a function of  $\text{CO}_2^+$ , and  
25  $\text{C}_2\text{H}_4\text{O}_2^+$  fragments, which were selected to describe the variability of the OOA and BBOA  
26 factors.

## 27 **1 Introduction**

28 Atmospheric aerosols affect climate (Lohmann et al., 2004, Schwarze et al., 2006), human  
29 health (Dockery et al., 2005, Laden et al., 2000), and ecosystems on a global scale.  
30 Quantification and characterization of the main aerosol sources are crucial for the  
31 development of effective mitigation strategies. The Aerodyne aerosol mass spectrometer  
32 (AMS, Canagaratna et al., 2007) and aerosol chemical speciation monitor (ACSM, Ng et al.,

1 2011, Fröhlich et al., 2013) have greatly improved air quality monitoring by providing real-  
2 time measurements of the non-refractory (NR) submicron aerosol (PM<sub>1</sub>) components.  
3 Analysis of organic mass spectra using positive matrix factorization (PMF, Paatero, 1997;  
4 Paatero and Tapper, 1994) has enabled the quantitative separation of OA factors, which can  
5 be subsequently related to major aerosol sources and formation processes (e.g. Lanz et al.,  
6 2007; Lanz et al., 2010; Zhang et al., 2011; Ulbrich et al., 2009; Elser et al., 2016 a). Despite  
7 its numerous advantages, AMS field deployment remains expensive and demanding, and  
8 therefore most of the studies are typically restricted to short-time periods and a single (or few)  
9 sampling site(s). The limited amount of long-term datasets suitable for OA source  
10 apportionment severely limits model testing and validation (Aksoyoglu et al., 2011;  
11 Aksoyoglu et al., 2014; Baklanov et al., 2014), as well as for the development of appropriate  
12 pollution mitigation strategies. AMS analysis of aerosol filter samples (Lee et al., 2011; Sun  
13 et al., 2011; Mihara and Mochida, 2011; Daellenbach et al., 2016), which are routinely  
14 collected at many stations worldwide, broadens the temporal and spatial scales available for  
15 AMS measurements.

16 In this study we present the application of the offline-AMS methodology described by  
17 Daellenbach et al. (2016) to yearly cycles of filter samples collected in parallel at three  
18 different locations in Lithuania between September 2013 and August 2014. The methodology  
19 consists of water extraction of filter samples, followed by nebulization of the liquid extracts,  
20 and subsequent measurement of the generated aerosol by high-resolution time-of-flight AMS  
21 (HR-ToF AMS). In this work, organic aerosol water extracts were nebulized in Ar, permitting  
22 direct measurement of the CO<sup>+</sup> ion (Fig. S1), which is typically not directly quantified in  
23 AMS data analysis due to interference with N<sub>2</sub><sup>+</sup>, but is instead estimated as being equal to  
24 CO<sub>2</sub><sup>+</sup> (Aiken et al., 2008). Direct measurement of CO<sub>2</sub><sup>+</sup> better captures the variability in the  
25 total OA mass and its elemental composition as well as potentially improving source  
26 apportionment of ambient aerosol. Aerosol elemental ratios and oxidation state are of  
27 particular relevance as they provide important constraints for understanding aerosol sources,  
28 processes, and for the development of predictive aerosol models (Canagaratna et al., 2015).

29 Aerosol composition in the south-east Baltic region has so far received little attention. To our  
30 knowledge the only investigation of OA sources in this area was during a five-day period of  
31 intense land clearing activity occurring in the neighboring Russian enclave of Kaliningrad  
32 (Ulevicius et al., 2016; Dudoitis et al., 2016), in which transported biomass burning emissions

1 dominated the aerosol loading. OA source contributions under less extreme conditions remain  
2 unstudied, with the most relevant measurements performed in Estonia with a mobile lab  
3 during March 2014 at two different locations (Elser et al., 2016b). On-road measurements  
4 revealed large traffic contributions with an increase of 20% from rural to urban environments.  
5 Also, residential biomass burning (BB) and oxygenated OA (OOA) contributions were found  
6 to be substantial.

7 In this study we present a complete source apportionment of the submicron OA fraction  
8 following the methodology described by Daellenbach et al. (2016) in order to quantify and  
9 characterize the main OA sources affecting the Lithuanian air quality. The three sampling  
10 stations were situated in the Vilnius suburb (urban background), Preila (rural coastal  
11 background), and Rūgšteliškis (rural terrestrial background), covering a wide geographical  
12 domain and providing a good overview of the most typical Lithuanian and south-eastern  
13 Baltic air quality conditions and environments. PMF analysis of offline-AMS measurements  
14 are compared with the results reported by Ulevicius et al. (2016) and with PMF analysis of  
15 chemical marker measurements obtained from the same filter samples.

## 16 **2 Sampling and offline measurements**

### 17 **2.1 Site description and sample collection**

18 We collected 24-h integrated PM<sub>1</sub> filter samples at 3 different stations in Lithuania from 30  
19 September 2013 to 2 September 2014 using 3 High-Volume samplers (Digitel DHA80, and  
20 DH-77) operating at 500 L min<sup>-1</sup>. In order to prevent large negative filter artifacts, the high  
21 volume were equipped with temperature control systems maintaining the filter storage  
22 temperature always below 25°C, which is lower or comparable to the maximum daily  
23 temperature during summer. The particulate matter was collected on 150-mm diameter quartz  
24 fiber filters (Pallflex Tissuquartz 2500QAT-UP / pure quartz, no binder) pre-baked at 800°C  
25 for 8 h. Filter samples were wrapped in pre-baked aluminum foils (400°C for 6 h), sealed in  
26 polyethylene bags and stored at -20°C after exposure. Field blanks were collected and stored  
27 following the same procedure.

28 Sampling was conducted at urban (Vilnius), rural terrestrial (Rūgšteliškis) and rural coastal  
29 (Preila) monitoring sites (Fig. 1). The rural terrestrial site of Rūgšteliškis serves as a baseline  
30 against which urban-specific sources in the major population center of Vilnius can be

1 compared. The rural coastal site of Preila provides an opportunity to distinguish terrestrial and  
2 marine sources.

3 The sampling station in Vilnius is located at the Center for Physical Sciences and Technology  
4 campus (54°38' N, 25°10' E, 165 m a.s.l.) 12 km southwest of the city center (population:  
5 535000) and is classified as an urban background site. The site is relatively far from busy  
6 roads, and surrounded by forests to the north/northeast, and by a residential zone to the  
7 south/east. It is ca. 350 km distant from the Baltic coast, and 98 km from the Rūgšteliškis  
8 station (Fig. 1).

9 The station in Preila (55°55' N, 21°04' E, 5 m a.s.l.) is a representative rural coastal  
10 background site, situated in the Curonian Spit National Park on the isthmus separating the  
11 Baltic Sea from the Curonian Lagoon. The monitoring station is located <100 m from the  
12 Baltic shore. The closest populated area is the village of Preila (population: 200 inhabitants),  
13 located 2 km to the south.

14 The rural terrestrial station of Rūgšteliškis (55°26' N and 26°04' E, 170 m a.s.l.) is located in  
15 the eastern part of Lithuania, about 350 km from the Baltic Sea. The site is surrounded by  
16 forest and borders the Utenas Lake in the southwest. The nearest residential areas are  
17 Tauragnai, Utena (12 km and 26 km west of the station, population: 32000 inhabitants) and  
18 Ignalina (17 km southeast of the station, population: 6000 inhabitants).

## 19 **2.2 Offline-AMS analysis**

20 The term *offline-AMS* will be used herein to refer to the methodology described by  
21 Daellenbach et al. (2016) and summarized below. For each analyzed filter sample, four 16-  
22 mm diameter filter punches were subjected to ultrasonic extraction in 15 mL of ultrapure  
23 water (18.2 MΩ cm at 25°C, total organic carbon (TOC) < 3 ppb) for 20 min at 30°C.

24 The choice of water instead of an organic solvent is motivated by two arguments:

- 25 - Water yields the lowest background and hence the highest signal to noise compared to  
26 other highly pure solvents (including methanol, dichloromethane and ethyl acetate).
- 27 - In contrast to the water extraction, the use of organic solvents precludes the  
28 quantification of the organic content in the extracts (e.g. by using a total OC analyzer),  
29 which in turn prevents a quantitative source apportionment.

1 Liquid extracts were then filtered and atomized in Ar ( $\geq 99,998$  % Vol. abs., Carbagas, CH-  
2 3073 Gümligen, Switzerland) using an Apex Q nebulizer (Elemental Scientific Inc., Omaha  
3 NE 68131 USA) operating at 60°C. The resulting aerosol was then dried by passing through a  
4 Nafion drier (Perma Pure, Toms River NJ 08755 USA), and subsequently analyzed by a HR-  
5 ToF-AMS. 12 mass spectra per filter sample were collected (AMS V-mode,  $m/z$  12-232, 30 s  
6 collection time per spectrum). A measurement blank was recorded before and after each  
7 sample by nebulizing ultrapure water for 12 minutes. Field blanks were measured following  
8 the same extraction procedure as the collected filter samples, yielding a signal not statistically  
9 different from that of nebulized milliQ water. Finally we registered the AMS fragmentation  
10 spectrum of pure gaseous CO<sub>2</sub> ( $\geq 99,7$  % Vol, Carbagas, CH-3073 Gümligen, Switzerland), in  
11 order to derive its CO<sub>2</sub><sup>+</sup>:CO<sup>+</sup> ratio.

12 Offline-AMS analysis was performed on 177 filter samples in order to determine the bulk  
13 water-soluble organic matter (WSOM) mass spectral fingerprints. In total, 63 filters from  
14 Rūgšteliškis, 42 from Vilnius, and 71 from Preila were measured in Ar. The reader is referred  
15 to DeCarlo et al. (2006) for a thorough description of the AMS operating principles and  
16 calibration procedures.

17 HR-ToF-AMS analysis software SQUIRREL (SeQUential Igor data RetRiEvaL, D. Sueper,  
18 University of Colorado, Boulder, CO, USA) v.1.53G and PIKA (Peak Integration by Key  
19 Analysis) v.1.11L for IGOR Pro software package (Wavemetrics, Inc., Portland, OR, USA)  
20 were utilized to process and analyze the AMS data. HR analysis of the AMS mass spectra was  
21 performed in the  $m/z$  range 12-115.

## 22 **2.3 Supporting measurements**

23 Additional offline analyses were carried out in order to validate and corroborate the offline-  
24 AMS source apportionment results. This supporting dataset was also used as input for PM<sub>1</sub>  
25 source apportionment as discussed below. The complete list of the measurements performed  
26 can be found in Table 1 and Table S1. Briefly, major ions were measured by ion  
27 chromatography (IC; Jaffrezo et al., 1998); elemental and organic carbon (EC, OC) were  
28 quantified by thermal optical transmittance following the EUSAAR2 protocol (Cavalli et al.,  
29 2010); water-soluble OC (WSOC) was measured by water extraction followed by catalytic  
30 oxidation and non-dispersive infrared detection of CO<sub>2</sub> using a total organic carbon analyzer  
31 (Jaffrezo et al., 2005). Organic markers were determined by gas chromatography-mass

1 spectrometry (GC-MS; Golly et al., 2015); high performance liquid chromatography (HPLC)  
 2 associated with a fluorescence detector (LC 240 Perkin Elmer) and HPLC-pulsed  
 3 amperometric detection (PAD; Waked et al., 2014) for 67 composite samples. Composites  
 4 were created merging two consecutive filter samples, but no measurements are available for  
 5 Vilnius during summer. Measurements included 18 polycyclic aromatic hydrocarbons  
 6 (PAHs), alkanes (C21-C40), 10 hopanes, 13 methoxyphenols, 13 methyl-PAHs (Me-PAHs), 6  
 7 sulfur-containing-PAHs (S-PAHs), 3 monosaccharide anhydrides, and 4 monosaccharides  
 8 (including glucose, mannose, arabitol, and mannitol). In this work ion concentrations always  
 9 refer to the IC measurements.

10 Table 1. Overview of supporting measurements. A complete list of measured compounds can  
 11 be found in table S1.

<b>Analytical Method</b>	<b>Measured compounds</b>	<b>Filters measured</b>
IC (Jaffrezo et al., 1998)	Ions EC/OC	All
Thermal optical transmittance using Sunset Lab Analyzer (Birch and Cary, 1996) using EUSAAR2 protocol (Cavalli et al., 2010)		All
TOC analyzer using persulphate oxidation at 100°C of the OM, followed by CO <sub>2</sub> quantification with a non-dispersive infrared spectrophotometer (Jaffrezo et al., 1998)	WSOC	All
HPLC associated with fluorescence detector (LC 240 Perkin Elmer) (Golly et al., 2015, Besombes et al., 2001)	PAHs (table S1)	67 composite samples
GC-MS (with and without derivatization step) (Golly et al., 2015)	S-PAHs, Me-PAHs, alkanes, hopanes, methoxyphenols, others	67 composite samples

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HPLC-PAD, (Waked et al., 2014)	Anhydrous sugars, sugars alcohols, monosaccharides	67 composite samples
Chemiluminescence (Environnement S.A., Model AC31M)	NO <sub>x</sub>	Online (Vilnius only)

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1 In the following, subscripts *avg*, and *med* will denote average and median values,  
2 respectively.

### 3 3 Source apportionment

4 Positive matrix factorization (PMF, Paatero and Tapper, 1994) is a bilinear statistical model  
5 used to describe the variability of a multivariate dataset as the linear combination of a set of  
6 constant factor profiles and their corresponding time series, as shown in Eq. (1):

$$7 \quad x_{i,j} = \sum_{z=1}^p (g_{i,z} \cdot f_{z,j}) + e_{i,j} \quad (1)$$

8 Here  $x$ ,  $g$ ,  $f$ , and  $e$  denote elements of data, factor time series, factor profiles and residual  
9 matrices, respectively, while subscripts  $i,j$  and  $z$  are indices for time, measured variables, and  
10 factor number. The value  $p$  represents the total number of factors chosen for the PMF  
11 solution. The PMF algorithm iteratively solves Eq. (1) by minimizing the objective function  
12  $Q$ , defined in Eq. (2) Only non-negative  $g_{i,z}$  and  $f_{z,j}$  values are permitted:

$$13 \quad Q = \sum_i \sum_j \left( \frac{e_{i,j}}{s_{i,j}} \right)^2 \quad (2)$$

14 Here the  $s_{i,j}$  elements represent entries in the input error matrix.

15 In this work the PMF algorithm was run in the robust mode in order to dynamically  
16 downweigh the outliers. The PMF algorithm was solved using the multilinear engine-2 (ME-  
17 2) solver (Paatero, 1999), which enables an efficient exploration of the solution space by *a*  
18 *priori* constraining the  $g_{i,z}$  or  $f_{z,j}$  elements within a certain variability defined by the scalar  $a$   
19 ( $0 \leq a \leq 1$ ) such that the modelled  $g_{i,z}$ ' and  $f_{z,j}$ ' satisfy Eq. (3):

$$20 \quad \frac{(1-a)f_{z,n}}{(1+a)f_{z,m}} \leq \frac{f_{z,n'}}{f_{z,m'}} \leq \frac{(1+a)f_{z,n}}{(1-a)f_{z,m}} \quad (3)$$



1 Here  $n$  and  $m$  are any two arbitrary columns (variables) in the normalized F matrix. The  
2 Source Finder toolkit (SoFi, Canonaco et al., 2013, v.4.9) for Igor Pro software package  
3 (Wavemetrics, Inc., Portland, OR, USA) was used to configure the ME-2 model and for post-  
4 analysis. PMF analysis was applied to two complementary datasets: (1) organic mass spectra  
5 from offline-AMS measurements for the apportionment of OM sources and (2) molecular  
6 markers for the apportionment of the measured  $PM_{10}$  mass. These two analyses are discussed  
7 separately below.

8

### 9 3.1 Offline-AMS PMF

10 In the following section we describe the offline-AMS source apportionment implementation,  
11 optimization and uncertainty assessment. Briefly, we selected the number of PMF factors  
12 based on residual analyses and solution interpretability; subsequently we explored the  
13 rotational uncertainty of our source apportionment model and discarded suboptimal solutions  
14 providing insufficient correlation of factor time series with external tracers. The offline-AMS  
15 source apportionment returns the water soluble PMF factor concentrations. Daellenbach et al.  
16 (2016) determined factor specific recoveries (including PMF factor extraction efficiencies),  
17 by comparing offline-AMS and online-ACSM OA source apportionments. In particular, the  
18 filter samples were collected for one year during an online-ACSM monitoring campaign  
19 conducted at the same sampling station. Briefly, the factor recoveries were determined as the  
20 ratio between the water soluble OA PMF-factor concentrations retrieved from offline-AMS  
21 source apportionment divided by the OA PMF factor concentrations obtained from ACSM  
22 OA source apportionment. Factor specific recoveries and corresponding uncertainties were  
23 determined for HOA, BBOA, COA, and OOA. Applying these recoveries enabled scaling the  
24 water soluble factor concentrations to the corresponding bulk OA concentrations. A  
25 sensitivity analysis of these recoveries was reported in Section 3.1.3, and the corresponding  
26 uncertainty was propagated to the source apportionment results.

27 A second selection step was carried out on the rescaled solutions as described in section 3.1.3.  
28 The offline-AMS source apportionment results presented in this study represent the average  
29 of the retained rescaled PMF solutions, while their variability represents our best estimate of  
30 the source apportionment uncertainty.

### 1 3.1.1 Inputs

2 The offline-AMS input matrices include in total 177 filter samples (62 filters from  
3 Rūgštelīškis, 42 from Vilnius, and 73 from Preila). Each filter sample was represented on  
4 average by 12 mass spectral repetitions to explore the effect of AMS and nebulizer stability  
5 on PMF outputs. A corresponding measurement blank was subtracted from each mass  
6 spectrum. The input PMF matrices included 269 organic fragments fitted in the mass range  
7 (12-115). The input error  $s_{i,j}$  elements include the blank variability ( $\sigma_{i,j}$ ) and the uncertainty  
8 related to ion counting statistic and ion-to-ion signal variability at the detector ( $\delta_{i,j}$ , Allan et  
9 al., 2003; Ulbrich et al., 2009):

$$10 \quad s_{i,j} = \sqrt{\delta_{i,j}^2 + \sigma_{i,j}^2} \quad (4)$$

11 We applied a minimum error to the  $s_{i,j}$  matrix elements according to Ulbrich et al. (2009), and  
12 a down-weighting factor of 3 to all fragments with an average signal to noise lower than 2  
13 (Ulbrich et al., 2009). Input data and error matrices were rescaled such that the sum of each  
14 row is equal to the estimated WSOM concentration, which is calculated as the product of the  
15 measured WSOC multiplied by the OM:OC<sub>i</sub> ratios determined from the offline-AMS PMF  
16 results.

### 17 3.1.2 Overview of retrieved factors and estimate of traffic exhaust OA (TEOA)

18 We used a 4-factor solution to represent the variability of the input data. The 4 separated OA  
19 factors included the following:

20 1/ a biomass burning OA (BBOA) factor highly correlated with levoglucosan originating from  
21 cellulose pyrolysis;

22 2/ a local OA (LOA) factor explaining a large fraction of N-containing fragments variability  
23 and contributing mostly in Vilnius during summer and spring;

24 3/ a background oxygenated-OA (B-OOA) factor showing relatively stable contributions at all  
25 seasons;

26 4/ a summer-OOA (S-OOA) factor showing increasing concentrations with the average daily  
27 temperature.

28 If the number of factors is decreased to 3, a mixed BBOA/B-OOA factor is retrieved, and  
29 significant structure appears in the residuals during winter (Fig. S2, S3, S4). Increasing the

1 number of factors to 5 and 6, leads to a splitting of OOA factors that cannot be interpreted in  
2 terms of specific aerosol sources/processes (Fig. S2, S3). The further separated OOA factor in  
3 the 5-factor solution possibly derived from the splitting of B-OOA; in fact the sum of the  
4 newly separated OOA and B-OOA in the 5-factor solution correlated well with the B-OOA  
5 time series from the 4-factor solution ( $R = 0.93$ ). Overall, a clear structure removal in the  
6 residual time-series was observed until a number of factors equal to 4 (Fig. S4, S5).

7 We also explored a 5-factor solution in which a hydrocarbon-like OA (HOA) profile from  
8 Mohr et al. (2012) was constrained to estimate the TEOA contribution. However, the water-  
9 soluble TEOA (WSTEOA) contribution to WSOM was estimated as  $0.2\%_{\text{avg}}$  (section 3.1.4),  
10 likely too small for PMF to resolve. We performed 100 PMF runs by randomly varying the  
11 HOA  $a$ -value. The obtained results showed a low TEOA correlation with hopanes ( $R_{\text{max}} =$   
12  $0.25$ ,  $R_{\text{min}} = -0.15$ ) with 45% of the PMF runs associated with negative Pearson correlation  
13 coefficients, supporting the hypothesis that this factor has too small contribution in the water  
14 extracts to be resolved. Therefore, we selected the 4-factor solution as our best representation  
15 of the data, while TEOA was instead estimated by a chemical mass balance (CMB) approach  
16 and not based on AMS mass spectral features.

17 TEOA concentrations are estimated using a CMB approach that assumes hopanes, present in  
18 lubricant oils engines, (Subramanian et al., 2006) to be unique tracers for traffic. However,  
19 hopanes can also be emitted upon combustion of different types of fossil fuel, in particular by  
20 coal combustion (Rutter et al., 2009), therefore the traffic contribution estimated here,  
21 although very small (as discussed in the result section) should be considered as an upper  
22 estimate. Still, the EC/hopanes ratio determined in this work ( $900 \pm 100$ ) is consistent with  
23 EC/hopanes for TE ( $1400 \pm 900$ : He et al., 2006; He et al., 2008; El Haddad et al., 2009; Fraser  
24 et al., 1998) and not with the coal EC/hopanes from literature profiles ( $300 \pm 200$ : Huang et al.,  
25 2014; supplementary information (SI)). To assess the traffic exhaust OC (TEOC) contribution  
26 we used the sum of the four most abundant hopanes (17a(H),21b(H)-norhopane,  
27 17a(H),21b(H)-hopane, 22S,17a(H),21b(H)-homohopane, and 22R,17a(H),21b(H)-  
28 homohopane (hopanes<sub>sum</sub>)). The TEOC contribution was estimated from the average  
29 hopanes<sub>sum</sub>/TEOC ratio ( $0.0012 \pm 0.0005$ ) from tunnel measurements reported by He et al.  
30 (2006), He et al. (2008), El Haddad et al. (2009), and Fraser et al. (1998), where the four  
31 aforementioned hopanes were also the most abundant. In order to rescale TEOC to the total  
32 TEOA concentration we assumed an OM:OC<sub>TEOA</sub> ratio of  $1.2 \pm 0.1$  (Aiken et al., 2008, Mohr

1 et al., 2012, Docherty et al., 2011, Setyan et al., 2012). The uncertainty of the estimated  
2 TEOA concentration was assessed by propagating the uncertainties relative to the  
3 OM:OC<sub>TEOA</sub> ratio (8.3%), the hopanes<sub>sum</sub>/TEOC ratio (41.7%), the hopane measurement  
4 repeatability (11.5%), and detection limits (7 pg m<sup>-3</sup>).

### 6 3.1.3. Source apportionment uncertainty

7 A common issue in PMF is the exploration of the rotational ambiguity, here addressed by  
8 performing 100 PMF runs initiated using different input matrices. We adopted a bootstrap  
9 approach (Davison and Hinkley, 1997) to generate the new input data and error matrices  
10 (Brown et al., 2015). Briefly, the bootstrap algorithm generates new input matrices by  
11 randomly resampling mass spectra from the original input matrices. As already mentioned,  
12 the input matrices contained ca. 12 mass spectral repetitions per filter sample; therefore the  
13 bootstrap approach was implemented in order to resample random filter sample mass spectra  
14 together with the corresponding measurement repetitions. Each newly generated PMF input  
15 matrix had a total number of samples equal to the original matrices (177 samples), although  
16 some of the original 177 filter samples are represented several times, while others are not  
17 represented at all. Overall we resampled on average 63±2% of the filter samples per bootstrap  
18 run. The generated data matrices were finally perturbed by varying each  $x_{i,j}$  element within  
19 twice the corresponding uncertainty ( $s_{i,j}$ ) assuming a normal distribution of the errors.  
20 Solutions were selected and retained according to three acceptance criteria based on PMF  
21 factor correlations with corresponding tracers: BBOA vs. levoglucosan, B-OOA vs. NH<sub>4</sub><sup>+</sup>,  
22 and S-OOA vs. average daily temperature. In order to discard suboptimal PMF runs, we only  
23 retained solutions associated with positive Pearson correlation coefficients for each criterion,  
24 for both the individual stations and the entire dataset. In total 95% of the solutions were  
25 retained following this approach. We note that no solution was discarded based on the first  
26 two criteria.

27 The offline-AMS PMF analysis provides the water-soluble contribution of the identified  
28 aerosol sources. In order to rescale the water-soluble organic carbon concentration of a  
29 generic factor  $z$  (WSZOC) to its total OC concentration (ZOC) we used the factor recoveries  
30 ( $R_z$ ) determined by Daellenbach et al. (2016) according to Eq. (5):

$$31 \quad ZOC_i = \frac{WSZOC_i}{R_z} \quad (5)$$

1 For each PMF factor (BBOA, W-OOA, and S-OOA), the water-soluble organic carbon  
 2 contribution was determined from the OM:OC ratio calculated from the (water-soluble) factor  
 3 mass spectrum (Aiken et al. 2008). For LOA, whose recovery was not previously reported,  
 4  $R_{LOA}$  was estimated from a single parameter fit according to Eq. (6)

$$5 \quad OC = TEOC + \frac{WS_{BBOA}}{(OM/OC)_{WS_{BBOA}} \cdot R_{BBOA}} + \frac{WS_{W-OOA}}{(OM/OC)_{WS_{W-OOA}} \cdot R_{OOA}} + \frac{WS_{S-OOA}}{(OM/OC)_{WS_{S-OOA}} \cdot R_{OOA}} + \frac{WS_{LOA}}{(OM/OC)_{LOA} \cdot R_{LOA}} \quad (6)$$

6 Here the water-soluble OA factor concentrations were converted to the corresponding water-  
 7 soluble OC concentrations to fit the measured OC. For each of the 95 retained PMF solutions,  
 8 Eq. (6) was fitted 100 times by randomly selecting a set of 100  $R_{BBOA}$ ,  $R_{OOA}$  value  
 9 combinations from those determined by Daellenbach et al. (2016). Each fit was initiated by  
 10 perturbing the input  $OC_i$  and  $TEOC_i$  within their uncertainties, assuming a normal distribution  
 11 of the errors. Additionally, in order to explore the effect of possible bulk extraction efficiency  
 12 (WSOC/OC) systematic measurement biases on our  $R_Z$  estimates, we also perturbed the OC,  
 13 WSOC (Daellenbach et al., 2016) inputs. Specifically, we assumed an estimated accuracy bias  
 14 of 5% for each of the perturbed parameters, which corresponds to the OC and WSOC  
 15 measurement accuracy. In a similar way, we also perturbed the input  $R_{BBOA}$  and  $R_{OOA}$   
 16 assuming an accuracy estimate of 5% deriving from a possible OC measurement bias in  
 17 Daellenbach et al. (2016) which could have affected the  $R_Z$  determination. In total  $9.5 \cdot 10^3$  fits  
 18 were performed (Eq. 6) and we retained only solutions (and corresponding perturbed  $R_Z$   
 19 combinations) associated with average OC residuals not statistically different from 0 within  
 20  $1\sigma$  for each station individually and for summer and winter individually ( $\sim 8\%$  of the  $9.5 \cdot 10^3$   
 21 fits, Fig. S6). The OC residuals of the accepted solutions did not manifest a clear correlation  
 22 with the LOA concentration (Fig. S7), indicating that the estimated  $R_{LOA}$  was properly fitted,  
 23 without compensating for unexplained variability of the PMF model or biases from the other  
 24  $R_z$ . Fig. S8 shows the probability density functions (PDF) of the retained perturbed  $R_z$  which  
 25 account for all uncertainties and biases mentioned above.  $R_{LOA,med}$  was estimated to be equal  
 26 to 0.66 (1<sup>st</sup> quartile 0.61, 3<sup>rd</sup> quartile 0.69, Fig. S8), while the retained  $R_{BBOA}$  and  $R_{OOA}$  values  
 27 ( $R_{BBOA,med}$  0.57, 1<sup>st</sup> quartile 0.55, 3<sup>rd</sup> quartile 0.60;  $R_{OOA,med}$  0.84, 1<sup>st</sup> quartile 0.81, 3<sup>rd</sup> quartile  
 28 0.88) were systematically lower than those reported by Daellenbach et al. (2016), reflecting  
 29 the lower bulk extraction efficiency (bulk EE = WSOC/OC) measured for this dataset  
 30 (median = 0.59, 1<sup>st</sup> quartile = 0.51, 3<sup>rd</sup> quartile = 0.72 vs. median = 0.74, 1<sup>st</sup> quartile = 0.66, 3<sup>rd</sup>  
 31 quartile 0.90 in Daellenbach et al. (2016)). All the retained  $R_k$  combinations are available at  
 32 DOI: doi.org/10.5905/ethz-1007-53.

1 Source apportionment uncertainties ( $\sigma_{S.A.}$ ) were estimated for each sample  $i$  and factor  $z$  as the  
2 standard deviation of all the retained PMF solutions ( $\sim 8\%$  of the  $9.5 \cdot 10^3$  fits). In addition to  
3 the rotational ambiguity of the PMF model (explored by the bootstrap technique) and  $R_Z$   
4 uncertainty, each PMF solution included on average 10 repetitions for each filter sample, and  
5 hence  $\sigma_{S.A.}$  accounted also for measurement repeatability. In this work, the statistical  
6 significance of a factor contribution is calculated based on  $\sigma_{S.A.,z,i}$  (Tables S2 and S3).

7 In general the recovery estimates reported in Daellenbach et al. (2016) represent the most  
8 accurate estimates available, being constrained to match the online-ACSM source  
9 apportionment results. The  $R_Z$  combinations reported by Daellenbach et al. (2016)  
10 demonstrated to positively apply to this dataset, enabling properly fitting the measured Bulk  
11 EE (WSOC/OC) with unbiased residuals and therefore providing a further confidence on their  
12 applicability (we note that in Eq. 6 we fitted OC as function of  $1/R_Z$  and  $WSOC_{Z,i}$ , therefore  
13  $R_Z$  fitted  $WSOC/OC = \text{Bulk EE}$ ). In general further  $R_Z$  determinations calculated comparing  
14 offline-AMS and online-AMS source apportionments would be desirable in order to provide  
15 more robust  $R_Z$  estimates. In absence of a-priori  $R_Z$  values for specific factors (e.g. for LOA in  
16 this study) we recommend constraining the  $R_Z$  combinations reported by Daellenbach et al.  
17 (2016) as a-priori information to fit the unknown recoveries, with the caveat that the  $R_Z$   
18 combinations reported by Daellenbach et al. (2016) were determined for filter samples  
19 extracted with water following a specific procedure; therefore we recommend adopting these  
20  $R_Z$  combinations for filter samples extracted in the same conditions. Nevertheless the  $R_Z$   
21 combinations reported by Daellenbach et al. (2016) should be tested also for filters water  
22 extracted in different conditions to verify whether they can properly fit the Bulk EE. In case  
23 the  $R_Z$  combinations reported by Daellenbach et al. (2016) would not apply for a specific  
24 location or extraction procedure (i.e. not enabling a proper fit of Bulk EE) we recommend a  
25  $R_Z$  redetermination by comparing the offline-AMS source apportionment results with well-  
26 established source apportionment techniques. In absence of data to perform a well-established  
27 source apportionment, we recommend to fit all the  $R_Z$  to match the bulk EE (i.e. fitting all the  
28 recoveries similarly as in Eq. 6 without constraining any a-priori  $R_Z$  value).

29 In general, the offline-AMS technique assesses less precisely the contribution of the lower  
30 water soluble factors. The higher uncertainty mostly stems from the larger PMF rotational  
31 ambiguity when separating a factor characterized by low concentration in the aqueous filter  
32 extracts. Nevertheless, the uncertainty is dataset dependent, as the separation of source

1 components with low water solubility can be improved in case of distinct time variability  
2 characterizing those sources in comparison with the other aerosol sources. The low aqueous  
3 concentration of scarcely water soluble sources in fact can be partially overcome by the large  
4 signal/noise characterizing the offline-AMS technique (170 on average for this dataset).

#### 6 3.1.4. Sensitivity of PMF to the un-apportioned TEOA fraction

7 Despite representing only a small fraction, the un-apportioned water-soluble TEOA  
8 (WSTEOA) contribution could in theory affect the apportionment of the other sources in the  
9 PMF model. To assess this, we performed a PMF sensitivity analysis by subtracting the  
10 estimated WSTEOA concentration from the input PMF data matrix, and by propagating the  
11 estimated WSTEOA uncertainty (section 3.1.2) in the input error matrices. To estimate the  
12 WSTEOA concentration we assumed  $R_{TEOA}$  of  $0.11 \pm 0.01$  (Daellenbach et al., 2016) and we  
13 used the HOA profile reported by Mohr et al. (2012) as surrogate for the TEOA mass spectral  
14 fingerprint. This approach is equivalent to constraining both the WSTEOA time series and  
15 factor profile. Overall the WSTEOA contribution to WSOM was estimated as  $0.2\%_{avg}$ ,  
16 making a successful retrieval of WSTEOA unlikely (Ulbrich et al., 2009). Consistently, PMF  
17 results obtained from this sensitivity analysis indicated that BBOA and B-OOA were robust,  
18 showing only 1% difference from the average offline-AMS source apportionment results,  
19 with BBOA increased and B-OOA decreased. S-OOA and LOA instead showed larger  
20 deviations from the average source apportionment results (S-OOA increased by 8% and LOA  
21 decreased by 15%), yet within our source apportionment uncertainties. These results highlight  
22 the marginal influence of the un-apportioned WSTEOA fraction on the other factors.

#### 24 3.2 Marker-PMF: measured $PM_1$ source apportionment

25 In the following section we describe the implementation of source apportionment using  
26 chemical markers (marker-PMF), as well as its optimization and uncertainty assessment. We  
27 discuss the number of factors and the selection of specific constraints to improve the source  
28 separation. Subsequently we discuss the source apportionment rotational uncertainty, and the  
29 sensitivity of our PMF results to the number of source specific markers, and to the assumed  
30 constraints.

### 1 3.2.1 Inputs

2 The marker-PMF yields a source apportionment of the entire measured PM<sub>1</sub> fraction (organic  
3 and inorganic). Measured PM<sub>1</sub> is defined here as the sum of EC, ions measured via IC, and  
4 OM estimated from OC measurements multiplied by the (OM:OC)<sub>*i*</sub> ratio determined from the  
5 offline-AMS PMF results by summing the factor profiles OM:OC ratios weighted by the time  
6 dependent factor relative contributions (rescaled by the recoveries). PMF was used to analyze  
7 a data matrix consisting of selected organic molecular markers, ions measured by IC, EC, and  
8 the remaining OM fraction (OM<sub>res</sub>) calculated as the difference between OM and the sum of  
9 the organic markers already included in the input matrix. OM<sub>res</sub> represented on average  
10 95±2% of total OM. The marker-PMF analysis is limited by the lack of elemental  
11 measurements (e.g. metals and other trace elements) typically used to identify mineral dust  
12 and certain anthropogenic sources. All markers showing concentrations above the detection  
13 limits for more than 25% of the samples were selected as input variables (72 in total). The  
14 PMF input matrices contain 67 composite samples (31 for Rūgšteliškis, 29 for Preila, and 7  
15 for Vilnius). The errors (*s*<sub>*i,j*</sub>) were estimated by propagating for each *j* variable the detection  
16 limits (DL) and the relative repeatability (*RR*) multiplied by the *x*<sub>*i,j*</sub> concentration according to  
17 Eq. (7) (Rocke and Lorenzato, 1995):

$$18 \quad s_{i,j} = \sqrt{(DL_j)^2 + (x_{i,j} \cdot RR_{i,j})^2} \quad (7)$$

### 19 3.2.2 Number of factors and constraints

20 We selected a 7-factor solution to explain the variability of the measured PM<sub>1</sub> components.  
21 The retrieved factors were biomass burning (BB), traffic exhaust (TE), primary biological  
22 organic aerosol (PBOA), SO<sub>4</sub><sup>2-</sup>-related secondary aerosol (SA), NO<sub>3</sub><sup>-</sup>-related SA, methane  
23 sulfonic acid (MSA)-related SA, and a Na<sup>+</sup>-rich factor explaining the variability of inorganic  
24 components typically related to resuspension of mineral dust, sea salt, and road salt.

25 We first tested an unconstrained source apportionment. This led to a suboptimal separation of  
26 the aerosol sources, with large mixings of PMF factors associated with contributions of  
27 markers originating from different sources. In particular we observed mixing of BB markers  
28 (e.g. levoglucosan) with fossil fuel combustion markers such as hopanes, as well as with  
29 inorganic ions such as NO<sub>3</sub><sup>-</sup> and Ca<sup>2+</sup>. All these markers, although related to different  
30 emission/formation processes, are characterized by similar seasonal trends, i.e. higher



1 concentrations during winter than in summer. Specifically, the BB tracers increase during  
2 winter because of domestic heating activity, hopanes presumably because of the accumulation  
3 in a shallower boundary layer and lower photochemical degradation,  $\text{NO}_3^-$  because of the  
4 partitioning into the particle phase at low temperatures, and  $\text{Ca}^{2+}$  because winter was the  
5 windiest season and therefore was associated with the most intense resuspension.

6 We subsequently exploited the markers' source-specificity to set constraints for the profiles  
7 output by our model: for each individual source, we treated the contribution of the unrelated  
8 source-specific markers as negligible (e.g. we assumed that TE, SA, Na-rich factor and PBOA  
9 do not contribute to levoglucosan). In contrast, the non-source specific variables (EC,  $\text{OM}_{\text{res}}$ ,  
10 (Me-)PAHs, S-PAHs, inorganic ions, oxalate, alkanes) were freely apportioned by the PMF  
11 algorithm. In a similar way we set constraints for primary markers (e.g.  $\text{K}^+$  and  $\text{Ca}^{2+}$ ) and  
12 combustion related markers (e.g. PAHs), which are not source-specific but the contribution of  
13 which can be considered as negligible in the SA factors. In this case the algorithm can freely  
14 apportion these markers to all the primary factors and combustion-related factors,  
15 respectively.

16 In details, EC, PAHs, and methyl-PAHs were constrained to zero in non-combustion sources,  
17 i.e. all profiles but TE and BB. While EC could partially derive from dust resuspension,  
18 literature profiles for this source suggest an EC contribution below 1% (Chow et al., 2003).  
19 This is expected to be also the case here given the distance of the three stations from  
20 residential areas and busy roads. Methoxyphenols and sugar anhydrides, considered to be  
21 unique BB markers, were constrained to zero in all sources but BB. Similarly, hopanes were  
22 constrained to zero in all factors but TE. We also assumed no contribution from glucose,  
23 arabitol, mannitol, and sorbitol to all secondary factors, and traffic exhaust. The  $\text{SO}_4^{2-}$   
24 contribution from primary traffic emissions was estimated to be negligible, given the use of  
25 desulfurized fuel for vehicles in Lithuania. Likewise, alkane contributions were assumed to be  
26 zero in the SA factors, similar to the contribution of  $\text{Ca}^{2+}$ ,  $\text{Na}^+$ ,  $\text{K}^+$  and  $\text{Mg}^{2+}$  in the SA factors  
27 and TE.

28 The number of factors was increased until no mixing between source-specific markers for  
29 different aerosol sources/processes was observed any more. Secondary sources instead were  
30 explained by three factors because of the distinct seasonal and site-to-site variability of MSA,  
31  $\text{NO}_3^-$  and  $\text{SO}_4^{2-}$ . Oxalate correlated well with  $\text{NH}_4^+$  ( $R=0.62$ ) and the latter well with the sum  
32 of  $\text{SO}_4^{2-}$  and  $\text{NO}_3^-$  equivalents ( $R=0.98$ ). Note that the aforementioned secondary tracers were

1 not constrained in any factor with the exception of  $\text{SO}_4^{2-}$  contributions which were assumed to  
2 be negligible in the TE factor. Moreover the 7-factor solution showed unbiased residuals  
3 (residual distribution centered at 0 within  $1\sigma$ ) for all the stations together and for each station  
4 individually, while lower order solutions showed biased residuals for at least one station or all  
5 the stations together.

6 PMF results obtained assuming only the aforementioned constraints returned suboptimal  
7 apportionments of  $\text{OM}_{\text{res}}$  and  $\text{Na}^+$  between the BB and the  $\text{Na}^+$ -rich factor, with unusually  
8 high  $\text{OM}_{\text{res}}$  fractional contributions in the  $\text{Na}^+$ -rich factor and unusually high  $\text{Na}^+$   
9 contributions in the BB profile in comparison with literature profiles (Chow et al., 2003;  
10 Huang et al., 2014 and references therein; Schauer et al., 2001). Similarly the  $\text{EC}/\text{OM}_{\text{res}}$  value  
11 for TE was substantially lower than literature profiles (El Haddad et al., 2013 and references  
12 therein). Other constraints were therefore introduced to improve the separation of these three  
13 variables. Specifically, EC and  $\text{OM}_{\text{res}}$  were constrained in the traffic profile to be equal to  
14 0.45 and 0.27 ( $a$ -value = 0.5) according to El Haddad et al. (2013), while EC:BB ratio was  
15 constrained to 0.1 ( $a$ -value = 1) according to Huang et al. (2014) and references therein.  $\text{Na}^+$   
16 was constrained to 0.2% ( $a$ -value = 1) in BB according to Schauer et al. (2001), while  $\text{OM}_{\text{res}}$   
17 was constrained to zero in the  $\text{Na}^+$ -rich factor to avoid mixing with BB. Although this  
18 represents a strict constraint, we preferred avoiding constraining  $\text{OM}_{\text{res}}$  to a specific value for  
19 the  $\text{Na}^+$ -rich factor which could not be linked to a unique source but possibly represents  
20 different resuspension-related sources (e.g. sea salt, mineral dust and road dust). However, we  
21 expect none of the aforementioned sources to explain a large fraction of the submicron  $\text{OM}_{\text{res}}$   
22 (the OC:dust ratio for dust profiles is 1-15% according to Chow et al., 2003). The sensitivity  
23 of our source apportionment to the constraints listed in this section is discussed in the next  
24 section.

25

### 26 3.2.3. Source apportionment uncertainty and sensitivity analyses

27 We explored the model rotational uncertainty by performing 20 bootstrap PMF runs, and by  
28 perturbing each input  $x_{i,j}$  element within  $2 \cdot s_{i,j}$  assuming a normal distribution of the errors.  
29 Results and uncertainties of the PMF model reported in this paper represent the average and  
30 the standard deviation of the bootstrap runs.

1 As discussed in section 3.2.2, we assumed the contribution of specific markers to be 0 in  
2 different factor profiles. Such assumptions preclude the PMF model to vary the contributions  
3 of these variables from 0 (Eq. 3). In order to explore the effect of such assumptions on our  
4 PMF results we loosened all these constraints assuming variable contributions equal to 50%,  
5 37.5%, 25%, and 12.5% of their average relative contribution to measured PM<sub>1</sub>. In all cases  
6 the  $a$ -value was set to 1. The average factor concentrations for the 12.5% case and the fully  
7 constrained average bootstrap PMF solutions were not statistically different (confidence  
8 interval of 95%, Fig. S9). Statistically significant differences arose for the of the SO<sub>4</sub><sup>2-</sup>-related  
9 SA in the 50% and 37.5% cases, and the Na<sup>+</sup>-rich factor in the 25% and 37.5% cases,  
10 indicating that loosening the constraints allowed additional rotational uncertainty in  
11 comparison to the uncertainty explored by the bootstrap approach. By contrast, the factors  
12 associated with large relative uncertainties from the marker source apportionment (TE and  
13 PBOA, Table S3) showed the best agreement in terms of concentrations (Fig. S9) with the  
14 fully constrained solution, suggesting that the variability introduced by loosening the  
15 constraints did not exceed that already accounted for by the bootstrap approach. As previously  
16 mentioned, the largest contribution discrepancies were observed for the SO<sub>4</sub><sup>2-</sup>-related SA and  
17 Na<sup>+</sup>-rich factor. Looser constraints increased the explained variability of primary components  
18 such as EC, arabinol, sorbitol, K<sup>+</sup>, Mg<sup>2+</sup>, and Ca<sup>2+</sup> by the (secondary) SO<sub>4</sub><sup>2-</sup>-related SA factor.  
19 The Na<sup>+</sup>-rich factor showed increasing contributions from OM<sub>res</sub> and from BB components  
20 such as methoxyphenols, and anhydrous sugars, which exhibited similar seasonal trends as the  
21 Na<sup>+</sup>-rich factor. None of the marker-PMF factors showed statistically different average  
22 contributions (confidence interval of 95%) when tolerating a variability of the constrained  
23 variables within 12.5% of their relative contribution to PM<sub>1</sub>. Note that with this degree of  
24 tolerance the contribution of OM to the Na<sup>+</sup>-rich was 28%, which is unrealistically high  
25 compared to typically reported values for OM:dust ratios (<15% Chow et al., 2003).  
26 Therefore, we consider the fully constrained PMF solution to represent best the average  
27 composition of the contributing sources.

28 The marker-PMF source apportionment depends strongly on the input variables (i.e. measured  
29 markers), as these are assumed to be highly source specific. That is, minor sources, such as  
30 MSA-related SA and PBOA, are separated because source-specific markers were used as  
31 model inputs. Meanwhile, more variables were used as tracers for TE and BB  
32 (methoxyphenols (5 variables), sugar anhydrides (3 variables), and hopanes (5 variables)),  
33 which gives more weight to these specific sources. We explored the sensitivity of the PMF

1 results to the number and the choice of traffic and wood burning markers, by replacing them  
2 with randomly selected input variables. In total 20 runs were performed and the average  
3 contribution of the different sources to  $OM_{res}$  was compared with the marker source  
4 apportionment average results, where bootstrap was applied to resample time points. Results  
5 displayed in Fig. S10 are in agreement the apportionment of  $OM_{res}$  from BB within  $11\%_{avg}$ ,  
6 highlighting its robustness. The agreement for TE was lower, which is not surprising given  
7 the lower contribution of this source and the smaller number of specific markers (hopanes).  
8 However, these uncertainties were within the marker source apportionment uncertainty (Fig.  
9 S10), implying that the results were not significantly sensitive to the number and the choice of  
10 input markers for BB and traffic exhaust.

11

## 12 **4 Results and Discussion**

### 13 **4.1 $PM_{10}$ composition**

14 An overview of the measured  $PM_{10}$  composition can be found in Fig. 1. Measured  $PM_{10}$   
15 average concentrations were in general low, with lower values detected at the rural terrestrial  
16 site of Rūgšteliškis ( $5.4 \mu\text{g m}^{-3}_{avg}$ ) than in Vilnius ( $6.7 \mu\text{g m}^{-3}_{avg}$ ) and Preila ( $7.0 \mu\text{g m}^{-3}_{avg}$ ).  
17 OM represented the major fraction of measured  $PM_{10}$  for all seasons and stations, with  $57\%_{avg}$   
18 of the mass. The average OM concentrations were higher during winter ( $4.2 \mu\text{g m}^{-3}$ ) than in  
19 summer ( $3.0 \mu\text{g m}^{-3}$ ) at all sites probably to a combination of domestic wood burning activity  
20 and accumulation of the emissions in a shallower boundary layer. For similar reasons, EC  
21 average concentrations showed higher values during winter ( $0.42 \mu\text{g m}^{-3}$ ) than in summer  
22 ( $0.25 \mu\text{g m}^{-3}$ ). During summer, the average EC concentration was  $\sim 5$  times higher in Vilnius  
23 ( $0.54 \mu\text{g m}^{-3}$ ) than in Preila and Rūgšteliškis ( $0.12$  and  $0.11 \mu\text{g m}^{-3}$ , respectively), indicating  
24 an enhanced contribution from combustion emissions. In the absence of domestic heating  
25 during this period, a great part of these emissions may be related to traffic. During winter, EC  
26 concentrations were comparable at all sites (only 25% higher in Vilnius than in Preila and  
27 Rūgšteliškis). This suggests that a great share of wintertime EC may be related to BB, the  
28 average contribution of which is significant at all stations within  $3\sigma$  (table S2). It should be  
29 noted that the highest measured  $PM_{10}$  concentrations were detected at the remote rural coastal  
30 site of Preila during three different pollution episodes. In particular, the early March episode  
31 corresponded to the period analyzed by Ulevicius et al. (2016) and Dudoitis et al. (2016), and

1 was attributed to regional transport of polluted air masses associated to an intense land  
2 clearing activity characterized by large scale grass burning in the neighboring Kaliningrad  
3 region.  $\text{SO}_4^{2-}$  represented the second major component of measured  $\text{PM}_{10}$  (20%<sub>med</sub>) at all sites  
4 and seasons. Its average concentration remained rather constant with only slightly higher  
5 concentrations in summer than in winter ( $1.2 \pm 0.7 \mu\text{g m}^{-3}$ , and  $1.1 \pm 0.6 \mu\text{g m}^{-3}$  respectively).  
6 Overall  $\text{SO}_4^{2-}$  concentrations did not show large differences from site-to-site, suggestive of  
7 regional sources. By contrast  $\text{NO}_3^-$  showed a clear seasonality with larger contributions in  
8 winter (average  $0.9 \pm 0.8 \mu\text{g m}^{-3}$  equivalent to 12% of measured  $\text{PM}_{10}$ ) than in summer  
9 ( $0.03 \pm 0.03 \mu\text{g m}^{-3}$ ), as expected from its semi-volatile nature.

#### 10 **4.2 OM source apportionment (Offline-AMS PMF)**

11 The apportioned PMF factors were associated to aerosol sources/processes according to their  
12 mass spectral features, seasonal contributions and correlations with tracers. The four  
13 identified factors were BBOA, LOA, B-OOA, and S-OOA, which are thoroughly discussed  
14 below. The TEOA contributions instead were determined using a CMB approach.

15 BBOA was identified by its mass spectral features, with high contributions of  $\text{C}_2\text{H}_4\text{O}_2^+$ , and  
16  $\text{C}_3\text{H}_5\text{O}_2^+$  (Fig. 2), typically associated with levoglucosan fragmentation from cellulose  
17 pyrolysis (Alfarra et al., 2007), accordingly the BBOA factor time series correlated well with  
18 levoglucosan (Pearson correlation coefficient:  $R=0.90$ , Fig. S11). BBOA contributions were  
19 higher during winter and lower during summer (Fig. 3a). We determined the biomass burning  
20 organic carbon (BBOC) concentration from the BBOA time series divided by the  
21  $\text{OM}:\text{OC}_{\text{BBOA}}$  ratio determined from the corresponding HR spectrum. The winter  
22 levoglucosan/BBOC ratio was  $0.16_{\text{med}}$ , consistent with values reported in continental Europe  
23 for ambient BBOC profiles (levoglucosan/BBOC range: 0.10-0.21, Zotter et al., 2014;  
24 Minguillón et al., 2011; Herich et al., 2014).

25 The second factor was defined as LOA because of its statistically significant contribution  
26 (within  $3\sigma$ ) only in Vilnius during summer (table S2), in contrast to other potentially local  
27 primary (e.g. BBOA) and secondary (S-OOA) sources which contributed at all sites. The  
28 LOA mass spectrum was characterized by a high contribution of N-containing fragments  
29 (especially  $\text{C}_5\text{H}_{12}\text{N}^+$ , and  $\text{C}_3\text{H}_8\text{N}^+$ ), with the highest N:C ratio (0.049) among the apportioned  
30 PMF factors (0.029 for BBOA, 0.013 for S-OOA, 0.023 for B-OOA). A similar factor was  
31 also observed by Byčenkienė et al. (2016) using an ACSM at the same station. In that work,

1 high LOA concentrations were associated with wind directions from N-NW, and the authors  
2 suggested the sludge utilization system of Vilnius (UAB Vilniausvandenys) situated 3.9 km  
3 NW from the sampling station as a probable source.

4 Two different OOA sources (S-OOA and B-OOA) were resolved and exhibited different  
5 seasonal trends. Separation and classification of OOA sources from offline-AMS is typically  
6 different from that of online AMS and ACSM measurements, mainly due to the different time  
7 resolution. Few online-AMS studies reported the separation of isoprene-related OA factor  
8 (Budisulistiorini et al., 2013; Hu et al., 2015, Xu et al., 2015) mostly driven by isoprene  
9 epoxides chemistry. Xu et al. (2015) showed that nighttime monoterpene oxidation by nitrate  
10 radical contributes to less-oxidized OOA. However, the large majority of online-AMS OOA  
11 factors are commonly classified based on their volatility (semi-volatile OOA and low-  
12 volatility OOA) rather than on their sources and formation mechanisms.

13 This differentiation is typically achieved only for summer datasets when the temperature  
14 gradient between day and night is sufficiently high, yielding a detectable daily partitioning  
15 cycle of the semi-volatile organic compounds and  $\text{NO}_3^-$  between the gas and the particle  
16 phases. Online AMS datasets have higher time resolution than filter sampling, but sampling  
17 periods typically cover only a few weeks. Therefore the apportionment is driven by daily  
18 variability rather than seasonal differences. By contrast, in the offline-AMS source  
19 apportionment, given the 24-h time resolution of the filter sampling and the yearly cycle time  
20 coverage, the separation of the factors is driven by the seasonal variability of the sources and  
21 by the site-to-site differences. Therefore, the offline-AMS source apportionment separates  
22 factors by seasonal trends rather than volatility.

23 The resolved B-OOA factor explained a higher fraction than S-OOA. It was associated with  
24 background oxygenated aerosols as no systematic seasonal pattern was observed. However,  
25 B-OOA correlated well with  $\text{NH}_4^+$  ( $R=0.69$ , Fig. S11), and had the highest OM:OC ratio  
26 among the apportioned PMF factors (2.21).

27 Unlike B-OOA, S-OOA showed a clear seasonality with higher contributions during summer,  
28 increasing exponentially with the average daily temperature (Fig. S12a). During summer the  
29 site-to-site S-OOA concentrations were not statistically different within a confidence interval  
30 of 95%, while during winter the site-to-site agreement was lower, possibly due to the larger  
31 model uncertainty associated with the low S-OOA concentrations. A similar S-OOA vs.  
32 temperature relationship was reported by Leitch et al. (2011) for a terpene dominated

1 Canadian forest using an ACSM and by Daellenbach et al. (2016) and Bozzetti et al. (2016)  
2 for the case of Switzerland (Fig. S12b), using a similar source apportionment model. This  
3 increase in S-OOA concentration with temperature is consistent with the exponential increase  
4 in biogenic SOA precursors (Guenther et al., 2006). Therefore, even though the behavior of S-  
5 OOA at different sites might be driven by several parameters, including vegetation coverage,  
6 available OA mass, air masses photochemical age and ambient oxidation conditions (e.g. NO<sub>x</sub>  
7 concentration), temperature seems to be the main driver of S-OOA concentrations. Overall  
8 more field observations at other European locations are needed to validate this relation. While  
9 the results indicate a probable secondary biogenic origin of the S-OOA factor, the precursors  
10 of the B-OOA factor are not identified. In section 4.4.2 more insights into the OOA sources  
11 will be discussed.

12 The S-OOA profile showed a CO<sub>2</sub><sup>+</sup>/C<sub>2</sub>H<sub>3</sub>O<sup>+</sup> ratio of 0.61<sub>avg</sub>, placing it in the region of semi-  
13 volatile SOA from biogenic emissions in the *f*<sub>44</sub>/*f*<sub>43</sub> space (Ng et al., 2011), as attributed by  
14 Canonaco et al. (2015). Despite the higher summer photochemical activity, the water-soluble  
15 bulk OA showed more oxidized mass spectral fingerprints during winter (O:C=0.61<sub>avg</sub>) than  
16 in summer (O:C=0.55<sub>avg</sub>), similar to the results presented by Canonaco et al. (2015) for  
17 Zurich. Accordingly, the S-OOA profile also showed a less oxidized water-soluble mass  
18 spectral fingerprint than B-OOA, with an O:C ratio of 0.40<sub>avg</sub>, in comparison with 0.80<sub>avg</sub> for  
19 B-OOA. Considering the sum of B-OOA and S-OOA, the median OOA:NH<sub>4</sub><sup>+</sup> ratios for  
20 Rūgšteliškis, Preila, and Vilnius were 3.2, 2.4, and 2.5 respectively, higher than the average  
21 but within the range of the values reported by Crippa et al. (2014) for 25 different European  
22 rural sites (2.0<sub>avg</sub>; minimum value 0.3; maximum 7.3).

23

#### 24 **4.3 PM<sub>1</sub> source apportionment (marker-PMF)**

25 The PMF factors in this analysis were associated with specific aerosol sources/processes  
26 according to their profiles, seasonal trends and relative contributions to the key variables. Fig.  
27 4 displays factor profiles, and the relative contribution of each factor to each variable. The  
28 Na<sup>+</sup>-rich factor explained a large part of the variability of Ca<sup>2+</sup>, Mg<sup>2+</sup>, and Na<sup>+</sup> (Fig. 4) and  
29 showed higher contributions during winter than in summer (Fig. 5), suggesting a possible  
30 resuspension of sand and salt typically used during winter in Lithuania for road de-icing. This  
31 seasonal trend is also consistent with wind speed, which showed the highest monthly values  
32 during December 2013 and January 2014. We cannot exclude the possibility that this factor

1 may include contributions from sea salt, although  $\text{Na}^+$  and  $\text{Cl}^-$  were not enhanced at the  
2 marine station in comparison with the other stations. The overall contribution of this  $\text{Na}^+$ -rich  
3 factor to measured  $\text{PM}_{10}$  was relatively small ( $1\%_{\text{avg}}$ ), but may be larger in the coarse fraction.

4 The BB factor showed a well-defined seasonality, with high contributions during winter. This  
5 factor explained a large part of the variability of typical wood combustion tracers such as  
6 methoxyphenols, sugar anhydrides (including levoglucosan, mannosan, and galactosan),  $\text{K}^+$ ,  
7  $\text{Cl}^-$ , EC, PAHs, and methyl-PAHs (Fig. 4). Using the  $\text{OM}:\text{OC}_{\text{BBOA}}$  ratio (1.88) calculated  
8 from offline-AMS, we estimated the levoglucosan:BBOC ratio to be  $0.18_{\text{avg}}$ , which is within  
9 the range of previous studies (Ulevicius et al., 2016 and references therein). Note that this  
10 factor explained also large fractions of variables typically associated with non-vehicular fossil  
11 fuel combustion, such as benzo(b)naphtho(2,1-d)thiophene (BNT[2,1]) and 6,10,14-trimethyl-  
12 2-pentadecanone (DMPT, Fig. 4, Manish et al., 2007; Subramanian et al., 2007), indicating a  
13 potential mixing of BB with fossil fuel combustion sources. However, the fossil fuel  
14 combustion contribution to BB is unlikely to be large, considering the low concentrations of  
15 fossil fuel tracers such as hopanes (66% of the samples below quantification limit ( $<\text{QL}$ )),  
16 BNT[2,1] (64% $<\text{QL}$ ), and DMPT (55% $<\text{QL}$ ). Moreover, the above mentioned agreement of  
17 the levoglucosan:BBOC ratio with previous studies corroborates the BB estimate from the  
18 marker-PMF.

19 The traffic exhaust factor explained a significant fraction of the alkane variability, with a  
20 preferential contribution from light alkanes (Fig. 4). Its contribution was never statistically  
21 significant within  $3\sigma$ . However on average the concentration was higher in Vilnius than at the  
22 other stations and in general higher in winter than in summer.

23 The PBOA factor explained the variability of the primary biological components, such as  
24 glucose, mannitol, sorbitol, arabitol, and alkanes with an odd number of carbon atoms  
25 (consistent with Bozzetti et al., 2016 and references therein). Highest PBOA concentrations  
26 were observed during spring, especially at the rural site of Rūgšteliškis. Overall the  
27 contribution of this factor was uncertain with an average relative model error of 160%  
28 probably due to the small PBOA contributions ( $0.6\%_{\text{avg}}$  of the total OM), which hampers a  
29 more precise determination by the model. In particular  $\text{OM}_{\text{res}}$  was the variable showing the  
30 highest mass contribution to the PBOA factor. However, the large contribution and the large  
31 uncertainty of  $\text{OM}_{\text{res}}$  to this factor ( $0.3\pm 0.4$ ) resulted in a large uncertainty in the PBOA  
32 estimated concentration.



1 The last three factors were related to SA, as indicated by the large contributions of secondary  
2 species such as oxalate,  $\text{SO}_4^{2-}$ , MSA, and  $\text{NO}_3^-$  to the factor profiles (Fig. 4). The three factors  
3 showed different spatial and temporal contributions.

4 The  $\text{NO}_3^-$ -related SA exhibited highest contributions during winter, suggesting temperature-  
5 driven partitioning of secondary aerosol components. Moreover the  $\text{NO}_3^-$ -related SA,  
6 similarly to BB and TE, showed the highest concentrations in Vilnius, and the lowest in  
7 Rūgštelėškis suggesting its possible relation with anthropogenic gaseous precursors (e.g.  
8  $\text{NO}_x$ ), as already reported in other studies (e.g. Xu et al., 2016; McMeeking et al., 2012).

9 The MSA-related SA factor manifested the highest concentrations at the marine site of Preila  
10 during summer, and in general larger contributions during summer than winter, suggesting its  
11 relation with marine secondary aerosol. MSA has been reported to be related to marine  
12 secondary biogenic emissions deriving from the photo-oxidation of dimethyl sulfide (DMS)  
13 emitted by the phytoplankton bloom occurring during the warm season (Li et al., 1993,  
14 Crippa et al., 2013 and references therein).

15 The last factor ( $\text{SO}_4^{2-}$ -related SA) showed higher contributions during summer than in winter  
16 without clear site-to-site variability, following the seasonal behavior of  $\text{SO}_4^{2-}$  showing slightly  
17 higher concentrations during summer than in winter, which is probably driven by the  
18 secondary formation from gaseous photochemical reactions and aqueous phase oxidation.  
19 This factor explained the largest part of the oxalate and  $\text{SO}_4^{2-}$  variability and represented  
20 48%<sub>avg</sub> of the measured  $\text{PM}_1$  by mass.

#### 21 **4.4 Comparison of the source apportionment methods**

22 In this section we compare the offline-AMS PMF and marker-PMF results. We begin with  
23 BBOA and TEOA emissions which were resolved by marker-PMF and offline-AMS (TEOA  
24 was not resolved by offline-AMS but determined through a CMB approach). The remaining  
25 OM fraction (Other-OA = OA – BBOA - TEOA) was apportioned by the offline-AMS source  
26 apportionment to B-OOA, S-OOA and LOA (Other-OA<sub>offline-AMS</sub>). However, the LOA  
27 contribution was statistically significant (within  $3\sigma$ ) only in Vilnius during summer (Table  
28 S2), while no data were available for these periods from the marker source apportionment.  
29 The marker source apportionment instead attributed the Other-OA mass fraction to 4 factors  
30 (Other-OA<sub>marker</sub>): PBOA, as well as to  $\text{SO}_4^{2-}$ ,  $\text{NO}_3^-$ , and MSA-related secondary organic  
31 aerosols (SOA, Fig. S13). The OA concentrations of the factors retrieved from the  $\text{PM}_1$

1 markers source apportionment were obtained by multiplying the factor time series by the sum  
2 of the organic markers and  $OM_{res}$  contributions to the normalized factor profiles. The PM  
3 concentrations from the marker PMF factors are displayed in Fig. 5.

#### 4 4.4.1 Primary OA sources

5 Offline-AMS and marker source apportionments provided comparable BBOA estimates, with  
6 concentrations agreeing within a 95% confidence interval (Fig. 6). Results revealed that  
7 BBOA contributed the largest fraction to the total OM during winter in Preila and Vilnius,  
8 while in Rūgšteliškis the largest OA source derived from B-OOA. The average winter BBOA  
9 concentration was  $1.1 \pm 0.8 \mu\text{g m}^{-3}$  in Rūgšteliškis and  $2 \pm 1 \mu\text{g m}^{-3}$  in Vilnius (errors in this  
10 section represent the standard deviation of the temporal variability). Overall the average  
11 BBOA concentrations were higher at the urban background site of Vilnius and lower at the  
12 rural terrestrial site of Rūgšteliškis. Preila showed higher values ( $3 \pm 3 \mu\text{g m}^{-3}$ ) driven by the  
13 grass burning episode occurred at the beginning of March (Ulevicius et al., 2016). Excluding  
14 this episode, the BBOA winter concentration was lower than in Vilnius ( $1.8 \mu\text{g m}^{-3}$ ). During  
15 winter, considering only the samples collected concomitantly, Preila and Vilnius showed well  
16 correlated BBOA time series ( $R = 0.91$ ) and significantly positive correlations were observed  
17 for also for Preila and Rūgšteliškis ( $R = 0.72$ ) and for Vilnius and Rūgšteliškis ( $R = 0.66$ )  
18 (offline-AMS BBOA time series). These results highlight the effect of regional  
19 meteorological conditions on the BBOA daily variability in the south east Baltic region.

20

21 By contrast, during summer BBOA concentrations were much lower, with 40% of the points  
22 showing statistically not significant contributions within  $3\sigma$  for the offline-AMS source  
23 apportionment and 100% for the marker source apportionment. Between late autumn and  
24 early March the offline-AMS source apportionment revealed three simultaneous episodes  
25 with high BBOA concentrations at the three stations, while the marker source apportionment  
26 which is characterized by lower time resolution did not capture some of these episodes. The  
27 first episode occurred between 19 and 25 December 2013 during a cold period with an  
28 average daily temperature drop to  $-9.7 \text{ }^\circ\text{C}$  as measured at the Rūgšteliškis station (no  
29 temperature data were available for the other stations). The third episode occurred between 5  
30 and 10 March 2014 and was associated with an intense grass burning episode localized mostly  
31 in the Kaliningrad region (Ulevicius et al., 2016, Dudoitis et al., 2016, Mordas et al., 2016).

1 The episode was not associated with a clear temperature drop, with the highest concentration  
2 ( $14 \mu\text{g m}^{-3}$ ) found at Preila on 10 March 2014, the closest station to the Kaliningrad region.  
3 Similarly, at the beginning of February high BBOA concentrations were registered at the  
4 three stations, without a clear temperature decrease. Other intense BBOA events were  
5 detected but only on a local scale, with intensities comparable to the regional scale episodes.  
6 Using the  $\text{OM:OC}_{\text{BBOA}}$  ratio calculated from the HR water-soluble BBOA spectrum (1.88),  
7 we estimated the  $\text{BBOC}_{\text{avg}}$  concentrations during the grass burning episode (5-10 March  
8 2014) to span between  $0.8$  and  $7.2 \mu\text{g m}^{-3}$ . On a daily basis our BBOC concentrations are  
9 consistent with the estimated ranges reported by Ulevicius et al. (2016) for non-fossil primary  
10 organic carbon ( $0.6$ - $6.9 \mu\text{g m}^{-3}$  during the period under consideration), showing also a high  
11 correlation ( $R=0.98$ ).

12 TEOA estimates obtained by CMB and marker-PMF always agreed with each other within  $3\sigma$   
13 (Fig. 6). The two approaches confirm that TEOA is a minor source at all three stations (Fig.  
14 6). Hopanes concentrations (used in this work as TEOA tracers) were below detection limits  
15 ( $7 \text{ pg m}^{-3}$ ) for 66% of the collected samples. Similarly to  $\text{NO}_x$ , hopanes, showed a clear spatial  
16 and seasonal variability with higher concentrations in Vilnius during winter, suggesting an  
17 accumulation of traffic emissions in a shallower boundary layer (Fig. 3b,  $\text{NO}_x$  data available  
18 only for Vilnius). During the grass burning event, we observed a peak in the total hopane  
19 concentration, and therefore also a peak of the estimated TEOA ( $2.4 \mu\text{g m}^{-3}$  maximum value).  
20 This relatively high concentration is most probably not due to a local increase of TE, but  
21 rather due to a regional transport of polluted air masses from neighboring countries (Poland  
22 and the Russian Kaliningrad enclave). By assuming an  $\text{OM:OC}_{\text{TEOA}}$  ratio of  $1.2 \pm 0.1$  (Aiken et  
23 al., 2008, Mohr et al., 2008, Docherty et al., 2011, Setyan et al., 2012), we determined the  
24 corresponding organic carbon content (TEOC). Our TEOC concentration was consistent  
25 within  $3\sigma$  with the average fossil primary OC over the whole episode estimated by Ulevicius  
26 et al. (2016), ( $0.4$ - $2.1 \mu\text{g m}^{-3}$ ) although on a daily basis the agreement was relatively poor.

27 Overall, offline-AMS source apportionment and marker-PMF returned comparable results for  
28 BBOA and similarly the TEOA estimate by markers-PMF and CMB were comparable,  
29 therefore not surprisingly the offline-AMS and marker-PMF approaches yielded OA  
30 concentrations also for the Other-OA fractions which agreed within  $3\sigma$ .

#### 1 4.4.2 Other-OA sources: offline-AMS and marker-source apportionment 2 comparison

3 The marker-source apportionment, in comparison to the offline-AMS source apportionment  
4 enables resolving well-correlated sources (e.g. BBOA and  $\text{NO}_3^-$ -related SOA) as well as  
5 minor sources (e.g. MSA-related SOA and PBOA) because source-specific markers were  
6 used as model inputs. By contrast, the offline-AMS source apportionment is capable of  
7 resolving OA sources for which no specific markers were available such as LOA, which was  
8 separated due to the distinct spatial and temporal trends of some N-containing AMS  
9 fragments. We first briefly summarize the Other-OA factor concentrations and their site-to-  
10 site differences retrieved by the two techniques; subsequently we compare the two source  
11 apportionment results.

12 The Other-OA<sub>offline-AMS</sub> factor time series are displayed in Fig. S13. The B-OOA factor  
13 showed relatively stable concentrations throughout the year with  $0.9 \pm 0.8_{\text{avg}} \mu\text{g m}^{-3}$  during  
14 summer and  $1.1 \pm 0.9_{\text{avg}} \mu\text{g m}^{-3}$  during winter. Although B-OOA concentrations were relatively  
15 stable throughout the year, higher contributions were observed in Preila and Rūgšteliškis  
16 compared to Vilnius. The extreme average seasonal concentrations were between 0.8 and 1.3  
17  $\mu\text{g m}^{-3}$  at Rūgšteliškis during fall and winter, between 0.9 and 1.1  $\mu\text{g m}^{-3}$  at Preila during  
18 spring and winter, and between 0.4 and 0.6  $\mu\text{g m}^{-3}$  in Vilnius during summer and winter.  
19 These values do not evidence clear seasonal trends, but highlight a site-to-site variability  
20 which will be further discussed in the following. S-OOA instead was the largest contributor to  
21 total OM during summer with an average concentration of  $1.2 \pm 0.8 \mu\text{g m}^{-3}$ , always agreeing  
22 between sites within a confidence interval of 95% (2 tails t-test). By contrast, during winter  
23 the S-OOA concentration dropped to an average value of  $0.3 \pm 0.2 \mu\text{g m}^{-3}$ , with 81% of the  
24 points not statistically different from 0  $\mu\text{g m}^{-3}$  within  $3\sigma$ . Finally, the LOA factor showed  
25 statistically significant contributions within  $3\sigma$  only during summer and late spring in Vilnius.  
26 Despite its considerable day-to-day variability this factor contributed  $1.0 \pm 0.8 \mu\text{g m}^{-3}_{\text{avg}}$  in  
27 Vilnius during summer.

28 The markers source apportionment instead attributed  $85\%_{\text{avg}}$  of the Other-OA<sub>marker</sub> mass to the  
29  $\text{SO}_4^{2-}$ -related SOA, while  $\text{NO}_3^-$ -related SOA, MSA-related SOA, and PBOA explained  
30 respectively  $9\%_{\text{avg}}$ ,  $5\%_{\text{avg}}$  and  $1\%_{\text{avg}}$  of the Other-OA<sub>marker</sub> mass (Fig. S13). The  $\text{SO}_4^{2-}$ -related  
31 SOA average concentration was  $2.4 \mu\text{g m}^{-3}$  during summer and  $1.7 \mu\text{g m}^{-3}$  during winter with

1 no significant differences from station to station, suggesting a regional origin of the factor.  
 2 The  $\text{NO}_3^-$ -related SOA concentration was  $0.4 \mu\text{g m}^{-3}_{\text{avg}}$  during winter, only  $0.03_{\text{avg}} \mu\text{g m}^{-3}$ ,  
 3 during summer, corresponding to  $10\%_{\text{avg}}$  and  $1\%$  of the OA, respectively. Moreover, the  $\text{NO}_3^-$   
 4 -related SOA during winter showed the highest average concentrations in Vilnius with  $0.5 \mu\text{g}$   
 5  $\text{m}^{-3}$  and the lowest in Rūgšteliškis with  $0.3 \mu\text{g m}^{-3}_{\text{avg}}$ . The MSA-related SOA instead  
 6 manifested the highest concentrations during summer with an average of  $0.12 \mu\text{g m}^{-3}_{\text{avg}}$ .  
 7 Higher values were observed during summer at the rural coastal site of Preila where the  
 8 average concentration was  $0.28 \mu\text{g m}^{-3}_{\text{avg}}$  corresponding to  $10\%_{\text{avg}}$  of the OM. Finally, the  
 9 PBOA factor exhibited the largest seasonal concentrations during spring at the rural terrestrial  
 10 site of Rūgšteliškis with an average of  $0.05 \mu\text{g m}^{-3}_{\text{avg}}$ , while the summer average  
 11 concentration was  $0.02 \mu\text{g m}^{-3}$  consistent with the low PBOA estimates reported in Bozzetti et  
 12 al. (2016) for the submicron fraction during summer.

13 Many previous studies reported a source apportionment of organic and inorganic markers  
 14 concentrations (Viana et al., 2008 and references therein). In these studies  $\text{SO}_4^{2-}$ ,  $\text{NO}_3^-$ , and  
 15  $\text{NH}_4^+$  were typically used as tracers for secondary aerosol factors commonly associated with  
 16 regional background and long-range transport; here we compare the apportionment of the  
 17 SOA factors obtained from the marker source apportionment and the OOA factors separated  
 18 by the offline-AMS source apportionment. Moreover, contrasting the two source  
 19 apportionments may provide insight into the origin of the OOA factors retrieved from the  
 20 offline-AMS source apportionment, and into the origin of the SOA factors resolved by the  
 21 offline-AMS source apportionment. To our knowledge an explicit comparison has not yet  
 22 been reported in the literature.

23

24 Table 2: Pearson correlation coefficients between Other-OA<sub>marker</sub> components from offline-AMS  
 25 and marker-source apportionment.

		Other-OA <sub>marker</sub>			
		SO <sub>4</sub> <sup>2-</sup> -related SOA	MSA-related SOA	NO <sub>3</sub> <sup>-</sup> -related SOA	PBOA
Other- OA <sub>offline- AMS</sub>	LOA	0.33	0.16	-0.08	0.10
	B-OOA	0.70	0.22	0.21	0.47
	S-OOA	0.60	0.45	-0.47	0.05

26

27 Table 2 reports the correlations between the time series of the Other-OA<sub>marker</sub> factors and the  
 28 Other-OA<sub>offline-AMS</sub> factors (Figs. 6 and S13). These correlations are mostly driven by seasonal

1 trends as none of these sources shows clear spikes except for LOA during summer in Vilnius.  
2 Using the correlations coefficients we can identify the mostly related factors from the two  
3 source apportionments.

4 The  $\text{SO}_4^{2-}$ -related SOA explained the largest fraction of the Other- $\text{OA}_{\text{marker}}$  mass (85%<sub>avg</sub>),  
5 and it was the only Other- $\text{OA}_{\text{marker}}$  factor always exceeding the individual concentrations of  
6 B-OOA and S-OOA, indicating that the variability explained by the  $\text{SO}_4^{2-}$ -related SOA in the  
7 marker-source apportionment is explained by both OOA factors in the offline-AMS source  
8 apportionment. Moreover, the  $\text{SO}_4^{2-}$ -related SOA seasonality seems consistent with the sum  
9 of S-OOA and B-OOA with higher concentrations in summer than in winter. This observation  
10 suggests that the OOA factors resolved by offline-AMS are mostly of secondary origin and  
11 the  $\text{SO}_4^{2-}$ -related SOA, typically resolved by the markersource apportionment, explains the  
12 largest fraction of the OOA factors apportioned by offline-AMS which includes both biogenic  
13 SOA and aged background OA.

14 The  $\text{NO}_3^-$ -related SOA and the PBOA were mostly related to the B-OOA factor as they  
15 showed higher correlations with B-OOA than with S-OOA. The B-OOA factor therefore may  
16 explain a small fraction of primary sources (PBOA), which however represents only 0.6%<sub>avg</sub>  
17 of the total OA. The  $\text{NO}_3^-$ -related SOA and the PBOA were mostly related to the B-OOA  
18 factor as they showed higher correlations with B-OOA than with S-OOA. The B-OOA factor  
19 therefore may explain a small fraction of primary sources (PBOA), which however represents  
20 only 0.6%<sub>avg</sub> of the total OA. In detail, the  $\text{NO}_3^-$ -related SOA correlation with B-OOA was  
21 poor ( $R = 0.21$ ), however the correlation with LOA and S-OOA was negative (Table 2),  
22 suggesting that the mass attributed to  $\text{NO}_3^-$ -related SOA by the markers source apportionment  
23 was fully attributed to the B-OOA factor in the offline-AMS source apportionment. This is  
24 also confirmed by the fact that the sum of LOA and S-OOA concentrations during winter  
25 (when the  $\text{NO}_3^-$ -related SOA substantially contributes) can't explain the  $\text{NO}_3^-$ -related SOA  
26 mass, which therefore has to be attributed to B-OOA.

27 The MSA-related SOA showed the highest correlation with the S-OOA factor, as the two  
28 sources exhibited the highest concentrations during summer, although the MSA-related SOA  
29 preferentially contributed at the rural coastal site of Preila. While we already discussed the  
30 probable secondary biogenic origin of S-OOA, the correlation with the MSA-related SOA  
31 suggests that the S-OOA factor, especially at the rural coastal site of Preila, explains also a  
32 large fraction of the marine biogenic SOA. The correlation between the two factors is

1 therefore not surprising as the precursor emissions (dimethyl sulfide, isoprene and terpenes)  
2 are strongly related to the temperature leading to higher summer MSA-related SOA and S-  
3 OOA concentrations. Assuming all the MSA-related SOA to be explained by the S-OOA  
4 factor, we estimate a marine biogenic SOA contribution to S-OOA of 27%<sub>avg</sub> during summer  
5 at Preila, while this contribution is lower at the other stations (12%<sub>avg</sub> in Rūgšteliškis during  
6 summer, 7% in Vilnius during spring, no summer data for Vilnius Fig. S13). As already  
7 mentioned, here we assume all the MSA-related SOA to be related to marine secondary  
8 biogenic emissions, however other studies also report MSA from terrestrial biogenic  
9 emissions (Jardine et al., 2015), moreover a certain fraction of the MSA-related SOA can also  
10 be explained by the B-OOA factor. Overall these findings indicate that the terrestrial sources  
11 dominate the S-OOA composition, nevertheless the marine SOA sources may represent a  
12 non-negligible fraction, especially at the marine site.

13 Another advantage obtained in coupling the two source apportionment results is the  
14 possibility to study the robustness of the factor analyses by evaluating the consistency of the  
15 two approaches as we already discussed for the primary OA and Other-OA fractions. Figure  
16 S14a displays the PMF modelled WSOC:measured WSOC PMF for the offline-AMS case,  
17 indicating a clear bias between Vilnius and the rural sites, with a WSOC overestimation of  
18 ~5% in Preila and Rūgšteliškis. While this overestimation is negligible for WSOC mass, it  
19 might have significant consequences on single factor concentrations. By contrast, OM  
20 residuals are more homogeneous for the case of markers PMF (Fig. S14b). As we show in  
21 Fig. S6, these residuals marginally affect the apportionment of combustion sources, as  
22 suggested by the well comparing estimates of BBOA and TEOA using the two methods.  
23 Therefore, these residuals are more likely affecting non-combustion sources (LOA, S-OOA  
24 and B-OOA). For the common days, the S-OOA concentration is not statistically different at  
25 the different stations during summer (confidence interval of 95%), indicating that the  
26 residuals are more likely affecting LOA and B-OOA, which instead show site-to-site  
27 differences. Now, the PMF WSOC residuals appear at all seasons, also during periods without  
28 significant LOA contribution in Vilnius. Therefore, we conclude that B-OOA is the factor  
29 most significantly affected by the difference in the WSOC residuals. We could best assess the  
30 residual effects by comparing the  $B-OOA_{\text{offline-AMS}}$  with that estimated using the other  
31 technique that seem to yield more homogeneous residuals:  $B-OOA_{\text{marker}}$ . Here  $B-OOA_{\text{marker}}$   
32 is estimated as  $\text{Other-OA}_{\text{markers}} - \text{LOA} - \text{S-OOA}$ . While  $B-OOA_{\text{offline-AMS}}$  shows site-to-site  
33 differences,  $B-OOA_{\text{markers}}$  did not show statistically different concentrations at all stations

1 within a confidence interval of 95%. Based on these observations, we conclude that observed  
2 site-to-site differences in B-OOA concentrations are likely to be related to model  
3 uncertainties.

#### 4 5 **4.5 $f\text{CO}^+$ vs. $f\text{CO}_2^+$**

6 Figure 7 displays the water-soluble  $f\text{CO}^+$  vs.  $f\text{CO}_2^+$  scatter plot. A certain correlation ( $R=0.63$ )  
7 is seen, with  $f\text{CO}_2^+$  values being higher than  $f\text{CO}^+$  ( $\text{CO}_2^+:\text{CO}^+$ : 1<sup>st</sup> quartile 1.50, median 1.75,  
8 3<sup>rd</sup> quartile 2.01), whereas a 1:1  $\text{CO}_2^+:\text{CO}^+$  ratio is assumed in standard AMS/ACSM analyses  
9 (Aiken et al., 2008; Canagaratna et al., 2007). Comparing the measured  $\text{CO}_2^+:\text{CO}^+$  values for  
10 the bulk WSOM and for pure gaseous  $\text{CO}_2$  might provide insight into the origin of the  $\text{CO}^+$   
11 fragment in the AMS. The fragmentation of pure gaseous  $\text{CO}_2$  returned a  $\text{CO}_2^+:\text{CO}^+$  ratio of  
12  $8.21_{\text{avg}}$  which is significantly higher than our findings for the water-soluble bulk OA  
13 ( $1.75_{\text{med}}$ ). Assuming thermal decarboxylation of organic acids as the only source of  $\text{CO}_2^+$  does  
14 not explain the observed  $\text{CO}_2^+:\text{CO}^+$  ratio of  $1.75_{\text{med}}$  and another large source of  $\text{CO}^+$  has to be  
15 assumed. Therefore, the carboxylic acid decarboxylation into  $\text{CO}_2$  can be considered as a  
16 minor source of  $\text{CO}^+$ .

17 Figure 7c and Fig. 8 show that not only does the water-soluble (WS)  $\text{CO}_2^+:\text{CO}^+$  ratio  
18 systematically differ from 1, but it also varies throughout the year with higher  $\text{CO}_2^+:\text{CO}^+$   
19 values associated with warmer temperatures (Fig. 7c). The lower  $\text{CO}_2^+:\text{CO}^+$  ratios in winter  
20 are primarily due to BB, as the WSBBOA factor profile showed the lowest  $\text{CO}_2^+:\text{CO}^+$  ratio  
21 ( $1.20_{\text{avg}}$ ) among all the apportioned WS factors ( $2.00_{\text{avg}}$  for B-OOA,  $2.70_{\text{avg}}$  for S-OOA, and  
22  $2.70_{\text{avg}}$  for LOA). We observed a seasonal variation of the  $\text{CO}_2^+:\text{CO}^+$  ratio also for the water-  
23 soluble OOA (S-OOA + B-OOA) mass spectral fingerprint. The  $\text{CO}_2^+:\text{CO}^+$  ratio was slightly  
24 lower for B-OOA than for S-OOA ( $2.00_{\text{avg}}$  for B-OOA, 2.70 for S-OOA). Nevertheless, given  
25 the low S-OOA relative contribution during winter (Fig. 3), we note that the total OOA  
26 showed a slightly lower  $\text{CO}_2^+:\text{CO}^+$  ratio during winter than in summer (Fig. S15), indicating  
27 that the OOA mass spectral fingerprint evolves over the year, possibly because of different  
28 precursor concentrations, and different photochemical activity.

29 Fig. 7a shows that most of the measured  $\{f\text{CO}^+;f\text{CO}_2^+\}$  combinations lies within the triangle  
30 defined by the BBOA, S-OOA and B-OOA  $\{f\text{CO}^+;f\text{CO}_2^+\}$  combinations. The LOA factor  
31  $\{f\text{CO}^+;f\text{CO}_2^+\}$  combination lies within the triangle as well, but is anyways a minor source and



1 thus unlikely to contribute to the  $\text{CO}_2^+/\text{CO}^+$  variability. We parameterized the  $\text{CO}^+$  variability  
2 as a function of the  $\text{CO}_2^+$ , and  $\text{C}_2\text{H}_4\text{O}_2^+$  fragment variabilities using a multi-parameter fit  
3 according to Eq. (8).  $\text{CO}_2^+$  and  $\text{C}_2\text{H}_4\text{O}_2^+$  were chosen as B-OOA and BBOA tracers,  
4 respectively, with B-OOA and BBOA being the factors that explained the largest fraction of  
5 the  $f\text{CO}^+$  variability (85% together).

$$6 \quad \text{CO}^+_i = a \cdot \text{CO}_2^+_i + b \cdot \text{C}_2\text{H}_4\text{O}_2^+_i \quad (8)$$

7 Although this parameterization is derived from the WSOM fraction  $\text{CO}_2^+$ ,  $\text{C}_2\text{H}_4\text{O}_2^+$ , and  $\text{CO}^+$   
8 originate from the fragmentation of oxygenated, i.e. mostly water-soluble compounds.  
9 Accordingly, this parameterization might also well represent the total bulk OA (as the offline-  
10 AMS recoveries of these oxygenated fragments are relatively similar:  $R_{\text{CO}_2^+}=0.74$ ,  
11  $R_{\text{C}_2\text{H}_4\text{O}_2^+}=0.61$ , Daellenbach et al., 2016). Note that this parameterization may represent very  
12 well the variation of  $\text{CO}^+$  in an environment impacted by BBOA and OOA, but should be  
13 used with caution when other sources (such as COA) may contribute to  $\text{CO}^+$ ,  $\text{CO}_2^+$  and  
14  $\text{C}_2\text{H}_4\text{O}_2^+$ . In order to check the applicability of this parameterization to a PMF output, we  
15 recommend monitoring the  $\text{CO}_2^+$  and  $\text{C}_2\text{H}_4\text{O}_2^+$  variability explained by the OOA and BBOA  
16 factors. In case a large part of the  $\text{CO}_2^+$  and  $\text{C}_2\text{H}_4\text{O}_2^+$  variability is explained by OOA and  
17 BBOA, the parameterization should return accurate  $\text{CO}^+$  values. The coefficients  $a$  and  $b$  of  
18 Eq. (8) were determined as 0.52 and 1.39 respectively, while the average fit residuals were  
19 estimated to be equal to 10% (Fig. S16). In contrast, parameterizing  $\text{CO}^+$  as proportional to  
20  $\text{CO}_2^+$  only (as done in the standard AMS analysis scheme with coefficients updated to the  
21 linear fit between  $\text{CO}^+$  and  $\text{CO}_2^+$  (1.75)) yielded 20%<sub>avg</sub> residuals, indicating that such a  
22 univariate function describes the  $\text{CO}^+$  variation less precisely.

23 An alternative parameterization is presented in the SI in which the contribution of moderately  
24 oxygenated species (such as S-OOA) to  $\text{CO}^+$  was also considered by using  $\text{C}_2\text{H}_3\text{O}^+$  as an  
25 independent variable. We show that the dependence of  $\text{CO}^+$  on  $\text{C}_2\text{H}_3\text{O}^+$  is statistically  
26 significant (Fig. 7c) as also suggested by the PMF results (S-OOA contributes 12% to the  
27  $\text{CO}^+$  variability). However, the parameter relating  $\text{CO}^+$  to  $\text{C}_2\text{H}_3\text{O}^+$  is negative, because the  
28  $\text{CO}^+:\text{CO}_2^+$  and  $\text{CO}^+:\text{C}_2\text{H}_4\text{O}_2^+$  ratios are lower in moderately oxygenated species compared to  
29 species present in BBOA and B-OOA. While this parameterization captures the variability of  
30  $\text{CO}^+$  across the seasons better compared to a 2-parameter fit for the present dataset, it may be  
31 more prone to biases in other environments due to the known contributions of other factors to  
32  $\text{C}_2\text{H}_3\text{O}^+$ . For example, cooking-influenced organic aerosol (COA) often accounts for a

1 significant fraction of  $C_2H_3O^+$ . For ambient datasets we propose the use of  $CO_2^+$  and  $C_2H_4O_2^+$   
2 only, which may capture less variation but is also less prone to biases. Although our results  
3 suggest that the available  $CO^+$  and O:C estimates (Aiken et al., 2008; Canagaratna et al.,  
4 2015) may not well capture the  $CO^+$  variability, our  $CO^+$  parameterization should not be  
5 applied to calculate the O:C ratios or recalculate the OA mass from AMS datasets, as those  
6 are calibrated assuming a standard fragmentation table (i.e.  $CO_2^+ = CO^+$ ).

7 In a recent work, Canagaratna et al. (2015) reported the Ar nebulization of water soluble  
8 single compounds to study the HR-AMS mass spectral fingerprints in order to improve the  
9 calculation of O:C and OM:OC ratios. Following the same procedure, we nebulized a subset  
10 of the same standard compounds including malic acid, azalaic acid, citric acid, tartaric acid,  
11 cis-pinonic acid, and D(+)-mannose. We obtained comparable  $CO_2^+ : CO^+$  ratios (within 10%)  
12 to those of Canagaratna et al. (2015) for all the analyzed compounds, highlighting the  
13 comparability of results across different instruments. With the exception of some  
14 multifunctional compounds (citric acid, malic acid tartaric acid, ketobutyric acid, hydroxyl  
15 methylglutaric acid, pyruvic acid, oxaloacetic acid, tartaric acid, oxalic acid and malonic  
16 acid), the water-soluble single compounds analyzed by Canagaratna et al. (2015) mostly  
17 showed  $CO_2^+ : CO^+$  ratios  $< 1$ , systematically lower than the  $CO_2^+ : CO^+$  ratios measured for the  
18 bulk WSOM in Lithuania (1<sup>st</sup> quartile 1.50, median 1.75, 3<sup>rd</sup> quartile 2.01), which represents a  
19 large fraction of the total OM (bulk EE: median = 0.59, 1<sup>st</sup> quartile = 0.51, 3<sup>rd</sup> quartile = 0.72).  
20 Considering the relatively high extraction efficiency, and considering that the  $CO^+$  and  $CO_2^+$   
21 fragmentation precursors tend to be more water soluble than the bulk OA, the aforementioned  
22 compounds could be representative of a large part of the  $CO^+$  and  $CO_2^+$  fragmentation  
23 precursors. This indicates that the selection of appropriate reference compounds for ambient  
24 OA is non-trivial, and the investigation of multifunctional compounds is of high importance.

25

## 26 **5 Conclusions**

27  $PM_1$  filter samples were collected over an entire year (November 2013 to October 2014) at  
28 three different stations in Lithuania. Filters were analyzed by water extraction followed by  
29 nebulization of the liquid extracts and subsequent measurement of the generated aerosol with  
30 an HR-ToF-AMS (Daellenbach et al., 2016). For the first time, the nebulization step was  
31 conducted in Ar, enabling direct measurement of the  $CO^+$  ion, which is typically masked by  
32  $N_2^+$  in ambient air and assumed to be equal to  $CO_2^+$  (Aiken et al., 2008).  $CO_2^+ : CO^+$  values  $> 1$

1 were systematically observed, with a mean ratio of  $1.7 \pm 0.3$ . This is likely an upper limit for  
2 ambient aerosol, as only the water-soluble OM fraction is measured by the offline-AMS  
3 technique.  $\text{CO}^+$  concentrations were parameterized as a function of  $\text{CO}_2^+$ , and  $\text{C}_2\text{H}_4\text{O}_2^+$ , and  
4 this two-variable parameterization showed a superior performance to a parameterization based  
5 on  $\text{CO}_2^+$  alone, because  $\text{CO}^+$  and  $\text{CO}_2^+$  show different seasonal trends.

6 PMF analysis was conducted on both the offline-AMS data described above and a set of  
7 molecular markers together with total OM. Biomass burning was found to be the largest OM  
8 source in winter, while secondary OA was largest in summer. However, higher concentrations  
9 of primary anthropogenic sources (biomass burning and hopanes here used as traffic markers)  
10 were found at the urban background station of Vilnius. The offline-AMS and marker-based  
11 analyses also identified local emissions and primary biological particles, respectively, as  
12 factors with low overall but episodically important contributions to PM. Both methods  
13 showed traffic exhaust emissions to be only minor contributors to the total OM; which is not  
14 surprising given the distance of the three sampling stations from busy roads.

15 The two PMF analyses apportioned SOA to sources in different ways. The offline-AMS data  
16 yielded factors related to regional background (B-OOA) and temperature-driven (likely  
17 biogenic-influenced) emissions (S-OOA), while the marker-PMF yielded factors related to  
18 nitrate, sulfate, and MSA. For the offline-AMS PMF, S-OOA was the dominant factor in  
19 summer and showed a positive exponential correlation with the average daily temperature,  
20 similar to the behavior observed by Leitch et al. (2011) in a Canadian boreal forest.  
21 Combining the two source apportionment techniques suggests that the S-OOA factor includes  
22 contributions from both terrestrial and marine secondary biogenic sources, while only small  
23 PBOA contributions to submicron OOA factors are possible. The analysis highlights the  
24 importance of regional meteorological conditions on air pollution in the southeastern Baltic  
25 region, as evidenced by simultaneous high BBOA levels at the three stations during three  
26 different episodes in winter and by statistically similar S-OOA concentrations across the three  
27 stations during summer.

28

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3

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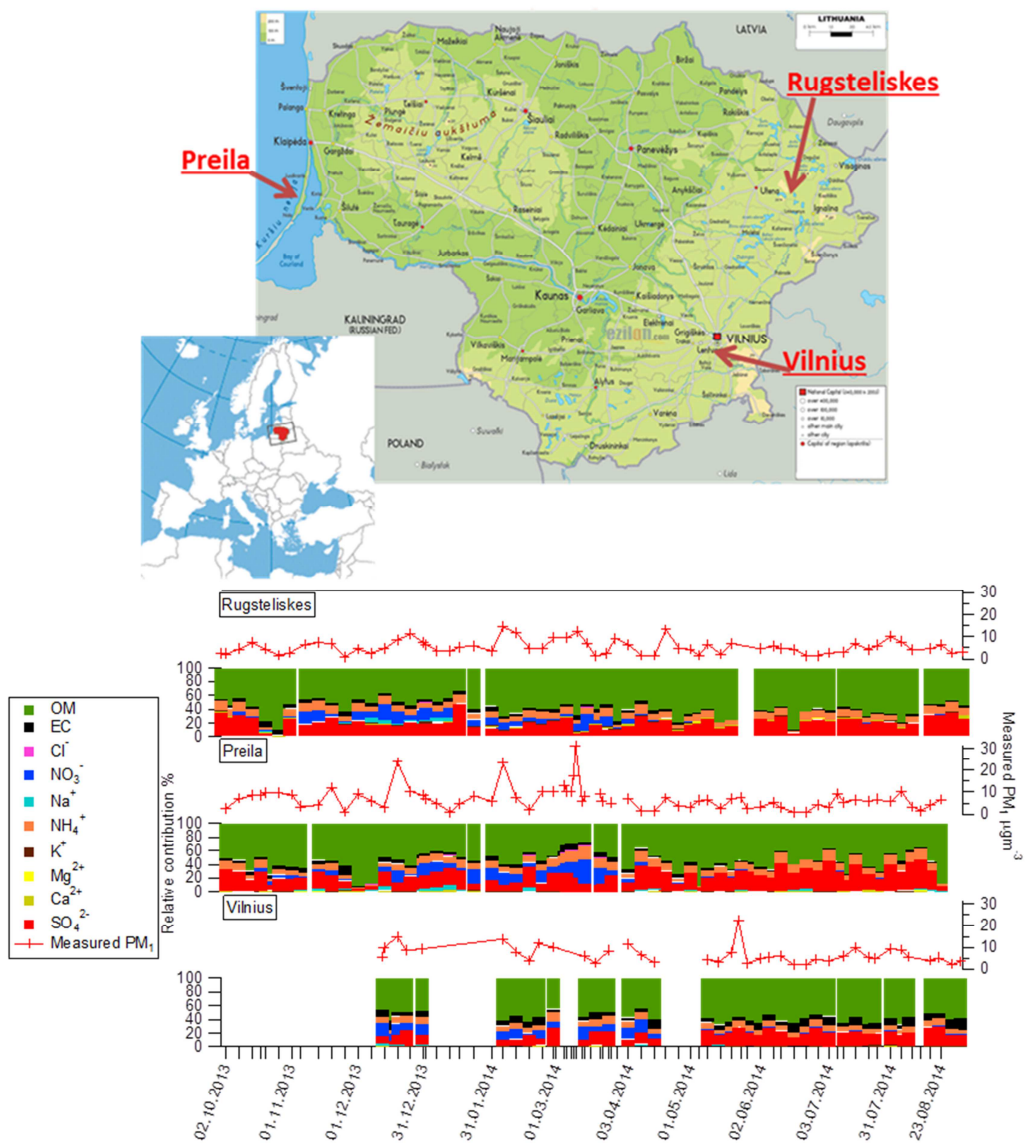
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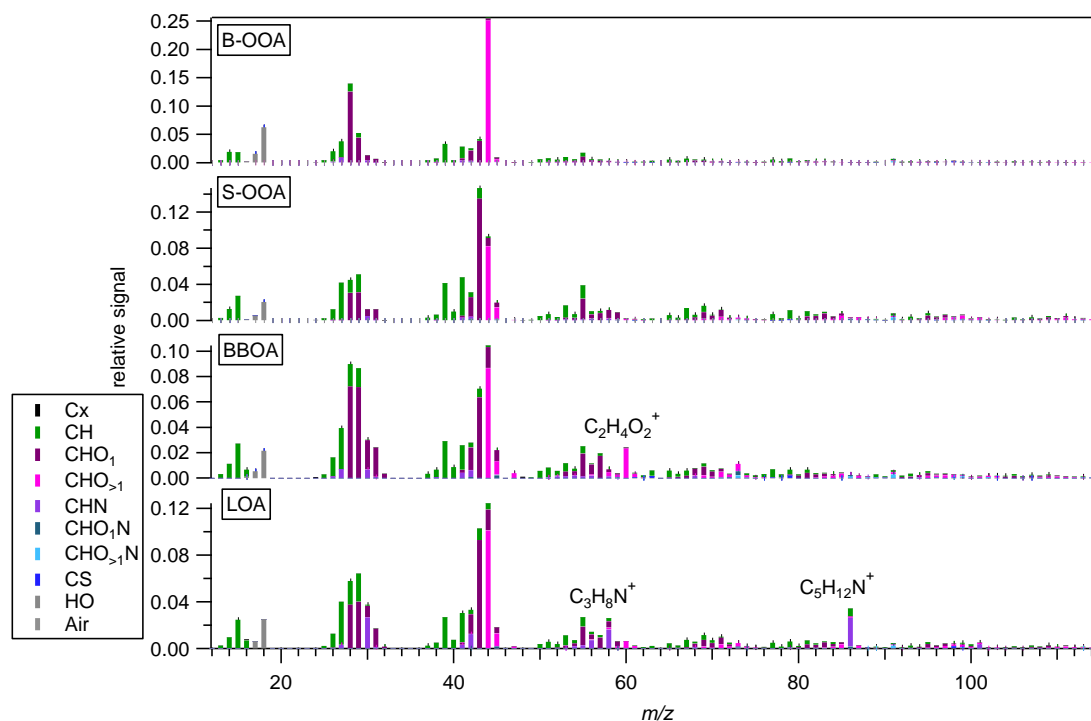
1 Figures main text



2

3 Figure 1. Sampling locations, and measured PM<sub>1</sub> composition. Ion concentrations from IC.

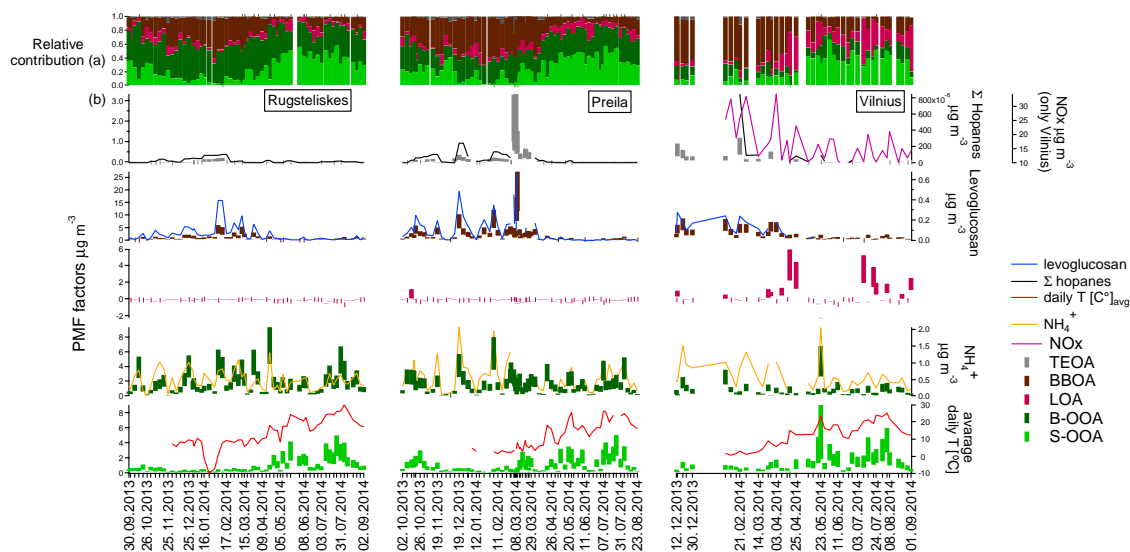




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2 Figure 2. Offline-AMS PMF factor profiles: background oxygenated OA (B-OOA), summer  
 3 oxygenated OA (S-OOA), biomass burning OA (BBOA), local OA (LOA).

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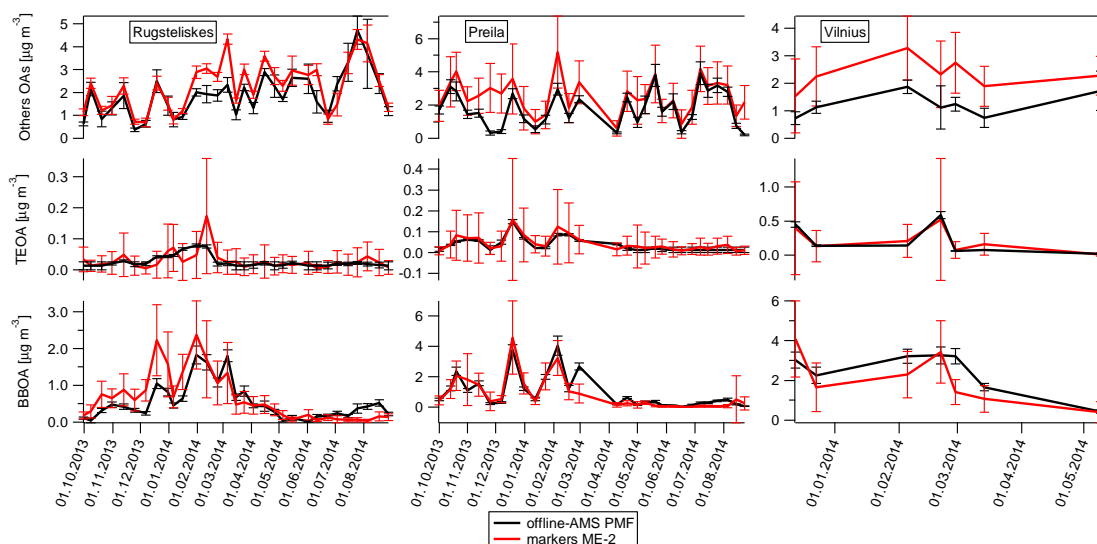


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6 Figure 3. a) Temporal evolutions of relative contributions to the OA factors; b) OA sources  
 7 and corresponding tracers: concentrations and uncertainties (shaded areas).

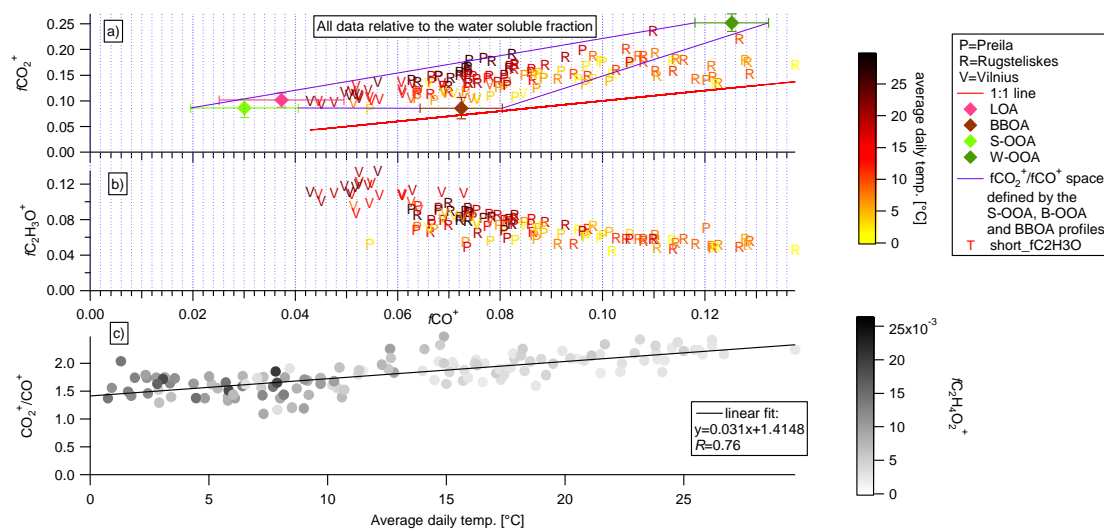
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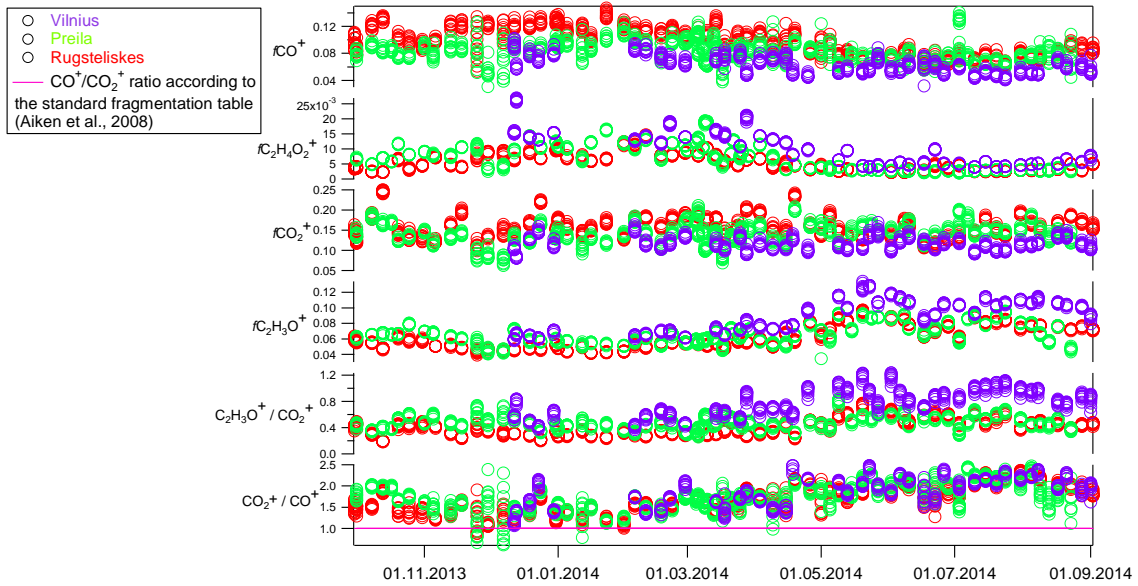
2 Figure 6. Marker-PMF and offline-AMS OM source apportionment comparison.



3

4 Figure 7. a) water-soluble  $f\text{CO}_2^+$  vs  $f\text{CO}^+$  scatter plot. Color code denotes the average daily  
 5 temperature [°C], diamonds indicate the  $f\text{CO}_2^+ / f\text{CO}^+$  ratio for different PMF factor profiles.  
 6 The 1:1 line is displayed in red. Few points from Rūgšteliškis lie outside the triangle,  
 7 suggesting they are not well explained by our PMF model. However, Fig. S5 displays flat  
 8 residuals for Rūgšteliškis, indicating an overall good WSOM explained variability by the  
 9 model. b) water-soluble  $f\text{C}_2\text{H}_3\text{O}^+$  vs  $f\text{CO}^+$  scatter plot. Color code denotes the average daily  
 10 temperature [°C] c) Scatter plot of the water-soluble  $\text{CO}_2^+$  to  $\text{CO}^+$  ratio vs. average daily  
 11 temperature. Grey code denotes  $f\text{C}_2\text{H}_4\text{O}_2^+$ .

12



1

2 Figure 8. Time-dependent fractional contributions ( $f$ ) of typical AMS tracers.

3