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Characterization of submicron aerosol chemical composition and sources in the coastal area of Central Chile

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Abstract

Chemical characteristics and the sources of submicron particles (< 1 μm in diameter) were investigated in Valle Alegre, the coastal area of Central Chile. The chemical composition of particles was studied by using a Soot Aerosol Mass Spectrometer and Multi-Angle Absorption Photometer. Submicron particles were dominated by organics (42% of mass) and sulfate (39% of mass) while the mass fractions of ammonium, nitrate and black carbon were much smaller (13, 2 and 4% of mass, respectively). Additionally, several metals (V, Zn, Fe, Cd, Cu, K, Na and Mg) were detected in submicron particles and also some of their inorganic salts (e.g. NaCl+, MgCl2−, CaCl2−, KCl+ and KNO3−). The sources of particles were examined by using Positive Matrix Factorization (PMF). Organic aerosol (OA) was divided into five factors by using PMF; hydrocarbon-like OA (HOA), biomass burning OA (BBOA), low-volatility oxygenated OA (LV-OOA), semi-volatile OA (SV-OOA) and marine oxygenated OA (MOOA). Oxygenated factors (LV-OOA; SV-OOA and MOOA) comprised 75% of total OA with LV-OOA being the dominant factor (38% of OA). Sulfate had two major sources in Valle Alegre; ~70% of sulfate was related to anthropogenic sources through the oxidation of gas phase SO2 whereas ~24% of sulfate was associated with biogenic origin related to the oxidation of dimethyl sulfide in the marine environment. Regarding total submicron particle mass (campaign-average 9.5 μg m−3), the contribution of anthropogenic sources was at least as large as that of biogenic origin.

1. Introduction

Global Burden of Disease studies have shown that the exposure to ambient particulate matter (PM) with aerodynamic diameter < 2.5 μm (PM2.5) is among the top ten risks faced by human beings today (WHO, 2016a; 2016b). On the global level, the exposure to outdoor air pollution, mostly to PM, causes 3.3 million premature deaths per year (Lelieveld et al., 2015). The relative contributions of different emission sources vary significantly but e.g. in densely populated western countries, 33–55% of premature deaths caused by air pollution are related to anthropogenic aerosol. In addition to adverse health effects, anthropogenic particles influence on climate. Due to their complex physical and chemical properties, atmospheric aerosols still cause the largest uncertainty to the total radiative forcing estimate (IPCC, 2013).

Sources of anthropogenic pollution have mostly been investigated in urban areas with rather well-known pollution sources such as traffic, residential burning (biomass/coal), cooking and energy production (e.g. Aiken et al., 2009; He et al., 2011; Elser et al., 2016). In rural areas, the contribution of biogenic sources is typically larger but also anthropogenic sources can be more diverse and significantly different from urban areas (Kortelainen et al., 2017). In rural locations, ambient particles can be e.g. strongly influenced by industrial sources as industrial activities are often situated near small towns or in sparsely populated areas. In case of industrial sources, emissions from point sources, such as from stacks, are usually regulated and the emissions can be determined by standard methods but there are also a lot of fugitive emissions that are much more difficult to control and quantify. Ideally, industrial facilities would be located far away from habitation, in order to minimize their air quality impact, but in most cases that is unfeasible because of the need of labor and efficient transportation.
system.

Chile is one of the countries suffering from air pollution episodes (Gramsch et al., 2006, 2016). In the past, the capital of Chile, Santiago, was one of the most polluted cities in the world, but due to the successful pollution control plan, the concentrations of air pollutants have been reduced (Koutrakis et al., 2005). Nowadays submicron particle mass concentrations in Santiago are not higher than in some other large cities (e.g. Huang et al., 2011; Carbone et al., 2013; Sun et al., 2015; Gramsch et al., 2016; Tagle et al., 2018). As Chile is wealth in minerals, mostly copper, it is one of the most highly industrialized Latin American countries. Currently, Chile has serious air quality concerns in rural areas where a large number of industrial activities are situated.

The largest industrial region in Chile is the Santiago-Valparaíso area. Valle Alegre is a rural valley in Valparaíso Region with a lot of agricultural activity but there is also a large industrial area with e.g. smelters, power plants, oil refineries and cement factories nearby. The industrial site also incorporates one of the most important port activities in the region. The area has been officially recognized by the state of Chile as “Saturated Area” for PM2.5 which means that mass concentrations exceed the national air quality standard for PM2.5 (annual limit of 20 μg m⁻³) and “Latent area” for PM10 which means concentration are close to the national air quality standard (over 80% of the annual limit of 50 μg m⁻³).

The aim of this study was to investigate the chemical characteristics and sources of submicron particles in Valle Alegre by using a Soot particle Aerosol Mass Spectrometer (SP-AMS). The SP-AMS enables the detection of non-refractory aerosol species (organics, sulfate, nitrate, ammonium and chloride) and refractory material e.g. refractory black carbon (rBC) and metals. In order to examine the sources of particles in Valle Alegre, Positive Matrix Factorization (PMF) was applied to the mass spectra (MS) of organic aerosol (OA). In addition to OA, PMF was also utilized conventionally by adding sulfate fragments to the OA data matrix to examine the origin of sulfate in Valle Alegre. The specific objective of this study was to assess the impact of anthropogenic and biogenic sources to particulate pollution. Although the sources and chemistry of submicron particles have been investigated extensively around the world, until now most of the studies have been carried out in the northern hemisphere. In the southern hemisphere, there are still areas that are lacking information related to local air quality and climate change that can be different from the northern hemisphere. To our knowledge, this is the most extensive study characterizing the chemical properties of submicron particles in the rural area of Central Chile.

2. Experimental methods

2.1. Measurement site and period

Measurement site was located in Valle Alegre in Central Chile (32°48’28.24”S, 71°26’12.85”W; Fig. 1a). Valle Alegre is a rural valley ~7 km inland from the bay of Quintero enclosed by the coastal mountain range at the east side. It is situated ~12 kilometers away from one of the largest industrial complex in Chile, Ventanas. The industrial park comprises a 5-terminal port and 14 industrial facilities: four coal-fired power plants, one copper sulfate smelter, one copper refinery, three gas and diesel storage facilities, two petrochemical factories, and two industries processing asphalt and cement. The industrial sites are mostly located north-west from the station, at the Pacific coast. Five largest industrial sites in the region are shown in Fig. 1a.

The measurements were carried out from 9 to 28 January 2015 (austral summer). Instruments were placed in a container at a corner of a private recreation area with some outdoor activities (e.g. pool, barbeque area). However, as the recreational area was still partially under construction, there was not much recreational or building activity during the measurement campaign. The other side on the container was bordered by a small forest. There was also some agricultural activity near the site as well as cattle and horses.

2.2. Instruments

2.2.1. Soot Particle Aerosol Mass Spectrometer

The chemical composition of submicron particles was measured by using a Soot Particle Aerosol Mass Spectrometer (Aerodyne Research Inc., Billerica, USA; Onasch et al., 2012). The SP-AMS is a high resolution Time-of-Flight aerosol mass spectrometer (HR-ToF-AMS; DeCarlo et al., 2006) that has an additional laser vaporizer. The SP-AMS is able to measure non-refractory species (organics, sulfate, nitrate, ammonium and chloride) but also rBC and some metals due to the laser vaporizer (ND:YAG 1064 nm). Typically, the SP-AMS operates in two modes, MS mode, in which the chemical composition of all particles in the size range of the instrument (~50–800 nm) is measured, and efficient Particle-time-of-flight (ePToF) mode that allows the determination of chemical species as a function of particle size. In the MS mode, the chopper is alternated between open and closed positions of which open allows the particle beam to pass through and closed blocks it. In the ePToF mode, the chopper is continuously rotating (~140 Hz) and as it has several slits that let the particle beam pass through. The SP-AMS data was recorded with one minute time-resolution of which half of the time the SP-AMS measured in the MS and half in the ePToF mode. The SP-AMS was equipped with both tungsten vaporizer (at 600 °C) and laser vaporizer. Typically, the SP-AMS operates by switching the laser on and off, however, during this campaign there was a malfunction with the laser power. Therefore, the laser was on continuously from 9 to 18 January after which the laser was off for the rest of the campaign. The SP-AMS had a cyclone (SCC 1.828, BG1 Inc., US) in the sample line outside the container in order to remove particles larger than 1 μm in diameter from the sample flow.

2.2.2. Multi Angle Absorption Photometer

A Multi-Angle Absorption Photometer (MAAP, Model 5012, Thermo Fisher Scientific, Waltham, USA) was used to determine the concentration of black carbon (BC). In the MAAP, particles are deposited on a quartz fiber filter and a continuous laser beam (670 nm) is passed perpendicular through the filter. The decrease in the transmitted light is due to two factors; absorption by the particles on the filter matrix. Absorbance of the particles is determined by subtracting scattering, measured by the additional detectors placed at selected angles, from the decrease in the light intensity (Petzold and Schönlinner, 2004). In this study, a default mass-specific aerosol absorption coefficient (σₐₐ₅₅) of 6.6 μm² g⁻¹ was used for the MAAP results. The data averaging time was one minute and the flow rate was 11 lpm. The MAAP had an inlet with the 50% cut-off at 1 μm placed outside the container. The comparison between rBC from the SP-AMS and BC from the MAAP is presented in Supplementary material (Fig. S1). The time series of rBC and BC followed similar trend but overall their correlation was rather poor. BC results presented in this article are mostly obtained from the MAAP. Only for the size-distribution of BC the results are taken from the SP-AMS but in that case the corresponding species is referred as rBC.

2.2.3. Air quality measurements, meteorology and backward trajectories

An air quality monitoring station operated by local industrial companies was situated next to the measurement container. Hourly averaged PM₁₀ concentration was measured with Continuous Ambient Particulate Monitor (FH62C14, Thermo Fisher Scientific, Waltham, USA), nitrogen oxides with NO-NO₂-NOx Analyzer (42i, Thermo Fisher Scientific, Waltham, USA), sulfur dioxide (SO₂) with SO₂ Analyzer (43i, Thermo Fisher Scientific, Waltham, USA) and ozone (O₃) with Ozone Analyzer (49i, Thermo Fisher Scientific, Waltham, USA). Unfortunately, there were neither PM2.5 nor PM1 measurement at the air quality monitoring station. Meteorological parameters (temperature, wind direction and wind speed) were monitored with Met One Sensors (Met One Instruments Inc, Grants Pass, USA), however, due to the malfunctioning of the temperature sensor in Valle Alegre, temperature was measured with a temperature sensor in the container (49i, Waltham, USA).

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obtained from the air quality monitoring station in Quintero (∼10 km from Valle Alegre station).

72-h air mass backward trajectories were calculated in order to investigate the origin and transport history of air masses that arrived in Valle Alegre. Trajectories were calculated by using the NOAA Hybrid Single-Particle Lagrangian Integrated Trajectory model (Draxler and Rolph, 2003; Stein et al., 2015) and global meteorological data from the Global Data Assimilation System (GDAS) archive with an arrival height of 500 m above the ground. Trajectories were calculated every 3 h for the period of 9-28 January 2015.

2.3. Laboratory measurements for reference mass spectra

Reference mass spectra for methane sulfonic acid (MSA) and ammonium sulfate were obtained by laboratory tests. MSA and ammonium sulfate standards were dissolved in Milli-Q water, and particles were produced by an aerosol generator (Model 3076, TSI Inc., Shoreview, USA). 300 nm particles were selected with the Differential Mobility Analyzer (DMA; Model 3082, TSI Inc., Shoreview, USA) and directed to the SP-AMS.

2.4. Data analysis

2.4.1. Relative ionization efficiency and collection efficiency for the SP-AMS

The SP-AMS data was analyzed with Igor 6.3 software using SQRL 1.56D and PIKA 1.15D. Elemental analysis of OA was accomplished with the method developed originally by Aiken et al. (2007, 2008) but extended to a wider range of OA species by Canagaratna et al. (2015). The ionization efficiency (IE) of nitrate was determined by calibrating the instrument by using monodisperse ammonium nitrate particles. Default relative ionization efficiencies (RIEs) were used for organics and chloride whereas for sulfate and ammonium the RIEs of 0.86 and 3.1 were used, respectively, based on the ammonium sulfate and ammonium nitrate calibrations carried out in the laboratory. For rBC, the RIE of 0.1 was obtained from the calibration with Regal black. For metals, the RIEs were taken from the study of Carbone et al. (2015). Since the exact RIE was not available for Cd, K and Mg, a default RIE of 1 was used for them. By using the values given in Carbone et al. (2015), it was assumed that metals were in the same particles with rBC. However, that may not always be true, especially in Valle Alegre where the sources of metals and rBC can be quite different. Mg and Zn were determined as their isotopes at m/z 25 (25Mg+) and 68 (68Zn+), respectively, as the main isotopes were overlapped by SO++ (at m/z 24) and SO2+ (at m/z 64) fragments. Size distribution (ePToF) data was analyzed only with unit mass resolution (UMR) because the tool for the high resolution ePToF data was not available.

The collection efficiency (CE) was calculated according to Middlebrook et al. (2012). Most of the time the CE was 0.45, which is the lower limit of the CE based on the equation of Middlebrook et al. (2012), but there were several events when the CE was as large as 0.8 (see Fig. S2). During those periods particles were acidic due to the larger amount of sulfate compared to that of ammonium. Relative humidity was not measured continuously from the sample air, and therefore, that could not be taken into account when calculating the CE. It should be noted that for the SP-AMS the CE can be different from the standard AMS due to the additional laser vaporizer. The factors affecting the CE in the SP-AMS have been discussed e.g. in Onasch et al. (2012), Willis et al. (2014) and Ahern et al. (2016). In this study the impact of the laser vaporizer was not taken into consideration when calculating the CE because the amount of refractory material was rather low, and therefore, most of the chemical species were estimated to be

Fig. 1. The location of the measurement site in Valle Alegre, Chile (a), the average wind direction pattern at the site (b), and the diurnal variation of meteorological parameters (c). Temperature and wind speed are presented as average ± stdv while in wind direction plot individual (hourly) values are shown. Temperature in (c) was obtained from the air quality monitoring station in Quintero ~10 km from the Valle Alegre station.
vaporized at the tungsten vaporizer. The uncertainty of the CE is discussed in supplemental material.

2.4.2. Positive matrix factorization

Sources of the particulate material were investigated by using Positive Matrix Factorization (PMF Tool 2.08D; Paatero and Tapper, 1994; Ulbrich et al., 2009). PMF was applied to the mass spectra of OA measured by the SP-AMS. PMF solutions were calculated up to eight factors (Fig. S3). It was found that a minimum of five factors were needed to explain the variation of OA in Valle Alegre. Based on their MS (Fig. S4a), these factors were identified as hydrocarbon-like OA (HOA), biomass burning OA (BBOA), semi-volatile oxygenated OA (SV-OOA), low-volatility oxygenated OA (LV-OOA) and marine oxygenated OA (MOOA). Compared to five factor solution, in four factor solution the typical fragments of BBOA and MOOA were mixed into the other factors whereas in the case of six, seven and eight factors HOA, LV-OOA and SV-OOA were split between several factors. PMF solutions with more than five factors did not give any additional information compared to five factor solution, and therefore, five factor solution was chosen to be presented and discussed in this article for OA (hereafter called OA-solution). Five factor solution was tested for the rotational freedom by varying speak, and also multiple seeds were calculated to confirm that the solution presented here was not a unique solution but could be achieved with various seeds.

In order to examine the sources of sulfate in Valle Alegre, PMF was also employed to the combined OA and sulfate MS (hereafter called OA + sulfate -solution). Sulfate fragments included in PMF were SO\textsuperscript{2−} (at m/z 48), HSO\textsuperscript{−} (at m/z 49), SO\textsubscript{2}O\textsuperscript{−} (at m/z 64), HSO\textsubscript{3}− (at m/z 65), H\textsubscript{2}SO\textsubscript{3}− (at m/z 66), SO\textsubscript{3}− (at m/z 80), HSO\textsubscript{4}− (at m/z 81), H\textsubscript{2}SO\textsubscript{4}− (at m/z 82) and H\textsubscript{2}SO\textsubscript{5}− (at m/z 98). For both OA and sulfate, CE and RIE of 1 were used. Errors for the sulfate fragments were calculated similar to the fragments of OA. CO\textsubscript{2} and sulfate fragments were downweighed. This approach has also been used in e.g. Kortelainen et al. (2017), Tiitta et al. (2016) and Hao et al. (2014), but instead of sulfate, they included nitrate fragments in the matrix of OA. Due to the small concentration of sulfate, nitrate was not included in PMF in this study. In addition to selected inorganic species, also a full mass spectrum encompassing all organic and inorganic components has been analyzed with PMF but that was performed in UMR mode (McGuire et al., 2014).

Similar to the PMF solution for OA, five factors, HOA, BBOA, SV-OOA, LV-OOA and MOOA, were obtained for OA + sulfate -solution (Fig. S4b). However, in OA + sulfate -solution there was an additional factor, called here as Sulfate factor, that consisted mostly of sulfate. Sulfate fragments were also found in MOOA and in smaller amount in LV-OOA and SV-OOA but in HOA and BBOA the amount of sulfate was very small.

PMF solutions for OA and OA + sulfate are compared for the MS in Fig. S4c and for the time series in Fig. S5a-b. Only organic fragments were included in the comparison. For HOA and BBOA, the MS and time series correlated well (R\textsuperscript{2} = 0.939−0.999) except that for BBOA the fragments CO\textsuperscript{2−} (at m/z 28) and CO\textsubscript{2}O\textsuperscript{−} (at m/z 44) were clearly smaller in the OA + sulfate -solution than in the OA-solution. Accordingly, for HOA the elemental composition of OA was very similar in the OA and OA + sulfate -solutions whereas for BBOA the oxygen to carbon ratio (O:C) was larger and the hydrogen to carbon ratio (H:C) smaller in the OA-solution than in the OA + sulfate -solution. For LV-OOA, the correlation between the time series of the OA and OA + sulfate -solutions was fairly good (R\textsuperscript{2} = 0.966), however, most of the time LV-OOA concentrations were larger in the OA-solution than in the OA + sulfate -solution. The oxidation state was clearly lower in the OA-solution than in the OA + sulfate -solution for LV-OOA. For MOOA, the concentrations and oxidation states were rather similar for the OA and OA + sulfate -solutions. SV-OOA had the greatest difference between the OA and OA + sulfate -solutions. The correlation between the time series of the OA and OA + sulfate -solutions was only modest (R\textsuperscript{2} = 0.725). Regarding the MS, the contributions of two main fragments, CO\textsuperscript{2−} and CO\textsubscript{2}O\textsuperscript{−}, were much larger, and accordingly SV-OOA was more oxidized, in the OA + sulfate -solution than in the OA-solution.

In terms of mass fractions (Fig. S5c), BBOA was equal in the OA- and OA + sulfate -solutions whereas SV-OOA was larger and HOA, LV-OOA and MOOA were smaller in the OA + sulfate -solution. The additional sixth factor (Sulfate) contributed 9% to total OA in the OA + sulfate -solution. Sulfate factor had most similar O:C and H:C to BBOA. The PMF solution for OA + sulfate is utilized in this article only to evaluate the sources of sulfate in Valle Alegre (Section 3.3.2.). The discussion on the sources of OA (Section 3.3.1.) is based on the OA-solution.

3. Results and discussion

3.1. Meteorological parameters, air quality monitoring data and air mass origin

Daily-averaged meteorological conditions were rather stable during the measurement campaign in Valle Alegre. However, there was a clear diurnal cycle for wind speed and direction (Fig. 1c) dominated by the topography of the area as the measurement site was located in the valley (Fig. 1a). At night-time and morning, the dominant wind direction was northeast, typically from the bottom of the valley, and wind speed was low (< 0.5 m/s). Wind direction turned to south-west (the sea) around midday and also wind speed increased at the same time (hourly average maximum ~1.5 m/s). Temperature was on average (± stdv) 16.9 (± 1.7) °C during the campaign being around 3 °C higher in daytime than during night and early morning.

For the inorganic gases, the average (± stdv) concentrations were 2.6 (± 3.3), 4.5 (± 3.8), 20 (± 9.3) and 5.4 (± 6.1) ppb for NO\textsubscript{2}, NO\textsubscript{3}, O\textsubscript{3} and SO\textsubscript{2}, respectively. The campaign-average (± stdv) PM\textsubscript{10}\textsubscript{−} concentration was 40 (± 16) μg m\textsuperscript{−3}. Of those species, only ozone displayed a clear diurnal trend (Fig. S6b). The concentration of ozone followed similar pattern to wind direction and speed increasing in the morning around 9 a.m. and having the maximum between 4 and 7 p.m. However, different from wind speed, ozone concentration remained at the elevated level later in the night while wind speed dropped at night-time approximately at 10 p.m.

According to the backward trajectories, air masses arrived in Valle Alegre mostly from south or south-west (Fig. S7). Three days before arriving in Valle Alegre, air masses were over the Pacific Ocean and from there they travelled to Valle Alegre by following loosely the Chilean coastline from south to north. Occasionally air masses also made a loop over the continental Chile before reaching Valle Alegre. That happened mostly during the night and early morning. For example on 20 January 2015, air masses arrived in Valle Alegre from south-east from midnight to 9 a.m (Fig. S7c), after which the air mass trajectories turned to the sea. In general, the origin of air masses was rather similar during the whole measurement period. Small differences observed in the backward trajectories were not seen in the results, probably because the concentrations of air pollutants were mostly governed by local meteorology. Therefore, the trajectories are not discussed in this paper in the following sections.

3.2. Chemical composition of submicron particles

The campaign-average (± stdv) concentrations of organics, sulfate, nitrate, ammonium, chloride and BC were 4.9 (± 3.9), 3.3 (± 2.5), 0.14 (± 0.22), 0.99 (± 0.50), 0.06 (± 0.06) and 0.34 (± 0.35) μg m\textsuperscript{−3}, respectively, measured by the SP-AMS and MAAP (10-min averages). The concentrations of chemical species varied noticeably during the campaign. For organics and nitrate, the concentration peaks were rather short in time, typically from an hour to few hours, whereas for sulfate and ammonium, the concentrations varied in a longer time-scale, from several hours to almost a day (Fig. 2a). This
suggested that sulfate and ammonium were mostly regionally dispersed whereas organics and nitrate originated also from the local sources such as vehicles or small-scale combustion. For BC, the concentration was rather stable throughout the campaign but there were some periods when BC was elevated. Regarding the total mass concentration of submicron particles, calculated by summing up all the species from the SP-AMS and MAAP, the campaign average concentration was 9.5 μg m⁻³. Organics and sulfate had roughly similar mass fractions (42 and 39% on average, respectively) whereas the campaign-average contributions of ammonium, BC, nitrate and chloride to total mass were 13, 3.6, 1.6 and 0.7%, respectively (Fig. 2b and Fig. S8a).

The submicron mass concentration was smaller in Valle Alegre (9.5 μg m⁻³) than that measured in rural (∼13–20 μg m⁻³) or urban (∼20–50 μg m⁻³) locations in Santiago Metropolitan Region (Carbone et al., 2013; Tagle et al., 2018). The most notable difference between Valle Alegre and Santiago was, however, the composition of submicron particles. In Valle Alegre, the mass fraction of sulfate (39%) was much larger than that in Santiago (< 10%) whereas the fraction of organics was larger in Santiago (∼55% in Santiago). The small fraction of sulfate in Santiago can be explained by the lack of industrial sites and ports in Santiago area that were abundant in proximity to Valle Alegre. A large fraction of sulfate in submicron particles has been detected earlier e.g. at the urban sites in Pittsburgh, New York and Houston, USA (Zhang et al., 2007). By contrast, the mass fraction of nitrate was much larger in Santiago (∼8-30%) than in Valle Alegre (∼2%) due to the larger amount of traffic in Santiago. In general, the mass concentration of submicron particles in Valle Alegre was at the same level with the mass concentrations measured typically in urban areas but larger than the concentrations measured at rural locations (Zhang et al., 2007; Crippa et al., 2014).

Regarding the diurnal variations, none of the measured chemical species had a distinctive diurnal trend (Fig. S9). The clearest diurnal pattern was observed for chloride that had smaller concentrations at daytime from ∼10 a.m. to 9 p.m. than in the other time of the day. Also the concentrations of organics and BC were slightly larger at night-time than daytime, however, the variation in concentrations was also greater during the night shown by the large standard deviations. Sulfate and ammonium concentrations followed loosely wind pattern (Fig. 1) being larger from midnight to midday when wind was from the valley and its speed was low resulting in the accumulation of pollutants in the boundary layer. After midday, the concentrations decreased as wind started to blow from the sea and the mixing of atmospheric species increased. For nitrate, there was a peak in concentrations between 7 and 9 a.m. that could be due to morning traffic, however, at that time of the day nitrate concentrations varied a lot shown by the large standard deviations.

In terms of mass size distributions, most of the chemical species had an accumulation mode with a maximum at ∼350 nm (Fig. 2c). Ammonium peaked at slightly smaller particle size than sulfate. The size distributions of organics and nitrate deviated from those of ammonium...
and sulfate. Organics had slightly wider accumulation mode than ammonium and sulfate having relatively more mass at < 350 nm whereas nitrate had an additional mode at ~100 nm. The time evolution of the size distributions (Fig. S10) revealed that organics and nitrate were found in small particles only in short time periods that is in agreement with the earlier speculation that nitrate and organics originated also from local sources, probably from traffic or small-scale burning. For BC, the mass size distribution was calculated by using unit mass resolution m/z 36 (that consists mostly of C3H3+ fragment) as a surrogate for rBC in the SP-AMS. The mass size distribution of rBC was similar to that of nitrate except that the accumulation mode extended to larger particle sizes for rBC. That indicates that a larger contribution of rBC was transported to site from distant sources supported by a more steady concentration of rBC compared to that of nitrate.

3.2.1. Organics

The average mass spectrum for organics is shown in Fig. S11. The dominant organic fragments were CO+ (at m/z 28) and CO2+ (at m/z 44) fragments indicating that organics were highly oxidized. In general, oxygenated fragments (CnH2n+Oz) constituted 54% of organics of which slightly more than half contained the fragments with one oxygen atom (e.g. CO+) whereas the other portion included the fragments with more than one oxygen atom (mostly CO2+). The rest of organics consisted of hydrocarbon fragments (CnH2n+ fragment) that had largest signal for CH3+ (at m/z 15), C2H3+ (at m/z 27), C3H5+ (at m/z 39) and C4H9+ (at m/z 41). Regarding the elemental composition of organics, the average (±stdev) H:C and O:C were 1.5 (±0.14) and 0.68 (±0.19), respectively, resulting in the average (±stdev) ratio of 2.0 (±0.25) for the organic matter to organic carbon ratio (OM:OC). O:C and OM:OC were slightly larger from 3 p.m. to around 9 p.m., and H:C accordingly smaller, than at the other times of the day (Fig. S12). The larger O:C and OM:OC resulted in the average (±stdev) ratio of 2.0 (±0.25) for (±stdev) H:C and O:C were 1.5 (±0.14) and 0.68 (±0.19), respectively, resulting in the average (±stdev) ratio of 2.0 (±0.25) for the organic matter to organic carbon ratio (OM:OC). O:C and OM:OC were slightly larger from 3 p.m. to around 9 p.m., and H:C accordingly smaller, than at the other times of the day (Fig. S12). The larger O:C and OM:OC accordingly.

3.2.2. Metals and inorganic salts

V, Zn, Fe, Cd, Cu, K, Na and Mg were detected in the particles in Valle Alegre. The concentrations of metals were calculated for the period when the laser vaporizer was on (9–18 January 2015). The

![Figure 3](image.png)

**Fig. 3.** Time series of metals detected in submicron particles using the SP-AMS; 10-min average concentrations for the period when the laser was on.
general assumption was that the metals were vaporized only with the laser, however, some metals (Zn, Cu, Na, K and Mg) were also observed during the period when the laser was off. Those metals were probably associated with some inorganics salts and/or vaporized with the tungsten vaporizer (Salcedo et al., 2012).

The average (± stdev) concentrations for V, Zn, Fe, Cd, Cu, K, Na and Mg were 0.65 (± 0.80), 1.6 (± 2.3), 0.48 (± 2.3), 0.10 (± 0.15), 0.53 (± 0.76), 70 (± 103), 22 (± 20) and 2.6 (± 2.2) ng m⁻³, respectively. Regarding the time evolution of metals, Fe had very short-lived concentration peaks whereas the concentrations of other metals changed more slowly (Fig. 3). Zn, Cd and Cu had some similarities in their time series but clear correlations between metals were not found. The largest concentration peak for Zn, Cd and Cu was observed on 10 January 2015 from midday to ~2 p.m. During that time also organics and sulfate concentration increased, but not ammonium (Fig. 2a), indicating that aerosol was acidic. Also gas phase NO and SO₂ concentrations were slightly elevated (Fig. S6a). It can be speculated that the increase in metal concentrations during that period can be due to the emissions from the industrial sites, however, that could not be verified since wind direction was between west and south-west while industrial sites were located north-west from the measurement site.

Diurnal variation was observed only for V and Mg (Fig. S14). They had slightly smaller concentrations in the afternoon, similar to sulfate (Fig. S9). Accordingly, the concentration of Mg somewhat correlated with that of sulfate (R² = 0.45). Mg and Ca were found as chloride salts (MgCl₂⁻ at m/z 94 and CaCl₂⁺ at m/z 110) but it was not possible to determine Ca itself due to the overlapping argon peak at m/z 40. MgCl₂⁻ and CaCl₂⁺ followed the time trend of Ca (Fig. S15) suggesting that their origin could be sea water as also Cd is found in sea water (Boyle et al., 1976). However, the time series of Na, chloride and NaCl⁺ (at m/z 58) were different from that of MgCl₂⁻, CaCl₂⁺ and Cd (Fig. S15) indicating that the source was probably other than sea-water.

K had a large concentration peak on 10–11 January 2015 from 10:30 p.m. to around 1 a.m. (Fig. 3). At the same time also PM₁₀ (Fig. 3). Zn, Cd and Cu had some similarities in their time series but clear correlations between metals were not found. The largest concentration peak for Zn, Cd and Cu was observed on 10 January 2015 from midday to ~2 p.m. During that time also organics and sulfate concentration increased, but not ammonium (Fig. 2a), indicating that aerosol was acidic. Also gas phase NO and SO₂ concentrations were slightly elevated (Fig. S6a). It can be speculated that the increase in metal concentrations during that period can be due to the emissions from the industrial sites, however, that could not be verified since wind direction was between west and south-west while industrial sites were located north-west from the measurement site.

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The sources and atmospheric processing of OA in Valle Alegre were investigated by using PMF. PMF for OA resolved different components than levoglucosan and BBOA (Saarikoski et al., 2012). The largest concentration peak for Zn, Cd and Cu was observed on 10 January 2015 from midday to ~2 p.m. During that time also organics and sulfate concentration increased, but not ammonium (Fig. 2a), indicating that aerosol was acidic. Also gas phase NO and SO₂ concentrations were slightly elevated (Fig. S6a). It can be speculated that the increase in metal concentrations during that period can be due to the emissions from the industrial sites, however, that could not be verified since wind direction was between west and south-west while industrial sites were located north-west from the measurement site.

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K had a large concentration peak on 10–11 January 2015 from 10:30 p.m. to around 1 a.m. (Fig. 3). At the same time also PM₁₀ (Fig. S6a), organics and BC (Fig. 2a), and KCl⁺ (at m/z 74; Fig. S15) had elevated concentrations. KCl⁺ followed more closely the time trend of chloride than that of K. Elevated concentration on 10 January were likely to be caused by biomass burning because KCl, a species that is found in fresh emissions (Zhang et al., 2005). In terms of vehicle emissions, the MS of HOA resembled to the standard MS profile given in Ng et al. (2011); R² = 0.917) but it was even more similar to the MS measured previously in Mexico City (Aiken et al., 2009; R² = 0.965), Helsinki (Carbone et al., 2014; R² = 0.946) and Xi’an and Beijing (Elser et al., 2016; R² = 0.931) (Fig. S16). HOA is typically associated with traffic emissions (Zhang et al., 2005). In terms of vehicle emissions, the MS of HOA in Valle Alegre was more similar to that of a gasoline vehicle (Timonen et al., 2017) than that of a diesel bus (Saarikoski et al., 2017). This finding is in agreement with the vehicle fleet in Chile that consist mostly (~80%) of gasoline cars. HOA was very lightly oxygenated having high H:C (1.95) and very low O:C (0.09). Compared to the previous studies, HOA was less oxidized in Valle Alegre than e.g. in Po Valley (O:C 0.20; Saarikoski et al., 2012) or Mexico City (O:C 0.16; Aiken et al., 2009) but close to that measured in the Pearl River Delta region in China (O:C 0.11; He et al., 2011). In general, the concentration of HOA was rather small most of the time but there were some periods, typically from an hour to few hours, when HOA concentration increased and HOA dominated OA (Fig. 4b and SBb). Regarding the diurnal variation, HOA displayed slightly larger concentrations during the night than at daytime, however, also the variation in HOA concentrations was large during the night. Even though HOA time trend somewhat followed to that of BC, HOA did not correlate with BC, or nitrogen oxides (NO, NO₂, NO₃) suggesting that there could be some additional sources for HOA besides vehicles.

The MS of BBOA had a large signal for CHO⁺ (at m/z 29) and C₂H₄O⁺ (at m/z 27) but also the oxygenated fragments C₃H₄O₂⁺ (at m/z 60) and C₄H₄O₂⁺ (at m/z 73), that have been earlier associated with biomass burning (Alfarra et al., 2007), were elevated. Oxygenated fragments C₃H₄O₂⁺ and C₄H₄O₂⁺ contributed 2.9 and 1.2%, to the total signal of BBOA, respectively. BBOA was clearly more oxidized than HOA having larger O:C (0.36) and smaller H:C (1.78) than HOA. BBOA was more oxidized in Valle Alegre than in Po Valley (O:C 0.23; Saarikoski et al., 2012), Mexico City (O:C 0.30; Aiken et al., 2009) or the Pearl River Delta region (O:C 0.32; He et al., 2011). Regarding the time series, BBOA concentration was elevated during four periods. The largest BBOA concentration was measured on 23 January after midnight but BBOA was also elevated on 10–11 January at the same time with K as discussed earlier in Section 3.2.2. In general, BBOA and K had some similarities in their time trends with the correlation coefficient R² = 0.47. BBOA correlated more clearly with BC than HOA (R² = 0.47) suggesting that the observed BC concentrations were probably partly associated with biomass combustion. BBOA had also a moderate correlation with chloride (R² = 0.43). In terms of diurnal variation BBOA had elevated concentrations at night-time, with a large variation in concentrations, similar to HOA.

SV-OOA had most pronounced signal for CHO⁺ (at m/z 43) followed by C₂H₄O⁺ (at m/z 27), C₃H₄O₂⁺ (at m/z 39) and C₄H₄O₂⁺ (at m/z 41). SV-OOA was slightly more oxygenated than BBOA with O:C and H:C ratios of 0.37 and 1.62, respectively. Regarding the diurnal trend, SV-OOA was slightly elevated in the morning (maximum at ~8–12 a.m.) after which it decrease towards the evening having minimum at ~7–10 p.m. The diurnal trend of SV-OOA somewhat agreed to that of nitrate (Fig. S9), however, nitrate concentration was elevated much shorter time period than SV-OOA concentration. Different from the time series of HOA and BBOA that had several short-term peaks, SV-OOA concentration was steadier throughout the campaign. SV-OOA had only one short-term maximum on 23 January after midnight together with BBOA. On 23 January SV-OOA seemed to be related to biomass burning. Previously, it has been suggested that biomass burning can produce OOA-type aerosol indicated by a stronger correlation between levoglucosan (marker compound for biomass burning) and OOA components than levoglucosan and BBOA (Saarikoski et al., 2012).

The largest fraction (38%) of OA consisted of LV-OOA. LV-OOA was clearly the most oxygenated factor having O:C and H:C of 1.19 and 1.22, respectively, and its MS dominated by CO₂⁺ (at m/z 28) and CO⁺ (m/z 29) fragments. LV-OOA had very flat diurnal trend with no
dependence on the time of the day but the LV-OOA concentrations were clearly larger at the beginning of the campaign than at the end. Also LV-OOA seemed to be somewhat associated with biomass burning as LV-OOA correlated stronger with K than BBOA ($R^2 = 0.51$). By looking at the time series of K, it was noticed that the largest K peaks were captured by BBOA whereas the general time-evolution of K was followed more closely by LV-OOA. As already mentioned, OOA can originate from biomass burning, however, it is also possible that the correlation of LV-OOA and K was related to the transport of regional/long-range transported pollutants.

On average, MOOA made 21% of total OA. In terms of diurnal variation, MOOA was clearly larger in the evening, which suggested it to be marine-related as wind blew from sea at that time of the day (Fig. S17). The maximum of MOOA (at ~7 p.m.) was probably related to the SOA formation peaking at the early evening as also the oxidants ($O_3 + NO_2$; Fig. S6b). The MS of MOOA was similar to that of LV-OOA having the largest signal for CO$_2^+$ (at m/z 44) and CO$^+$ (at m/z 28) and displaying the second largest O:C ratio (1.03) after LV-OOA. MOOA had a clear signal for the fragments CH$_2$SO$_2^+$ (at m/z 78) and CH$_3$SO$_2^+$ (at m/z 79) with their fractions in MOOA being 0.32 and 0.94%, respectively. Also CH$_2$SO$^+$ (at m/z 62) and CH$_3$SO$^+$ (at m/z 63) were present in the MS of MOOA but with smaller amounts (< 0.2%). As already mentioned these fragments are typical for MSA that is primarily derived from the oxidation of DMS in marine environment, and therefore, can be considered as a tracer for marine SOA. Also sulfur-containing organic fragments without oxygen atoms CH$_2$S$^+$ (at m/z 45) and CH$_3$S$^+$ (at m/z 47) were associated with MOOA having the contributions of 0.26 and 0.05%, respectively.

Sulfur-containing organic fragments found in MOOA were compared to the reference MS of MSA obtained in the laboratory tests (Fig. S18). The pattern of CH$_3$SO$^+$ (at m/z 62), CH$_2$SO$^+$ (at m/z 63), CH$_2$SO$_2^+$ (at m/z 78) and CH$_3$SO$_2^+$ (at m/z 79) was very similar to MOOA and MSA standard whereas the relative contributions of CH$^+$ (at m/z 45) and CH$_4$SO$_3^+$ (at m/z 96) were larger for MSA than for MOOA. One reason for the difference can be that CH$_2$S$^+$ and CH$_3$SO$_2^+$ are difficult to separate from overlapping peaks at same m/z when they are present in small quantities (in MOOA). Based on the ratio of CH$_3$SO$_2^+$ to the total organic signal in the reference MS of MSA (0.11), the contribution of MSA to MOOA was estimated to be 8.5%.

Due to the presence of sulfur-containing fragments in MOOA, S:C was several times larger for MOOA ($6.7 \times 10^{-3}$) than that for the other PMF factors ($0.24$–$1.7 \times 10^{-3}$). Also N:C was largest for MOOA ($1.0 \times 10^{-2}$). That is in line with the previous finding that biogenic amines are an important source of marine secondary aerosol (Facchini et al., 2008). In addition to MOOA, nitrogen-containing fragments were observed in BBOA and SV-OOA.

### 3.3.2. Source apportionment for sulfate

The sources of sulfate in Valle Alegre were investigated by adding sulfate fragments to the data matrix analyzed with PMF. PMF for OA + sulfate resulted in six factors of which five factors were similar to those without the sulfate fragments whereas sixth factor was dominated by sulfate (called Sulfate factor). The comparison between OA and OA + sulfate factors was presented earlier in Section 2.4.2. (Experimental methods).

The distribution of sulfate between the PMF factors is shown in Fig. 4. PMF factors for organic aerosol. Mass spectra (a), time series with tracers (a) and diurnal trends (median ± 25/75%) (c). CS species in (a) refer both to C$_m$H$_n$O$_z$S$^+$ and C$_m$H$_n$S$^+$ fragments.
can be considered to be the only atmospheric source of MSA, MSA can
salt sulfate (nss-sulfate) in order to distinguish it from sulfate originated
HSO+ (31%) and smallest to H2SO3
the opposite was detected to MOOA having the largest contribution to
eral. The lack of correlation between SO2 and sulfate can be due to the
evolution of Sulfate factor followed that of SO2 at times (Fig. S19a) but
also partly associated with LV-OOA (11%), and SO+ and SO2
there was no clear correlation between Sulfate factor and SO2 in gen-
smelter and coal-
clearly attributed to speci
measurement site. The maximum for sulfate in MOOA in the evening
(Alegre. On average, 70% of sulfate was related to Sulfate factor that was
assumed to be related to the oxidation of anthropogenic SO2. The time
evolution of Sulfate factor followed that of SO2 at times (Fig. S19a) but
there was no clear correlation between Sulfate factor and SO2 in general.
The lack of correlation between SO2 and sulfate can be due to the
fact that SO2 is a primary component while sulfate is formed from the
oxidation of SO2. Therefore, their concentration ratio depends on the
oxidation time of SO2 as well as the removal efficiencies for both SO2
and sulfate. Sulfate factor, however, somewhat correlated with the
acidity of particles (Fig S19b).

The second largest fraction of sulfate was associated with marine-
related OA (MOOA; 24%). Sulfate in MOOA was assumed to be related
to the oxidation of DMS in the marine environment because, in addition
to MSA, the oxidation of DMS produces SO2 which is oxidized to sulfate
in the atmosphere. A small portion of sulfate was also associated with
LV-OOA and SV-OOA (4 and 2%, respectively) whereas HOA and BBOA
contained only a negligible fraction of total sulfate (< 1%; not shown in
Fig. 5a).

The origin of sulfate depended clearly on the time of the day
(Fig. 5b). From midnight until ~ 4 p.m. the largest fraction of sulfate
was attributed to Sulfate factor, its contribution being especially large
between 6 a.m. and 4 p.m., whereas in the evening a greater portion of
sulfate was associated with MOOA. The diurnal trend for sulfate in
Sulfate factor followed that of SV-OOA (Fig. 4c) suggesting that they
were both products of local secondary aerosol formation. In the Valle
Alegre area, the major industrial sources of SO2, the copper sulfate
smelter and coal-fired power plants, are located ~ 12 km from the
measurement site. The maximum for sulfate in MOOA in the evening
coincided with the maximum concentration of oxidants (56b).

Regarding different sulfate fragments, none of the fragments
was clearly attributed to specific PMF factor. All the fragments were mostly
related to the Sulfate-factor (Fig. 5c). Sulfate factor had the largest
contribution to H2SO3+ (85%) and smallest to HSO− (56%) whereas
the opposite was detected to MOOA having the largest contribution to
HSO− (31%) and smallest to H2SO3+ (13%). HSO− and H2SO3+ were
also partly associated with LV-OOA (11%), and SO+ and SO2 were
associated with SV-OOA (~ 4%).

Sulfate from the oxidation process of DMS is often called non-sea
salt sulfate (nss-sulfate) in order to distinguish it from sulfate originated
from sea-salt in the marine environment. Since the oxidation of DMS
can be considered to be the only atmospheric source of MSA, MSA can
be used as a tracer to separate marine biogenic sulfate from the other
sulfate sources (Davis et al., 1999; Legrand et al., 1991). The ratio of
MSA to nss-sulfate, however, depends on the ambient condition such as
temperature (Bates et al., 1992). In this study, for MOOA, the ratio of
MSA to nss-sulfate was 0.11 when MSA was calculated by using the
ratio of CH3SO2+ to the total organic signal in the reference MS of MSA
and nss-sulfate was calculated by using the measured Na concentration
(> 99% of total sulfate in MOOA was nss-sulfate). This ratio was in the
same magnitude, or slightly larger, than that observed earlier in South
Pacific (0.010–0.210), however, modelled MSA to nss-sulfates ratios
were typically larger (0.093–0.205; Gondwe et al., 2004). The ratio of
MSA to nss-sulfate obtained in this study suggested that most of sulfate
in MOOA was likely to be associated with marine origin, DMS oxida-

Besides sulfate from the oxidation of DMS, the fragmentation
of MSA produces inorganic fragments similar to sulfate. Therefore, the
pattern of sulfate fragments detected for MOOA and Sulfate factor was
compared to the pattern observed for MSA and ammonium sulfate in
the laboratory (Fig. S18). It is clear that the sulfate pattern for MOOA
and Sulfate-factor resembled the pattern of ammonium sulfate more
than that of MSA. MSA had relatively larger contribution of HSO2+ (at
m/z 65) than ammonium sulfate, MOOA or Sulfate-factor whereas
SO3+ (at m/z 80) was almost absent in the MS of MSA. It can be con-
cluded that most of sulfate in MOOA and Sulfate-factor was related to
inorganic sulfate (ammonium sulfate, ammonium bisulfate, sulfuric
acid) not organic sulfur, like MSA. That was also confirmed by calcu-
ling the amount of sulfate formed from the fragmentation of MSA by
using the ratio of total sulfate to CH3SO2+ obtained from the reference
MS of MSA (Fig. S18). Based on this calculation, ~17% of sulfate in
MOOA (and 2.5% in Sulfate-factor) was attributed to the fragmentation
of MSA. As already mentioned sulfur-containing organic and inorganic
fragments may also originate from the fragmentation of organosulfates
(Farmer et al., 2010) but their contribution could not be taken into
account in this study.

4. Conclusions

The aim of this study was to investigate the chemical composition
and sources of submicron particles in Valle Alegre, the coastal area of
Central Chile. The specific feature of the measurement location was the
vicinity of large natural particle source (Pacific Ocean) and several
anthropogenic sources (industrial area). In general, Valle Alegre is

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**Fig. 5.** Distribution of sulfate between the PMF factors in terms of total mass (a) diurnal hour (b), and individual fragments (c).
considered rather polluted as the mass concentrations often exceed the national air quality standards for PM_{10}, PM_{2.5} and SO_{2}. However, the average mass concentration for submicron particles during the three-week campaign was only 9.5 \mu g m^{-3} that corresponds with the mass concentrations measured typically in urban areas (e.g. Zhang et al., 2007). The reason for the rather small concentration was likely to be the location of the measurement site that was in a corner of agricultural and recreational area not supposed to represent the most polluted location in the valley.

Particles were dominated by organic compounds (42% of mass). Organic aerosol were largely influenced by secondary OA that was mostly long-range transported, or regionally distributed highly-oxygenated material (LV-OOA), or marine-related (MOOA). The specific feature of MOOA was that, in addition to oxygenated organic fragments, it contained organic fragments with sulfur. Sulfur-containing fragments were associated with the fragmentation of MSA (CH_{3}SO_{2}^{+}, CH_{2}SO_{2}^{+} and CH_{3}SO_{4}^{+}). Additionally, a factor probably representing local SOA production was detected (SV-OOA). A smaller fraction of OA was composed of primary OA (HOA and BBOA) that probably originated from local combustion sources, traffic and biomass burning. Direct industrial-related OA could not be identified in this study. However, based on the PMF solution for OA and sulfate, 9% of OA was attributed to the PMF factor related to anthropogenic sulfate. This is higher OA fraction than that found in a previous study conducted in an industrialized environment in Marseille, France, where < 5% of total OA was associated with industry (El Haddad et al., 2013).

Submicron particles contained a significant fraction of sulfate (39% of mass). Sulfate was partly neutralized by ammonium but there were periods when particles were clearly acidic. Sulfate had two major sources, or formation pathways, in Valle Alegre. 70% of sulfate seemed to be related to the oxidation of anthropogenic SO_{2} whereas 24% of sulfate was associated with the oxidation of DMS in the marine environment. The most important industrial sources of sulfate at the area were copper smelter and coal-fired power plants. The concentrations of other inorganics species, nitrate, BC and chloride, were rather small peaking only in few cases, their sources being possibly quite local.

Due to the laser vaporizer, the SP-AMS also allowed to detect metals (V, Zn, Cd, Cu, K, Na and Mg) in submicron particles. The time series of Zn, Cd and Cu resembled each other suggesting same origin that was speculated to be industry as also sulfate and SO_{2} concentration were elevated at the same time. However, separating the sources of particles was a challenge because the industrial sites and sea were located at the same direction from the measurement site. Some of the metals were also found as inorganic salts of chloride and nitrate (NaCl^{+}, MgCl_{2}^{+}, CaCl_{2}^{+}, KCl^{+} and KNO_{3}^{+}) likely to be related to both natural and anthropogenic origin.

Overall, secondary species constituted 84% of total submicron particle mass in Valle Alegre when BBOA, HOA and BC were considered as primary components. Large contribution of secondary mass can be explained by high concentration of atmospheric oxidants, since the measurements were conducted in summertime, but also by several industrial sources in the region that emit sulfate precursor SO_{2}. The impact of anthropogenic and biogenic sources on total mass was somewhat equal when sulfate from anthropogenic SO_{2} oxidation, BC, nitrate, ammonium, chloride and primary OA (HOA and BBOA) were assumed to be anthropogenic, and sulfate from DMS oxidation and all OOA fractions biogenic. However, if OOA originated also from anthropogenic sources, anthropogenic particle mass exceeded biogenic particle mass. Purely marine-related was ~19% of mass (MOOA and sulfate from DMS oxidation) and ~18% of mass was from combustion sources (HOA, BBOA and BC).

This study showed that the contribution of anthropogenic and biogenic (marine) sources to organic and inorganic particulate matter can be identified and separated. The results of this study provided valuable, novel information on the characteristics and sources of submicron particles in the rural location influenced by industrial and marine sources especially abundant of sulfur-containing inorganic and organic species. Although the measurements were carried out in a single location for a rather short time, the results and the applied measurement and data analysis methods are also applicable to other similar sites to some extent. To our knowledge this is the most extensive study characterizing the chemical properties of submicron particles in coastal Chile.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.atmosenv.2018.11.040.

References


