Semi-continuous gas and inorganic aerosol measurements at a Finnish urban site: comparisons with filters, nitrogen in aerosol and gas phases, and aerosol acidity

U. Makkonen¹, A. Virkkula¹², J. Mäntykenttä¹, H. Hakola¹, P. Keronen², V. Vakkari², and P. P. Aalto²

¹Finnish Meteorological Institute, 00560, Helsinki, Finland
²University of Helsinki, 00560, Helsinki, Finland

Correspondence to: U. Makkonen (ulla.makkonen@fmi.fi)

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Abstract. Concentrations of 5 gases (HCl, HNO₃, HONO, NH₃, SO₂) and 8 major inorganic ions in particles (Cl⁻, NO₃⁻, SO₄²⁻, NH₄⁺, Na⁺, K⁺, Mg²⁺, Ca²⁺) were measured with an online monitor MARGA 2S in two size ranges, \(D_p<2.5\ \mu m\) and \(D_p<10\ \mu m\), in Helsinki, Finland from November 2009 to May 2010. The results were compared with filter sampling, mass concentrations obtained from particle number size distributions, and a conventional SO₂ monitor. The MARGA yielded lower concentrations than those analyzed from the filter samples for most ions. Linear regression yielded the following MARGA vs. filter slopes: 0.72 for Cl⁻, 0.90 for NO₃⁻, 0.85 for SO₄²⁻, 0.91 for NH₄⁺, 0.49 for Na⁺, 3.0 for Mg²⁺, and 3.0 for Ca²⁺ and 0.90 for the MARGA vs. SO₂ monitor. For K⁺ there were not enough data points to calculate a statistically significant linear regression. There were clear seasonal cycles in the concentrations of the nitrogen-containing gases: the median concentrations of HNO₃, HONO, and NH₃ were 0.09 ppb, 0.37 ppb, and 0.01 ppb in winter, respectively, and 0.15, 0.15, and 0.14 in spring, respectively. The gas-phase fraction of nitrogen decreased roughly with decreasing temperature, so that in the coldest period from January to February the median contribution was 28 % but in April to May was 53 %. There were also large fractionation variations that temperature alone cannot explain. HONO correlated well with NOₓ but a large fraction of the HONO-to-NOₓ ratios were larger than published ratios in a road traffic tunnel, suggesting that a large amount of HONO had other sources than vehicle exhaust. Aerosol acidity was estimated by calculating ion equivalent ratios. The sources of acidic aerosols were studied with trajectory statistics that showed that continental aerosol is mainly neutralized and marine aerosol acidic.

1 Introduction

Concentrations of major inorganic ions in aerosols have been measured for decades by sampling on filters and subsequently analyzing them with ion chromatography (e.g., Mulik et al., 1976; Stevens et al., 1978; Mulik and Sawicki, 1979). These kind of measurements are an integral part of monitoring networks such as the European Monitoring and Evaluation Programme (EMEP) (EMEP, 2007). With filter sampling, time resolution is low – from some hours to days – depending on concentrations. In addition, filter sampling suffers from both negative and positive artifacts (e.g., Lipfert, 1994). In order to study atmospheric processes, methods have been developed for measuring aerosol chemical composition at a higher time resolution and to avoid the artifacts associated with filter sampling.

A method with a good time resolution, in the order of seconds/minutes, is the Aerosol Mass Spectrometer (AMS) (Jayne et al., 2000; Jimenez et al., 2003), which provides size-resolved chemical composition of submicron aerosols. The AMS measures the concentrations of major aerosol constituents: organics, sulphate, nitrate, ammonium, and chloride. The disadvantage of it is that it does not detect some other important elements such as sodium, one of the main constituents of sea salt, and calcium and magnesium, significant constituents of both both soil dust and sea salt. To
analyze these elements, semicontinuous methods involving ion chromatography have been developed. In these methods, the sample is first taken for a period of some minutes to hours and then analyzed. An example is the Particle Into Liquid Sampler (PILS) that can be connected, for instance, to an ion chromatograph (IC) or an instrument for analyzing the concentration of water-soluble organic carbon (Weber et al., 2001; Orsini et al., 2003; Sullivan et al., 2004). The PILS combines two aerosol technologies: particle growth in a mixing condensation particle counter, and droplet collection by a single jet inertial impactor. It takes advantage of the principle of steam-jet aerosol collector (SJAC) (e.g., Khlystov et al., 1995). Another semicontinuous method that uses a SJAC is the ambient ion monitor (AIM, URG Corporation, USA) that determines the concentrations of several anions and cations in aerosol (e.g., Wu and Wang, 2007; Nie et al., 2010) and exists in four different configurations (URG 9000A–D) that have different performances. Yet another method that is based on the same principle is the Dionex Gas Particle Ion Chromatography (GPIC) system that measures concentrations of water-soluble inorganic aerosol constituents (Cl\(^-\), NO\(_3\)\(^-\), SO\(_4\)\(^2-\), and NH\(_4\)\(^+\)) and their associated precursor gases (HCl, SO\(_2\), HNO\(_3\), NH\(_3\)) (Godri et al., 2009). The ability to measure both water-soluble aerosol and the precursor gas concentrations at a high time resolution has recently been shown to be especially valuable for evaluating models, for instance the secondary inorganic aerosol formation (Schaap et al., 2011) and the gas-aerosol partitioning of ammonium nitrate (Aan de Brugh et al., 2012).

For measuring ammonia, several methods have been developed based on very different principles: wet chemistry, optics, optoacoustics and mass spectrometric techniques. An intercomparison of eleven instruments for measuring atmospheric ammonia at ambient concentrations was presented by von Bobrutzki et al. (2010). It was shown in that paper that wet chemistry instruments had good long-term stability and provided a reliable differentiation between gas-phase NH\(_3\) and aerosol NH\(_4\)\(^+\), which could not be validated for the other instruments.

The instrument for Measuring AeRosols and GAses (MARGA) (ten Brink et al., 2007) has a SJAC (Slanina et al., 2001) and it is connected to two ion chromatographs, one for anions and another for cations. In addition, it also measures the concentrations of water-soluble gases that produce ions observable with an IC, using the principles presented, e.g., by Wyers et al. (1993). Trebs et al. (2004) who used basically the same method, even though they did not call it MARGA then, at a rural site in the Amazon Basin. The instrument consisted of a wet rotating annular denuder in combination with a SJAC, followed by an IC for measuring anions and flow injection analysis (FIA) for measuring ammonium; other cations were not measured. The MARGA was used to monitor the size distribution of nitrate, ammonium, sulphate and chloride in aerosol at the top of a 200-m meteotower in Cabauw, Netherlands (ten Brink et al., 2007). Four parallel impactors with cut-offs of 0.18, 0.32, 0.56 and 1.0\(\mu\)m were attached to the MARGA inlet. In a clean background environment, a MARGA was used at an EMEP supersite in Scotland (Cape, 2009).

A commercially available MARGA (Applikon Analytical BV, Netherlands) was operated by the Finnish Meteorological Institute Air Quality Research (FMI/AQR) at the urban background station in Kumpula, Helsinki, from November 2009 to May 2010. These were the first semi-continuous measurements of sulphur dioxide (SO\(_2\)), nitric acid (HNO\(_3\)), nitrous acid (HNO\(_2\) or HONO), hydrochloric acid (HCl) and ammonia (NH\(_3\)) using a rotating wet annular denuder in combination with an on-line ion chromatograph in Finland. In addition, the major anion (Cl\(^-\), NO\(_3\)\(^-\), SO\(_4\)\(^2-\)) and cation (NH\(_4\)\(^+\), Na\(^+\), K\(^+\), Mg\(^{2+}\), Ca\(^{2+}\)) concentrations in aerosols in two size ranges, \(D_p < 2.5 \mu m\) \((PM_{2.5})\) and \(D_p < 10 \mu m\) \((PM_{10})\), were measured with the instrument.

The most important goal of this study was to investigate whether the MARGA could be used to replace the traditional EMEP filter pack method (EMEP, 2007). Furthermore, the Clean Air for Europe (CAFE) Directive (EU 2008/50/EC) demands that the member states measure also the chemical composition of PM\(_{2.5}\). For monitoring networks it is important that the measurement methods give comparable results. There are published comparisons between filter sampling and semicontinuous methods (e.g., Nie et al., 2010) but not with the commercially available MARGA. Trebs et al. (2008) compared Amazonian ammonium, nitrate, chloride, and sulphate concentrations analyzed from filter samples with those measured with an instrument similar to MARGA. In the comparison, ammonium was the only cation, and in that instrument it was analyzed with FIA. In the instrument used in the present work, five cations were analyzed, all with an IC. To our knowledge, there are no published comparisons between a MARGA that analyzes the concentrations of all the above-mentioned three anions and five cations and conventional filter sampling methods.

In this paper the performance of the MARGA 2S instrument is first evaluated by comparing it with independent methods: filter sampling, mass concentrations obtained from particle number size distributions, and a conventional SO\(_2\) monitor. Next, the observations on nitrogen compounds (both in the gas and aerosol phases) are discussed, and finally the aerosol data are used to study aerosol acidity and its sources.

2 Experimental

2.1 Measurement site

The Station for Measuring Ecosystem-Atmosphere Relationships (SMEAR III) is located next to the main building of the Finnish Meteorological Institute (FMI) on a rocky hill 26 m a.s.l., about 5 km northeast of the center of Helsinki. The station consists of a measurement container...
and meteorological measurements on ground and in a tower. The container is air conditioned and kept at a constant temperature at 20 °C. A comprehensive description of the SMEAR III station can be found in Järvi et al. (2009).

2.2 MARGA

Concentrations of 5 gases (HCl, HNO\textsubscript{3}, HONO, NH\textsubscript{3}, SO\textsubscript{2}) and 8 major inorganic ions in particles (Cl\textsuperscript{-}, NO\textsubscript{3}\textsuperscript{-}, SO\textsubscript{4}\textsuperscript{2-}, NH\textsubscript{4}\textsuperscript{+}, Na\textsuperscript{+}, K\textsuperscript{+}, Mg\textsuperscript{2+}, Ca\textsuperscript{2+}) were measured with a MARGA 2S from 1 November 2009 to 24 May 2010. The MARGA 2S ADI 2080 (Applikon Analytical BV, Netherlands) consists of two identical sample boxes and one analytical box. Ambient air was drawn at the flow rate of 2 m\textsuperscript{3} h\textsuperscript{-1} through a PM\textsubscript{10}-inlet (Teflon coated, URG-2000-30DBN-TC, 2 m\textsuperscript{3} h\textsuperscript{-1}) and divided into the PM\textsubscript{10} sample box and into the PM\textsubscript{2.5} sample box through a PM\textsubscript{2.5} cyclone (1 m\textsuperscript{3} h\textsuperscript{-1} Teflon coated inlet, URG-2000-30ENB). However, the first part of the measurements (1 November 2009–19 January 2010) was carried out without any specific inlet – there was just a rain shield on the inlet tubing. The PM\textsubscript{10}-inlet and the PM\textsubscript{2.5} cyclone were installed on 19 January 2010.

Sample air was first drawn through the Wet Rotating Denuder (WRD) where water-soluble gases diffused to the absorption solution, then particles were collected in a steam-jet aerosol collector (SJAC) (Slanina et al., 2001). Diluted hydrogen peroxide solution (10 ppm) was used as the absorption solution to prevent microbiological growth. Absorption solutions were drawn from the WRD and the SJAC to syringes (25 ml) in the analytical box. Each hour after the syringes had been filled, samples were injected into Metrohm anion (250 µl loop) and cation (500 µl loop) chromatographs with the internal standard (LiBr). Cations were separated in a Metrosep C4 (100/4.0) cation column using 3.2 mmol l\textsuperscript{-1} HNO\textsubscript{3} eluent. For anions a Metrosep A Supp 10 (75/4.0) column and Na\textsubscript{2}CO\textsubscript{3}-NaHCO\textsubscript{3} (7/8 mmol l\textsuperscript{-1}) eluent was used. Components were detected by conductivity measurements. For anions a chemical suppressor (H\textsubscript{3}PO\textsubscript{4} for regeneration) was used. The detection limits for all the components were 0.1 µg m\textsuperscript{-3} or better, except for K\textsuperscript{+} (0.16 µg m\textsuperscript{-3}), Mg\textsuperscript{2+} (0.12 µg m\textsuperscript{-3}) and Ca\textsuperscript{2+} (0.21 µg m\textsuperscript{-3}). The detection limits and the repeatability of the instrument studied from parallel measurements of the two sample boxes (neither the PM\textsubscript{10}-inlet nor PM\textsubscript{2.5} cyclone was used) are presented in Table 1.

Field blanks for gases were checked by installing filters in the sampling line before the denuder of the instrument. An oxalic-acid-impregnated filter was used to remove ammonia and a NaOH impregnated filter to remove acidic gases. The blank values for gases were insignificant, except for nitric acid. The instrument blank for nitrate and nitric acid was remarkable, of the order of 0.4 µg m\textsuperscript{-3}, and the blank-value caused by the instrument was subtracted from the results. The high nitrate blank was caused by small amounts of cation eluent leaking continuously to the anion injector.

### Table 1. The detection limits and the repeatability calculated from real air samples collected using the two parallel sample boxes of the MARGA instrument (1 November 2009–18 January 2010).

| Compound | Detection limit µg m\textsuperscript{-3} | Repeatability % |
|----------|----------------------------------------|-----------------|
| HCl      | 0.02                                   | 30              |
| HNO\textsubscript{3} | 0.03                                   | 3.3             |
| SO\textsubscript{2} | 0.04                                   | 4.9             |
| HNO\textsubscript{3} | 0.05                                   | 1.1             |
| NH\textsubscript{3} | 0.05                                   | 3.9             |
| Cl\textsuperscript{-} | 0.02                                   | 4.5             |
| NO\textsubscript{3} | 0.04                                   | 1.0             |
| SO\textsubscript{4} | 0.03                                   | 1.1             |
| Na\textsuperscript{+} | 0.02                                   | 1.2             |
| NH\textsubscript{4} | 0.03                                   | 1.9             |
| K\textsuperscript{+} | 0.01                                   | 13              |
| Mg\textsuperscript{2+} | 0.01                                   | 8.7             |
| Ca\textsuperscript{2+} | 0.01                                   | 1.1             |

2.3 Filter sampling

The aerosol results of MARGA were compared with the results of daily sampling of PM\textsubscript{10} (Digitel PM\textsubscript{10} sampling inlet, 1 m\textsuperscript{3} h\textsuperscript{-1}) and PM\textsubscript{2.5} (Digitel PM\textsubscript{2.5} inlet. 1 m\textsuperscript{3} h\textsuperscript{-1}) on Teflon filters. The filters were extracted in ultrapure water (Milli-Q) and analyzed by ion chromatographs (Waters). The procedure is described in detail in the EMEP Manual (EMEP, 2007). Daily averages were calculated from the MARGA results and compared with the results of the filter method from 5 February to 5 May 2010.

2.4 Other measurements

The routine measurements of the SMEAR III station include particle number size distribution measurements with a Twin DMA (TDMPS) in the size range 3–950 nm: SO\textsubscript{2} measurements with a Horiba APSA 360 monitor; O\textsubscript{3} measurements with a TEI 49 monitor; and NO\textsubscript{x} measurements with a TEI 42S monitor. There is also other instrumentation at the site (see Järvi et al. (2009) for details) but those mentioned above are used here. The number size distributions were used for estimating submicron aerosol mass concentration and comparing that with the sum of the analyzed ions. The SO\textsubscript{2} concentrations could be used as such since the MARGA also measures it. The NO\textsubscript{x} and ozone concentrations were used for studies of nitric and nitrous acid.

3 Results and discussion

A statistical summary of the MARGA data is presented in Table 2, including averages, standard deviations, selected
percentiles of the cumulative distribution of concentrations, and the fraction of time data are obtained for each ion or gas. When calculating the averages, data values below detection limit (DL) were taken into account by giving them the value of 0.5 × DL. The concentrations of sulphate, nitrate, ammonium, and calcium were most of time higher than the detection limits. The lowest fraction of concentrations above detection limit in PM$_{10}$ was for potassium.

The gas-phase concentrations presented below are in ppb and the aerosol concentrations in µg m$^{-3}$ at 20°C and 1013 mbar. The time of the measurements in the figures is the local winter time (UTC+2 h).

### 3.1 Temporal variations of aerosol concentrations

Several observations can be made from the time series of the concentrations of the ions analyzed from the aerosol phase in both the PM$_{2.5}$ and PM$_{10}$ size ranges (Fig. 1). The data for which the MARGA software automatic fitting routine to the chromatograms does not yield any concentrations are given as a discontinuity in the graph. This is an effective detection limit. Some of the breaks in the time series are due to the fact that the instrument was not running because of some technical issue such as maintenance operations.

The MARGA software does not always recognize a small sodium peak next to a larger ammonium peak or vice versa, which leads to discontinuities in these data. For chloride there is an additional reason for the discontinuities: before the anion column gets exhausted, the chloride peak slowly disappears to the water dip in the beginning of the chromatogram.

The two size ranges measured with the MARGA followed each other well, especially for sulphate, ammonium and nitrate but worse for calcium and magnesium. This is reasonable as it is well known that the latter two exist both in soil dust and sea salt and that the size distributions of both of these aerosol types are dominated by coarse particles. The size fractions will be discussed more below.

The variation of the aerosol concentrations has two types: short peaks that are probably due to some local source in Helsinki, and slower, several-day-long variations that are due to long-range transport. The latter will be discussed below by using back trajectories. One interesting peak can be mentioned. The highest potassium concentrations were measured during the New Year Eve celebrations: K$^+$ concentrations rose above 1 µg m$^{-3}$ at 19:00 p.m. on 31 December 2009, reached the peak 8.1 µg m$^{-3}$ at 21:00–22:00 and decreased again below 1 µg m$^{-3}$ at 03:00 a.m. on 1 January 2010. In
this same fireworks-related peak, the concentrations of the other ions also increased clearly.

The sum of the analyzed ions follows well the temporal variation of the mass concentration calculated from the DMPS (Fig. 2). Here the mass concentration from the DMPS data was calculated assuming the density of 1.5 g cm$^{-3}$ for spherical particles to generate an independent estimate for aerosol mass concentration. Figure 2 shows that the high and the low concentrations of the two methods occurred mainly simultaneously. However, the ion concentrations obviously explain only part of the mass: even if we compare PM$_{10}$ from the DMPS and the sum of ions in PM$_{10}$ from the MARGA, PM$_{1}$ is 97% of time larger than the sum of the ions. This means that a large fraction of aerosol mass has not been analyzed. This is in line with earlier studies, for instance by Timonen et al. (2008) who observed that most of aerosol mass often consists of organics also in Helsinki.

3.2 Comparisons of aerosol data with ions analyzed from filter samples

The concentrations measured with the MARGA after the PM$_{10}$ inlet was installed were compared with those analyzed from the Teflon filters by plotting them in the time series (Fig. 1) and by calculating linear regressions (Fig. 3). For the regressions the hourly MARGA data were first averaged over the sampling periods of the filters. Values below detection limit (DL) were taken into account by giving them the value of 0.5 $\times$ DL. For the regressions, only those filter sampling periods were used, during which at least 50% of the MARGA data of the ion to be analyzed were $>$ DL, in order to reduce the weight of the more uncertain data. The reason is that for some ions there were several sampling periods during which there were no hourly data $>$ DL. The extreme example is K$^+$ that did not have any single filter sampling period during which all hourly data were $>$ DL, and there were only 5 sampling periods out of 88 during which at least 50% of the hourly data were $>$ DL (Table 3). If K$^+$ data from all the 88 filter sampling periods had been used for the regressions, most of the data points used for the fitting would have been given the same value, and 0.5 $\times$ DL and the regression would have been meaningless. On the other hand, if only those sampling periods had been taken into account during which concentrations during all 24 h had been $>$ DL, the amount of data points for the regression would have been reduced an unnecessary amount. For instance, the concentration of an ion may have been $<$ DL during two hours of some sampling period and it makes sense to replace these two values by 0.5 $\times$ DL, calculate the sampling-period average, and use that value in the regression. The approach used now is a compromise between these two: using either all data even if most of them were $<$ DL or not using any of the data where any hourly value was $<$ DL.

Fig. 1. Hourly-averaged concentrations of inorganic ions in aerosols at SMEAR III from 1 November 2009 to 25 May 2010. MARGA data in PM$_{10}$ (black line) and PM$_{2.5}$ (blue line); concentrations analyzed from 24-h PM$_{10}$ filter samples (red line). Note: before 19 January 2010, there was only a total aerosol inlet in use and no PM$_{2.5}$ cyclone.

Fig. 2. Sum of inorganic ion concentrations in aerosols measured with MARGA in the size range $D_p < 10\mu m$ (noting that before 19 January 2010 there was only a total aerosol inlet in use) and the mass concentration of particles estimated from the particle number size distributions ($m(DMPS)$) in the size range $D_p < 1 \mu m$ and assuming the density of 1.5 g cm$^{-3}$. 
Fig. 3. Comparisons of concentrations of ions in the size range $D_p < 10 \mu m$ measured with the MARGA and analyzed from Teflon filters. The MARGA data were averaged over the filter sampling periods. Unit: $\mu g m^{-3}$. The linear regressions with the black lines and equations were calculated by fitting both slopes and offsets and those with the red dashed lines and red equations by forcing the offset to zero.

The linear regressions were calculated both by fitting both slope and offset and by forcing the offset to zero. Regressions calculated by using the MS Excel 2010 linest function also yield the uncertainties of the slope and offset as standard errors (stderr). From these the respective standard deviations were calculated from $\text{stdev} = \sqrt{n \cdot \text{stderr}}$, where $n$ is the number of observations. The slopes, offsets and their uncertainties are given in Table 3.

The slopes for sulphate, ammonium, and nitrate were all close to each other, ranging from 0.85 to 0.91 (Table 3, Fig. 3). From these comparisons alone it is not possible to determine whether the MARGA or the filter sampling results are closer to the true ones. However, the filter sampling method is routine and the laboratory follows strict EMEP protocols (EMEP, 2007) so this suggests that the concentrations given by the MARGA were underestimated. For ammonium it should also be noted that there is the possibility of a positive artifact in the filter samples. If there is ammonia in the laboratory air that reacts with sulphate in the filters and forms ammonium, the resulting ammonium concentrations will be higher than they were originally. Filters may also result in a negative artifact: ammonium nitrate is semivolatile and if the filters are stored at elevated temperatures, both ammonium and nitrate may volatilize from the filters. The filter handling was done at room temperature so this effect can be considered to be negligible.

The MARGA software does not always recognize small ammonium and sodium peaks which eluate next to each other. This leads to underestimation of both compounds and it is obviously more severe for sodium (Table 3, Fig. 3). For Mg$^{2+}$ and Ca$^{2+}$ the slopes were clearly above one, i.e., 3.0. It was discovered during the following campaign that all these were artifacts that could be significantly reduced by replacing the loop with a concentration column. It was thus shown that the reason for high Mg$^{2+}$ and Ca$^{2+}$ was either a dirty loop or sensitivity or the MARGA software could not calculate the small peak areas to concentrations correctly. The

| Ion  | N  | $k \pm \text{stderr}(k)$ | $C_0 \pm \text{stderr}(C_0)$ | $r^2$ |
|------|----|--------------------------|-----------------------------|-------|
| Cl$^-$ | 38 | 0.72 ± 0.33              | 0.03 ± 0.11                 | 0.83  |
| NO$_3^-$ | 84 | 0.90 ± 0.24              | 0.46 ± 0.69                 | 0.93  |
| SO$_4^{2-}$ | 87 | 0.85 ± 0.12              | 0.24 ± 0.30                 | 0.98  |
| Na$^+$ | 28 | 0.49 ± 0.46              | −0.03 ± 0.20                | 0.55  |
| NH$_4^+$ | 85 | 0.91 ± 0.42              | −0.30 ± 0.54                | 0.83  |
| K$^+$ | 5  | −0.26 ± 0.70             | 0.13 ± 0.13                 | 0.19  |
| Mg$^{2+}$ | 69 | 3.03 ± 2.04              | −0.02 ± 0.09                | 0.69  |
| Ca$^{2+}$ | 82 | 3.03 ± 1.26              | 0.09 ± 0.23                 | 0.86  |
| SO$_2$ | 3701 | 0.90 ± 0.31           | −0.08 ± 0.68                | 0.89  |

Table 3. Coefficients of the linear regressions of the concentrations measured with the MARGA data vs. those analyzed from the PM$_{10}$ filters and the SO$_2$ measured with a Horiba APSA 360 monitor: $C(\text{MARGA}) = k \cdot C(\text{comparison method}) + C_0$. $N$ = number of observations used in the regression. For explanation of the uncertainties of $k$ and $C_0$ see Sect. 3.2. For K$^+$ the $r^2$ statistic is negative for the model without a constant, indicating that the model is not appropriate for the data.
of each ion to the sum was calculated. On average, sulphate was the most abundant ion with 50.4% of inorganic ion mass after K⁺. The relative contributions of the other ions are shown in Fig. 4. The calculated ratios did not exceed the range of 0.1 to 10. The median ratio was 1 for all ions except for the ratio of K⁺ to PM10, which was 0.98.

The $r^2$ statistics for the K⁺ regression were 0.19, indicating a weak correlation. The negative $r^2$ values suggest that the model with no offset is not appropriate for K⁺. For all other ions, the $r^2$ values were positive and greater than 0.98, indicating a strong correlation.

3.3 Aerosol concentration in the two size ranges

The MARGA aerosol data were used for estimating their contribution to the sum of the analyzed ions and for studying their size distribution in the time period when both the PM10 inlet and the PM2.5 cyclone were in use, from 19 January to 24 May 2010 (Fig. 4). The size distribution is of course just a crude estimation, since there were only two size ranges. At each hour the concentrations were summed up and the ratio of each ion to the sum was calculated. On average, sulphate is responsible for most (50.4%) of inorganic ion mass after the PM2.5 inlet and slightly less (46.4%) after the PM10 inlet.

3.4 Temporal variation of the gases and their relationships

The hourly-averaged concentrations of gases measured with the MARGA (HNO3, HNO2, NH3, and SO2) at SMEAR III from 1 November 2009 to 25 May 2010. The red line in the SO2 time series is the data measured with a TEI 43iTL monitor.
and in the scatter plot (Fig. 6). For the other gases, we can only compare our data with values measured in other sites just to see whether the values are in a reasonable range.

### 3.4.1 HNO$_3$ and HNO$_2$

To get some estimate of the general level of HNO$_3$ concentrations, the data from the SMEAR II station in Hyytiälä in the southwestern part of central Finland were used for comparison. The average HNO$_3$ concentration (0.16 ppb) of the whole measurement period at SMEAR III was almost the same as the average HNO$_3$ measured using an EMEP filter pack at SMEAR II (0.15 ppb) in central Finland during the same period (unpublished data, analyzed at FMI/AQR). The yearly average concentration 0.05 ppb measured with MARGA at the EMEP supersite in south-eastern Scotland in 2007 was much lower (Cape, 2009). So, in principle the order of magnitude of the HNO$_3$ concentrations is reasonable. However, we cannot exclude losses in our inlet system and underestimation due to that. To evaluate inlet losses our present data are not sufficient.

A major pathway in the formation of HNO$_3$ is the reaction NO$_2$ + OH$^-$ + M → HNO$_3$ + M, where the hydroxyl radical OH$^-$ is originated from photochemical reactions. Nitrous acid HNO$_2$ formed in the reaction NO + OH$^-$ + M → HNO$_2$ + M, on the other hand, is dissociated by solar radiation: HNO$_2$ + h$\nu$ → OH$^+$ + NO. Ozone is produced in photochemical reactions in sunlight and it is also transported downward from ozone rich air from above during daytime when mixing is efficient. At night it is destroyed when it reacts rapidly with NO according to NO + O$_3$ → NO$_2$ + O$_2$ and also due to dry deposition. In locations where NO source is large, ozone can be totally destroyed (Finlayson-Pitts and Pitts, 2000). In other words, when there is sunlight, the expected HNO$_3$ and O$_3$ concentrations are higher and the HONO concentrations are lower than in the darkness. This is in agreement with our observations. Concentrations of the gases relevant to the above discussion are presented during a dark period in December (Fig. 7) and a light period in May (Fig. 8), when there is already a clear diurnal cycle. The concentration of nitrate does not follow the temporal variation of nitric acid, which suggests its sources are further away and it is transported over long distances. This is supported by the trajectory statistical analysis presented below in Sect. 3.6.

During the darkest months (November–January) when sunlight was very limited in the northern latitudes, the concentration of nitric acid was mostly below 0.1 ppb and stayed stable throughout day and night. At the beginning of February when the amount of daylight increased and there was still snow cover on the ground, the variation of HNO$_3$ increased and peaks up to 1–1.5 ppb were detected. In March and April the concentrations varied below 0.5 ppb. In May HNO$_3$ concentration increased again and peaked up to 1 ppb. In the spring days there is enough light for the photochemical reactions to form OH-radicals, which react with NO$_2$ forming nitric acid. Also the snow cover affects the amount of radiation.

The diurnal cycle was calculated both for the winter and for the spring periods. Only the latter is shown in Fig. 9 because in winter there was no clear daily pattern. Also the diurnal cycle of nitrate in aerosol (PM$_{10}$) is shown because nitric acid is its precursor and because ammonium nitrate particles are semivolatile and may evaporate into ammonia
and nitric acid when temperature increases (e.g., Seinfeld and Pandis, 1998). Even though it was discussed above that the nitrate sources in general are far away from our measurement site, it did also have a diurnal cycle in spring, which can be observed from 3-month statistics. In March–May the HNO$_3$ concentration had a minimum at 07:00–09:00 and a clear maximum at 12:00–14:00 LT. Nitrate, on the other hand, had its maximum average concentration two hours earlier, at 10:00 (Fig. 9). These diurnal cycles resemble those observed at other sites, for instance Zürich (Fisseha et al., 2006), Pittsburgh (Wittig et al., 2004), and Oensingen, Switzerland (Wolff et al., 2010): the peak nitrate concentration is a few hours earlier than that of HNO$_3$. The time lag can be explained by the temperature dependent partitioning of ammonium nitrate between the gas and aerosol phase (e.g., Fisseha et al., 2006). The times of the diurnal maxima are not quite the same at the sites. The reason is that the diurnal cycles depend both on the site and the time of the year. For example, in Oensingen in the late summer of 2006, the maximum HNO$_3$ concentration was clearly later in the afternoon (16:00–17:00) (Wolff et al., 2010), which is about 4 h later than in our spring period in Helsinki. In our study the HNO$_3$ concentration decreased in the afternoon, probably due to dry deposition and heterogeneous formation of nitrate, as suggested for example by Fisseha et al. (2006) for the similar cycle in Zürich. The day-to-day variation of HNO$_3$ was large in our data, which is due to variation in both sources and removal. HNO$_3$ is typically removed both by dry deposition and by below-cloud scavenging. Precipitation data have not been used for the analysis to explain the variation.

Contrary to nitric acid, the HONO amount in air was higher in winter and decreased in spring. The highest concentrations were measured in December and January. The maximum (4.3 ppb) was measured on 18 December. From the middle of February, HONO concentrations varied below 1 ppb. Between 21 April and 18 May, HONO concentration was at the lowest, staying below 0.3 ppb. In winter HONO was higher in the daytime and lower at night, having a minimum at 04:00–05:00 LT. In spring concentrations rose in the early mornings, reaching maximum at 07:00–08:00, and decreased in the afternoons (minimum at 13:00–16:00, Fig. 9). This again resembles the cycle in Zürich, Switzerland, where HONO reached maximum at about 09:00 and decreased during the day until it reached minimum at 17:00 in March 2003 (Fisseha et al., 2006).

The amount of HONO in air depends on the NO$_x$ concentration. About half of the NO$_x$ emitted from the Earth’s surface annually arises from fossil fuel combustion and the remainder from biomass burning and emissions from soil (Mosier, 2001). There are indications that HONO may be...
formed heterogeneously from NO$_2$ on ground or airborne surfaces, for instance aerosol particles, especially soot, or cloud droplets (e.g., Gutzwiller et al., 2002). Positive correlations between NO$_x$ and HONO have been found. Kurtenbach et al. (2001) made extensive investigations of emissions and heterogeneous formation of HONO in a road traffic tunnel and found that the mean HONO-to-NO$_x$ ratio was 0.008 ± 0.001. We have plotted this ratio and the lower and higher values 0.007 and 0.009 as blue lines together with our data (Fig. 10). It is interesting that when a regression line was fit through the data by forcing the offset to zero, the slope is the same as was observed by Kurtenbach et al. (2001), suggesting that also at our site traffic is a major source of HONO. There are not many data points below the line. This means that most of the time there was at least as much HONO as the Kurtenbach et al. (2001) HONO-to-NO$_x$ ratio predicts. On the other hand, a large amount of the data points are clearly above the line, even by an order of magnitude. This suggests that traffic is not the only source of HONO at our site.

3.4.2 Ammonia

As the major base in the atmosphere, ammonia plays a key role for the neutralization of acidic gases and the formation of particulate matter (Asman et al., 1998; Kirkby et al., 2011). Furthermore, the effects of ammonia deposition can lead to eutrophication and acidification of soils (Fern, 1998; Erisman et al., 2001; Butterbach-Bahl et al., 2011). The main sources of ammonia (NH$_3$) are animal waste, ammonification of humus followed by emission from soils, losses of ammonium-containing fertilizers from soils, industrial emissions, oceans, biomass burning and crops (Asman et al., 1998; Seinfeld and Pandis, 1998). About 50 to 75% of NH$_3$ from terrestrial systems is emitted from animal and crop-based agriculture, from animal excreta and synthetic fertilizer application (Mosier, 2001). In winter when land is covered by snow and the sea is frozen, it may be expected that the ammonia concentrations are low. The agriculture-related and soil-related sources are strongest in summer, which as such already leads to a seasonal cycle.

We observed a seasonal cycle that is consistent with the above. Most NH$_3$ concentrations were below detection limit in January and February. After that concentrations rose slightly, and as expected, highest peaks (1–3 ppb) were detected later in spring in April and May. Usually ammonia concentrations increase in May when the agricultural activities start (Ruoho-Airola et al., 2010).

The additional explanation for the seasonal cycle of ammonia is related to temperature in another way: at warm temperatures at daytime, ammonium nitrate particles may volatilize, and at cold temperatures the other way round, so the reaction NH$_3$+HNO$_3$(g) ↔ NH$_4$NO$_3$(s) has a temperature-dependent balance. These are also in agreement with our observations: the concentrations of ammonium and nitrate were the highest in winter.

3.4.3 SO$_2$

The anthropogenic sources of SO$_2$ are primarily the combustion of sulphur-containing fossil fuels in power and heating plants, as well as industry and traffic, especially ships (e.g., Corbett et al., 1999, 2010; Stern, 2005; Vestreng et al., 2007). There are reasons to expect a seasonal cycle in SO$_2$ concentrations: for instance heating plants operate at a higher power in winter, and in winter there is less light and thus lower concentrations of hydroxyl radical that oxidises SO$_2$ (e.g., Seinfeld and Pandis, 1998). There was some seasonality in the SO$_2$ concentrations, with higher concentrations in winter in January–February. The highest peak (18 ppb) was measured on 20 February. There was also some diurnal cycle in the SO$_2$ concentration. In winter (November–February) they were not so clear: the average ± standard deviation (median) SO$_2$ concentrations were 1.6 ± 2.1 ppb (0.75 ppb) at noon and 1.2 ± 1.4 ppb (0.66) at midnight. In spring (March–May) they were clearer: the respective concentrations were 1.3 ± 2.2 ppb (0.62 ppb) at noon, and 0.56 ± 0.50 ppb (0.37 ppb) at midnight. The most plausible explanation for this cycle is that in daytime the emissions from ships at the harbors of Helsinki are stronger than at night, and ships are well-known sources of SO$_2$ (e.g., Corbett et al., 1999, 2010). This interpretation was supported by calculating SO$_2$ diurnal cycles using data measured at an air quality measurement station at the harbor of Helsinki. There the cycle was clearer and the concentrations higher (P. Anttila, senior scientist at FMI, personal communication, 2012). A further analysis of the SO$_2$ concentration is out of the scope of the present paper.

3.4.4 HCl

The hydrochloric acid (HCl) data is the most discontinuous of the gases. Its concentrations were above the detection limit most of February and March, and the rest of the time they were mainly below it. The highest peaks, up to 0.5–2 ppb,
were measured in February and March. HCl stayed close to its detection limit in November–January. In spring it had a maximum at noon. HCl may be formed when some acid, for instance sulphuric or nitric acid, replaces chloride from sea-salt aerosols. Anthropogenic sources include coal combustion, biomass burning, and waste incineration (Knipping and Dabdub, 2003). Now in our data the highest concentrations were observed in February and March when the Baltic Sea is still frozen. This suggests that the source is not sea salt.

3.5 Nitrogen in gas and aerosol phases

The fractionation of nitrogen into gas (excluding NOx) and aerosol phases was studied by converting the concentrations into ppb units, adding them up $N_{tot} = NO_3^- + NH_4^+ + HNO_3 + HONO + NH_3$, and calculating the contribution of each of them to $N_{tot}$ (Fig. 11). The variation is large but one observation can be made: in the coldest months (January and February), most nitrogen is in the aerosol phase, and the balance turns to the gas phase when temperature increases. The median and the 95 percent range (2.5th to 97.5th percentiles) of the contribution of gas-phase nitro-

\[ y = 2.86x + 1.11 \]

\[ R^2 = 0.06 \]

Fig. 12. Relationship of nitrate and nitric acid.

\[ f(N_{gas}) = (HNO_3 + HONO + NH_3)/N_{tot} \] was 28 % (9–70 %) in January–February, and 53 % (11–81 %) in April–May. This is a general trend but large variations from this are also obvious in the time series. It can be observed that the contribution of nitrate is largest in March, which is not in any of the extremes either for temperature or solar radiation. No good explanation could be given.

Nitrate is formed when nitric acid gets into the aerosol phase. The scatter plot of nitric acid vs. nitrate (Fig. 12) shows that these species do not correlate well, which can be seen also for winter and summer in Figs. 7 and 8, respectively. A probable explanation is that the nitrate observed is more from long-range transport whereas the nitric acid is more from local sources. Also, the temperature-dependent fractionation into gas and aerosol phases affects the relationship.

3.6 Aerosol acidity and source area analysis

The aerosol data were next used for studies on the acidity of aerosols. The major acidifying and long-range transported aerosol components are sulphate, ammonium and nitrate. The ion equivalent sum of anions was calculated from

\[ SO_4^{2-} + NO_3^- + Cl^- = \frac{m(SO_4)}{48} + \frac{m(NO_3)}{62} + \frac{m(Cl)}{35.5} \]

and that of the cations from

\[ \frac{m(NH_4)}{18} + \frac{m(Na)}{23} + \frac{m(K)}{39} + \frac{m(Ca)}{20} + \frac{m(Mg)}{12.5} \]

where m stands for mass concentration. The ratio of these is $<1$ for acidic aerosols and $>1$ for basic. This calculation was done for the PM$_{10}$ particles. The ratio was calculated with and without taking calcium into account because we wanted to see the effect of soil dust. In winter the aerosol is mainly acidic but in March and April basic (Fig. 13, top panel). It is
Fig. 13. Studies on the acidity of aerosols in PM$_{10}$. Upper panel: ratio of sums of ion equivalent concentrations of cations to that of anions, including and excluding Ca$^{2+}$ concentrations; middle panel: ratio of sums of ion equivalent concentrations of cations to that of anions excluding Ca$^{2+}$ concentrations, the ratio of ammonium to non-sea salt sulphate ion equivalent concentrations, and the ratio of ammonium to the sum of non-sea salt sulphate and nitrate ion equivalent concentrations; lower panel: nitrate concentrations.

an indication of the well-known urban air quality problem in all Finnish cities in spring: road dust that is brushed from the roads during the spring road cleaning.

On the other hand, if the cation-to-anion ratio is calculated using only ammonium, non-sea salt sulphate (nss SO$_4^{2-}$), and sodium (Na$^+$), where 0.25 is the sulphate-to-sodium ratio in sea water from the average sea salt composition in Seinfeld and Pandis, (1998), and nitrate, the ratio is essentially the same as when using all of the ions (Fig. 13, middle panel), whereas when only ammonium and nss sulphate is used, the ratio is often much higher than one. This means that there is more ammonium than is needed for neutralizing sulphuric acid into ammonium sulphate. Then the rest of ammonium is neutralizing nitric acid into ammonium nitrate. However, there are long periods when the ratio is < 1.

To study the origin of the acidic aerosol, the concentration data were combined with back trajectories.

HYPLIT4 (HYbrid Single-Particle Lagrangian Integrated Trajectory) trajectories (Draxler and Hess, 1998; Heinzerling, 2004; HYPLIT-web) were calculated for an arrival height of 100 m with hourly interval, 96 h back in time. The trajectory data were used in a statistical way, presented earlier by Vakkari et al. (2011). At each time step the measured value of the chosen parameter was assigned to the grid cells ($0.5 \times 0.5^{\circ}$) along the corresponding back trajectory so that the arrival time of the trajectory was equal to the measurement time. The geometric mean of values accumulated to each grid cell was calculated. The end result is a concentration field that suggests for each cell passed by air masses on the way to the measurement site, whether it contributed to relatively high or low values monitored at the receptor site. In order to ensure the statistical significance of the result, the geometric mean was calculated only if a minimum number of trajectories, set to 10 in this work, crossed a grid cell.

The analysis was done for nss sulphate, ammonium and nitrate. It shows clearly that these ions come from continental Europe (Fig. 14). When it was done for cation-to-anion ratios that were calculated using all ions, the result is far from clear (Fig. 15). This is because calcium comes with soil dust from near-by sources. But when the calculation is done using only ammonium, sulphate, and nitrate, the difference between continental and marine air from the Norwegian Sea becomes obvious: the continental aerosol is neutralized and the marine aerosol not.

4 Summary and conclusions

The first deployment of a MARGA instrument in Finland was presented here. Its performance was analyzed by comparing it with independent methods and the data were used to demonstrate some of the possibilities it gives for atmospheric chemistry studies.

The concentrations measured by the MARGA were mainly lower than those measured with standard Teflon filters. Linear regression yielded the following MARGA vs. filter slopes: 0.72 for Cl$^-$, 0.90 for NO$_3^-$, 0.85 for SO$_4^{2-}$, 0.91 for NH$_4^+$, 0.49 for Na$^+$, 3.0 for Mg$^{2+}$, and 3.0 for Ca$^{2+}$ and 0.90 for the MARGA vs. SO$_2$ monitor. For K$^+$ there were not enough data points to calculate a statistically significant linear regression. One of the goals with online measurements
is to use them to replace filter sampling within the EMEP network. From the results of this study this cannot be recommended, at least with the MARGA setup used during the period presented here. However, part of the discrepancies were later in a following campaign proven out to be solvable by using a concentration column for the cations. This needs to be studied rigorously.

However, the method is promising and it gives very valuable information on both gas and aerosol phases at time resolutions high enough for process studies. For instance, in spring we observed clear diurnal cycles of nitric and nitrous acids that could not have been observed with some impregnated filter method.

There were clear seasonal cycles of the nitrogen containing gases: the median concentrations of HNO$_3$, HONO, and NH$_3$ were 0.09 ppb, 0.37 ppb, and 0.01 ppb in winter, respectively; and 0.15, 0.15, and 0.14 in spring, respectively. The HNO$_3$ and HONO seasonal cycles were driven by the amount of solar radiation: when there was sunlight, the HNO$_3$ concentrations were higher and the HONO concentrations were lower than in darkness. This was also the case with the diurnal cycles of the same gases in spring. Comparison with published HONO-to-NO$_x$ ratios from a road traffic tunnel suggests that at our site traffic is a major source of HONO. On the other hand, a large amount of the HONO-to-NO$_x$ ratios were clearly larger than these, even by an order of magnitude. This shows that traffic is not the only source of HONO at our site.

The seasonal cycle of ammonia was such that in the coldest months the concentrations were mainly below the detection limit and they increased with increasing temperature. This can be explained by agriculture-related and soil-related sources that are low when land is frozen and covered with snow.

The fractionation of nitrogen into gas and aerosol phases was studied by converting all concentrations into ppb units, summing then up and calculating the contribution of each of them to the sum. The gas-phase fraction of nitrogen decreased roughly with decreasing temperature, so that in the coldest period from January to February the median contribution was 28 % but in April to May 53 %. However, in all months there were large fractionation variations that temperature alone cannot explain.

The analysis of ion balances showed that in winter aerosol is mainly acidic. In spring it gets basic due to soil dust. The trajectory statistics shows that even though the measurement site is in Helsinki and in the middle of the most densely populated parts of Finland, clear differences can be seen in the degree of neutralization as a function of source area. The long-range transported aerosol is acidic when it comes from the northern sectors and neutralized by ammonia when it comes from Central and Eastern Europe.
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