1	Variability of Black Carbon mass concentration in surface snow at Svalbard						
2	Michele Bertò ^{1#} , David Cappelletti ^{2,7} , Elena Barbaro ^{1,3} , Cristiano Varin ¹ , Jean-Charles Gallet ⁴ , Krzysztof						
3	Markowicz ⁵ , Anna Rozwadowska ⁶ , Mauro Mazzola ⁷ , Stefano Crocchianti ² , Luisa Poto ^{1,3} , Paolo Laj ⁸ ,						
4	Carlo Barbante ^{1,3} and Andrea Spolaor ^{1,2*} .						
5							
6	¹ Ca' Foscari University of Venice, Dept. Environmental Sciences, Informatics and Statistics, via Torino,						
7	155 - 30172 Venice-Mestre, Italy;						
8	² Università degli Studi di Perugia, Dipartimento di Chimica, Biologia e Biotecnologie, Perugia, Italy;						
9	³ CNR-ISP, Institute of Polar Science – National Research Council –via Torino, 155 - 30172 Venice-						
10	Mestre, Italy;						
11	⁴ Norwegian Polar Institute, Tromsø, Norway.						
12	⁵ University of Warsaw, Institute of Geophysics, Warsaw, Poland;						
13	⁶ Institute of Oceanology, Polish Academy of Sciences, Sopot, Poland;						
14	⁷ CNR-ISP, Institute of Polar Science – National Research Council – Via Gobetti 101, Bologna;						
15	⁸ Univ. Grenoble-Alpes, CNRS, IRD, Grenoble-INP, IGE, 38000 Grenoble, France						
16							
17	[#] Now at Laboratory of Atmospheric Chemistry, Paul Scherrer Institute, 5232 Villigen PSI, Switzerland						
18							
19	Correspond to: Andrea Spolaor, andrea.spolaor@cnr.it; Michele Bertò, michele.berto@gmail.it						
20							

21 Abstract

Black Carbon (BC) is a significant forcing agent in the Arctic, but substantial uncertainty remains 22 23 to quantify its climate effects due to the complexity of the different mechanisms involved, in particular related to processes in the snowpack after deposition. In this study, we provide detailed and unique 24 information on the evolution and variability of BC content in the upper surface snow layer during the 25 spring period in Svalbard (Ny-Ålesund). Two different snow-sampling strategies were adopted during 26 spring 2014 and 2015, providing the refractory BC (rBC) mass concentration variability on a 27 28 seasonal/daily and daily/hourly time scales. The present work aims to identify which atmospheric 29 variables could interact and modify the mass concentration of BC in the upper snowpack, the snow layer 30 which BC particles affects the snow albedo. Atmospheric, meteorological, and snow-related physicalchemical parameters were considered in a multiple linear regression model to identify the factors that 31 32 could explain the variations of BC mass concentrations during the observation period. Precipitation

events were the main drivers of the BC variability. Snow metamorphism and activation of local sources
 during the snow melting periods appeared to play a non-negligible role (wind resuspension in specific
 Arctic areas where coal mines were present). The statistical analysis suggests that the BC content in the
 snow is not directly associated to the atmospheric BC load.

37 **1. Introduction**

In the last two decades, the Arctic region has been exposed to dramatic changes in terms of 38 atmospheric temperature rise, sea ice decrease, and increase of air mass transport from lower latitudes 39 40 bringing warmer and humid air masses containing pollutants and anthropogenic derived compounds (Law 41 and Stohl, 2007; Comiso et al., 2008; Screen and Simmonds, 2010; Eckhardt et al., 2013; Schmale et al., 2018; Maturilli et al., 2019). Long-range transport and local emissions of combustion generating aerosols 42 43 like black carbon (BC) can influence the radiative budget of the Arctic atmosphere, especially the impacts 44 of atmospheric aging on the mixing state of BC particles (Eleftheriadis et al., 2009; Bond et al., 2013; 45 Zanatta et al., 2018). When deposited over snow, numerous aerosol species directly increase the quantity 46 of solar radiation absorbed by the snowpack, thus favouring snow aging processes and the decrease of the snow albedo (Hansen and Nazarenko, 2004; Flanner et al., 2007; Hadley and Kirchstetter, 2012; Skiles et 47 48 al., 2018; Skiles and Painter, 2019).

Among these light-absorbing aerosols, black carbon (BC) particles are the most effective in 49 absorbing the visible and near infrared solar radiation. These primarily emitted, insoluble, refractory and 50 51 carbonaceous particles originate from natural and anthropogenic sources such as open fires or diesel engine exhausts. Currently, the anthropogenic emissions are higher compared to the natural ones 52 (Moosmüller et al., 2009; Bond et al., 2013). In 2000, the energy production sector (including fossil fuels 53 and solid residential fuels combustion) generated approximately 59% of the total global BC emissions 54 55 while the remaining came from biomass burning (Bond et al., 2013). BC particles are characterized by a 56 mass size distribution peaking around 100-250 nm (or mass equivalent diameter), e.g. 240 nm in the Svalbard area in spring (Bond et al., 2013; Laborde et al., 2013; Zanatta et al., 2016; Motos et al., 2019). 57 58 The impact of BC particles absorbing the incoming solar radiation has indeed a non-negligible role in the 59 Arctic region, which is already threatened by a two-fold temperature increase compared to the midlatitude areas, the so-called "Arctic Amplification" (Bond et al., 2013; Cohen et al., 2014; Serreze and 60 Barry, 2011). BC has an atmospheric lifetime of about seven days and has been directly targeted in 61 62 important international mitigation agreements (AMAP, 2015). Theoretical and experimental results showed that the cryosphere is affected both by the BC-induced warming of the atmosphere and by direct 63 64 and indirect BC effects on the snow once deposited over it (Flanner, 2013),

65 Atmospheric BC measurements in the Arctic regions are still rare, despite an extraordinary effort 66 done by the international scientific community to evaluate the sources, transport paths, concentration, and 67 climate impact (Eleftheriadis et al., 2009; Pedersen et al., 2015; Ferrero et al., 2016; Ruppel et al., 2017; Osmont et al., 2018; Zanatta et al., 2018; Laj et al., 2020). BC mass concentrations can be directly 68 measured by using incandescent or thermal techniques and indirectly, by absorption measurements using 69 an appropriate mass absorption cross-section (Petzold, 2013). Various terms such as refractory black 70 71 carbon (rBC) for incandescent measurements, elemental carbon (EC) using thermal techniques, or 72 equivalent black carbon (eBC) based on optical technique are used. Forsström et al. (2009) reported 73 measurements performed in Arctic snow in the past and new measurements of EC in snow surface using 74 filters and a thermo-optical method. The geographical and seasonal eBC variability was investigated in 75 the Arctic region by Doherty et al. (2010). Other BC measurement in snow samples from the Arctic 76 region can be found in Aamaas et al. (2011), Forsström et al. (2013), Pedersen et al. (2015), Gogoi et al. 77 (2016), Khan et al. (2017) and Mori et al. (2019). Intercomparison of different techniques agree within a 78 factor of 2 uncertainty at Alert (Sharma et al., 2017), Ny-Ålesund, and Barrow (Sinha et al., 2017).

A complex combination of processes are involved in the BC particles transfer from the 79 80 atmosphere to the surface snow. Via a modelling approach, Liu et al. (2011) found that approximately 81 50% of BC's total burden in the Arctic atmosphere is removed through wet deposition-related processes. 82 Yasunari et al. (2013) estimated the intensity of BC dry deposition on the Himalayan glaciers; they found 83 that the surface roughness and the surface wind speed are critical parameters in order to retrieve realistic 84 results. In a recent study, Jacobi et al. (2019) confirmed the previous estimates suggesting that 85 approximately 60% of the BC particles are deposited on the surface snow via wet deposition in spring in 86 the Svalbard Arctic area. Models are still not fully able to describe the actual deposition and transport 87 processes in Svalbard, resulting in underestimating the BC concentration in the snowpack (Eckhardt, S. et 88 al 2015, Stohl, A. et al. 2013). Although wet deposition is suggested to be the main driver of BC concentration in the snow, little is known about other environmental processes potentially affecting the 89 90 BC particles concentration once deposited, i.e. physical post-depositional processes.

91 In this study we present two unique experiments performed in a clean area close to the town of Ny-Ålesund (Svalbard) at the Gruvebadet Aerosol Laboratory (78.91734 N, 11.89535 E, 40 m a.s.l.), 92 during spring 2014 and 2015. Daily and hourly time resolution samplings were performed on the snow 93 94 surface to investigate which atmospheric variables could directly or indirectly modify the BC mass 95 concentration in the surface snow once deposited. The daily sampling lasted for approximately 85 days to assess the intra-seasonal variability covering the transition from a cold period (April) to the melting 96 97 period in late June. The hourly time resolution experiment was performed to investigate the existence of 98 potential processes affecting the BC concentration over the diurnal cycle.

99

100 2. Experimental Methods

101 2.1 Study Area

Both experiments were conducted in the proximity of the Ny-Ålesund research station (78.5526 102 N, 11.5519 E, 25 m a.s.l.), located on the Spitzbergen Island in Svalbard archipelago. Along the west 103 coast, Svalbard is characterized by a maritime climate with an annual average temperature of -3.9°C in 104 Ny-Ålesund (between 1994 and 2017) (Maturilli et al., 2019). On average, the snowpack starts building 105 up in September and melts away at the end of May (Førland et al. 2011). Ny-Ålesund has become one of 106 107 the reference locations for conducting Arctic climate studies focusing on atmospheric composition and physics. Long-term monitoring of atmospheric aerosols is performed at the Gruvebadet station (Feltracco 108 et al., 2019, 2020, 2021a, 2021b; Moroni et al., 2018; Ferrero et al., 2016; Bazzano et al., 2015; Moroni et 109 al., 2015; Zangrando et al., 2013; Scalabrin et al., 2012, Turetta er al., 2021), and at the Zeppelin 110 observatory (475 m a.s.l.) (Eleftheriadis et al., 2009; Tunved et al., 2013; Lupi et al., 2016, and reference 111 112 therein).

113

114 **2.2** Snow Sampling

There are no standardized methods for sampling, filtering and analytical protocols for detecting atmospheric carbon deposited in snow, even if a few protocols have been developed (Ingersoll et al., 2009; Gallet et al., 2018; Meinander et al., 2020). In the present work, two different sampling strategies were adopted regarding the thickness of the sampled layer and the temporal sampling frequency.

119 Snow samples were collected during two field campaigns: The first campaign was carried out in Spring 2014, from April 1st to June 24th for a total of 85 days, it consists of daily sampling and it is 120 referred hereafter as the "85-days experiment". The second campaign was conducted in Spring 2015 from 121 April 28th to May 1st. During these three days, measurements were collected with hourly sampling. This 122 123 second campaign is hereafter referred as the "3-days experiment". Snow samples were collected about 1 km North-West of Ny-Ålesund (Figure 1). The area is a dedicated clean site for aerosols and snow 124 125 sampling, with no fuel engine traffic. The wind at the site is usually blowing from east to west, and rarely 126 from North to South, minimizing the emission of the town reaching the sampling area. The main wind pattern during the experiment is presented in Figures 1 and 2. The samples for both experiments were 127 128 kept frozen until the lab analyses. The samples were collected using neck nylon gloves to avoid any 129 contamination.

The two experiments aim to capture the rBC mass concentration on a daily basis in the surface snow (upper 10 cm) during the seasonal change and on an hourly basis on a thinner surface snow layer (upper 3 cm) during a daily cycle. Although wet and dry deposition are the main sources of BC in the Artic snow, the aim of our experiments was to evaluate if other atmospheric parameters could contributeto the snow surface rBC mass concentration variability.

135 In the 85-days experiment, the first 10 cm of surface snow were collected on a daily basis (approximately at 11.00 am, GMT+2) in the same area, using a 5 cm diameter and 10 cm long Teflon 136 tube. The samples were collected following a straight line leaving about 15 cm between the sampling 137 points to minimize the spatial variability. The collected snow was homogenized in a pre-cleaned plastic 138 139 bag and then, without melting, 50 mL was transferred into vial (Falcon[™] 50mL Conical Centrifuge Tubes) for BC, coarse mode particles number (mix of soil, mineral coarse mode and possibly coal coarse 140 mode) concentration and electrical conductivity analyses. The 85-days experiment was designed with the 141 142 aim to investigate the BC presence in the upper snow layer, where most of the snow-radiation interaction 143 takes place and where BC particles' presence can decrease the snow albedo (Doherty et al., 2010). Snow 144 albedos increased rapidly and asymptotically as the snow depth increased. Visible albedos reached 0.9 for a snow depth of only 5 cm (Perovich et al. 2007). Moreover, this sampling strategy allowed to evaluate 145 146 the variation of BC on a seasonal basis and to capture the impacts of wind, precipitation or melting.

147 During the 3-days experiment, the first 3 cm of surface snow were collected on an hourly basis in 148 pre-cleaned vials in a delimited area of 2 x 2 m using the same sampling tools as above (Spolaor et al., 149 2019). In this case the samples were collected following a straight line leaving about 5 cm between the 150 sampling points. The aim of the 3-days experiment was to investigate the potential daily cycle of surface 151 BC concentration; therefore, we foresaw that small variations could derive from the impact of the daily 152 variation of short-wave radiation (SWR) and subsequent induced snow metamorphism at the surface of 153 the snowpack, often at cm scale. To avoid dilution of the signal, we reduced the vertical sampling 154 thickness to 3 cm to enhance our chances of observing variation in the rBC mass concentration, if such 155 variation exists.

The temperature at the surface of the snowpack (at 7 cm for 85-days and at 3 cm for 3-days experiment) was always measured. The daily/hourly snow accumulation was determined by measuring the emerging part of 4 poles placed around the sampling area. The average standard deviation calculated from the four poles provides us a reasonable estimate of the variability in snow accumulation\depletion within the sampling area. The standard deviation obtained ranges from 2 to 4 cm for the entire periods, indicating a limited spatial variability.

162

163 2.3 Atmospheric Optical Measurements

164 **2.3.1** Aethalometer (AE-31)

165 In this study, the equivalent BC (eBC) concentration in the Boundary Layer (around 3 m a.s.l) 166 was measured by an AE-31 aethalometer (Gundel et al., 1983), during the 3-day campaign. The device is 167 equipped with 7-wavelengths (370, 470, 520, 590, 660, 880, 950 nm). It determines the attenuation 168 coefficient by using the light attenuation ratio through a sensing spot and a referenced clean spot, both on a quartz fiber filter substrate. The sampling and reference spots surface areas are 0.5 cm², while the 169 volumetric flow rate is 4 L min⁻¹. The flow rate was calibrated with a TetraCal (BGI Instruments) 170 volumetric airflow before and after the field campaign. A 5 minutes temporal resolution was used for data 171 acquisition. However, due to the low background concentration in the Arctic, the signal/noise ratio is 172 173 high, so that data were hourly averaged. The data presented in this study were processed according to Segura et al. (2014) methodology. For this purpose the multiple scattering and filter loading effect 174 (Weingartner et al., 2003) was corrected with new values of mass absorption cross section (MAC) and 175 multiple scattering factor (C=3.1), reported by Zanatta et al. (2018). The MAC value was derived using 176 observations and observationally constrained Mie calculations in spring at the Zeppelin Arctic station 177 (Svalbard, 78°N). Zanatta et al. (2018) estimated the MAC at 550 nm (9.8 m² g⁻¹) and at 880 nm (6.95 m² 178 g^{-1}), which we used to estimate MAC at 520 nm (10.2 m² g⁻¹). 179

180

181 **2.3.2** Particle Soot Absorption Photometer (PSAP)

182 During the 85-days sampling period the aerosol absorption coefficient was also measured by 183 means of a 3-wavelengths PSAP (this instrument was not available during the 3-days experiment period). 184 It measures the variation of light transmission through a filter where particles are continuously deposited 185 with constant airflow. A second filter identical to the first one remains clean and is used as a reference to 186 take into account possible variations of the light source, i.e. a 3-color LED (blue, green and red with wavelength centred around 470, 530 and 670 nm, respectively). The correction developed by Bond et al. 187 188 (1999) was applied to consider the filter loading effect. The complete eBC mass concentration time series for the 85-days experiment was retrieved using the Aethalometer (first period) and the PSAP (second 189 190 period), with an overlapping period with simultaneous measurements of 5 days. For the retrieved eBC mass concentration from the two instruments to be equal during the overlapping period, the PSAP eBC 191 was calculated dividing the absorption measurements (at 530 nm) with a MAC equal to 7.25 $m^2 g^{-1}$ 192 (keeping the AE31 data as reference). Daily averages were calculated from the 1-minute data to compare 193 194 with the rBC daily data obtained from the snow.

195

196 2.4 Surface Snow measurements

197 2.4.1 Coarse Mode Particles Number Concentration

The snow samples were melted at room temperature before the on-line coarse-mode particles and conductivity measurements (the water was pumped from the vials by a 12 channels peristaltic pump, ISMATECH, type ISM942). Specifically, the number concentration of coarse mode particles in the surface snow was measured with a Klotz Abakus laser sensor particle counter. This instrument optically counts the total number of particles and measures each particle's size in a liquid constantly flowing through a laser beam cavity (LDS 23/23). The measurements size range of the instrument is from 0.8 to about 80 μ m with 32 dimensional bins (Table SI 1), not overlapping with that of the SP2. Only the 32nd bin has a dimensional range above 15.5 μ m, i.e. of 80 μ m. The data were recorded by a LabView® based software obtaining a sufficient number of data points in order to have a standard deviation less than 5% of the mean value. The particles number concentration was calculated using the constant water flow value.

208

209 2.4.2 rBC Measurement – SP2

210 The rBC mass concentration and mass size distribution were measured following the methods described in Lim et al. (2014). The snow samples were melted at room temperature prior to the analyses. 211 212 The vials with the melted snow were sonicated for ten minutes at room temperature. The samples were nebulized before the injection in the Apex-Q desolvation system (APEX-Q, Elemental Scientific Inc., 213 Omaha, USA). The nebulization efficiency was evaluated daily by injecting Aquadag® solutions with 214 different mass concentrations, ranging from 0.1 to 100 ng g⁻¹, obtaining an average value of 61%, that 215 was used to correct all the BC mass concentrations reported in this manuscript. More details on the 216 217 method can be found in Lim et al. (2014) and in Wendl et al. (2014).

The SP2 data were analyzed using the IGOR based toolkit from M. Gysel (Laboratory of Atmospheric Chemistry, Paul Scherrer Institute, Switzerland). The large amount of signals derived from every single particle are elaborated achieving rBC mass and number concentrations and size distributions.

221

222 2.5 Meteorological Parameters

223 Meteorological parameters, in addition to the atmospheric and snow ancillary measurements, 224 were used in the statistical exercise to study the variability of rBC mass concentration in surface snow samples as a function of the atmospheric conditions. BC particles are deposited on the snowpack 225 226 following a combination of wet and dry deposition. However, once deposited on/in the snowpack other 227 processes can potentially induced a significant variability in the surface BC content. The wind direction and its velocity can modify the BC distribution in the upper snowpack due to snow-mobilization. The 228 solar radiation and relative humidity may enhance snow sublimation and surface hoar formation thus 229 230 modifying the relative BC concentration in the upper snow layer by removing or adding "water" mass to 231 the snow surface.

Air temperature and relative humidity at 2 meter height have been retrieved from a meteorological station
located about 800 meters north of the sampling site, using a ventilated PT-100 thermo-couple by Thies
Clima and a HMT337 humicap sensor by Vaisala, respectively. Wind speed and direction at 10 meter

235 height were obtained from a Combined Wind Sensor Classic by Thies Clima (see Maturilli et al., 2013). 236 At about 50 m distance, the radiation measurements for the Baseline Surface Radiatio Network (BSRN) 237 provide among others the downward solar radiation detected by a Kipp&Zonen CMP22 pyranometer (Maturilli et al., 2015). Both meteorological and surface radiation measurements are available in a 1-238 minute time resolution via the PANGAEA data repository (Maturilli et al., 2020). The daily/hourly mean 239 240 values of the meteorological parameters were used in the statistical analyses of the 85-days/3-days 241 experiment and in Figures 2 and 3 (the physical-chemical parameters from the snow samples are punctual 242 values).

243

244 **2.6 Parameters considered** in the statistical analysis

245 The snowpack evolution is primarily driven by meteorological parameters, which are responsible for 246 adding/removing mass to the annual snowpack. Wind can affect the snow pack evolution in several ways: 1) by snow redistribution, 2) favouring the ablation/sublimation, and 3) lifting particles from nearby 247 248 sources and areas. Surface snow and air temperatures are two fundamental parameters required to fully 249 understand the varying conditions of the snow pack. In our study, the temperature variables are proxies 250 for the melting episodes and for the presence of liquid water potentially affecting the concentration of 251 impurities. The air and snow temperatures do not have a direct effect in the rBC concertation in surface 252 snow, but they are fundamental indicators to identify the spring warming events (T > 0° C, called also the 253 Rain on Snow events - ROS) that yield the snow melting. Moreover, air and snow temperature could be 254 relevant to evaluate possible snow metamorphism and the response of the upper snowpack to the 255 meteorological conditions. Snow and air temperatures can be used during the 3-days experiment to 256 evaluate the daily scale frequency and be helpful to investigate the daily scale variability of rBC in the 257 surface snow.

The SWR is not expected to be directly linked to the surface mass concentration of rBC, however the 258 259 surface process could affect it indirectly by favouring sublimation (water mass removal), as well as hoar 260 formation (water mass addition) during the colder parts of the day (night/early morning). The relative 261 humidity gives an idea of the amount of water present in the atmosphere and the high RH might favour 262 the deposition of BC suspended by the formation of water droplets through the cloud condensation nuclei. 263 This parameter is especially significant for the selected sampling location, nearby to the shore. Indeed, 264 relative humidity values close or higher than 90% could be associated to fog or low cloud conditions and 265 not directly to wet or dry precipitations. The last meteorological parameter considered is the precipitation amount. This aspect is important to understand the wet deposition processes able to transfer BC particles 266 267 from the atmosphere to the snow surface.

268 The additional selected parameters are 1) the atmospheric eBC mass concentration, to investigate 269 the possible link between eBC particles present in the atmosphere and the rBC in snow surface, 2) the 270 coarse mode particles that could have a similar transport pathways to the black carbon and gives an idea of the amount of total impurities deposition and 3) the total water conductivity, an indirect measurement 271 of the salinity content of the snow. It is important to note that the eBC and the rBC mass concentrations 272 273 are not the same physical quantities: the former is obtained from an absorption measurement assuming a 274 constant MAC, whereas the second is obtained via a laser-induced-incandescence method with an SP2 empirically calibrated with a reference material (Petzold et al., 2013). Considering the location of the 275 sampling site (<1 km from the coastline), the contribution of the ocean emissions to the snowpack 276 277 chemical composition is significant. We considered the total conductivity as an indication of sea spray 278 deposition, and to investigate common deposition patterns and/or similarities to the behaviour of BC 279 (although BC is not emitted from ocean surface). The conductivity was also considered to determine if 280 there was a large sea-spray aerosol event which could, potentially affecting the SP2 measurements (see 281 supplementary material).

282

283 2.7 Statistical Analysis

284 Multiple linear regression was carried out to evaluate the relationship between the observed 285 surface snow rBC mass concentration and the selected set of covariates consisting of the meteorological 286 and snow physical-chemical parameters that could have a direct effect on controlling snowpack dynamics 287 as well on the BC concentration as discussed in Section 2.6. All the atmospheric parameters described in 288 the previous section (wind, snow and air temperature, incoming solar radiation, relative humidity, and 289 snow precipitation amount) were initially considered as covariates to be included in the multiple linear regression. However, wind speed and direction, as well as the atmospheric stability, expressed as vertical 290 291 wind speed, were removed because preliminary statistical analyses indicate that none of them is associated with the observed variations in snow rBC mass concentrations. This does not mean that such 292 293 parameters do not play a role in controlling the BC concentration, but that no statistically significant 294 associations were found with the data collected in our study and thus these parameters were no longer 295 considered in the statistical analyses discussed below.

Multiple linear regression models were fitted on the logarithm scale because the distribution of rBC concentrations in both experiments is characterized by a significant skewness. Coarse mode particles number concentrations and conductivity were also log-transformed to linearize their relationships with log(rBC). The regression model fitted on the two experiments is

$$log(rBC) = \beta_0 + \beta_1 log(dust) + \beta_2 eBC + \beta_3 temp + \beta_4 snow + \beta_5 swr + \beta_6 log(cond) + \epsilon.$$

301

302 In the above model, 'dust' indicates coarse mode particles number concentrations, 'temp' is the snow 303 temperature at 7 cm depth for the 85-days experiment (daily resolution) and at 2 cm dept for the 3-days experiment (hourly resolution), 'snow' is a binary indicator for the presence of solid precipitation, 'swr' is 304 solar incoming shortwave radiation, 'cond' is the conductivity and ε is a zero-mean normal error. 305 Graphical inspection of residuals plots and normal probability plots confirmed that after the logarithm 306 transformations, the regression models meet the assumptions of linearity, constant error variance (called 307 homoschedasticity in the statistical literature) and normal errors. The statistical analyses were performed 308 309 with the statistical language R (R Core Team, 2020).

310

311 **3. Results and Discussions**

312 **3.1 Seasonal BC variability in surface snow**

313 **3.1.1** Atmospheric eBC and atmospheric condition

During the experimental period, the atmospheric eBC concentration ranged between from 80 ng m⁻³ to <314 5 ng m⁻³ (Figure 2) with an average of 34 ± 23 ng m⁻³. The highest concentrations were measured at the 315 beginning of the campaign, especially from April 15th to 27th, followed by a general decreasing trend 316 characterized by the presence of several concentration peaks (on May 8th, 17th and 24th) potentially due to 317 Eurasian fires, as already suggested from Feltracco et al., 2020 (Figure S1). The ammonia daily 318 319 concentration time series (the only available biomass burning tracer for that period in the area) measured 320 at the Zeppelin station is plotted together with the Gruvebadet atmospheric BC measurements in Figure 321 S3. Biomass burning is a significant source of atmospheric ammonia (Andreae and Merlet, 2001), often affecting the Arctic region (Moroni et al. 2020). As shown in Figure S3, both time series have a similar 322 behaviour at the very beginning of the campaign, from April 3rd to 8th and during the period between May 323 7th and 21st. This suggests that the BC detected in the atmosphere could be originated from biomass 324 burning episodes during these two time periods. During the 85-days sampling period, wind was 325 characterized by the following median values (25th and 75th percentiles) for direction and speed: 205° 326 (152°, 257°) and 2.7 (1.9, 3.7) m s⁻¹, respectively, therefore mostly coming from south-west (Figure 2). 327 Daily air temperature at 3 m increased during the campaign from -15°C to about +5°C (Figure 2) 328 following the seasonal variation of incoming solar energy: from 100 to 300 W m⁻² with an average of 185 329 \pm 75 W m⁻² (Figure 2, orange line). The snow precipitation episodes are presented as daily-accumulated 330 331 values (Figure 2, blue bars) ranging from zero to 12 cm.

332

333 **3.1.2 Surface Snow Conditions**

Over the 85 days experiment, the snow rBC mass concentration varies from 0.2 to 6 ng g^{-1} (Figure 2), 334 with an average of 1.4 ± 1.3 ng g⁻¹, in agreement with results available in the literature (Mori et al., 2019; 335 Jacobi et al., 2019; Aamaas et al., 2011). An increasing trend can be observed for the rBC mass 336 concentration in the surface snow across the sampling period. The median of the rBC mass equivalent 337 diameter in the snow is 313 ± 35 nm (Figure 2), similar to what obtained in other studies (e.g. Schwarz et 338 al., 2013). The rBC mass equivalent diameter show high variability, ranging from 200 to 500 nm. 339 340 However, since the rBC concentrations were low, the evaluation of the geometric mean of the particles diameter for the biggest sizes, above 300 to 400 nm, has been considered as qualitative information due to 341 the high signal noise. 342

The number of coarse mode particles (Figure 2, blue line) shows a constant concentration in the first half 343 of the campaign $(1^{st} \text{April} - \text{May } 11^{th} - \text{average concentration of } 3435 \pm 1824 \# \text{ml}^{-1})$ whereas it increases in 344 the second half (12^{th} of May to 27^{th} of June - average concentration of $7782\pm5683 \text{ # ml}^{-1}$), especially after 345 the 1st of June (1st of June to 27th of June - average concentration of $9352\pm6741 \text{ # ml}^{-1}$), in concomitance 346 with the onset of the snow melting period. The conductivity (Figure 2, green line) also shows an 347 increasing trend at the end of the sampling campaign when snow is melting, with an overall average value 348 of $30 \pm 8 \mu$ S. The spatial variability of rBC, calculated in the same manner as proposed by Spolaor et al. 349 350 (2019) for other species, was obtained from six surface snow samples collected in the four corners of the 351 sampling area and two surface snow samples in the centre right before the beginning of the experiment. The following rBC mass concentrations were obtained: a) 3.95 ng g⁻¹; b) 4.92 ng g⁻¹ c) 4.20 ng g⁻¹d) 3.10 352 ng g⁻¹e) 3.82 ng g-1 f) 3.58 ngg⁻¹, resulting in a rBC spatial variability of 16% in the surface snow of the 353 354 considered sampling area.

355

356 3.1.3 Statistical Results

The fitted multiple linear regression model for the 85-days experiment data explains the 69% of the variance of the logarithm of the snow rBC mass concentration ($R^2 = 0.69$). The fitted model indicates the presence of strongly statistically significant associations of the (log transformed) snow rBC mass concentration with the coarse-mode particles number concentration (p < 0.001) and the snow temperature (p < 0.001). A weaker association is found with the occurrence of snow precipitations (p = 0.03). The statistical associations of rBC mass concentration with the other covariates considered in the model are non-significant. See Table 1 for the estimated coefficients and the corresponding p-values.

In order to interpret the statistical results, the description of the 85-days campaign is split into two periods identified as the transition from the "cold" to the "melting" state. The first period occurred before the end of May: the rBC mass concentration often increases with snowfall episodes (April 9th/10th/11th and 17th;

May 17th, 22nd and 27th/28th; June 1st) as suggested by previous studies, with exceptions for April 24th and 367 368 May 7^{th} . Over the sampling period, a weakly statistically significant positive association (p = 0.03) was 369 found between snow rBC mass concentration in surface snow and the occurrence of snow precipitations. 370 BC wet deposition processes are estimated to remove 50% - 60% of the total atmospheric BC burden in 371 the Arctic (Liu et al., 2011; Jacobi et al., 2019). In our study, the wet deposition impacts could be partially 372 masked due to the sampling frequency and the wind snow. In Kongsfjord, a strong wind is often present 373 during the precipitation events (Figure 2). Consequently, the freshly deposited snow is frequently 374 removed from the surface before being able to sample it. Interestingly, our observations show that, on a daily scale, the precipitation episodes are not clearly related to a decrease in the atmospheric eBC mass 375 376 concentration (Figure 2). A possible explanation is that the precipitation amounts were small so that the 377 precipitation events did not significantly alter the atmospheric BC reservoir.

In the second period, from the beginning of June, the atmospheric temperature increases, causing the 378 379 snow-melting season's onset. At the beginning of June, the snow rBC mass concentration increases up to approximately 5 ng g⁻¹, and a simultaneous increase was detected in the coarse mode particles number 380 concentration (peaks between June 4th and 7th). As suggested in previous studies, the surface melting 381 process could explain the observed increase in rBC and coarse mode particles concentrations. However, 382 we also have to consider that rBC can be dry deposited, as it has been recently suggested (up to 50-60%; 383 Liu et al., 2011; Jacobi et al., 2019). Very few field validation data exist for estimating the amount of dry 384 385 deposition at the snow surface, and this process is often used as an ancillary information since most 386 models underestimate the BC in the Arctic snowpack compared to field measurements.

Our data support the hypothesis related to local sources' activation in enhancing the dry deposition impacts in an old mining town as Ny-Alesund. Especially during poor snow cover conditions, as during the snow-melting season, coarse mode particles as residuals of carbon extraction mining activities are available for wind lift\suspension (Vecchiato et al. 2018). The possible effect of local sources' activation is further supported by a recent analysis of the Brøggerbreen glacier and Ny-Ålesund annual snowpack. This analysis shows the presence of retene (an organic compound frequently used to track the presence of coal), most likely due to local sources (Vecchiato et al., 2018).

The simultaneous increase of rBC mass and coarse mode particle number concentrations during the second part of the experiment (e.g. visible between June 3rd and June 7th-8th) could be explained via similar post-depositional processes: snow melting and sublimation. The episodes of snow surface melting can significantly affect the snow particulate content and we hypothesize that the hydrophobicity of pure BC particles, and of several species in the coarse mode particles, might affect its physical location in the 399 snowpack (in the literature, the response of the BC particles is still debated): the hydrophobicity of the 400 particles can cause the surface concentration to increase while losing water mass through percolation. 401 This could lead into a positive feedback process: the increase of BC concentration can thus enhance snow 402 sublimation (water evaporation) resulting in a further increase of BC concentration in surface snow, and 403 so on.

404 In this study, the estimated statistical association between snow rBC mass concentration and the daily snow temperature is negative and strongly significant (p < 0.001). During the 85-days experiment, we can 405 distinguish two events where the temperature appeared to play a role in the BC concentration. Both of 406 407 them show an increase in rBC mass concentration during melting/refreezing episodes, in agreement with other studies (Aamaas et al., 2011; Xu et al., 2006; Doherty et al., 2013; Doherty et al. 2016). The first 408 event occurred between May 5th to May 12th and the second event after May 20th, when the proper snow 409 melting began (Figure 2). The first event was characterized by a rapid rise of the daily air temperature 410 (from $-6^{\circ}C$ to $-1^{\circ}C$) in concomitance to a snow precipitation event, followed by a rapid temperature 411 decrease to -6 °C. The surface snow (10 cm) mirrored this behaviour, first rising from -6 °C to 0 °C, and 412 then cooling down to -6 °C. During this warm event, the upper snow strata underwent a melting episode 413 with surface water percolation (although limited), making the surface BC concentration to increase. The 414 second event started approximately on May 20th and lasted until the end of the experiment (Figure 2). 415 During this period, the atmospheric temperature increased constantly, and the snowpack started to melt 416 consequently. Moreover, surface BC concentration increased almost continuously from May 25th to its 417 maximum observed on June 6th. Afterward, the upper snow rBC mass concentration tended to decrease 418 419 following the rapid snowpack decline.

- 420
- 421 **3.2 Diurnal variation of rBC in surface snow**

422 3.2.1 Surface Snow/Atmospheric Aerosol Content and Atmospheric Conditions

The 3-days experiment was performed at the end of April 2015, during the Arctic spring. The samples were collected on an hourly basis over 3 days achieving a high-resolution sampling frequency. The atmospheric concentration of eBC ranged from 2 to 50 ng m⁻³, decreasing during the sampling period and not showing any particular diurnal pattern (Figure 3). The mean value of the atmospheric eBC mass concentration is 34 ± 23 ng m⁻³, similar to the average of the 85-days experiment.

The surface snow rBC mass concentration undergoes to daily time scale change of surface concentration showing up to 2-fold hourly increases (Figure 3, bottom panel, smoothed dark blue line). rBC mass concentrations of approximately 15 ng g⁻¹ were measured in the snow samples from the beginning of the sampling to the end of the second day. Later, from the beginning of the third day until

the end of experiment, rBC mass concentrations show an average concentration of about 5 ng g^{-1} (Figure 432 3). The average value over the whole sampling period is 9 ± 5 ng g⁻¹ (approximately 6 times higher than 433 434 during the 85-days experiment). The rBC mass size distribution was characterized by a median value of the geometric means of about 230 ± 32 nm, significantly lower than that which was measured during the 435 85-days, and still in agreement with previous studies (Sinha et al., 2018; Schwarz et al., 2013). The 436 concentrations of EC and OC measured in parallel snow samples (not of the same volume) are reported 437 438 and described in Figure S4; the interpretation of the differences between the rBC and the EC 439 measurements in snow samples was beyond this manuscript's objectives.

440 The number concentration of coarse mode particles remains stable in the first half of the 441 experiment, until the end of April, and shows an average value over the three days of $26642 \pm 9261 \ \text{mL}^{-}$ 442 ³. The water conductivity shows a similar behaviour, and it is characterized by an average of $39 \pm 9 \ \mu\text{S}$ 443 (30% higher than during the 85-days experiment).

All the measured snow impurities show two common features (see supplementary material and 444 Figure S4): first, a decrease in the absolute values detected between 4 and 8 a.m. of April 30th, despite the 445 absence of precipitations or any other particular meteorological episode (Figure 3); second, the impact of 446 the snow precipitation event from approximately 4 p.m. to midnight of the April 30th, where the 447 448 concentrations of aerosols in the snow slightly increased at the very beginning whereas decreasing at the 449 end of the event. Only the BC core diameter remained above the average when the other aerosol snow 450 content decreased (up to approximately 400 nm of mass equivalent diameter), consequently returning to 451 the average value. The spatial variability of BC, calculated as proposed by Spolaor et al. (2019) for other 452 species, was obtained by the analysis of 5 surface snow samples, collected in the four corners of the sampling area and one in the centre obtaining the following concentrations: a) 10.17 ngg⁻¹, b) 10.64 ng g⁻¹ 453 ¹, c) 7.04 ngg^{-1} , d) 11.98 ngg^{-1} , and e) 11.91 ngg^{-1} , thus resulting in a spatial variability of 19%. Clear 454 sky conditions where observed for the duration of the sampling period except for the snowfall occurred at 455 456 the end of the third day.

457

458 **3.2.2 Statistical Results**

The multiple linear regression model for the 3-days experiment explains the 78% of the snow rBC mass concentration variance, a percentage higher than the 85-days experiment, likely due to the more stable atmospheric conditions and the greater interaction with the atmosphere of the upper 3 cm of the snow pack compared with the depth resolution used during the seasonal experiment. Similar for the 85days experiment we evaluate (Figure S4) the 10 days back-trajectory during the 3 days of the experiment. The result suggests that the air mass arriving in Ny-Ålesund during the experiment were mainly originated from the Arctic Ocean. 466 The fitted multiple linear regression model indicates a statistically significant association between 467 the logarithm of the rBC mass concentration in the snow and the logarithm of the conductivity (p < p468 0.001), the logarithm of the number concentration of coarse-mode particles (p < 0.001) and the occurrence of snow precipitations (p < 0.001). The estimated coefficients of the covariates are reported in 469 470 Table 1. In Figure 4 are displayed the 95% and 90% confidence intervals for the estimated coefficients of 471 regression models fitted to two experiments (85-days and the 3-days). Since the covariates considered in 472 the two experiments have quite different unit scales, Figure 4 shows the confidence intervals for the 473 standardized covariates. The standardization simplifies the comparison among the estimated effects of the 474 different covariates and between the two experiments, in this way allowing a visual comparison of the 475 estimated statistical associations between the logarithm of the snow rBC mass concentration and the 476 considered parameters.

477 The association between the logarithm of the coarse-mode particles number concentration and the 478 logarithm of the snow rBC mass concentration is positive and strongly significant (p < 0.001), similarly to 479 what observed for the 85-days experiment, confirming the similar behaviour of these types of particles 480 also in the surface snow pack (3 cm). The association between the logarithm of conductivity and the logarithm of the snow rBC mass concentration is positive and strongly significant (p < 0.001). Snow 481 482 conductivity is mostly influenced by the presence of sea salt ions (mainly coming from sea spray aerosol 483 considering the location of the experimental site) in the snow samples. Sea spray aerosol is not considered 484 a source of rBC and a direct effect of the sea spray emission on the rBC snow concentration is here 485 consider negligible. However the positive association between rBC and conductivity can be explained by 486 the fact that both sea spray aerosol and BC particles (as well dust) undergoes to similar dry deposition 487 process (when concentration increase) favoured by the stable atmospheric condition occurred during the 488 experiment (with the exception of the snow event during the third day) as well from similar physical 489 removal process (concentration decrease) from the snow surface. Considering we are exploring the rBC 490 concentration change in the upper 3 cm, we explore the possible existence of a daily cycle. The BC 491 particles are known to be non-volatile and not photo-chemically active, therefore the decrease/increase in 492 their concentration observed during the experiment can only be driven by physical process such as wind 493 erosion and snow deposition. However and additional process that might drive the rBC concentration 494 change in the upper snow pack is the condensation of water vapour on the top of the snow crystals and the 495 formation of surface hoar as well the sublimation. The formation of surface hoar has the effect to adding 496 "water" mass without BC particles in the snow surface causing a relative rBC dilution, while sublimation 497 has the effect remove "water" mass causing a relative concentration increase. Surface hoar and 498 sublimation are depending mainly by the temperature and solar radiation, two parameters that exhibits the 499 diurnal cycle (Figure 4). From the statistical analysis no associations were found on rBC with the

incoming solar radiation (at hour resolution) and the snow temperature during the sampling period. These
 results indicate that the rBC mass concentration in the surface snow does not undergo to diurnal changes
 and this process are negligible in controlling the rBC snow surface concentration.

The occurrence of snow precipitations is negatively associated with the logarithm of the rBC 503 504 mass concentration in the snow (p < 0.001). As previously remarked, the aerosol scavenging intensity is 505 not measurable with snow sampling strategies based on the sampling of a constant snow thickness from 506 the surface (3 cm in this case). We tentatively explain the negative relation observed in this study with the 507 high frequency sampling, being able to follow the evolution of the BC particles scavenged during a snow episode (from 3 to 12 p.m. of the 30th April 2015). The beginning of the precipitation episodes appeared 508 to remove the highest amount of BC particles, leaving the atmosphere cleaner as reflected by the lower 509 510 BC mass concentration revealed in subsequent samples. The snow collected at 18:00 of April 30 showed 511 a higher amount of rBC as well as the highest coarse mode particles number concentration and conductivity. In the next few hours, from 9 to 12 p.m., the snow precipitations were depleted in terms of 512 513 aerosol content and rBC mass concentration.

514

515 4. Conclusions and Future Perspectives

516 The seasonal and daily experiments (85- and 3-days long, respectively) suggest that the rBC 517 concentration in the upper snow layer is not only driven by a cumulative process, as it happens when the 518 entire annual snow pack is evaluated, but it is a rather more complex process involving atmospheric, 519 meteorological and snowpack conditions. Our results based on a multiple linear regression models 520 suggest that the amount of BC in the surface snow is not associated to the BC atmospheric load. This 521 finding suggests that, despite the potentially high atmospheric BC concentrations (as in the case of long-522 range transport of biomass burning plumes), this parameter does not seem to be the primary driver of the 523 variations in the surface snow rBC over the experiment periods. In both experiments, the coarse mode 524 particles are positively associated with the snow BC mass concentration, suggesting that the BC and 525 coarse mode particles deposition undergo similar deposition and, in case, to post-depositional processes in 526 the upper snowpack. Specifically, before the beginning of the melting season, the wet deposition episodes 527 appeared to have major impacts, whereas the activation of common local sources favour the wind 528 suspension from uncovered areas enhancing the intensity of dry deposition processes, might lead to an 529 accelerated snow melting.

530 Our results also suggest that in order to explain the observed BC mass concentration variability 531 during seasonal and diurnal time ranges other processes than wet and dry depositions should be 532 considered. Surface melting episodes enrich the BC content in the surface layer not because of an 533 enhanced deposition but mainly because of water mass loss. In particular, the snow mass loss is stronger during the snow-melting season, where an increase in the rBC concentration could significantly alter the snow albedo and further enhance the radiative absorption, hence promoting a positive feedback. The proposed processes and the rBC concentration determined in Ny-Ålesund could be influenced by local emission in particular at the begging and at the end of the snow season when the snowpack does not cover homogeneously the surface. However, the process described by our results could occur in other Arctic sites although with different magnitudes and impacts.

The remarkable diurnal and daily variability, as well as the complex interdependent mechanisms affecting the rBC mass concentration in the Arctic surface snow, makes the results of albedo-based radiative impact model of the active layer a potential source of erroneous conclusions: the impacts of long distance biomass burning episodes might be overestimated, whereas the impact of local sources and dry deposited impurities during the melting season might be underestimated. Additional empirical studies are therefore necessary in order to improve our understanding of the involved physical mechanisms and to better constrain modelling studies.

547

548 Acknowledgements

549 This work was part of the PhD (in "Science and Management of Climate Change") of Michele Bertò at 550 the Ca' Foscari University of Venice that was partly funded with the Early Human Impact ERC project. 551 Thanks to Giuseppe Pellegrino for helping collecting the samples. Thanks to Jacopo Gabrieli and the 552 technicians of the Ca'Foscari University of Venice for the precious help in building up the coarse mode 553 particles and conductivity measurement apparatus. We acknowledge the use of data and imagery from 554 LANCE FIRMS operated by the NASA/GSFC/Earth Science Data and Information System (ESDIS) with 555 funding provided by NASA/HQ. We want to thank Paolo Laj and the LGGE (Grenoble, France) for 556 lending us the SP2 and Marco Zanatta for transferring the SP2 know-how on instrumental functioning and 557 data analyses. Thanks to Martin Gysel-Beer, PSI, for the IGOR based SP2 Toolkit for SP2 data analyses. We thank Marion Maturilli and AWI for providing us with the meteorological data. Thanks to Giorgio 558 559 Bertò for checking and correcting the language of this manuscript. This paper is an output of the AMIS project in the framework of "Project MIUR - Dipartimenti di Eccellenza 2018-2022". This project has 560 561 received funding from the European Union's Horizon 2020 research and innovation programme under 562 grant agreement No 689443 via project iCUPE (Integrative and Comprehensive Understanding on Polar 563 Environments).

564

565

567	Data Availability							
568	Meteorological and surface radiation data are available at the PANGAEA database (Maturilli, 2015a;							
569	2015b; 2015c; 2016a; 2016b; 2018a; 2018b; 2018c; 2018d; 2018e). The data for precipitation amount at							
570	Ny-Ålesund can be accessed via the eKlima database of MET Norway. The BC data are available upon							
571	request.							
572								
573	Author Contributions							
574	Author contributions. AS, EB, DC and MB conceived the experiments; AS, EB, DC, and LP collected the							
575	samples; MB measured the samples; KM and MMaz provided the atmospheric eBC concentrations; SC							
576	and DC provided the back-trajectories analyses; CV performed the statistical analyses with inputs from							
577	MB and AS. MB prepared the manuscript mainly with inputs from AS, J-C. G and DC (in the methods							
578	section from AS, KM, MMaz) and all co-authors contributed to the interpretation of the results as well as							
579	manuscript review and editing.							
580								
581	Data repository							
582	Maturilli, Marion (2020): Basic and other measurements of radiation and continuous meteorological							
583	observations at station Ny-Ålesund (April, May 2014 and April, May, June 2015), reference list of 10							
584	datasets. Alfred Wegener Institute - Research Unit Potsdam, PANGAEA,							
585	https://doi.pangaea.de/10.1594/PANGAEA.913988 (DOI registration in progress)							
586								
587								
507								
588								
589								
505								
590								
501								
551								
592								
592								
594								

596 FIGURES

- 597 Figure 1. a) Experimental sampling site location (dark grey rectangle), in proximity of the Gruvebadet
- 598 Aerosol Laboratory. b) Gruvebadet area (black square), close to the Ny-Ålesund research village. From:
- 599 Spolaor et al., 2019 (maps from <u>https://toposvalbard.npolar.no/</u>). The red arrow points to the North.



612 Figure 2. The 85-days experiments daily snow samples rBC mass concentration (light blue), eBC mass 613 concentration in the atmosphere (black), geometric mean mass equivalent diameter (purple), number of 614 coarse mode particles (Cmp - blue), total conductivity (green), meteo/snow parameters used in the statistical exercise: wind speed color coded for wind direction, solar radiation (orange line), air and 615 surface snow temperatures (blue bars and green line respectively), amount of fresh snow ("snow 616 precipitations", light blue bars) and the snow accumulation ("Neg. accumulation"; the values where 617 multiplied by -1 in order to show the similar trend of the snow lost and of the air/snow temperature during 618 619 the melting period at the end of the campaign).

620



Figure 3. The 3-days experiments snow samples hourly rBC mass concentration and smoothed line (light blue bars), atmospheric eBC mass concentration in the atmosphere (black), geometric mean mass equivalent diameter (purple), the number concentration of coarse mode particles (Cmp - blue) and the total conductivity (green), meteo/snow parameters used in the statistical exercise: wind speed color coded for wind direction, solar radiation (Orange line), Air and surface snow temperature (blue bars and green line respectively), amount of fresh snow ("snow precipitations", light blue bars). The yellow bars are centered on the midnight hours for the three sampling days.

630



631

633 Figure 4. Estimated coefficients of the standardized covariates of the multiple linear regression models 634 fitted to the 3 days and 85 days experiments. The segments correspond to 95% confidence intervals about 635 the corresponding estimated coefficients. The internal thicker segments correspond to 90% confidence intervals. Intervals, that do not include the zero, correspond to statistically significant covariates. If a 636 confidence interval consists of positive values, then there is a significant positive association between the 637 corresponding covariate and the logarithm of the snow rBC mass concentration given the remaining 638 639 covariates. Vice versa, if the confidence interval consists of negative values, then the association is 640 negative. The abbreviations used in the plot are: "log(cond)" - logarithm of the water conductivity time 641 series, "log(dust)" - logarithm of the coarse mode particles number concentration time series, "eBC" equivalent black carbon atmospheric concentration, "snow" - presence of snow precipitation episodes, 642 "swr" - short wave radiation, "temp" - the snow temperature. The plot is produced with the R package (R 643 Core Team, 2020) jtools (Long, 2020). 644

645



646

647

648

650 TABLES

Table 1. Estimated coefficients, 95% confidence intervals and the corresponding p-values for the covariates of the multiple linear regression model fitted to the 85 days and the 3-days experiments. The last rows of the table report the number of observations, the multiple coefficient of determination (R^2) and its adjusted version.

	85-days			3-days		
Covariates	Estimates	CI	р	Estimates	CI	р
(Intercept)	-6.74	-8.744.74	<0.001	1.39	-2.30 - 5.09	0.453
Cond[log]	-0.02	-0.51 - 0.48	0.950	1.38	0.89 - 1.87	<0.001
Dust[log]	1.29	1.03 - 1.55	<0.001	0.74	0.39 - 1.09	<0.001
eBC	-0.01	-0.01 - 0.00	0.074	0.00	-0.00 - 0.01	0.272
Snow[TRUE]	0.29	0.02 - 0.55	0.033	-0.77	-1.150.38	<0.001
SWR	0.00	-0.00 - 0.00	0.613	0.00	-0.00 - 0.00	0.468
Temp	-0.10	-0.140.05	<0.001	0.02	-0.04 - 0.08	0.535
Observation	72			68		
R2 /R2 adjusted	0.688/0.6590			0.779 /0.758		

667 References

- Aamaas, B., Bøggild, C. E., Stordal, F., Berntsen, T., Holmèn, K. and Strùm, J.: Elemental carbon
 deposition to Svalbard snow from Norwegian settlements and long-range transport, Tellus B Chem.
 Phys. Meteorol., 63(3), 340–351, doi:10.1111/j.1600-0889.2011.00531.x, 2011.
- AMAP, A. M. and A.: ARCTIC MONITORING AND ASSESSMENT PROGRAMME (AMAP): Work
 Plan 2015–2017., Working Paper, Arctic Monitoring and Assessment Programme (AMAP). [online]
 Available from: https://oaarchive.arctic-council.org/handle/11374/1443 (Accessed 6 May 2020), 2015.
- Andreae, M. O. and Merlet, P.: Emission of trace gases and aerosols from biomass burning, Glob.
 Biogeochem. Cycles, 15(4), 955–966, doi:10.1029/2000GB001382, 2001.
- Bazzano, A., Ardini, F., Becagli, S., Traversi, R., Udisti, R., Cappelletti, D. and Grotti, M.: Source
 assessment of atmospheric lead measured at Ny-Ålesund, Svalbard, Atmos. Environ., 113, 20–26,
 doi:10.1016/j.atmosenv.2015.04.053, 2015.
- Bond, T. C., Anderson, T. L. and Campbell, D.: Calibration and Intercomparison of Filter-Based
 Measurements of Visible Light Absorption by Aerosols, Aerosol Sci. Technol., 30(6), 582–600,
 doi:10.1080/027868299304435, 1999.
- Bond, T. C., Doherty, S. J., Fahey, D. W., Forster, P. M., Berntsen, T., DeAngelo, B. J., Flanner, M. G.,
 Ghan, S., Kärcher, B., Koch, D., Kinne, S., Kondo, Y., Quinn, P. K., Sarofim, M. C., Schultz, M. G.,
 Schulz, M., Venkataraman, C., Zhang, H., Zhang, S., Bellouin, N., Guttikunda, S. K., Hopke, P. K.,
 Jacobson, M. Z., Kaiser, J. W., Klimont, Z., Lohmann, U., Schwarz, J. P., Shindell, D., Storelvmo, T.,
 Warren, S. G. and Zender, C. S.: Bounding the role of black carbon in the climate system: A scientific
 assessment: BLACK CARBON IN THE CLIMATE SYSTEM, J. Geophys. Res. Atmospheres, 118(11),
 5380–5552, doi:10.1002/jgrd.50171, 2013.
- Cohen, J., Screen, J. A., Furtado, J. C., Barlow, M., Whittleston, D., Coumou, D., Francis, J., Dethloff,
 K., Entekhabi, D., Overland, J. and Jones, J.: Recent Arctic amplification and extreme mid-latitude
 weather, Nat. Geosci., 7(9), 627–637, doi:10.1038/ngeo2234, 2014.
- Comiso, J. C., Parkinson, C. L., Gersten, R. and Stock, L.: Accelerated decline in the Arctic sea ice cover,
 Geophys. Res. Lett., 35(1), doi:10.1029/2007GL031972, 2008.
- DeMott, P.J., Hill, T.C., McCluskey, C.S., Prather, K.A., Collins, D.B., Sullivan, R.C., Ruppel, M.J.,
 Mason, R.H., Irish, V.E., Lee, T. and Hwang, C.Y.: Sea spray aerosol as a unique source of ice
 nucleating particles. Proceedings of the National Academy of Sciences, 113(21), pp.5797-5803, doi:
 10.1073/pnas.1514034112, 2016.
- Doherty, S. J., S. G. Warren, T. C. Grenfell, A. D. Clarke, and R. E. Brandt.: Light-absorbing impurities
 in Arctic snow. Atmospheric Chem. Phys.10, no. 23: 11647, doi: 10.5194/acp-10-11647-2010, 2010.
- Doherty, S. J., T. C. Grenfell, S. Forsström, D. L. Hegg, S. G. Warren and R. Brandt, Observed vertical redistribution of black carbon and other light-absorbing particles in melting snow, J. Geophys. Res., 118(11), 5553-5569, doi:10.1002/jgrd.50235, 2013.
- Doherty, S. J., D. A. Hegg, P. K. Quinn, J. E. Johnson, J. P. Schwarz, C. Dang and S. G. Warren, Causes of variability in light absorption by particles in snow at sites in Idaho and Utah, J. Geophys. Res. Atmos., 121, doi:10.1002/2015JD024375, 2016.

Eckhardt, S., Hermansen, O., Grythe, H., Fiebig, M., Stebel, K., Cassiani, M., Baecklund, A. and Stohl,
A.: The influence of cruise ship emissions on air pollution in Svalbard - a harbinger of a more polluted
Arctic?, Atmospheric Chem. Phys., 13(16), 8401–8409, doi: 10.5194/acp-13-8401-2013, 2013.

Eckhardt, S., Quennehen, B., Oliviè, D. J. L., Berntsen, T. K., Cherian, R., Christensen, J. H., W. Collins
et al.:Current model capabilities for simulating black carbon and sulfate concentrations in the Arctic
atmosphere: a multi-model evaluation using a comprehensive measurement data set. Atmospheric
Chem. Phys., 15, no. 16: 9413-9433, doi: 10.5194/acp-15-9413-2015, 2015.

- Eleftheriadis, K., Vratolis, S. and Nyeki, S.: Aerosol black carbon in the European Arctic: Measurements
 at Zeppelin station, Ny-Ålesund, Svalbard from 1998–2007, Geophys. Res. Lett., 36(2),
 doi:10.1029/2008GL035741, 2009.
- Feltracco, M., Barbaro, E., Kirchgeorg, T., Spolaor, A., Turetta, C., Zangrando, R., Barbante, C. and
 Gambaro, A.: Free and combined L- and D-amino acids in Arctic aerosol, Chemosphere, 220, 412–421,
 doi:10.1016/j.chemosphere.2018.12.147, 2019.
- 719 Feltracco, M., Barbaro, E., Tedeschi, S., Spolaor, A., Turetta, C., Vecchiato, M., Morabito, E., 720 Zangrando, R., Barbante, C. and Gambaro, A.: Interannual variability of sugars in Arctic aerosol: 721 burning and biogenic Total Environ., 136089. Biomass inputs. Sci. 706. 722 doi:10.1016/j.scitotenv.2019.136089, 2020.
- Feltracco, M., Barbaro, E., Spolaor, A., Vecchiato, M., Callegaro, A., Burgay, F., Vardè, M., Maffezzoli,
 N., Dallo, F., Scoto, F., Zangrando, R., Barbante, C., Gambaro, A.: Year-round measurements of sizesegregated low molecular weight organic acids in Arctic aerosol. Sci. Total Environ., 763, 142954, doi:
 10.1016/j.scitotenv.2020.142954, 2021a.
- Feltracco, M., Barbaro, E., Hoppe, C. J., Wolf, K. K., Spolaor, A., Layton, R., Keuschnig, C., Barbante,
 C., Gambaro, A., Larose, C.: Airborne bacteria and particulate chemistry capture Phytoplankton bloom
 dynamics in an Arctic fjord, Atmos. Environ., 256, 118458, doi: 10.1016/j.atmosenv.2021.118458,
 2021b.
- Ferrero, L., Cappelletti, D., Busetto, M., Mazzola, M., Lupi, A., Lanconelli, C., Becagli, S., Traversi, R.,
 Caiazzo, L., Giardi, F., Moroni, B., Crocchianti, S., Fierz, M., Mocnik, G., Sangiorgi, G., Perrone, M.
 G., Maturilli, M., Vitale, V., Udisti, R. and Bolzacchini, E.: Vertical profiles of aerosol and black carbon
 in the Arctic: a seasonal phenomenology along two years (2011-2012) of field campaign, Atmospheric
 Chem. Phys., 16, 12601–12629, doi: 10.5194/acp-16-12601-2016, hdl:10013/epic.48736, 2016.
- Flanner, M. G.: Arctic climate sensitivity to local black carbon, J. Geophys. Res. Atmospheres, 118(4),
 1840–1851, doi:10.1002/jgrd.50176, 2013.
- Flanner, M. G., Zender, C. S., Randerson, J. T. and Rasch, P. J.: Present-day climate forcing and response
 from black carbon in snow, J. Geophys. Res. Atmospheres, 112(D11), doi:10.1029/2006JD008003,
 2007.
- Forsström, S., Ström, J., Pedersen, C. A., Isaksson, E. and Gerland, S.: Elemental carbon distribution in
 Svalbard snow, J. Geophys. Res. Atmospheres, 114(D19), doi:10.1029/2008JD011480, 2009.
- Forsström, S., Isaksson, E., Skeie, R. B., Ström, J., Pedersen, C. A., Hudson, S. R., Berntsen, T. K.,
 Lihavainen, H., Godtliebsen, F. and Gerland, S.: Elemental carbon measurements in European Arctic
 snowpacks, J. Geophys. Res. Atmospheres, 118(24), 13,614-13,627, doi:10.1002/2013JD019886, 2013.

- Gallet JC, Björkman M, Larose C, Luks B., Martma T. and Zdanowics C. (eds). Protocols and
 recommendations for the measurement of snow physical properties, and sampling of snow for black
 carbon, water isotopes, major ions and micro-organisms. Norwegian Polar Institute. Kortrapport / Brief
 Report no. 046, ISBN 978-82-7666-415-7 (printed), www.npolar.no, 2018.
- Gogoi, M. M., Babu, S. S., Moorthy, K. K., Thakur, R. C., Chaubey, J. P. and Nair, V. S.: Aerosol black
 carbon over Svalbard regions of Arctic, Polar Sci., 10(1), 60–70, doi:10.1016/j.polar.2015.11.001, 2016.
- Gundel, L. A., Dod, R. L., Rosen, H. and Novakov, T.: Relationship between optical attenuation and
 black carbon concentration for ambient and source particles, Lawrence Berkeley Lab., CA (USA).
 [online] Available from: https://www.osti.gov/biblio/5653266 (Accessed 7 May 2020), 1983.
- Hadley, O. L. and Kirchstetter, T. W.: Black-carbon reduction of snow albedo, Nat. Clim. Change, 2(6),
 437–440, doi:10.1038/nclimate1433, 2012.
- Hansen, J. and Nazarenko, L.: Soot climate forcing via snow and ice albedos, Proc. Natl. Acad. Sci.,
 101(2), 423–428, doi:10.1073/pnas.2237157100, 2004.
- Ingersoll, G.P., Don Campbell, M. Alisa Mast, David W. Clow, Leora Nanus, and Brent Frakes. 2009.
 Snowpack Chemistry Monitoring Protocol for the Rocky Mountain Network; Narrative and Standard
 Operating Procedures. United States Geological Service (USGS), Reston, Virginia. Administrative
 Report, 2009
- Jacobi, H.-W., Obleitner, F., Da Costa, S., Ginot, P., Eleftheriadis, K., Aas, W. and Zanatta, M.:
 Deposition of ionic species and black carbon to the Arctic snowpack: combining snow pit observations
 with modeling, 10361-10377, doi:10.5194/acp-19-10361-2019, 2019.
- Khan, A. L., Dierssen, H., Schwarz, J. P., Schmitt, C., Chlus, A., Hermanson, M., Painter, T. H. and
 McKnight, D. M.: Impacts of coal coarse mode from an active mine on the spectral reflectance of
 Arctic surface snow in Svalbard, Norway, J. Geophys. Res. Atmospheres, 122(3), 1767–1778,
 doi:10.1002/2016JD025757, 2017.
- Laborde, M., Crippa, M., Tritscher, T., Jurányi, Z., Decarlo, P. F., Temime-Roussel, B., Marchand, N.,
 Eckhardt, S., Stohl, A., Baltensperger, U., Prévôt, A. S. H., Weingartner, E. and Gysel, M.: Black
 carbon physical properties and mixing state in the European megacity Paris, Atmospheric Chem. Phys.,
 13(11), 5831–5856, doi:10.5194/acp-13-5831-2013, 2013.
- 774 Laj, P., Bigi, A., Rose, C., Andrews, E., Lund Myhre, C., Collaud Coen, M., Wiedensohler, A., Schultz, 775 M., Ogren, J. A., Fiebig, M., Gliß, J., Mortier, A., Pandolfi, M., Petäjä, T., Kim, S.-W., Aas, W., Putaud, 776 J.-P., Mayol-Bracero, O., Keywood, M., Labrador, L., Aalto, P., Ahlberg, E., Alados Arboledas, L., 777 Alastuey, A., Andrade, M., Artíñano, B., Ausmeel, S., Arsov, T., Asmi, E., Backman, J., Baltensperger, U., Bastian, S., Bath, O., Beukes, J. P., Brem, B. T., Bukowiecki, N., Conil, S., Couret, C., Day, D., 778 779 Dayantolis, W., Degorska, A., Santos, S. M. D., Eleftheriadis, K., Fetfatzis, P., Favez, O., Flentje, H., 780 Gini, M. I., Gregorič, A., Gysel-Beer, M., Hallar, G. A., Hand, J., Hoffer, A., Hueglin, C., Hooda, R. K., Hyvärinen, A., Kalapov, I., Kalivitis, N., Kasper-Giebl, A., Kim, J. E., Kouvarakis, G., Kranjc, I., 781 Krejci, R., Kulmala, M., Labuschagne, C., Lee, H.-J., Lihavainen, H., Lin, N.-H., Löschau, G., Luoma, 782 K., Marinoni, A., Meinhardt, F., Merkel, M., Metzger, J.-M., Mihalopoulos, N., Nguyen, N. A., 783 Ondracek, J., Peréz, N., Perrone, M. R., Petit, J.-E., Picard, D., Pichon, J.-M., Pont, V., Prats, N., Prenni, 784 A., Reisen, F., Romano, S., Sellegri, K., Sharma, S., Schauer, G., Sheridan, P., Sherman, J. P., Schütze, 785 786 M., Schwerin, A., Sohmer, R., Sorribas, M., Steinbacher, M., Sun, J., Titos, G., Tokzko, B., et al.: A

- global analysis of climate-relevant aerosol properties retrieved from the network of GAW near-surface
 observatories, Atmospheric Meas. Tech. Discuss., 1–70, doi: 10.5194/amt-2019-499, 2020.
- Law, K. S. and Stohl, A.: Arctic Air Pollution: Origins and Impacts, Science, 315(5818), 1537–1540,
 doi:10.1126/science.1137695, 2007.
- Lim, S., Fain, X., Zanatta, M., Cozic, J., Jaffrezo, J. L., Ginot, P. and Laj, P.: Refractory black carbon
 mass concentrations in snow and ice: method evaluation and inter-comparison with elemental carbon
 measurement, Atmospheric Meas. Tech., 7(10), 3307–3324, doi:10.5194/amt-7-3307-2014, 2014.
- Liu, J., Fan, S., Horowitz, L. W. and Levy, H.: Evaluation of factors controlling long-range transport of
 black carbon to the Arctic, J. Geophys. Res. Atmospheres, 116(D4), doi:10.1029/2010JD015145, 2011.
- Long J.A.. jtools: Analysis and Presentation of Social Scientific Data (2020). URL: https://cran.r project.org/package=jtools
- Lupi, A., Busetto, M., Becagli, S., Giardi, F., Lanconelli, C., Mazzola, M., Udisti, R., Hansson, H.-C.,
 Henning, T., Petkov, B., Ström, J., Krejci, R., Tunved, P., Viola, A. P. and Vitale, V.: Multi-seasonal
 ultrafine aerosol particle number concentration measurements at the Gruvebadet observatory, NyÅlesund, Svalbard Islands, Rendiconti Lincei, 27(1), 59–71, doi:10.1007/s12210-016-0532-8, 2016.
- Maturilli, M., Herber, A. and König-Langlo, G.: Climatology and Time Series of Surface Meteorology in
 Nylesund, Svalbard, Earth Syst. Sci. Data, 5, 155–163, doi: 10.5194/essd-5-155-2013, 2013.
- Maturilli, M., Herber, A. and König-Langlo, G.: Surface radiation climatology for Ny-Ålesund, Svalbard
 (78.9° N), basic observations for trend detection, Theor. Appl. Climatol., 120(1), 331–339,
 doi:10.1007/s00704-014-1173-4, 2015.
- Maturilli, M., Hanssen-Bauer, I., Neuber, R., Rex, M. and Edvardsen, K.: The Atmosphere Above NyÅlesund: Climate and Global Warming, Ozone and Surface UV Radiation, in The Ecosystem of
 Kongsfjorden, Svalbard, edited by H. Hop and C. Wiencke, pp. 23–46, Springer International
 Publishing, Cham., 2019.
- Meinander, O.; Heikkinen, E.; Aurela, M.; Hyvärinen, A. Sampling, Filtering, and Analysis Protocols to 811 812 Detect Black Carbon, Organic Carbon, and Total Carbon in Seasonal Surface Snow in an Urban 813 Background and Arctic Finland (>60° Atmosphere 923. N). 2020. 11. https://doi.org/10.3390/atmos11090923, 2020a. 814
- Moosmüller, H., Chakrabarty, R. K. and Arnott, W. P.: Aerosol light absorption and its measurement: A
 review, J. Quant. Spectrosc. Radiat. Transf., 110(11), 844–878, doi:10.1016/j.jqsrt.2009.02.035, 2009.
- Mori, T., Goto-Azuma, K., Kondo, Y., Ogawa-Tsukagawa, Y., Miura, K., Hirabayashi, M., Oshima, N.,
 Koike, M., Kupiainen, K., Moteki, N., Ohata, S., Sinha, P. R., Sugiura, K., Aoki, T., Schneebeli, M.,
 Steffen, K., Sato, A., Tsushima, A., Makarov, V., Omiya, S., Sugimoto, A., Takano, S. and Nagatsuka,
 N.: Black Carbon and Inorganic Aerosols in Arctic Snowpack, J. Geophys. Res. Atmospheres, 124(23),
 13325–13356, doi:10.1029/2019JD030623, 2019.
- Moroni, B., Becagli, S., Bolzacchini, E., Busetto, M., Cappelletti, D., Crocchianti, S., Ferrero, L., Frosini,
 D., Lanconelli, C., Lupi, A., Maturilli, M., Mazzola, M., Perrone, M. G., Sangiorgi, G., Traversi, R.,
 Udisti, R., Viola, A. and Vitale, V.: Vertical Profiles and Chemical Properties of Aerosol Particles upon
 Ny-Ålesund (Svalbard Islands), Adv. Meteorol., 2015, e292081, doi: 10.1155/2015/292081, 2015.

- 826 Moroni, B., Arnalds, O., Dagsson-Waldhauserová, P., Crocchianti, S., Vivani, R. and Cappelletti, D.:
- 827 Mineralogical and Chemical Records of Icelandic Coarse mode Sources Upon Ny-Ålesund (Svalbard
- 828 Islands), Front. Earth Sci., 6, doi:10.3389/feart.2018.00187, 2018.
- Moroni, B., Ritter, C., Crocchianti, S., Markowicz, K., Mazzola, M., Becagli, S., et al.. Individual particle
 characteristics, optical properties and evolution of an extreme long-range transported biomass burning
- event in the European Arctic (Ny-Ålesund, Svalbard Islands). Journal of Geophysical Research:
- Atmospheres, 125, e2019JD031535, doi:10.1029/2019JD031535, 2020.
- 833
- Motos, G., Schmale, J., Corbin, J. C., Modini, R. L., Karlen, N., Bertò, M., Baltensperger, U. and GyselBeer, M.: Cloud droplet activation properties and scavenged fraction of black carbon in liquid-phase
 clouds at the high-alpine research station Jungfraujoch (3580 m a.s.l.), Atmospheric Chem. Phys., 19(6),
 3833–3855, doi:10.5194/acp-19-3833-2019, 2019.
- Osmont, D., Wendl, I. A., Schmidely, L., Sigl, M., Vega, C. P., Isaksson, E. and Schwikowski, M.: An
 800-year high-resolution black carbon ice core record from Lomonosovfonna, Svalbard, Atmospheric
 Chem. Phys., 18(17), 12777–12795, doi: 10.5194/acp-18-12777-2018, 2018.
- Pedersen, C. A., Gallet, J.-C., Ström, J., Gerland, S., Hudson, S. R., Forsström, S., Isaksson, E. and
 Berntsen, T. K.: In situ observations of black carbon in snow and the corresponding spectral surface
 albedo reduction, J. Geophys. Res. Atmospheres, 120(4), 1476–1489, doi: 10.1002/2014JD022407,
 2015.
- Perovich, D. : Light reflection and transmission by a temperate snow cover. *Journal of Glaciology*,
 53(181), 201-210. doi:10.3189/172756507782202919, 2007
- 847
- Petzold, A., Ogren, J. A., Fiebig, M., Laj, P., Li, S.-M., Baltensperger, U., Holzer-Popp, T., Kinne, S.,
 Pappalardo, G., Sugimoto, N., Wehrli, C., Wiedensohler, A. and Zhang, X.-Y.: Recommendations for
 reporting "black carbon" measurements, Atmospheric Chem. Phys., 13(16), 8365–8379, doi:
 10.5194/acp-13-8365-2013, 2013.
- R Core Team. R: A language and environment for statistical computing. R Foundation for Statistical
 Computing, Vienna, Austria, (2020). URL: https://www.R-project.org/
- Ruppel, M. M., Soares, J., Gallet, J.-C., Isaksson, E., Martma, T., Svensson, J., Kohler, J., Pedersen, C.
 A., Manninen, S., Korhola, A. and Ström, J.: Do contemporary (1980–2015) emissions determine the
 elemental carbon deposition trend at Holtedahlfonna glacier, Svalbard?, Atmospheric Chem. Phys.,
 17(20), 12779–12795, doi: 10.5194/acp-17-12779-2017, 2017.
- Scalabrin, E., Zangrando, R., Barbaro, E., Kehrwald, N. M., Gabrieli, J., Barbante, C. and Gambaro, A.:
 Amino acids in Arctic aerosols, Atmospheric Chem. Phys., 12(21), 10453–10463, doi:10.5194/acp-1210453-2012, 2012.
- Schmale, J., Arnold, S. R., Law, K. S., Thorp, T., Anenberg, S., Simpson, W. R., Mao, J. and Pratt, K. A.:
 Local Arctic Air Pollution: A Neglected but Serious Problem, Earths Future, 6(10), 1385–1412,
 doi:10.1029/2018EF000952, 2018.
- Schwarz, J. P., Gao, R. S., Perring, A. E., Spackman, J. R. and Fahey, D. W.: Black carbon aerosol size in snow, Sci. Rep., 3(1), 1–5, doi:10.1038/srep01356, 2013.

- Screen, J. A. and Simmonds, I.: The central role of diminishing sea ice in recent Arctic temperature
 amplification, Nature, 464(7293), 1334–1337, doi:10.1038/nature09051, 2010.
- Segura, S., Estellés, V., Titos Vela, G., Lyamani, H., Utrilla Navarro, P., Zotter, P., Prévot, A. S. H.,
 Močnik, G., Alados-Arboledas, L. and Martínez-Lozano, J. A.: Determination and analysis of in situ
 spectral aerosol optical properties by a multi-instrumental approach, doi:10.5194/amt-7-2373-2014,
 2014.
- 872 Serreze, M. C. and Barry, R. G.: Processes and impacts of Arctic amplification: A research synthesis,
 873 Glob. Planet. Change, 77(1), 85–96, doi:10.1016/j.gloplacha.2011.03.004, 2011.
- Sharma, S., W. Richard Leaitch, Lin Huang, Daniel Veber, Felicia Kolonjari, Wendy Zhang. An
 evaluation of three methods for measuring black carbon in Alert, Canada. Atmos. Chem. Phys., 17,
 15225-15243, https://doi.org/10.5194/acp-17-15225-2017, 2017.
- Sinha, P. R., Y. Kondo, M. Koike, J. Ogren, A. Jefferson, T. Barrett, R. Sheesley, S. Ohata, N. Moteki, H.
 Coe, D. Liu, M. Irwin, P. Tunved, P. K. Quinn, and Y. Zhao, Evaluation of ground-based black carbon
 measurements by filter-based photometers at two Arctic sites, J. Geophys. Res., 122,
 doi:10.1002/2016JJD025843, 2017.
- Sinha, P. R., Kondo, Y., Goto-Azuma, K., Tsukagawa, Y., Fukuda, K., Koike, M., Ohata, S., Moteki, N.,
 Mori, T., Oshima, N., Førland, E. J., Irwin, M., Gallet, J.-C. and Pedersen, C. A.: Seasonal Progression
 of the Deposition of Black Carbon by Snowfall at Ny-Ålesund, Spitsbergen: Deposition of Black
 Carbon in Spitsbergen, J. Geophys. Res. Atmospheres, 123(2), 997–1016, doi:10.1002/2017JD028027,
 2018.
- Skiles, S. M. and Painter, T. H.: Toward Understanding Direct Absorption and Grain Size Feedbacks by
 Coarse mode Radiative Forcing in Snow With Coupled Snow Physical and Radiative Transfer
 Modeling, Water Resour. Res., 55(8), 7362–7378, doi:10.1029/2018WR024573, 2019.
- Skiles, S. M., Flanner, M., Cook, J. M., Dumont, M. and Painter, T. H.: Radiative forcing by light-absorbing particles in snow, Nat. Clim. Change, 8(11), 964–971, doi:10.1038/s41558-018-0296-5, 2018.
- Spolaor, A., Angot, H., Roman, M., Dommergue, A., Scarchilli, C., Vardè, M., Del Guasta, M., Pedeli,
 X., Varin, C., Sprovieri, F., Magand, O., Legrand, M., Barbante, C. and Cairns, W. R. L.: Feedback
 mechanisms between snow and atmospheric mercury: Results and observations from field campaigns on
 the Antarctic plateau, Chemosphere, 197, 306–317, doi:10.1016/j.chemosphere.2017.12.180, 2018.
- Spolaor, A., Barbaro, E., Cappelletti, D., Turetta, C., Mazzola, M., Giardi, F., Björkman, M. P.,
 Lucchetta, F., Dallo, F., Pfaffhuber, K. A., Angot, H., Dommergue, A., Maturilli, M., Saiz-Lopez, A.,
 Barbante, C. and Cairns, W. R. L.: Diurnal cycle of iodine, bromine, and mercury concentrations in
 Svalbard surface snow, Atmospheric Chem. Phys., 19(20), 13325–13339, doi: 10.5194/acp-19-133252019, 2019.
- Stephens, M., Turner, N. and Sandberg, J.: Particle identification by laser-induced incandescence in a solid-state laser cavity, Appl. Opt., 42(19), 3726–3736, doi:10.1364/AO.42.003726, 2003.
- Stohl, A., Klimont, Z., Eckhardt, S., Kupiainen, K., Shevchenko, V. P., Kopeikin, V. M., and Novigatsky,
 A. N.: Black carbon in the Arctic: the underestimated role of gas flaring and residential combustion
 emissions., Atmospheric Chem. Phys., 13(17), 8833-8855, doi: 10.5194/acp-13-8833-2013, 2013.

Tunved, P., Ström, J. and Krejci, R.: Arctic aerosol life cycle: linking aerosol size distributions observed
between 2000 and 2010 with air mass transport and precipitation at Zeppelin station, Ny-Ålesund,
Svalbard, Atmospheric Chem. Phys., 13(7), 3643–3660, doi: 10.5194/acp-13-3643-2013, 2013.

908 Turetta, C., Feltracco, M., Barbaro, E., Spolaor, A., Barbante, C., & Gambaro, A.: A Year-Round
909 Measurement of Water-Soluble Trace and Rare Earth Elements in Arctic Aerosol: Possible Inorganic
910 Tracers of Specific Events, Atmosphere, 12(6), 694, doi: 10.3390/atmos12060694, 2021.

- 911 Vecchiato, M., Barbaro, E., Spolaor, A., Burgay, F., Barbante, C., Piazza, R. and Gambaro, A.,
 912 Fragrances and PAHs in snow and seawater of Ny-Ålesund (Svalbard): Local and long-range
 913 contamination. Environmental Pollution 242, 1740-1747, doi: 10.1016/j.envpol.2018.07.095, 2018.
- Weingartner, E., Saathoff, H., Schnaiter, M., Streit, N., Bitnar, B. and Baltensperger, U.: Absorption of
 light by soot particles: determination of the absorption coefficient by means of aethalometers, J. Aerosol
 Sci., 34(10), 1445–1463, doi:10.1016/S0021-8502(03)00359-8, 2003.
- Wendl, I. A., Menking, J. A., Färber, R., Gysel, M., Kaspari, S. D., Laborde, M. J. G. and Schwikowski,
 M.: Optimized method for black carbon analysis in ice and snow using the Single Particle Soot
 Photometer, Atmospheric Meas. Tech., 7, 2667–2681, doi:10.5194/amt-7-2667-2014, 2014.
- **920** Xu, B., T. Yao, X. Liu, and N. Wang, Elemental and organic carbon measurements with a two-step
- heating gas chromatography system in snow samples from the Tibetan Plateau, Ann. Glaciol., 43, 257–
 262, doi: 10.3189/172756406781812122, 2006.
- Yasunari, T. J., Tan, Q., Lau, K.-M., Bonasoni, P., Marinoni, A., Laj, P., Ménégoz, M., Takemura, T. and
 Chin, M.: Estimated range of black carbon dry deposition and the related snow albedo reduction over
 Himalayan glaciers during dry pre-monsoon periods, Atmos. Environ., 78, 259–267,
 doi:10.1016/j.atmosenv.2012.03.031, 2013.
- Zanatta, M., Gysel, M., Bukowiecki, N., Müller, T., Weingartner, E., Areskoug, H., Fiebig, M., Yttri, K.
 E., Mihalopoulos, N., Kouvarakis, G., Beddows, D., Harrison, R. M., Cavalli, F., Putaud, J. P., Spindler,
 G., Wiedensohler, A., Alastuey, A., Pandolfi, M., Sellegri, K., Swietlicki, E., Jaffrezo, J. L.,
 Baltensperger, U. and Laj, P.: A European aerosol phenomenology-5: Climatology of black carbon
 optical properties at 9 regional background sites across Europe, Atmos. Environ., 145, 346–364,
 doi:10.1016/j.atmosenv.2016.09.035, 2016.
- Zanatta, M., Laj, P., Gysel, M., Baltensperger, U., Vratolis, S., Eleftheriadis, K., Kondo, Y., Dubuisson,
 P., Winiarek, V., Kazadzis, S., Tunved, P. and Jacobi, H.-W.: Effects of mixing state on optical and
 radiative properties of black carbon in the European Arctic, Atmospheric Chem. Phys., 18(19), 14037–
 14057, doi: 10.5194/acp-18-14037-2018, 2018.
- Zangrando, R., Barbaro, E., Zennaro, P., Rossi, S., Kehrwald, N. M., Gabrieli, J., Barbante, C. and
 Gambaro, A.: Molecular Markers of Biomass Burning in Arctic Aerosols, Environ. Sci. Technol.,
 47(15), 8565–8574, doi:10.1021/es400125r, 2013.