



1	Aerosol optical properties calculated from size distributions, filter samples and absorption photometer data
2	at Dome C, Antarctica and their relationships between seasonal cycles of sources
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21	Abstract
22	Optical properties of surface aerosols at Dome C, Antarctica in 2007-2013 and their potential source areas are
23	presented. Scattering coefficients ( $\sigma_{sp}$ ) were calculated from measured particle number size distributions with a
24	Mie code and from filter samples using mass scattering efficiencies. Absorption coefficients ( $\sigma_{ap}$ ) were
25	determined with a 3-wavelength Particle Soot Absorption Photometer (PSAP) and corrected for scattering by
26	using two different algorithms. The scattering coefficients were also compared with $\sigma_{sp}$ measured with a
27	nephelometer at the South Pole Station (SPO). The minimum $\sigma_{\sf ap}$ was observed in the austral autumn and the
28	maximum in the austral spring, similar to other Antarctic sites. The darkest aerosol, i.e., the lowest single
29	scattering albedo $\omega_o \approx$ 0.91 was observed in September and October and the highest $\omega_o \approx$ 0.99 in February and
30	March. The uncertainty of the absorption Ångström exponent $lpha_{ap}$ is high. The lowest $lpha_{ap}$ monthly medians were
31	observed in March and the highest in August – October. The equivalent black carbon (eBC) mass concentrations
32	were compared with eBC measured at three other Antarctic sites: the SPO and two coastal sites, Neumayer and





1 Syowa. The maximum monthly median eBC concentrations are almost the same ( $^{3} \pm 1$  ng m<sup>-3</sup>) at all these sites 2 in October-November. This suggests that there is no significant difference in eBC between the coastal and 3 plateau sites. The seasonal cycle of the eBC mass fraction exhibits a minimum f(eBC) ≈ 0.1% in February-March 4 and a maximum ~4-5% in August-October. Source areas were calculated using 50-day FLEXPART footprints. The 5 highest eBC concentrations and the lowest  $\omega_o$  were associated with air masses coming from South America, 6 Australia and Africa. Vertical simulations that take BC particle removal processes into account show that there 7 would be essentially no BC particles arriving at Dome C from north of latitude 10°S at altitudes < 1600 m. The 8 main biomass-burning regions Africa, Australia and Brazil are more to the south and their smoke plumes have 9 been observed at higher altitudes than that so they can get transported to Antarctica. The seasonal cycle of BC 10 emissions from wildfires and agricultural burning and other fires in South America, Africa and Australia were 11 calculated from data downloaded from the Global Fire Emissions Database (GFED). The maximum total emissions 12 were in August-September but the peak of monthly average eBC concentrations is observed 2 – 3 months later 13 in November not only at Dome C but also at SPO and the coastal stations. The air mass residence-time-weighted 14 BC emissions from South America are approximately an order of magnitude larger than from Africa and Oceania 15 suggesting that South American BC emissions are the largest contributors to eBC at Dome C. At Dome C the 16 maximum and minimum scattering coefficients were observed in austral summer and winter, respectively. At 17 SPO  $\sigma_{sp}$  was similar to that observed at Dome C in the austral summer but there was a large difference in winter, 18 suggesting that in winter SPO is more influenced by sea spray emissions than Dome C. The seasonal cycles of  $\sigma_{sp}$ 19 at Dome C and at the SPO were compared with the seasonal cycles of secondary and primary marine aerosol 20 emissions. The  $\sigma_{sp}$  measured at SPO correlated much better with the sea-spray aerosol emission fluxes in the 21 Southern Ocean than  $\sigma_{sp}$  at Dome C. The seasonal cycles of biogenic secondary aerosols were estimated from 22 monthly average phytoplankton biomass concentrations obtained from the CALIOP satellite sensor data. The 23 analysis suggests that a large fraction of the biogenic scattering aerosol observed at Dome C has been formed in 24 the polar zone but it may take a month for the aerosol to be formed, grown and get transported from the sea 25 level to Dome C.

26

# 27 1. Introduction

The Antarctic interior region has scarce observations of atmospheric constituents and many aspects of the atmospheric properties are underdetermined. The Antarctic dome or the polar vortex, which is much stronger than its northern counterpart and present throughout the year (Karpetchko et al., 2005), at most times efficiently prevents transport into the Antarctic troposphere from lower latitudes. However, wildfires and agricultural burning emissions from Africa, South America and Australia do affect vast regions of the southern hemisphere,





including Antarctica. For instance, Hara et al. (2010) found that haze episodes at Syowa Station, during which
visibility can drop to 10 km for periods of ~30 h, were caused by biomass burning aerosol from South America
transported to the Antarctic coast via the eastward approach of cyclones. At the Neumayer station large-scale
meridional transport of biomass-burning derived black carbon, preferentially from South America, seems to
determine the BC burden and causes a distinct and consistent spring / early summer concentration maximum
(Weller et al., 2013).

7

8 Concordia station lies on Dome C (75°06'S, 123°23'E), at 3233 m above sea level (.a.s.l.) on the East Antarctic 9 plateau, about 1100 km from the nearest coastline, the Ross Sea. The base is French / Italian operated, with 10 research fields within astronomy and glaciology as well as atmospheric sciences. The atmospheric 11 instrumentation is located in a small cabin southwest of the main base (at the site described by Udisti et al., 12 2012) where it is upwind of the base at the prevailing wind directions. Concordia is one of only three permanent 13 year-round stations operated on the Antarctic Plateau, the others being the American Amundsen-Scott 14 observatory (South Pole (SPO), 2835 m.a.s.l., about 1300 km from the nearest open sea, 1600 km away from Dome C) and the Russian Vostok station (78°28'S, 106°51'E, 3488 m.a.s.l., 600km away). Thus, there are large 15 16 spatial distances between the continuous atmospheric observation. However, properties of the Antarctic 17 atmosphere tend to extend both over longer temporal and spatial scales than elsewhere (Fiebig et al., 2014) 18 suggesting that the scarce observations that exist can be assumed to be representative of larger areas than 19 typical in other climate regions. This would imply that Dome C is an important indicator for the entire Antarctic 20 inland. Though measurement conditions are harsh the continuous long-term monitoring provided here can be a 21 baseline for the aerosol optical properties of the Antarctic inland and may provide indications of changes in 22 atmospheric constituents and aerosol levels.

23

24 There are several studies on the aerosol chemical composition at Dome C (e.g., Jourdain et al., 2009; Becagli et 25 al.,2012; Udisti et al., 2012; Legrand et al., 2016, 2017a, 2017b), and also the aerosol optical depth (AOD) has 26 been measured there (Tomasi et al., 2007). However, in situ surface aerosol scattering and absorption 27 coefficients at Dome C have not been presented. The light absorption coefficient and particle number size 28 distributions (PNSD) have been measured continuously with a 3-wavelength Particle Soot Absorption 29 Photometer (PSAP) and a differential mobility particle sizer (DMPS) since 2007. The PNSD data have already been 30 used in several papers. Järvinen et al. (2013) analyzed the seasonal cycle and modal structure of PNSD measured 31 with the DMPS, Chen et al. (2017) analyzed number size distribution of air ions measured with an Air Ion 32 Spectrometer (AIS) and the PNSD measured with the DMPS and Lachlan-Cope et al. (2020) used the Dome C





- DMPS data for comparing with the PNSD measured at the coastal site Halley. The PSAP data, however, have not been presented in detail. Caiazzo et al. (2021) used some of the PSAP data mainly for evaluating elemental carbon (EC) sample contamination. Grythe (2017) used the data from 2007-2013 as part of his PhD thesis but in the present paper we will analyze that period in more detail. Here we will describe the methods for measuring absorption and calculating scattering from the size distributions and filter samples.
- 6
- 7 The goals of the paper are to present descriptive statistics of extensive and intensive aerosol optical properties 8 at Dome C in 2007 – 2013, their seasonal cycles and the relationships between the seasonal cycles of major 9 sources of absorbing and scattering aerosols. The AOPs will be compared with other observations from other 10 Antarctic sites, in most detail the scattering coefficients measured at the South Pole.
- 11

# 12 **2. Methods**

# 13 2.1 Sampling site

- 14 Concordia station is a permanently operated French / Italian Antarctic research base on East Antarctic plateau. 15 The observations are performed in isolated sites around the main base. The Dome C sampling site is the same as 16 used by Udisti et al. (2012), Becagli et al. (2012), and Järvinen et al. (2013). It is located about 1 km southwest of 17 the station main buildings, upwind in the direction of the prevailing wind. The northeastern direction (10°-90°) 18 has been declared as the contaminated sector. Below the validity of the contaminated sector will be analyzed by 19 using the absorption photometer data. For in situ aerosol instrumentation the sample air was taken at the 20 flowrate of 5 Liters Per Minute (LPM) from the roof of the cabin with a straight 2-m long 25-mm diameter 21 stainless steel tube inlet. It was covered with a protective cap to protect against snow fall and ice buildup.
- 22

## 23 2.2 Instruments

# 24 2.2.1 Aerosol measurements

25 Light absorption by particles was measured with a Radiance Research  $3\lambda$  PSAP at three wavelengths,  $\lambda = 467$  nm, 26 530 nm, and 660 nm. There was no nephelometer measuring scattering coefficient so it was calculated from 27 particle size distributions and filter sample data as described below. Particle number size distributions were 28 measured at 10-minute time resolution in the size range 10 - 620 nm with a custom-built differential mobility 29 particle sizer (DMPS) as described by Järvinen et al. (2013) and in the size range  $0.3 - 20 \,\mu$ m with a Grimm model 30 1.108 optical particle counter (OPC) in 2007 – 2009. In addition to the in-situ instruments, PM<sub>1</sub> and PM<sub>10</sub> filter 31 samples were collected for chemical analyses by ion chromatography. The length of the sampling period of the 32 PM<sub>1</sub> and PM<sub>10</sub> samples was 3 or 4 days and 1 day, respectively.





- The data coverage for the PSAP, the DMPS, the OPC and the PM<sub>1</sub> and PM<sub>10</sub> filter sample data are presented in
   Fig. 1. The number of hours of accepted data and the number of samples are shown in parentheses for the
- 3 continuous instruments and the filter samplers, respectively. The filtering criteria will be presented below
- 4 (section 2.4).
- 5

# 6 2.2.2 Meteorological measurements

- 7 Ambient air temperature (t), relative humidity (RH), wind speed (WS) and wind direction (WD) data were from
- 8 the routine meteorological observation at Station Concordia as part of the IPEV/PNRA Project a collaborative
- 9 project between "Programma Nazionale di Ricerche in Antartide" (PNRA) and Institut Polaire Français Paul-Emile
- 10 Victor (IPEV) (<u>www.climantartide.it</u>).
- 11

# 12 2.3 Data processing

# 13 2.3.1 Mass concentrations from size distributions

- 14 60-minute average size distributions  $n(D_p)$  were first calculated from the original 10-minute data and corrected
- 15 for STP (p = 1013 hPa, T = 273.15 K). Mass concentration were calculated from the number size distributions
- 16 measured with the DMPS from

17 
$$m(DMPS) = \rho_p V(DMPS) = \rho_p \int_{10nm}^{620nm} \frac{\pi}{6} D_p^3 n(D_p) dD_p$$
 (1)

where the density  $\rho_p = 1.7 \text{ g cm}^{-3}$  was used. For particle density of 1.7 g cm<sup>-3</sup> the particle diameter 620 nm corresponds to the aerodynamic diameter  $D_a = \sqrt{\rho_p / \rho_0} D_p = \sqrt{1.7} \times 620 \text{ nm} \approx 808 \text{ nm}$ , where  $\rho_0 = 1 \text{ g cm}^{-3}$ . To be consistent with the definitions of filter-sample size ranges that typically show the upper aerodynamic diameter of a sampler inlet the mass concentration calculated from Eq. (1) will be referred to as m(DMPS,PM<sub>0.8</sub>) and the volume concentration as V(DMPS,PM<sub>0.8</sub>).

23

In December 2007 – July 2009 there was also the Grimm 1.108 OPC that measures number concentrations of particles in the D<sub>p</sub> range of 0.3 – 20  $\mu$ m. The three largest channels of the OPC measure the number concentrations in D<sub>p</sub> range of 7.5 – 20  $\mu$ m. For an assumed density  $\rho_p = 1.7$  g cm<sup>-3</sup> the diameter D<sub>p</sub> = 7.5  $\mu$ m corresponds to the aerodynamic diameter D<sub>a</sub> = 9.8  $\mu$ m. Assuming that  $\rho_p$  is constant over the whole size range the mass concentration of particles smaller than D<sub>a</sub> = 10  $\mu$ m is calculated from the number size distributions by excluding the three largest particle OPC channels as

30 
$$m(n(D_p), PM_{10}) = \rho_p V(n(D_p), PM_{10}) = \rho_p (V(DMPS, PM_{0.8}) + V(OPC, PM_{0.8-10}))$$
 (2)





1 The fraction of volume concentration measured by the DMPS equals

2 
$$fV(DMPS) = \frac{V(DMPS, PM_{0.8})}{V(n(D_p), PM_{10})}$$
 (3)

3 This fraction was calculated from data collected during the simultaneous operation of the DMPS and the OPC.

- 4 The monthly average fV(DMPS) values presented in Table 1 were used for the period 2008 2013 to calculate
- 5 mass concentrations in the size range  $D_a < 10 \ \mu m$  from

6 
$$m(DMPS, PM_{10}) = \frac{\rho_p V(DMPS, PM_{0.8})}{fV(DMPS)} = \frac{m(DMPS, PM_{0.8})}{fV(DMPS)}$$
(4)

In other words, the variable names m(DMPS,PM<sub>0.8</sub>) and m(DMPS,PM<sub>10</sub>) will be used below to emphasize that
 these mass concentrations were calculated from DMPS data. The mass concentrations m(DMPS,PM<sub>0.8</sub>) and
 m(DMPS,PM<sub>10</sub>) can be considered to be the lower and upper estimates of m.

10

### 11 2.3.2 Scattering coefficients from the size distributions

12 Scattering coefficients were calculated using the 60-minute average size distributions from

13 
$$\sigma_{\rm sp}({\rm m},\lambda) = \int Q_{\rm s}({\rm D}_{\rm p},{\rm m},\lambda)\frac{\pi}{4}{\rm D}_{\rm p}^2{\rm n}({\rm D}_{\rm p}){\rm d}{\rm D}_{\rm p}$$
(5)

where  $Q_s$  is the scattering efficiency calculated using the Mie code by Barber and Hill (1990), m is the refractive index,  $\lambda$  is the wavelength and  $n(D_p)$  is the particle number size distribution. Analogous to the mass concentrations the scattering coefficients were determined from the simultaneous DMPS and OPC measurements in December 2007 – July 2009 from

18 
$$\sigma_{sp}(n(D_p), PM_{10}) = \sigma_{sp}(DMPS, PM_{0.8}) + \sigma_{sp}(OPC, PM_{0.8-10})$$
 (6)

19 where  $\sigma_{sp}(OPC, PM_{0.8})$  and  $\sigma_{sp}(OPC, PM_{0.8-10})$  are the scattering coefficient calculated from the particle number size 20 distributions in the size ranges measured by the DMPS and the OPC, respectively. For  $\sigma_{sp}$ (OPC,PM<sub>0.8</sub>) the 21 refractive index of sulfuric acid (SA, H<sub>2</sub>SO<sub>4</sub>, m<sub>r</sub> = 1.426 + 0i, Seinfeld and Pandis, 1998) was used. This refractive 22 index is slightly lower than that estimated for submicron aerosols at two low-altitude Antarctic stations Aboa, 23 and Neumayer in Queen Maud Land. Virkkula et al. (2006) measured particle number size distributions in the 24 size range  $D_p < 800$  nm with a DMPS and light scattering of submicron particles with a nephelometer at the 25 Finnish a site about 130 km inland from the open Weddell Sea in January 2000. With an iteration procedure 26 matching nephelometer-measured and size-distribution-derived scattering coefficients the real refractive indices 27 were 1.43  $\pm$  0.07 and 1.45  $\pm$  0.04 at  $\lambda$  = 550 nm for all data and excluding new particle formation, respectively. 28 Jurányi and Weller (2019) measured size distributions with an SMPS and a laser aerosol spectrometer (LAS) for 29 a full year at the coastal site Neumayer and by fitting data of the two instruments in the overlapping range of





1 120 - 340 nm obtained  $m_r$  = 1.44 ± 0.08. Considering that both Aboa and Neumayer are closer to sources of 2 ammonia that neutralizes aerosol and increases the refractive index above that of pure sulfuric acid (1.426) it 3 was assumed here that the use of 1.426 for the calculation of  $\sigma_{sp}$  from the size range measured with the DMPS 4 is reasonable. For the larger particle size range,  $\sigma_{sp}(OPC, PM_{0.8-10})$  the refractive index of NaCl (m<sub>r</sub> = 1.544, Seinfeld 5 and Pandis, 1998) was used. This value is in line with the average refractive index of 1.54 with a range from 1.50 6 to 1.58 in the particle size range 0.3 - 12  $\mu$ m in impactor samples taken at the South Pole (Hogan et al., 1979) 7 and with the supermicron particle refractive index of  $1.53 \pm 0.02$  calculated from the chemical composition of 8 12-stage impactor samples taken at the coastal site Aboa (Virkkula et al., 2006).

9

10 The fraction of scattering coefficient measured by the DMPS was calculated from

11 
$$f\sigma_{sp}(DMPS,\lambda) = \frac{\sigma_{sp}(DMPS, PM_{0.8}, \lambda)}{\sigma_{sp}(n(D_p), PM_{10}, \lambda)}$$
(7)

12 The wavelengths of  $\lambda$  = 467 nm, 530 nm, and 660 nm were used to match the PSAP data. Similar to fV(DMPS), 13 fo<sub>sp</sub>(DMPS, $\lambda$ ) was calculated from data collected during the simultaneous operation of the DMPS and the OPC, 14 the seasonal monthly statistics were calculated (Table 1) and the respective monthly averages were applied to

15 the period 2008 – 2013 to calculate  $\sigma_{sp}$  in the size range  $D_a < 10 \ \mu m$  from

16 
$$\sigma_{\rm sp}(\rm DMPS, \rm PM_{10}, \lambda) = \frac{\sigma_{\rm sp}(\rm DMPS, \rm PM_{0.8}, \lambda)}{f\sigma_{\rm sp}(\rm DMPS, \lambda)}$$
(8)

17 The wavelength symbol  $\lambda$  will be used below only when necessary. The variable names  $\sigma_{sp}(DMPS,PM_{0.8})$  and 18  $\sigma_{sp}(DMPS,PM_{10})$  will be used to emphasize that these scattering coefficients were calculated from DMPS data in 19 the aerodynamic particle size ranges  $D_a < 0.8 \mu m$  and  $D_a < 10 \mu m$ . The scattering coefficients  $\sigma_{sp}(DMPS,PM_{0.8})$ 20 and  $\sigma_{sp}(DMPS,PM_{10})$  can also be considered to be the lower and upper estimates of  $\sigma_{sp}$  at the given wavelength. 21

22 Figure 2 shows the average particle number, volume and scattering size distributions at  $\lambda$  = 530 nm in the size 23 range 10 nm - 10  $\mu$ m and the respective normalized cumulative size distributions in the size range of 10 nm -24 7.5 μm during the period from 14 December 2007 to 14 July 2009 in summer and in winter. Fig. 2a and 2b show 25 that for the number concentrations the OPC size range plays an insignificant role whereas the larger particles 26 contribute significantly to both total particle volume concentration (Figs. 2c and 2d) and scattering coefficients 27 (Figs. 2e and 2f) and that this contribution varies seasonally. The contributions of fV(DMPS) and  $f\sigma_{sp}$ (DMPS, $\lambda$ ) 28 were calculated for hourly-averaged size distributions from Eqs. (3) and (7), the monthly seasonal statistics were 29 calculated and presented in Table 1 and as a boxplot in Fig. 3. Both the table and the boxplot show that both 30 fV(DMPS) and  $f\sigma_{sp}$ (DMPS, $\lambda$ ) have maxima in summer and minima in winter. They also show that the ranges are





- 1 large. Consequently the use of the monthly averages presented in Table 1 for calculating m(DMPS,PM<sub>10</sub>) and 2  $\sigma_{sp}$ (DMPS,PM<sub>10</sub>), Eqs. (4) and (8), creates an additional uncertainty to the results. Another important result is that
- 3 the wavelength dependency of  $f\sigma_{sp}(\text{DMPS},\!\lambda)$  is clear and it also has a seasonal cycle.
- 4

5 The wavelength dependency of the scattering coefficient can be described by the scattering Ångström exponent

$$6 \qquad \alpha_{sp} = -\frac{\ln(\sigma_{sp}(\lambda_1)/\sigma_{sp}(\lambda_2))}{\ln(\lambda_1/\lambda_2)}$$
(9)

9

#### 10 2.3.3 Absorption coefficients and equivalent black carbon concentrations

The PSAP data were first corrected for flow and spot size. The flow was calibrated 37 times during 2007 - 2013 with a TSI flow meter. The slopes and offsets of the calibrations were interpolated for each hour and the PSAP flows were corrected accordingly. All absorption coefficients were corrected to STP (1013.25 hPa and 273.15 K).

The PSAP measures signal and reference detector counts and the respective sums,  $\sum SIG$  and  $\sum REF$  are used for calculating non-scattering-corrected absorption coefficient, here  $\sigma_{ap,nsc}$ , from

17
$$\sigma_{ap,nsc} = f(Tr) \frac{A}{Q\Delta t} \ln \left( \frac{\left( \sum_{\Delta t} SIG / \sum_{\Delta t} REF \right)_{t,\Delta t}}{\left( \sum_{\Delta t} SIG / \sum_{\Delta t} REF \right)_{t}} \right) = f(Tr)\sigma_{0}$$
(10)

18 where A is the filter spot area, Q the flow rate,  $Tr = (\Sigma SIG/\Sigma REF)_t/(\Sigma SIG/\Sigma REF)_{t=0}$  is the transmittance, f(Tr) the 19 loading correction function and  $\Delta t$  the count integration time. The PSAP reports  $\sigma_{ap,nsc}$  with a 0.1 Mm<sup>-1</sup> resolution 20 at a 1-second time resolution. Averaging the 1-sec data is not good enough since at Dome C absorption 21 coefficients are most of the time clearly lower than 0.1 Mm<sup>-1</sup>. Therefore the signal and reference counts  $\Sigma SIG$  and 22  $\Sigma REF$  were used in (10) with  $\Delta t = 60$  min. Manufacturer-cut spots of the standard filter material Pallflex E70-23 2075W were used in the PSAP. The spot diameter was measured to be 4.9 ± 0.1 mm, so the spot area A was 18.9 24 ± 0.6 mm<sup>2</sup>. The uncertainty of A is ~3%.

25

Transmittance is reduced mainly by light absorption but also also due to scattering aerosol which results in the so-called apparent absorption and has to be taken into account in the data processing. There are different algorithms for processing PSAP data, e.g. by Bond et al. (1999), Virkkula et al. (2005), Müller et al. (2014), and Li





1 et al. (2020). Here we will use both the algorithm presented by Bond et al. (1999) (here B1999) with the

2 adjustment presented by Ogren (2010):

$$3 \qquad \sigma_{ap} = \frac{1}{1.22} \left( \frac{0.97 \cdot 0.873}{1.0796 \cdot Tr + 0.71} \sigma_0 - 0.02 \cdot \sigma_{sp} \right) = \frac{1}{1.5557 \cdot Tr + 1.0227} \sigma_0 - 0.0164 \cdot \sigma_{sp} = \sigma_{ap,nsc} - 0.0164 \cdot \sigma_{sp}$$
(11)

and the algorithm presented by Virkkula et al. (2005) with the constants updated by Virkkula (2010) (here
 V2010):

$$6 \qquad \sigma_{ap} = \left(k_0 + k_1(h_0 + h_1\omega_0)\ln(Tr)\right)\sigma_0 - s \cdot \sigma_{sp} \tag{12}$$

7 where

$$\omega_0 = \sigma_{sp} / (\sigma_{sp} + \sigma_{ap}) \tag{13}$$

9 is the single-scattering albedo and  $k_0$ ,  $k_1$ ,  $h_0$ ,  $h_1$ , and s are wavelength-dependent constants. In the rest of the 10 paper the symbol  $\sigma_{ap,nsc}$  will be used to present the non-scattering-corrected absorption coefficient, corrected 11 with the constants and formula in Eq. (11) excluding the subtraction of  $\sigma_{sp}$ .

12

8

13 Since there are the above-explained size-dependent uncertainties of scattering coefficient, additional absorption 14 coefficient estimates were calculated by using both algorithms. The upper estimates of absorption coefficients 15  $\sigma_{ap}(\sigma_{sp}(DMPS,PM_{0.8}))$  were calculated by using the lower estimate of scattering coefficient  $\sigma_{sp} = \sigma_{sp}(DMPS,PM_{0.8})$ 16 in the scattering corrections in Eqs. (11) and (12) and the lower estimates of absorption coeffcient 17  $\sigma_{ap}(\sigma_{sp}(DMPS,PM_{10}))$  were calculated by using the upper estimate of scattering coefficient  $\sigma_{sp} = \sigma_{sp}(DMPS,PM_{10})$ 18 in the scattering corrections. Consequently the lower and upper estimates of  $\omega_o$  are denoted as 19  $\omega_o(\sigma_{sp}(DMPS,PM_{0.8}))$  and  $\omega_o(\sigma_{sp}(DMPS,PM_{10}))$ , respectively. They were calculated by using both Eqs. (11) and (12) 20 for calculating  $\sigma_{ap}$ .

21

Considering that the period with the simultaneous measurements with the DMPS and the OPC showed that the DMPS size range always leads to an underestimation of both aerosol mass and scattering coefficient, it is likely that  $\sigma_{ap}$  corrected for scattering with  $\sigma_{sp}$ (DMPS, PM<sub>10</sub>) is closer to the true  $\sigma_{ap}$  than that corrected with  $\sigma_{sp}$ (DMPS, PM<sub>0.8</sub>). In the results both  $\sigma_{ap,nsc}$ ,  $\sigma_{ap}$ ( $\sigma_{sp}$ (DMPS,PM<sub>0.8</sub>)) and  $\sigma_{ap}$ ( $\sigma_{sp}$ (DMPS,PM<sub>10</sub>)) will be presented to evaluate the effect of using only the size range mesured with the DMPS for the scattering correction.

27

Similar to  $\sigma_{sp}$ , the wavelength dependency of light absorption by particles can roughly be described by the absorption Ångström exponent:





1 
$$\alpha_{ap} = -\frac{\ln(\sigma_{ap}(\lambda_1)/\sigma_{ap}(\lambda_2))}{\ln(\lambda_1/\lambda_2)}$$
(14)

that was calculated by using  $\lambda = 467$  nm and 660 nm for  $\sigma_{ap,nsc}$ ,  $\sigma_{ap}(\sigma_{sp}(DMPS,PM_{10}))$  and both Eqs. (11) and (12). The variable names  $\alpha_{ap}(\sigma_{ap,nsc})$ ,  $\alpha_{ap}(\sigma_{sp}(DMPS,PM_{10}),B1999)$  and  $\alpha_{ap}(\sigma_{sp}(DMPS,PM_{10}),V2010)$ , respectively, will be used to denote the  $\alpha_{ap}$  calculated in different ways. These calculations were conducted to study the uncertainty of  $\alpha_{ap}$  due to scattering corrections.

6

7 The absorption coefficient was used to estimate the concentration of equivalent black carbon, eBC (Petzold et8 al. 2013) from:

9 
$$eBC = \frac{\sigma_{ap}}{MAC}$$
(15)

10 where MAC is the mass absorption coefficient. For freshly-emitted BC the MAC value is approximately 7.5  $m^2 g^{-1}$ 11 <sup>1</sup> at  $\lambda$  = 550 nm (Bond et al., 2013). By assuming a wavelength-dependency of  $\lambda^{-1}$  this corresponds to MAC  $\approx$  7.8 12  $m^2 g^{-1}$  at  $\lambda$  = 530 nm. This can be considered to yield an upper estimate for eBC concentrations since for coated 13 BC particles MAC is larger (Bond et al., 2013). eBC was calculated by using  $\sigma_{ap,nsc}$  and  $\sigma_{ap}(\sigma_{sp}(DMPS, PM_{10}))$ 14 calculated with both algorithms, Eq. (11) and (12). The corresponding variable names  $eBC(\sigma_{ap,nsc})$  and 15  $eBC(\sigma_{sp}(DMPS,PM_{10}))$  will be used below for them. The scattering-corrected  $eBC(\sigma_{sp}(DMPS,PM_{10}))$  can be 16 considered to be closer to the true eBC concentration. The reason for also presenting eBC( $\sigma_{ap,nsc}$ ) is that often an 17 estimate of BC concentrations is needed even if it is known that it is an upper estimate (Caiazzo et al., 2021). It 18 is also comparable with the eBC often presented from Aethalometer measurements. Presenting both yields a 19 quantitative estimate of the bias due to not correcting the data for scattering.

20

21 The eBC mass fractions in the two size ranges  $D_a < 0.8 \,\mu$ m and  $D_a < 10 \,\mu$ m were calculated from

22 
$$feBC(m(DMPS, PM_{0.8})) = 100\% \frac{eBC}{m(DMPS, PM_{0.8})}$$
 (16)

23 
$$feBC(m(DMPS, PM_{10})) = 100\% \frac{eBC}{m(DMPS, PM_{10})}$$
(17)

where the mass concentrations m(DMPS,PM<sub>0.8</sub>) and m(DMPS,PM<sub>10</sub>) were defined in Eq. (4) and eBC calculated from Eq. (15). Mass fractions were calculated for eBC( $\sigma_{ap,nsc}$ ) and eBC( $\sigma_{sp}$ (DMPS,PM<sub>10</sub>)).

- 26
- 27
- 28





# 1 2.3.4 Noise of scattering and absorption coefficients and eBC

2 The uncertainty of scattering coefficients should in principle be calculated from the error propagation formula  $(\delta\sigma_{sp})^2 = \sum (\partial\sigma_{sp}/\partial x_i)^2 (\delta x_i)^2$ , where  $\delta x_i$  is the uncertainty of variable  $x_i$  in calculating  $\sigma_{sp}$  (e.g., Sherman et al., 3 4 2015). That would require taking into account all uncertainties of the size distribution measurements and Mie 5 modeling. However, a simplified approach was used here. The  $\sigma_{sp}$  calculated from the size distribution data and 6 the uncertainty of the size distribution range were used for calculating lower and upper estimates of  $\sigma_{sp}$  as 7 explained above. In addition to that the noise of  $\sigma_{sp}$  was estimated from the average of the absolute differences 8 of all two consecutive hourly-averaged scattering coefficients  $\delta \sigma_{sp}(average,1h) = average(|\Delta \sigma_{sp}(1h)|) =$ 9 average( $|\sigma_{sp}(t_{i+1}) - \sigma_{sp}(t_i)|$ ). The average noise of 24-h averages was calculated from  $\delta\sigma_{sp}(24h) =$  $\delta\sigma_{sp}(average, 1h)/\sqrt{24}$ . The noises were calculated for both  $\sigma_{sp}(DMPS, PM_{0.8})$ ) and  $\sigma_{sp}(DMPS, PM_{10})$ . The noises 10 11 are presented in Table 2. Note that the difference  $|\sigma_{sp}(t_{i+1}) - \sigma_{sp}(t_i)|$  is not only due to random noise so higher 12  $|\Delta\sigma_{sp}|$  values are observed when  $\sigma_{sp}$  is in reality increasing or decreasing so the true random noise is slightly 13 lower. When  $\sigma_{sp}$  is used in calculating the scattering correction of  $\sigma_{ap}$  in B1999 (Eq. (11))  $\sigma_{sp}$  is multiplied by 14 0.0164. Consequently, the  $\sigma_{sp}$  noise for the 24-h averages results in a 0.0164 $\sigma_{sp}$  noise for  $\sigma_{ap}$ . These noises are 15 also presented in Table 2.

16

17 The uncertainty of the absorption coefficient should also be calculated from the error propagation formula, 18 similar to Sherman et al. (2015). ). However, here only the uncertainty of the spot size (~3%) and the statistical 19 noise are taken into account. The noise of the non-scattering-corrected hourly  $\sigma_{ap,nsc}$  was estimated from the 20 average of the absolute differences of all two consecutive absorption measurements  $\delta\sigma_{ap,nsc}$  (average) = 21 average( $|\Delta \sigma_{ap,nsc}|$ ) = average( $|\sigma_{ap,nsc}(t_i) - \sigma_{ap,nsc}(t_i)|$ ) similar to the noise estimate of  $\sigma_{sp}$ . The noise of 24-hour averages was estimated from  $\delta\sigma_{ap,nsc}(24h) = \delta\sigma_{ap,nsc}(average, 1h)/\sqrt{24}$ . The noise in the scattering-22 23 corrected absorption cofficients were calculated from  $\delta \sigma_{ap} = \delta \sigma_{ap,nsc} + 0.0164 \delta \sigma_{sp}$  for both  $\sigma_{ap}(\sigma_{sp}(DMPS,PM_{0.8}))$ 24 and  $\sigma_{ap}(\sigma_{sp}(DMPS,PM_{10}))$  and for 1h and 24h averages (Table 2). The noise determined this way is formally correct 25 only for  $\sigma_{ap}$  calculated with the B1999 formula, Eq. (11), not for V2010. However, calculated directly fom the 26 absolute differences, the average  $|\Delta \sigma_{ap}(B1999)| \approx average |\Delta \sigma_{ap}(V2010)|$ , but the contribution of scattering to 27 the noise was only determined for B1999 as explained above. For V2010, Eq. (12) a formal error propagation 28 calculation is more complicated due to the iterative form of the procedure and it is out of the scope of the present 29 paper. The noise of eBC was calculated from  $\delta(eBC(\sigma_{ap})) = \delta \sigma_{ap}/MAC$  for both non-scattering-corrected and 30 scattering-corrected eBC. The detection limits were defined as  $2\times\delta(eBC(\sigma_{ao}))$ . The results are presented in Table 31 3.





1 The largest uncertainty factor for  $\sigma_{ap}$ ,  $\omega_{o}$ ,  $\alpha_{ap}$ , and eBC is not related to noise. It is due to the uncertainty of the 2 refractive index and size distributions used for calculating  $\sigma_{sp}$  and the algorithm. This was evaluated by calculating 3  $\sigma_{ap}$  by using the lower and upper estimates of  $\sigma_{sp}$  in both scattering correction algorithms. These four values 4 were used then for calculating  $\omega_{o}$ ,  $\alpha_{ap}$ , and eBC and they are presented below in relevant tables and figures.

5

# 6 2.4 Filtering and preprocessing the in situ data

Both PSAP absorption and DMPS-derived scattering coefficient data were filtered manually by removing rapidly changing values since they can be assumed to result from contamination from the station or from some technical problem. The PSAP transmittance data were used to filter out data measured at Tr < 0.7 following recommendations in WMO/GAW Report No. 227 (2016) and the PSAP handbook (Springston, 2018). During most of 2010 the PSAP flow was extremely unstable so practically the whole year was removed.

12

All major sources of light absorbing aerosol other than the Dome C base are so far away that rapid variations in  $\sigma_{ap,nsc}$  are due to either instrument malfunction or influence from the base, for instance emissions from vehicles. Further filtering of the data was done by removing data in which 10-minute averages of  $\sigma_{ap,nsc}$  were more than 10 times larger than the hourly  $\sigma_{ap,nsc}$ . This was done to remove short events that are local but do not appear to come directly from the base, based on wind direction. In all roughly 13% of the data were deemed contaminated.

- 19 Additionally, wind data were used to remove clear contamination from the station. The sampling site is located 20 upwind of the base itself by the prevailing wind directions. The base has a year-round diesel generator and 21 vehicles operated within the base-area move around the base from November to February. Fig. 4 shows the 22 distribution of  $\sigma_{ap,nsc}$  in 5° wind direction (WD) sectors at wind speed WS > 2 m s<sup>-1</sup>. The generator at the base is 23 clearly observed as a pronounced peak of in  $\sigma_{ap,nsc}$  at WD 60°. If the 75<sup>th</sup> percentile of the  $\sigma_{ap,nsc}$  cumulative 24 distribution is used as the criterion for the contaminated sector data when sector data when the wind direction 25 was between 30°<WD<90° would be filtered which is 6 % of the of data. If the 99<sup>th</sup> percentile of  $\sigma_{ao,nsc}$  is used 26 the contamination sector is wider, 20°<WD<110°, and 10% of the data would be filtered. Here the latter, i.e., the 27 stricter criterion was used. The distribution of  $\sigma_{ap,nsc}$  in the same WD sectors at several wind speed intervals are 28 shown in the supplement. It is obvious that at low wind speeds contaminated air can come from all directions. 29 Therefore, when WS < 2 m s<sup>-1</sup> all data were filtered out, regardless of WD.
- 30

Since the size distribution and absorption measurements are done in the same cabin, the DMPS and OPC data were also removed when the PSAP observations indicated contamination. Fig. 1 shows the instruments'





operational time in hours. The DMPS measurements had more gaps than the PSAP. The three instruments
 required for a valid measurement were not always operational at the same time. After filtering, altogether 15815
 hours of data remained for the statistical analyses. No filtering was applied to the PM<sub>1</sub> and PM<sub>10</sub> filter samples.
 The contamination is mainly BC so it was be assumed that the effect on ion concentrations was not significant.

5

The calculations were done using hourly-averaged data. These data were filtered to remove contaminated data as explained above. The filtered data were then averaged over 24 hours to reduce noise and improve detection limits. In the discussions below, the running 24-hour averages were used, centered at each hour, i.e.  $\sigma_{ap}(t,24H)$ = average( $\sigma_{ap}(t-12,1H),...,\sigma_{ap}(t+11,1H)$ ) which means, for instance, that at noon  $\sigma_{ap}(t=12,24H)$  = average( $\sigma_{ap}(t=0,1H), ..., \sigma_{ap}(t=23,1H)$ ) so the noon average represents all absorption coefficients measured during that day. If, during any period to be averaged, there were less than 12 hours of non-contaminated data then that 24-hour average was excluded from further analysis.

13

## 14 **2.5** Filter sample analyses and data processing

There were two samplers in the immediate vicinity of the cabin where the other in situ measurements were made. There was a  $PM_{10}$  sampling head operating following the CSN EN 12341 European Standard. The  $PM_1$ samples were collected on the backup filter of a Dekati  $PM_{10}$  impactor. In both of these particles were sampled on Teflon filters (Pall-Gelman, 47-mm diameter, 2-µm nominal porosity).  $PM_{10}$  and  $PM_1$  load is obtained by summing the mass of the ions determined on Teflon filters. Note that this can be considered to be the lower estimate since there could be unidentified compounds, such as organic carbon on the filters.

21

22 Just before the analysis, half of each filter was extracted with 10 mL of ultrapure water (18 M $\Omega$  Milli-Q) in 23 ultrasonic bath for 20 min. Every filter manipulation was carried out under a class-100 laminar-flow hood, to 24 minimize contamination risks. Inorganic anions and cations, as well as selected organic anions, were 25 simultaneously measured by using a three Thermo Scientific Dionex ion-chromatography system, equipped with 26 electrochemical-suppressed conductivity detectors. The sample handling during the IC injection was minimized 27 by using a specifically-designed Flow-Injection Analysis (IC-FIA) device (Morganti et al. 2007). Cations (Na<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, 28 K<sup>+</sup>, Mg<sup>2+</sup> and Ca<sup>2+</sup>) were determined by using a Thermo Scientific Dionex CS12A-4 mm analytical column with 20 29 mM H<sub>2</sub>SO<sub>4</sub> eluent. Inorganic anions (Cl<sup>-</sup>, NO<sub>3</sub><sup>-</sup>, SO<sub>4</sub><sup>2-</sup> and C<sub>2</sub>O<sub>4</sub><sup>2-</sup>) were measured by a Thermo Scientific Dionex 30 AS4A-4 mm analytical column with a 1.8 mM Na<sub>2</sub>CO<sub>3</sub>/1.7 mM NaHCO<sub>3</sub> eluent. F<sup>-</sup> and some organic anions 31 (acetate, glycolate, formate and methanesulfonate) were determined by a Thermo Scientific Dionex AS11 32 separation column by a gradient elution (0.075-2.5 mM Na2B4O7 eluent). Further details on the ion





- 1 chromatographic measurements are reported in Udisti et al. (2004) and Becagli et al. (2011). All concentrations
- 2 were corrected to STP (1013.25 hPa and 273.15 K)
- 3
- 4 In addition to calculating scattering coefficients from the DMPS data, PM1 and PM10 mass concentrations were 5 also used to calculate the scattering coefficients. The scattering coefficients were calculated by multiplying the 6 mass concentrations with mass scattering efficiencies (MSE) presented by Hand and Malm (2007). The PM<sub>10</sub> mass 7 concentrations were multiplied by the mass scattering efficiency of 1.9 m<sup>2</sup> g<sup>-1</sup> and the PM<sub>1</sub> concentrations were 8 multiplied by 3.6 m<sup>2</sup> g<sup>-1</sup>. These are the MSE for "total mixed" aerosol and "fine mixed" aerosol in the Table 5 in 9 Hand and Malm (2007), respectively. It has to be kept in mind that the MSE values in the above-mentioned paper 10 were derived from measurements in the continental USA so they most likely have a high uncertainty when 11 applied to the Dome C aerosol. The MSE values presented by Quinn et al. (2002) were used for calculating 12 scattering coefficient of nss sulfate in PM<sub>1</sub> filters.
- 13

# 14 **2.6 Scattering data from the South Pole**

- At the South Pole Station (SPO) light scattering coefficient has been measured for more than 40 years. An integrating nephelometer was installed in 1979 and used to measure  $\sigma_{sp}$ , at four wavelengths (450, 550, 700, 850 nm). This nephelometer (Meteorology Research Inc. (MRI), Altadena, CA) was used until its failure in 2002, and a TSI Model 3563 3-wavelength nephelometer ( $\lambda$  = 450 nm, 550 nm, and 700 nm) replaced it in November 2002. (Sheridan et al., 2016). Running 24-hour averages of  $\sigma_{sp}$ (550 nm) were calculated for the years 2007 – 2013 the same way as was done for the Dome C data. The data were used for comparisons with  $\sigma_{sp}$  calculated from the Dome C data.
- 22

# 23 **2.7 Source area analyses**

24 The airmass history and transport of aerosols to Dome C were calculated with the Lagrangian dispersion model 25 FLEXPART (Stohl et al., 2005; Pisso et al, 2020). ECMWF reanalysis meteorology was used to run 60000 26 trajectories every 6 hour 50 days backwards from Dome C to make a statistical sampling of the air measured 27 there. The FLEXPART trajectories follow the mean flow of the atmosphere plus random perturbations to account 28 for turbulence. In backward mode, the FLEXPART output is emission sensitivity or residence time within a grid 29 (S). When coupled with emissions, FLEXPART emission sensitivity creates a concentration at the release point 30 that is equivalent to forward simulations from emissions, except for some small numerical differences (Seibert 31 and Frank, 2004). One advantage of using a backward simulation in a case like this is that the emission sensitivity





- 1 fields can be used not only to simulate concentrations but also directly to quantitatively describe exactly where
- 2 the air that reaches Dome C originates, and, thus, potential emissions influences.
- 3
- To investigate the role of removal processes during transport, for all model runs, two different tracers were used, one atmospheric tracer with no removal and simulated BC particles with a lognormal size distribution (geometric mean diameter = 150 nm, geometric standard deviation 1.5) experiencing both dry and wet deposition. All tracers were run backwards for 50 days, in most cases sufficient for the aerosol tracer to have less than 1e-12 of the emission sensitivity of the inert air tracer, meaning any emission prior to this would have been removed by the time of arrival at Dome C. The wet removal differentiates removal within and below clouds, also considering
- 10 the water phase of the clouds and the precipitation type. The FLEXPART removal parameters are the efficiency
- 11 of aerosols to serve as cloud condensation nuclei (CCN<sub>eff</sub>) and ice nuclei (IN<sub>eff</sub>). The values used for them were
- 12 CCN<sub>eff</sub> = 0.9 and IN<sub>eff</sub> = 0.1 as in Table 4 of Grythe et al. (2017). The FLEXPART below-cloud scavenging is a scheme
- 13 based on Laakso et al. (2003) and Kyrö et al. (2009), both described in Grythe et al. (2017). The model includes a
- 14 realistic distribution of clouds by incorporating three-dimensional cloud information from ECMWF. For a detailed
- 15 description see Grythe et al. (2017).
- 16

# 17 2.7.1 Footprint difference calculations

A statistical analysis was applied to differentiate types of air pathways using a method derived from Hirdman et al. (2010). With the main aim to investigate the different pathways to Dome C, each 6hr interval was given a rank in regards to eBC concentration and single-scattering albedo. The emission sensitivity of the 50-day transport for an aerosol tracer was sorted according to its relative type. The emission sensitivities of the highest (S<sub>H</sub>) and the lowest (S<sub>L</sub>) 10% of eBC concentration and  $\omega_o$  were calculated by averaging their emission sensitivities for a given grid cell i, j, n by:

24 
$$S_* = \frac{1}{M} \sum_{m=1}^{M} S(i, j, m)$$
(18)

Where M is the number of measurements, and S<sub>\*</sub> can be any of the sorting criteria. The relative difference between two emission sensitivities in % is then calculated as:

27 
$$RD_{1,2}(i,j) = 100\% \frac{S_1(i,j) - S_2(i,j)}{S_1(i,j) + S_2(i,j)}$$
(19)

where  $S_1$  and  $S_2$  are the two footprints. This analysis of the footprint can be used to differentiate between different influencing factors on the airmass. This can be either the influence of transport, or removal or combination of these (transport efficiency) or the emission strength.





#### 1 2.7.2 Emissions used for interpreting the footprint statistics and observed seasonal cycles

2 The Global Fire Emissions Database (GFED) is a satellite information-based fire activity map. Monthly gridded 3 burned area and emissions from fires are included in the product (http://www.globalfiredata.org). Emitted BC is 4 calculated based on emission factors, which depend on the type of vegetation that is burning. Satellites give 5 snapshots collected to give pseudo global coverage and not continuous coverage. GFED v3.1 is based on the area 6 burned, which is derived by coupling Moderate resolution Imaging Spectroradiometer (MODIS) fire pixel counts 7 with surface reflectance images (Giglio et al., 2006, 2009, 2010). This widely used emission inventory has 8 uncertainties that arrive both from the emission factors and also from the amount of burnt material. A 9 comparison of this bottom-up inventory with top-down inventories found large regional differences, and top-10 down estimates were about 30% higher (Bond et al 2013). 11 12 For the scattering aerosol two sources were considered. An off-line tool (FLEX-SSA) developed by Grythe et al. 13 (2014) and Grythe (2017) to simulate sea spray aerosol (SSA) with FLEXPART was used. It uses inputs from the 14 ECMWF model. These inputs are the wind speed at 10 m above the surface (U10) and the sea surface

15 temperature (SST). The tool takes into account the sea ice fraction which is important to the Southern Ocean SSA 16 emissions. The other major marine scattering aerosols discussed below are biogenic secondary aerosols. 17 Behrenfeld et al. (2016) estimated monthly average phytoplankton biomass (C<sub>phyto</sub>) concentrations in 2007 – 18 2015 from the Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP) satellite sensor data in three zones: 19 zone 1 =  $45^{\circ}$  -  $55^{\circ}$ S, zone 2 =  $55^{\circ}$  -  $65^{\circ}$ S and zone 3 =  $65^{\circ}$ - $75^{\circ}$ S. The data provided by Behrenfeld (2021, personal 20 communication) were used for calculating seasonal monthly  $C_{phyto}$  averages in the three zones in 2008 – 2013. 21 C<sub>phyto</sub> can be used as a proxy of biological activity and emissions of dimethyl sulfide (DMS), a precursor of 22 secondary biogenic aerosols.

- 23
- 24
- 25





1 3. Results and discusion

#### 2 3.1 Overview of the data

- 3 The time series of  $\sigma_{sp}$  calculated from the size distributions and from the PM<sub>1</sub> and PM<sub>10</sub> concentrations and the
- $\sigma_{ap,nsc}$  at Dome C and  $\sigma_{sp}$  measured with the nephelometer at the SPO are presented in Fig. 5. For the DMPS-5 derived  $\sigma_{sp}$  only the upper estimate,  $\sigma_{sp}$  (DMPS, PM<sub>10</sub>), Eq. (8) is shown. The descriptive statistics of aerosol optical
- 6 properties and mass concentrations in the whole period are presented in Tables 4 and 5.
- 7

4

8 Several observations can be made from the time series in Fig. 5. First, the scattering coefficients calculated from 9 the size distributions and the filter samples follow each other relatively well. There is a clear seasonal cycle of 10 both  $\sigma_{sp}$  and  $\sigma_{ap,nsc}$ . It is clearly seen that  $\sigma_{ap,nsc}$  follows the temporal variations of  $\sigma_{sp}$  (DMPS), the high and low 11 values occur mainly simultaneously which is good, considering that these two AOPs were measured with 12 independent instruments. Since the PSAP and other filter-based absorption photometers are sensitive not only 13 to absorbing but also to scattering aerosol and since Dome C is far from BC sources it is possible that the good 14 correlation is due to the apparent absorption only. Below this will be studied simply by using Eqs. (11) and (12) 15 to account for the scattering artifact on the absorption measurement.

16

17 The  $\sigma_{sp}$  at Dome C and SPO agree better in austral summer than in winter. However, many high-concentration 18 episodes are observed also in winter almost simultaneous at Dome C and SPO. As an example, a four-month 19 period in May-August 2011 is presented in more detail in Fig. 6. The figure shows 24-hour running averages of 20  $\sigma_{sp}$ ,  $\sigma_{ap}$  and  $\omega_o$  at  $\lambda$  = 530 nm at Dome C and  $\sigma_{sp}$  at  $\lambda$  = 550 nm at SPO. Fig. 6a shows the upper estimate  $\sigma_{sp}$  = 21  $\sigma_{sp}$ (DMPS,PM<sub>10</sub>), the corresponding lower estimate of  $\sigma_{ap} = \sigma_{ap}(\sigma_{sp}(DMPS,PM_{10}))$  (corrected according to B1999, 22 Eq. 11) and the upper estimate of  $\omega_o$ . Fig. 6b presents the lower estimate of  $\sigma_{sp} = \sigma_{sp}(DMPS, PM_{0.8})$ , the 23 corresponding upper estimate of  $\sigma_{ap} = \sigma_{ap}(\sigma_{sp}(DMPS,PM_{0.8}))$  and the lower estimate of  $\omega_0$ . In both Fig. 6a and 6b 24 also the non-scattering-corrected absorption coefficient  $\sigma_{ap,nsc}$  is shown.

25

26 In Fig. 6a, the period denoted by (1) shows an episode in which  $\omega_0$  decreases significantly for several days, being 27 an example of long-range-transported eBC. Episodes (2) and (4) are examples of periods when  $\sigma_{sp}$  is 28 approximately an order of magnitude higher at SPO than at Dome C. There are also events such as episode (3) 29 when  $\sigma_{sp}$  is approximately the same at both sites. The peaks often seem to appear slightly earlier at SPO than at 30 Dome C, suggesting transport from SPO to Dome C rather than the other way around. An example of this is shown 31 in the footprint (Fig. 7) calculated for the episode denoted by (3) in Fig. 6. The footprint shows that the air masses 32 came from the direction of the Antarctic peninsula via SPO to Dome C. Air flow from the direction of the Weddell





1 Sea to SPO and then to Dome C is consistent with a very long known winter-time circulation pattern (Alt et al., 2 1959) as reviewed by Shaw (1979). During the event denoted by episode (3)  $\sigma_{ap,nsc}$  was also high. However, when 3 the scattering correction (Eq. 11) was applied the resulting  $\sigma_{ap}$  was not especially high and  $\omega_o$  was in the range 4 of 0.98 – 1.00 for both the upper and lower estimates of  $\sigma_{sp}$ , which indicates that non-scattering-corrected 5 absorption coefficients may be considerably overestimated when  $\sigma_{sp}$  is high.

6

7 The scattering coefficients calculated from the size distributions, averaged over the filter sampling periods 8 correlate positively with the scattering coefficients calculated from the PM<sub>1</sub> and PM<sub>10</sub> filters (Fig. 8). According 9 to the slopes 0.78  $\pm$  0.02 and 0.76  $\pm$  0.04 of the regression lines in Figd. 8a and 8c  $\sigma_{sp}$ (DMPS, PM<sub>0.8</sub>) seems to be 10 the lower estimate of  $\sigma_{sp}$  also when it is compared with the filter-sample-derived  $\sigma_{sp}$ . According to the slope of 11 1.29 ± 0.04 in Fig. 8b  $\sigma_{sp}$ (DMPS, PM<sub>10</sub>) is an upper estimate of  $\sigma_{sp}$  compared with  $\sigma_{sp}$ (PM<sub>10</sub>) but when  $\sigma_{sp}$ (DMPS, 12  $PM_{10}$ ) it is compared with  $\sigma_{sp}(PM_1)$  the slope is 1.01 ± 0.06 which appears to be somewhat controversial. There 13 are also other peculiarities in the scatter plots. The scatter plot of  $\sigma_{sp}$  (DMPS, PM<sub>0.8</sub>) vs.  $\sigma_{sp}$  (PM<sub>10</sub>) (Fig. 8a) have 14 data points where  $\sigma_{sp}$  (DMPS) is low, in the range of ~0.02 – 0.03 Mm<sup>-1</sup> but  $\sigma_{sp}$  (PM<sub>10</sub>) varies in a much larger range 15 from ~0.02 to ~0.9 Mm<sup>-1</sup>. This also occurs when  $\sigma_{sp}$  (DMPS, PM<sub>10</sub>) is compared with  $\sigma_{sp}$  (PM<sub>10</sub>) (Fig. 8b). The 16 pattern could be explained by too low values of of both  $\sigma_{so}(DMPS, PM_{0.8})$  and  $\sigma_{so}(DMPS, PM_{10})$  or by too high 17 values of  $\sigma_{sp}$  (PM<sub>10</sub>). Similar suspicious pattern is not observed in the comparison with the PM<sub>1</sub> filters (Fig. 8c and 18 8d) suggesting the problem may be with  $\sigma_{so}$  (PM<sub>10</sub>). It is clear that this is not a calibration of either the size-19 distribution-derived or the filter-sample-derived  $\sigma_{sp}$  but the main message of the regressions is that the values 20 are in the same order of magnitude and that there is a statistically significant positive correlation between them 21 which increases confidence in the results. When the regressions are compared with each other it has to be kept 22 in mind that the sampling periods and the number of samples of the  $PM_1$  and  $PM_{10}$  data were not the same.

23

24 Other reasons for the wide scatter of the data points are the mass scattering efficiencies (MSE) used for 25 calculating scattering coefficients from the filter samples (see section 2.5), uncertainties in ion analyses from the 26 filters and uncertainties in calculating scattering coefficient from the size distributions, especially the estimation 27 of  $\sigma_{so}$  (DMPS, PM<sub>10</sub>) from size distributions measured with the DMPS only. In spite of all these uncertainties the 28 statistical values (averages and percentiles of the cumulative distributions) of the scattering coefficients are 29 reasonably similar. For instance, the medians of  $\sigma_{sp}$  (PM<sub>10</sub>,  $\lambda$ =550 nm),  $\sigma_{sp}$  (PM<sub>1</sub>,  $\lambda$ =550 nm),  $\sigma_{sp}$  (DMPS, PM<sub>0.8</sub>, 30  $\lambda$ =530 nm) and  $\sigma_{sp}$  (DMPS, PM<sub>10</sub>,  $\lambda$ =530 nm), were 0.24 Mm<sup>-1</sup>, 0.24 Mm<sup>-1</sup>, 0.15 Mm<sup>-1</sup>, and 0.22 Mm<sup>-1</sup>, respectively 31 (Table 4). The fact that the medians of  $\sigma_{sp}(PM_{10})$  and  $\sigma_{sp}(PM_1)$  are the same is somewhat suspicious, it would be 32 expected that  $\sigma_{sp}(PM_1) < \sigma_{sp}(PM_{10})$ . At this point it is worth paying attention to the statistics of the mass





- 1 concentrations calculated from the size distributions and from the sum of ions in the filter samples (Table 5). The 2 median mass concentrations of the PM1 and PM10 filters were 66 ng m<sup>-3</sup> and 126 ng m<sup>-3</sup>, respectively, in the 3 expected order. These mass concentrations are also in reasonably good agreement with median m(DMPS, PM<sub>0.8</sub>) 4 of 70 ng m<sup>-3</sup> and median m(DMPS, PM<sub>10</sub>) of 108 ng m<sup>-3</sup> (Table 5). This suggests that the MSE values used for 5 calculating scattering coefficients from the filter masses were not correct. As it was written in section 2.5 the 6 MSE values were taken from Hand and Malm (2007) who derived them from measurements conducted mainly 7 in US national parks. Considering this, the agreement of the filter-sample-derived with the size-distribution-8 derived  $\sigma_{sp}$  is reasonable.
- 9

### 10 **3.2 Seasonal cycles of AOPs**

# 11 3.2.1 Seasonal cycles of scattering and absorption coefficients

12 The seasonal cycles of scattering and absorption coefficients are presented in Fig. 9. The SPO scattering 13 coefficients presented in Fig. 9a-d were measured using the TSI nephelometer and the Dome C scattering 14 coefficients were calculated using the PM<sub>1</sub> (Fig. 9a) and PM<sub>10</sub> (Fig. 9b) filter sample data as explained in section 15 2.5 and and from the number size distributions (Figs. 9c-d). The maximum and minimum monthly average and 16 median scattering coefficients were observed in austral summer and winter, respectively. At SPO the scattering 17 coefficient was similar to that at Dome C in austral summer but there was a large difference in austral winter. At 18 SPO the maximum monthy average scattering coefficients were observed in austral winter but at Dome C in 19 austral summer. This suggests that in austral winter SPO is more influenced by sea spray emissions than Dome 20 C. However, even though the averages and medians are lower at Dome C high scattering coefficients are also 21 occasionally observed there in austral winter, as is shown by the 95<sup>th</sup> percentiles in Fig. 9c and 9d and above in 22 the time series of winter 2011 (Fig. 6). The data does not explain the reasons of the difference between Dome C 23 and SPO in austral winter. It may either be due to different geographical locations, different size ranges measured 24 by the instruments or both.

- 25
- The minimum monthly means and medians of  $\sigma_{ap}$  at Dome C were observed in austral autumn (MAM) and the maximum monthly means and medians in austral spring (SON), which is different than the seasonal cycle of  $\sigma_{sp}$ . (Fig 9e and 9f, Tables S2 and S4). As a result, the seasonal cycle of the single-scattering albedo  $\omega_o$  is such that the darkest aerosol, i.e., the lowest  $\omega_o$  is observed in September and October and the highest  $\omega_o$  in February and March (Fig. 9g and 9h, Table S5). When the lower estimate for  $\sigma_{sp}$  (i.e.,  $\sigma_{sp}$ (DMPS, PM<sub>0.8</sub>)) is used for the scattering correction (Eqs. (11) and (12)) the October monthly medians of  $\omega_o$  are 0.862 and 0.868 when using the B1999 and V2010 algorithms, respectively, and when the upper estimate  $\sigma_{sp}$ (DMPS, PM<sub>10</sub>) is used for the scattering





1 corrections the October monthly medians of  $\omega_o$  are 0.909 and 0.914 when using the B1999 and V2010 algorithms, 2 respectively (Table S5). The highest monthly median single-scattering albedos are ~0.98 and > 0.99 with both 3 algorithms when using the  $\sigma_{sp}$  lower and upper estimates for the scattering corrections, respectively. These 4 results show that when  $\sigma_{sp}$  is not measured but calculated from the size distributions the  $\sigma_{ap}$  and  $\omega_o$  are clearly 5 less sensitive to the selection of the algorithm (B1999 or V2010) than to the scattering coefficient used for the 6 scattering correction. But as was noted in section 2.3.4, it is likely that  $\sigma_{ap}(\sigma_{sp}(DMPS,PM_{10}))$  is closer to the true 7 absorption coefficient than  $\sigma_{ap}(\sigma_{sp}(DMPS,PM_{0.8}))$  so we can also consider the seasonal cycles presented in Figs. 8 9d, 9f and 9h to be the closest to the true ones.

9

# 10 3.2.2 Seasonal cycles of scattering and absorption Ångström exponents

11 The wavelength dependency of both scattering and absorption have clear seasonal cycles. The average scattering 12 Ångström exponent of particles in the DMPS size range,  $\alpha_{sp}$ (DMPS,PM<sub>0.8</sub>) varies from ~2.6 in austral summer 13 (DJF) to ~2.1 in austral winter (JJA) indicating that in austral summer the size distributions are dominated by 14 smaller particles than in winter (Fig. 10a, Table S3). This cycle is much clearer, when  $\alpha_{sp}$  is calculated from the 15 upper estimate of scattering: average  $\alpha_{sp}(DMPS,PM_{10})$  varies from ~1.9 in austral summer to ~0.8 in winter. The 16 seasonal cycle of  $\alpha_{sp}$  (DMPS,PM<sub>10</sub>) is actually strikingly similar to the seasonal cycle of  $\alpha_{sp}$  of  $\sigma_{sp}$  measured at SPO. 17 This supports the use of the wavelength-dependent formula (Eq. 8) for calculating  $\sigma_{sp}$ (DMPS,PM<sub>10</sub>, $\lambda$ ) from 18  $\sigma_{sp}$ (DMPS,PM<sub>0.8</sub>, $\lambda$ ). The range of  $\alpha_{sp}$  is much larger at SPO than at Dome C, however. The main reason is probably 19 that when  $\sigma_{sp}$  (DMPS,PM<sub>10</sub>, $\lambda$ ) was calculated with Eq. (8) only the monthly averages of  $f\sigma_{sp}$  (DMPS, $\lambda$ ) (Eq. 7) were 20 used but the  $f\sigma_{sp}(DMPS,\lambda)$  range is actually quite large (Fig. 3). The SPO values were calculated from direct PM<sub>10</sub> 21 scattering measurements from a nephelometer.

22

The absorption Ångström exponent  $\alpha_{ap}$  was calculated for the non-scattering corrected absorption coefficient  $\sigma_{ap,nsc}$  and for the scattering-corrected  $\sigma_{ap}(\sigma_{sp}(DMPS,PM_{10}))$  with the two algorithms. Close to the  $\sigma_{ap}$  detection limit the ratios of  $\sigma_{ap}$  at two wavelengths are very noisy so Fig. 10b, Table 4 and Table S6 present  $\alpha_{ap}$  statistics of absorption coefficients for  $\sigma_{ap} > 3 \times \delta \sigma_{ap}$  where  $\delta \sigma_{ap}$  is the wavelength-dependent 24-h average noise at  $\lambda = 467$ nm and  $\lambda = 660$  nm (Table2). Note that the number of accepted data points is lower for the scattering-corrected than for the non-scattering-corrected  $\alpha_{ap}$  (Table 4). The reason is that the scattering correction often decreases  $\sigma_{ap}$  below  $3 \times \delta \sigma_{ap}$ .

30

The first observation that can be made from looking at the statistics (Fig. 10b, Table 4 and Table S6) is that  $\alpha_{ap}(\sigma_{sp}(DMPS,PM_{10}),V2010)$  is always larger than  $\alpha_{ap}(\sigma_{ap,nsc})$  and  $\alpha_{ap}(\sigma_{sp}(DMPS,PM_{10}),B1999)$ . The main





- 1 explanation of this is that the constants in the V2010 algorithm (Eq. 12) depend on wavelength but the B1999 2 algorithm (Eq. 11) uses the same constants for all wavelengths. The differences between the  $\alpha_{ap}$  obtained from 3 different algorithms were also discussed by Backman et al. (2014) and Luoma et al. (2019).
- 4

5 The seasonal cycles of  $\alpha_{ap}(\sigma_{ap,nsc})$  and  $\alpha_{ap}(\sigma_{sp}(DMPS,PM_{10}),B1999)$  are qualitatively similar: the lowest medians 6 are observed in March and the maxima in August – October. This cycle is approximately anticorrelated with the 7  $\omega_{o}$  seasonal cycle: in March the median  $\omega_{o}$  is the highest and the lowest in August – October. In March the median 8  $\alpha_{ap}(\sigma_{ap,nsc})$  and  $\alpha_{ap}(\sigma_{sp}(DMPS,PM_{10}),B1999)$  were ~0.6 and 0.37 and in August-September 0.96 and ~0.92-0.95, 9 respectively (Table S6), essentially the value generally used for pure BC. The seasonal cycle of 10  $\alpha_{ap}(\sigma_{sp}(DMPS,PM_{10}),V2010)$  is a little bit different: the minimum median of ~1.2 is in February and the maximum 11 of ~1.7 occurs in June (Table S6).

12

13 The interpretation of  $\alpha_{ap}$  is complicated. The  $\alpha_{ap}$  is related to the dominant absorbing aerosol type but physical 14 properties of the particles also affect it. For externally mixed BC particles it is generally assumed to be around 1 15 (Hegg et al., 2002; Bond and Bergstrom, 2006; Bond et al., 2013a) and higher for some organic aerosol from 16 biomass smoke and mineral dust (Kirchstetter et al., 2004; Russell et al., 2010; Devi et al, 2016). However, α<sub>ap</sub> 17 also depends on the size of BC cores and coating thickness. It is easy to show with Mie models that for single 18 non-coated BC particles with  $D_p < 20$  nm  $\alpha_{ap}$  is indeed close to 1, but when  $D_p \approx 100$  nm  $\alpha_{ap} \approx 1.3$  depending 19 on the wavelength pair used for the calculation and < 1 when  $D_p$  > ~150 nm. For BC particle size distributions the 20 width and the dominant particle size affect  $\alpha_{ap}$ . Coating of BC cores affects  $\alpha_{ap}$  even more: when BC particles are 21 coated either with a light-absorbing shell or even with a light-scattering shell  $\alpha_{ap}$  can be clearly larger than 1 22 (e.g., Gyawali et al., 2009; Lack and Cappa, 2010; Virkkula 2021). Core-shell simulations of size distributions of 23 BC particles coated with a light-scattering shell show that  $\alpha_{ab} > -1.4$  for the wavelength pair of 470/950 nm could 24 be obtained for BC particle size distributions when the shell volume fraction is > ~89 - 90% and the geometric 25 mean diameter of the BC particles is in the range of  $\sim$ 70 – 100 nm (Virkkula, 2021). Higher  $\alpha_{ap}$  would be obtained 26 also by coating with a light-absorbing shell such as brown carbon. In the present work such  $\alpha_{ap}$  values were 27 obtained for  $\alpha_{ao}(\sigma_{so}(DMPS,PM_{10}),V2010)$  for the wavelength pair 467/660 nm. So, if these values are closer to 28 the truth it seems that the BC particles that are observed at Dome C are thickly coated and their dominant particle 29 size is < ~100 nm. On the other hand, if the average  $\alpha_{ap} \approx 0.8$  obtained for  $\alpha_{ap}(\sigma_{ap,nsc})$  and 30  $\alpha_{a0}(\sigma_{sp}(DMPS,PM_{10}),B1999)$  is closer to the truth, the core-shell simulation of Virkkula (2021) suggests that BC 31 particle size distributions would dominated by thinly-coated particles in the size range > 100 nm.

32





#### 1 3.3 Seasonal cycles of mass concentrations, eBC mass concentrations and mass fractions

2 The seasonal cycles of the mass concentrations m(DMPS,PM<sub>0.8</sub>) and m(DMPS,PM<sub>10</sub>), see section 2.3.2, the mass 3 concentrations of the PM1 and PM10 filter samples, the mass fraction of the sum of secondary sulfur ions, the 4 eBC mass concentrations and the eBC mass fractions feBC(m(DMPS,PM<sub>0.8</sub>)) and feBC(m(DMPS,PM<sub>10</sub>)) (Eqs. 16 5 and 17) presented in Fig. 11 and in Tables S1, S7, S8, and S9. Some corresponding published Antarctic data are 6 also plotted in Fig. 11 for comparison. The m(DMPS,PM0.8) and the m(PM1) are consistent with each other in that 7 the minimum median mass concentrations are observed in May and June and maximum medians in February. 8 This cycle is very similar to that observed at the Norwegian station Troll in 2007-2011 (Fiebig et al., 2014). The 9 monthly average volume concentrations of particles in the size range 33 - 830 nm in Fig. 9 of Fiebig et al. (2014) 10 were digitized and multiplied by the same particle density  $\rho = 1.7$  g cm<sup>-3</sup> that was used for the Dome C data and 11 plotted in Fig 11a. The average (± standard deviation) of the ratio m(DMPS,PM<sub>0.8</sub>,Dome C)/m(DMPS,Troll) of the 12 monthly averages is ~0.6 ± 0.2, i.e., about 40% lower at Dome C. Fiebig et al. (2014) reasoned that the seasonal 13 cycle of particles in the size range measured by the DMPS, i.e., m(DMPS) is controlled by photo-oxidation-limited 14 aerosol formation. This is obviously true for Dome C also. In February, when the maximum monthly average PM<sub>1</sub> 15 and PM<sub>10</sub> concentrations were observed also the contribution of the sum of secondary sulfur ions (nss SO<sub>4</sub><sup>2-</sup> + 16 MSA<sup>-</sup>) was the highest (Fig. 11b): the average (± standard deviation) contributions to sum of ions in the PM<sub>1</sub> and 17 PM<sub>10</sub> filters was then ~81 ± 12% and ~ 61 ± 23%, respectively. The concentrations and the contributions of nss 18  $SO_4^{2-}$  + MSA<sup>-</sup> were the lowest in July, ~9 ±5 % and ~ 5 ± 5% for PM<sub>1</sub> and PM<sub>10</sub>, respectively. 19

20 The seasonal cycle of larger particles ( $m(PM_{10})$  and  $m(DMPS,PM_{10})$ ) is much weaker (Fig. 11a) than the  $m(PM_1)$ 21 and m(DMPS,PM<sub>0.8</sub>) cycle. The explanation is that the contribution of sea salt to aerosol mass is the highest in 22 winter (Fig. 11b) and that a large fraction of sea-salt particles is in the supermicron size range, in line with other 23 studies of aerosols at Dome C (e.g., Jourdain et al., 2009; Udisti et al., 2012; Legrand et al., 2017a, 2017b). Note 24 that the seasonal cycle of the mass fraction of secondary sulfur ions is qualitatively similar to the seasonal cycle 25 of the scattering Ångström exponent  $\alpha_{sp}$  (Fig. 10a): both have the highest values in the austral summer and the 26 lowest values in the austral winter. This is especially clear for the PM<sub>10</sub> filters. The small insert in Fig. 11b shows 27 the scatter plot of the monthly average  $\alpha_{so}$  (DMPS,PM<sub>10</sub>) vs. (nss SO<sub>4</sub><sup>2-</sup> + MSA<sup>-</sup>)/PM<sub>10</sub>. The relationship is essentially 28 linear and the correlation coefficient is high,  $r^2 = 0.93$ . Since the usual interpretation of the size dependence of 29  $\alpha_{sp}$  is that it is inversely proportional to dominating particle size it indicates that when the mass fraction of 30 secondary aerosol is the highest the dominating particle size is the smallest. As such this is not a surprising 31 observation but it is an additional piece of information that links the chemical composition and aerosol optical 32 properties.





The estimated m(PM<sub>10</sub>) values are consistent with the concentrations measured gravimetrically by Annibaldi et al. (2011) in December 2005 – January 2006. The average PM<sub>10</sub> mass concentration they obtained was 134 ± 12 ng m<sup>-3</sup> at p = 1013 hPa and T = 298 K which equals 146 ± 12 ng m<sup>-3</sup> at p = 1013 hPa and T = 273 K used in the present paper. The average (and median) PM<sub>10</sub> mass concentrations in the present work were 167 ng m<sup>-3</sup> (140 ng m<sup>-3</sup>) and 167 ng m<sup>-3</sup> (143 ng m<sup>-3</sup>) in December and January, respectively (Table S1), in a good agreement with the gravimetric measurement of Annibaldi et al. (2011) even though their measurements were not conducted in the same period as ours.

8

9 In the austral summer the mass concentration calculated from the size distributions (m(DMPS,PM0.8) and 10 m(DMPS,PM<sub>10</sub>)) were ~100 ng m<sup>-3</sup> higher than the sums of ions in the PM<sub>1</sub> and PM<sub>10</sub> filters (Table S1). Part of 11 the explanation could in principle be that the density 1.7 g cm<sup>-3</sup> used for calculating mass concentrations from 12 the size distibutions was too high but it cannot explain all of it. Another possible explanation is that there were 13 organic compounds not observed with ion chromatography. Caiazzo et al. (2021) took filter samples at Dome C 14 in a different period, December 2016 – January 2018, analyzed them for organic and elemental carbon with an OC/EC analyzer. The average OC concentration was 86 ± 29 ng m<sup>-3</sup>, approximately the concentration difference 15 16 between the size distribution-derived and the sums of ions in the filter samples.

17

18 The eBC concentrations eBC( $\sigma_{ap,nsc}$ ) and eBC( $\sigma_{sp}$ (DMPS,PM<sub>10</sub>)) were calculated from Eq. (15). For the scattering-19 corrected  $\sigma_{av}$  the two algorithms, Eq. (11) ad (12) yielded essentially the same absorption coefficients at  $\lambda = 530$ 20 nm. Therefore only one of them is shown in the seasonal cycle plot in Fig 11c but both are presented in the 21 supplement Table S7. On the other hand, eBC( $\sigma_{ap,nsc}$ ) is also plotted to show how much the scattering correction 22 affects the calculated eBC concentrations in different seasons. For comparison, published monthly median eBC 23 seasonal cycles at three other Antarctic sites are plotted in Fig 11c: at Neumayer, a coastal site in Queen Maud 24 Land, using two two methods, an Aethalometer and a MAAP (Weller et al., 2013), at Syowa, another Queen 25 Maud Land coastal site using an Aethalometer (Hara et al., 2019), and at SPO using an Aethalometer (Sheridan 26 et al., 2016). The maximum median eBC concentrations are observed in October-November at all sites. The 27 maximum eBC in October-November is  $^{3} \pm 1$  ng m<sup>-3</sup>, quite similar at all sites. For eBC It appears that there is no 28 significant difference between the coastal and plateau sites. The highest monthly median eBC concentrations 29 are those measured with the MAAP at Neumayer in October but, for the same month, the median Aethalometer-30 derived eBC at Neumayer is the lowest. The lowest monthly median eBC concentrations are observed in April-31 May at Neumayer, SPO and Dome C and three months earlier in February at Syowa. The lowest monthly medians, 32 ~0.2 ng m<sup>-3</sup> and ~0.3 ng m<sup>-3</sup> were observed at Dome C and SPO in May, respectively. The minima were higher at





1 the coastal sites. Note, however, that the eBC concentrations measured with the Aethalometer in Fig. 11b were 2 not corrected for scattering. This correction was done only for the PSAP data from Dome C and automatically for 3 the MAAP data from Neumayer. After the corrections the Dome C monthly median eBC( $\sigma_{sp}$ (DMPS,PM<sub>10</sub>)) ranged 4 from ~0.2 in May to ~3 ng m<sup>-3</sup> in October-November, i.e., approximately by an order of magnitude and 5 approximately the same as at SPO. The range is smaller at the coastal sites. This might be due to not correcting 6 for the scattering artifact even though the range of MAAP-derived eBC concentrations at Neumayer is also 7 smaller than on the plateau sites.

8 9 The seasonal cycle of eBC is somewhat different from that of the mass concentration. Consequently, the 10 minimum eBC mass fractions in both size ranges (feBC(m(DMPS,PM<sub>0.8</sub>) and feBC(m(DMPS,PM<sub>10</sub>)), Eqs. (16) and 11 (17), were in February-March and the maxima in August-October (Fig. 11d, Tables S8 and S9). The eBC mass 12 fractions during this peak were actually quite high. Especially, if it is assumed that all eBC is in the size range 13 measured with the DMPS even for the scattering-corrected eBC monthly medians and averages of feBC varied 14 around 4-5% and the 75<sup>th</sup> percentiles around 6-7% by using both algorithms (Table S8). These are BC mass 15 fractions typically observed in urban locations (e.g., Liu et al., 2014; Shen et al., 2018), in airborne measurements 16 over Europe (McMeeking et al., 2010) and in biomass burning plumes (Pratt et al., 2011), suggesting that in 17 these periods a large fraction of aerosol was long-range transported aerosol from other continents or highly 18 processed air with larger more scattering aerosol preferentially removed. The highest eBC monthly average and 19 median mass concentrations were observed in November but then feBC was lower than its maximum. This can 20 be explained by the increase of the amount of new, non-absorbing natural secondary particles and 21 condensational growth of BC cores by compounds originating from the sea austral during spring and summer. 22 Järvinen et al. (2013) classified new particle formation (NPF) events observed at Dome C and the highest fraction 23 of new particle formation events was in November while in austral spring the particle growth rate was also the 24 highest. The minimum feBC monthly averages were < ~0.5% and medians < ~0.3% in February-March (Tables S8 25 and S9). This minimum occurs simultaneously also with the minimum eBC concentrations. This suggests that 26 during this time of the year the amount of long-range transported aerosol from other continents is at minimum 27 at the same time when the biogenic aerosol production from the oceans is still high.

28

The seasonal cycles of single-scattering albedo (Fig 9) and eBC mass fraction (Fig. 11d) are anticorrelated with each other. It is logical: the lower the feBC is, the higher is the fraction of scattering aerosol and  $\omega_0$ . Their relationships can be used for assessing whether their observed seasonal cycles could be explained by internal mixing of BC particles and scattering components. Linear regressions of monthly average and median  $\omega_0$  vs. feBC





yield high correlation coefficients but the regression lines would yield negative values at feBC = 100%. So an exponential function of the form of  $\omega_0$ (feBC) =  $\omega_0$ (feBC=0)exp(-k·feBC) was fitted with the data (Fig. 12). The correlation coefficients were slightly worse, ~0.89 ± 0.01 for  $\omega_0$  vs. feBC(m(DMPS,PM<sub>0.8</sub>) (Fig. 12a) than the ~0.98 ± 0.01 for  $\omega_0$  vs. feBC(m(DMPS,PM<sub>10</sub>) (Fig. 12b). If the fitted exponential functions were valid up to feBC = 100% the  $\omega_0$ (feBC(m(DMPS,PM<sub>0.8</sub>) would predict that the average  $\omega_0 \approx 0.2$  and the  $\omega_0$ (feBC(m(DMPS,PM<sub>10</sub>) would predict that  $\omega_0$  (feBC=100%)  $\approx 0.06$ . These are reasonable values for pure BC: it has been measured that for fresh pure BC  $\omega_0$  is approximately 0.2 ± 0.1 (e.g., Bond and Bergstrom, 2006; Mikhailov et al, 2006; Bond et al., 2013).

9 To evaluate whether these relationships could be explained by coated spherical BC particles, lognormal size 10 distributions n(D<sub>p</sub>, GMD, GSD) were generated where GMD is the geometric mean diameter and GSD the 11 geometric standard deviation and  $D_p$  range is 3 nm – 10  $\mu$ m. In the simulations all particles had a BC core and a 12 scattering shell and the core volume fraction was the same for all particles in the size distribution. The core-shell 13 model N-Mie (Voshchinnikov and Mathis, 1999) that is based on a recursive algorithm of Wu and Wang (1991) 14 was used for calculating scattering and absorption efficiencies Q<sub>s</sub> and Q<sub>a</sub> and scattering and absorption 15 coefficients. See Virkkula (2021) for a detailed description of the simulations. The BC core volume fraction varied 16 from 0.25 to 4% to cover the range of feBC(m(DMPS,PM10)) shown in Fig. 12b. The simulated ω<sub>o</sub> vs. feBC(PM<sub>10</sub>) 17 agrees best with the observation-derived values when GSD = 1.8 and GMD = 200 nm. Smaller GMD and GSD yield 18 lower  $\omega_o$  vs. feBC(PM<sub>10</sub>). A rigorous error-minimizing procedure was not done since the main goal of this 19 simulation was to show that core-shell geometry can explain the observed  $\omega_o$  vs. feBC(PM<sub>10</sub>) relationship. 20 However, a deeper analysis of the modal structure of the size distributions was considered to be out of the scope 21 of the present paper.

22

# 23 **3.4** Aerosol sources and transport and their seasonal cycles

Source areas were modeled by using FLEXPART as described in section 2.7. The 60000 50-day backtrajectories were used for estimating vertical transport routes and the source areas of the observed aerosols. Sources of individual plumes are not analyzed, the main goal is to find explanations of the observed seasonal cycles of aerosol optical properties.

28

## 29 **3.4.1 Vertical and horizontal transport**

The average residence time of the trajectories in each grid cell in the altitude range 50 – 14400 m a.g.l. south of the Equator is shown for the inert tracer and for the simulated BC aerosol tracer (Fig. 13). The average altitude of both the inert tracer and the BC tracer decreases sharply as the latitude approaches 75°S which shows that

25





on the average air masses descend over the high plateau. This is in line with several studies that show that stratospheric air is brought down into the lower troposphere by descending air masses above the Antarctic continent (Ito, 1989; James, 1989; Parish and Bromwich, 1991; King and Turner, 1997; Krinner and Genthon, 2003; Stohl and Sodemann, 2010). Even though the average altitude of the trajectories shows descending air masses above to Dome C for the inert tracer the low-level residence times are high also at low levels (<1000 m a.g.l.) at latitudes south of about 40°S indicating that it is not uncommon that air masses flow near the surface up to Dome C.

8

9 When transport was modeled with FLEXPART using in-cloud and below-cloud scavenging of aged BC particles as 10 described by Grythe et al. (2017), for the BC tracer the residence times are lower than for the inert tracer except 11 near Dome C. Additionally, the average trajectory altitude is clearly different than for the inert tracer. Far from 12 Antarctica the BC tracer residence times are clearly smaller than for the inert tracer. This is due to the removal 13 of BC particles by in-cloud and below-cloud scavenging and dry deposition. As a result there would be essentially 14 no BC particles arriving at Dome C from north of latitude 10°S at altitudes < 1600 m which is indicated by the 15 respective area in Fig. 13b just white. Consequently, those BC particles that do get lifted above the clouds 16 increase the average altitude of the BC tracer backtrajectories compared with the inert tracer average altitude 17 as is shown by the respective black and white lines in Fig 13b. At latitudes ~20°S – 50 °S the BC tracer average 18 altitude is approximately 1000 m higher than that of the inert tracer.

19

20 Between Figs. 13a and 13b there are grey text boxes that show the approximate latitude range of some major 21 BC emitting regions. Indonesia is north of 10°S so BC particles should rise to about 2 km altitude in order to get 22 transported to Dome C. The Indonesian smoke plumes are generally lower than that (Tosca et al., 2011) so its 23 contribution to eBC observed at Dome C is probably negligible. The other three main biomass-burning regions 24 Africa, Australia and Brazil are more to the south and the smoke plume heights have been observed at clearly 25 higher altitudes (Pereira et al., 2016; Remy et al., 2017; Gonzalez-Alonso et al., 2019). Smoke plumes were 26 recently observed at a height of 4 km over Brazil and even higher, at 6 km over Australia (Shikwambana and 27 Kganyago, 2021). Comparison with Fig. 13b shows that from these altitudes it is likely that BC particles reach also 28 Dome C. All this is in agreement with Fiebig et al. (2009) who established a source-receptor relationship between 29 biomass burning events in Central Brazil and the aerosol observed at the Norwegian station Troll.

30

Near Dome C and over all continental Antarctica the BC tracer average altitude is lower than that of the inert tracer. When BC has a lower average altitude than tracer it means that the air at that point has undergone more





- removal above than below the average altitude. This suggests that near Dome C most removal takes place in the
   clouds, higher up than the release point. This implies that there is more in-cloud removal than precipitation
- 3 scavenging or dry deposition over the plateau.
- 4

# 5 3.4.2 Source areas of eBC and high and low $\omega_o$ using footprint differences

- 6 Source areas were next estimated by using FLEXPART and calculating the footprint differences as described in
- 7 section 2.7.1. The relative difference (RD) (Eq. 19) of the emission sensitivities in the two lowest layers (>1000 m
- 8  $\,$  a.g.l.) was calculated for the highest and the lowest 10% of eBC concentration and  $\omega_{o}.$  As it was shown in section
- 9 3.3 the highest and lowest  $\omega_o$  percentiles correspond to the lowest and the highest eBC mass fraction percentiles.
- 10

11 The highest eBC concentrations have an increased signal from the surrounding continents Australia, Africa and 12 South America (Fig. 14a). However, high RD values are also associated with large areas over the Pacific Ocean, 13 the Indian Ocean and the Atlantic Ocean. These high RD values on the oceans may in principle be due to shipping 14 emissions but it is more likely that they are due to BC emissions from the continents where the emissions are 15 considerably higher than from shipping. The highest  $\omega_0$  are obviously related to emissions from the Southern 16 Ocean (Fig. 14b).

17

# 18 **3.4.3** Seasonal cycles of air mass transport from different regions

Seasonal cycles of meteorological processes and conditions also affect properties of aerosols observed at Dome C. The most evident is solar radiation that affects new particle formation (e.g., Ito, 1989; Fiebig et al., 2014; Järvinen et al., 2013) but important are also the seasonal cycles of transport from the surrounding continents and from the marine boundary layer of the Southern Ocean (Stohl and Sodemann, 2010). The seasonal contributions of air masses from different regions were calculated as the fraction of time the FLEXPART trajectories spent over Antarctica, the surrounding oceans and continents (Fig. 15).

25

In the austral summer (DJF) the contribution of continental air other than Antarctica is low and its minimum is in April. This is true also for marine air masses other than the Southern Ocean. In summer the contribution of Antarctic air is at the highest. Then in May the air mass transport from other continents increases clearly and it reaches a maximum in July, a smaller fraction in August and a second maximum in September. The seasonal cycle of transport from all the surrounding continental areas is qualitatively similar. The next question is how well do the seasonal cycles of the aerosol optical properties observed at Dome C follow the emissions of major absorbing and scattering aerosols in different regions.





## 1 3.4.4 Relationships of seasonal cycles of BC emissions in the surrounding continents and eBC at Dome C

2 Monthly BC emissions from wildfires and agricultural burning and other fires in South America, Africa and 3 Oceania (Australia, New Zealand, Melanesia, Micronesia and Polynesia) in 2006-2012 were downloaded from 4 the Global Fire Emissions Database (GFED, V3.1). The seasonal cycles are presented in Fig. 16a together with the 5 monthly average eBC concentrations – corrected for scattering – observed at Dome C. It is obvious in Fig 16a 6 that the maximum of the sum of the emissions occurs three months earlier than the maximum eBC concentration 7 at Dome C.

8

9 The seasonal cycles of air mass residence time over each of the major source areas plays a crucial role in 10 explaining the relationship between eBC at DomeC and BC emissions. The BC emissions were multiplied with the 11 fraction of time (=f(t)) the FLEXPART trajectories spent over each of the three major source areas, shown in Fig. 12 15. These quantities (f(t)×BC emissions) will be called residence-time-weighted BC emissions, RTW BC emissions. 13 They are plotted in Fig. 16b together with the normalized eBC concentrations. Linear regressions between the 14 eBC concentration and RTW BC emissions were calculated but only the respective squared correlation 15 coefficients R<sup>2</sup> are shown. The correlation between eBC concentrations and RTW BC emissions is weak, R<sup>2</sup> < 0.3 16 (Fig. 16b). But the main reason is that the residence times shown in Fig. 15 present the fraction of time that 17 trajectories arriving at Dome C at any given month have spent over each of the regions before arrival at Dome C. 18 So next it was assumed that the transport time from each of the source areas to Dome C is one month so f(t) of 19 each month was multiplied with the BC emission one month earlier. This shifts the RTW BC emission peaks of all 20 the source areas so that all R<sup>2</sup> increase somewhat (Fig. 16c). When it is assumed that the transport time is 2 21 months R<sup>2</sup> increase clearly, 0.688 for Oceania and 0.665 for South America, for Africa correlation still remains 22 low (Fig. 16d). In October-November, when the eBC concentrations at Dome C are the largest the 2-month-23 shifted RTW BC emissions from South America are approximately an order of magnitude larger than from Africa 24 and Oceania suggesting that South American BC emissions are the largest contributor to eBC at Dome C.

25

The highest correlation between the eBC concentrations and RTW BC emissions with the 2-month time shift suggests further that transport time is considerably longer than the 30 days Stohl and Sodemann (2010) obtained in a 5.5-year climatology of atmospheric transport into the Antarctic troposphere. Fig. 16 also shows that the observed concentration varies by an order of magnitude only although the emissions vary by two orders of magnitude. This suggests that at the observed seasonal minimum eBC in March-April the sources are something else than wildfires and agricultural burning BC emissions in the GFED data. Possible sources can be other anthropogenic emissions such as traffic including shipping, industrial emissions and heating.





#### 1 3.4.5 Relationships of seasonal cycles of scattering aerosol emissions and $\sigma_{sp}$ at Dome C and SPO

2 The seasonal cycles of scattering coefficient at Dome C and at the SPO were compared with the seasonal cycles 3 of secondary and primary marine aerosol emissions. The sea-spray aerosol (SSA) emissions calculated with the 4 FLEX-SSA offline tool (section 2.7.2) show that SSA emissions from the Southern Ocean peak in the austral winter 5 (Fig. 17a). Secondary marine aerosols are formed from the oxidation of emission products of phytoplankton. The 6 phytoplankton concentration maximum is in summer and minimum in winter in the satellite data analysis of 7 Behrenfeld et al. (2016) (Fig. 17a). However, phytoplankton concentration alone does not explain the seasonal 8 cycle of secondary aerosol scattering coefficient. The DMS oxidation is a photochemical process so a simplified 9 method was used for estimating solar radiation intensity. Global radiation intensity I<sub>T</sub> (W m<sup>-2</sup>) was calculated by 10 using a clear sky model for direct and diffuse insolation (Bird and Hulstrom, 1981). IT was calculated at the surface 11 (p =1000 mbar) for each hour of the year at the prime meridian (0°) at three latitudes: 50°S, 60°S, and 70°S 12 corresponding to the central latitudes of zone 1 (45°-55°S), zone 2 (55°-65°S), and the polar zone, zone 3 (65°-13 75°S). The monthly averages of  $I_T$  at 10:00 – 14:00 in each day of a month were calculated. The reasoning for 14 using this time is that new particle formation is typically, although not exclusively, a daytime phenomenon at 15 numerous locations around the world, also in Antarctica (e.g., Weller et al., 2015; Kerminen et al., 2018; Kim et 16 al., 2019; Brean et al, 2021). The monthly product C<sub>phytol</sub>T, was then used as a proxy of biogenic secondary aerosol 17 emissions. Behrenfeld et al. (2016) also presented the photosynthetically active radiation (PAR) in the polar zone 18 obtained from the MODIS Aqua ocean colour sensor acquired from http://oceancolor.gsfc.nasa.gov/cms. In 19 addItion to the product C<sub>phyto</sub>I<sub>T</sub>, also the monthly product C<sub>phyto</sub>PAR was calculated for the zone 3 and used as a 20 proxy for biogenic secondary aerosol emissions. In Zone 3 the two emission proxies CphytolT and CphytoPAR agee 21 well (Fig. 17a).

22

23 The normalized seasonal cycles of the SSA flux and the secondary aerosol emission proxy C<sub>phyto</sub>PAR in the polar 24 zone are plotted together with the normalized seasonal cycles of the scattering coefficients at Dome C (Fig. 17b) 25 and at SPO (Fig. 17c). At Dome C both  $\sigma_{sp}$  (DMPS,PM<sub>0.8</sub>) and  $\sigma_{sp}$  (DMPS,PM<sub>10</sub>) approximately follow the seasonal 26 cycle of the secondary aerosol emissions in October - April. In the austral winter the effect of SSA is visible 27 especially in the high monthly  $\sigma_{sp}$  averages. The medians are cleary lower, they follow the secondary aerosol 28 emission proxy even in May. Also the normalized seasonal cycle of  $\sigma_{sp}$ (nss-SO<sub>4</sub><sup>2-</sup>,PM<sub>1</sub>), the scattering coefficient 29 calculated from non-seasalt sulfate concentrations in PM<sub>1</sub> filter samples is presented in Fig. 17b. It also follows 30 approximately the seasonal cycle of C<sub>phyto</sub>PAR for the whole year. However, there appears to be a time lag. 31  $C_{phyto}$  PAR grows fast in October but  $\sigma_{sp}$  (nss-SO<sub>4</sub><sup>2-</sup>, PM<sub>1</sub>) in November. When  $C_{phyto}$  PAR is shifted by one month they 32 agree clearly better. The linear regression of of  $\sigma_{sp}(nssSO_4^2; PM_1)$  vs.  $C_{phyto}PAR$  yields the correlation coefficient





1  $R^2$  = 0.65 when there is no C<sub>phyto</sub>PAR time shift and  $R^2$  = 0.92 when C<sub>phyto</sub>PAR is shifted by one month (Fig. 18). 2 This suggests that on the average it takes approximately a month for the secondary aerosol to be formed, grown, 3 get mixed in the upper atmospheric layers and be transported to the upper plateau. There is one thing that the 4 time shift cannot explain. The maximum  $\sigma_{sp}$  – and mass concentrations as discussed in Section 3.3 – was in 5 February. This cannot be explained by the seasonal cycle of the biogenic secondary aerosol emission proxy: the 6 maximum C<sub>phyto</sub>PAR was in December so even a one month transport time does not make February the 7 maximum. Above it was assumed that the proxy is simply the product C<sub>phyto</sub>PAR. However, the underlying 8 hypothesis of this proxy is that the emissions of the precursor gases of new particle formation, mainly DMS, are 9 linearly related to the phytoplankton mass concentrations. This is probably not the case, there are other factors. 10 Further analyses of that relationship are out of the scope of the present paper, however. Note that  $\sigma_{sp}(nssSO_4^{2-}$ 11 , PM<sub>1</sub>) correlates well also with  $C_{phyto}I_T$  of zones 1 and 2 shown in Fig. 17a which means the secondary marine 12 aerosol may have been formed and grown also further to the north than in the polar zone. 13

At SPO  $\sigma_{sp}$  is more strongly affected by the SSA emissions than at Dome C, especially in the austral winter and spring (Fig. 17c). Note that at SPO the seasonal cycle of the normalized median  $\sigma_{sp}$  follows roughly the normalized SSA emission flux in July – December. The seasonal cycle of the normalized average  $\sigma_{sp}$  is different because averages are affected more by individual high transport events and medians represent better the prevailing background aerosol of the season. As explained above, the difference between Dome C and SPO may either be due to geographical locations, different size ranges measured by the instruments or both.

20

## 21 **4.** Summary and conclusions

22 Aerosol optical properties have been measured at several Antarctic sites but scattering and absorption data 23 measured at Dome C have not been not examined in detail earlier. This work fills that gap using light absorption 24 from a 3 $\lambda$  PSAP and light scattering coefficients ( $\sigma_{sp}$ ) calculated from particle number size distributions measured 25 with a Differential Mobility Particle Sizer (DMPS) and an optical particle counter (OPC). Additionally, single 26 scattering albedo ( $\omega_o$ ), absorption Ångström exponent ( $\alpha_{ap}$ ), scattering Ångström exponent ( $\alpha_{sp}$ ) and equivalent 27 black carbon (eBC) concentrations were calculated. The sources of the aerosol were estimated by calculating 28 footprints with FLEXPART and by calculating seasonal cycles of transport of both scattering and absorbing 29 particles from different source areas. 30

Aerosol light scattering coefficient was calculated from the DMPS size distributions using two different refractive indices and the contribution of scattering due to particles larger than those measured with the DMPS to provide





1 a lower and upper estimate of  $\sigma_{sp}$ . Light scattering was also estimated using mass scattering efficiencies in 2 conjunction with mass concentrations obtained from PM<sub>1</sub> and PM<sub>10</sub> filter samples. The two most frequently used 3 algorithms to calculate aerosol absorption coefficients ( $\sigma_{ap}$ ) from PSAP measurements require scattering 4 coefficients. Both algorithms were applied and both the upper and lower estimate of  $\sigma_{sp}$  were used in order to 5 provide understanding of the differences between the algorithms and the impact of the scattering adjustment. 6 The absorption coefficient calculated using the  $\sigma_{sp}$  upper estimate was considered to be the best  $\sigma_{ap}$  estimate.

7

8 There were clear seasonal cycles of  $\sigma_{sp}$  and  $\sigma_{ap}$  at Dome C. The maximum and minimum of  $\sigma_{sp}$  were observed in 9 austral summer and winter, respectively. The Dome C scattering coefficients were also compared with  $\sigma_{sp}$ 10 measured with a nephelometer at the South Pole (SPO). At SPO the scattering coefficient was similar to that 11 measured at Dome C in austral summer but there was a large difference in the austral winter. At SPO the 12 maximum monthly averages were observed in austral winter. This suggests that, in winter, SPO is more 13 influenced by sea spray emissions than Dome C. At Dome C the  $\sigma_{ap}$  exhibited a different seasonal cycle of than 14  $\sigma_{sp}$  - the minimum  $\sigma_{ap}$  was observed in the austral autumn and the maximum in spring. As a result the seasonal 15 cycle of the single-scattering albedo  $\omega_o$  is such that the darkest aerosol, i.e., the lowest  $\omega_o \approx 0.91$  is observed 16 October and the highest  $\omega_{0} > 0.99$  in February and March.

17

18 The scattering Ångström exponent  $\alpha_{sp}$  calculated from the  $\sigma_{ap}$  lower estimate varied from ~2.6 in austral summer 19 to ~2.1 in austral winter indicating that in austral summer the size distributions are dominated by smaller 20 particles than in winter. For the  $\sigma_{sp}$  upper estimate  $\alpha_{sp}$  varied from ~1.9 in austral summer to ~0.8 in winter. This 21 seasonal cycle is quite similar to the seasonal cycle of  $\alpha_{sp}$  of  $\sigma_{sp}$  measured at SPO. The uncertainty of the 22 absorption Ångström exponent  $\alpha_{ap}$  is high, particularly in the clean conditions existing in Antarctica. However, 23 despite the high uncertainties the seasonal cycles of  $\alpha_{ap}$ , with and without the scattering correction, are 24 qualitatively similar: the lowest monthly medians are observed in March while the maxima occur in August -25 October. This cycle – even that of the non-scattering corrected  $\alpha_{ap}$  – is anticorrelated with the  $\omega_{o}$  seasonal cycle. 26

20

The eBC mass concentrations were compared with eBC measured at three other sites: the South Pole and two coastal sites Neumayer and Syowa. The maximum monthly median eBC concentrations are almost the same (~3  $\pm$  1 ng m<sup>-3</sup>) at all these sites in October-November. This suggests that, as far as eBC is concerned, there is no significant difference between the coastal and plateau sites. The seasonal cycle of eBC is slightly different from the mass concentration calculated from the number size distributions measured with the DMPS. Consequently, the seasonal cycle of the eBC mass fraction (f(eBC)) is such that f(eBC) minimum is in February-March and the





1maximum is in August-October, anticorrelating with  $\omega_o$ . The eBC mass fractions in this peak are actually quite2high and vary around 4-5% with the 75<sup>th</sup> percentiles > 6-7%, only slightly depending on PSAP correction algorithm.3These levels of eBC mass fractions are typically observed in polluted air, suggesting that in these periods a large4fraction of aerosol is long-range transported aerosol from other continents.

5

Source areas were calculated with 50-day FLEXPART footprints. The relative differences of the footprints calculated for the highest and lowest 10% of eBC concentrations and  $\omega_0$  showed that the highest eBC concentrations and the lowest  $\omega_0$  were associated with air masses coming from South America, Australia and Africa. Vertical simulations that take BC particle removal processes into account show that there would be essentially no BC particles arriving at Dome C from north of latitude 10°S at altitudes < 1600 m. The main biomassburning regions Africa, Australia and Brazil are more to the south and their smoke plumes have been observed at higher altitudes than that so they can get transported to Antarctica.

13

14 The seasonal cycle of BC emissions from wildfires and agricultural burning and other fires in South America, Africa 15 and Australia were calculated from data downloaded from the Global Fire Emissions Database (GFED). The 16 maximum total emissions were in August-September but the peak of monthly average eBC concentrations is 17 observed November, 2 – 3 months later not only at Dome C but also at the South Pole and the coastal stations. 18 This is considerably longer than the 30 days presented in an earlier study. If this peak eBC concentration is really 19 due to the peak emissions from the above-mentioned fires in the surrounding continents it means that the 20 aerosol from these fires remains in air for several months and gets mixed essentially over the entire Southern 21 Hemisphere. The seasonal contributions of air masses from different regions were calculated as the fraction of 22 time the FLEXPART trajectories spent over Antarctica, the surrounding oceans and continents. The BC emissions 23 were multiplied with the fraction of time the trajectories spent over each of the three major source areas. In 24 October-November, when the eBC concentrations at Dome C are the largest the 2-month-shifted residence-time-25 weighted BC emissions from South America are approximately an order of magnitude larger than from Africa 26 and Oceania suggesting that South American BC emissions are the largest contributor to eBC at Dome C.

27

The seasonal cycles of scattering coefficient at Dome C and at the SPO were compared with the seasonal cycles of secondary and primary marine aerosol emissions. The seasonal cycles of sea spray aerosol (SSA) emissions were simulated with the FLEX-SSA offline tool. The seasonal cycles of biogenic secondary aerosols were estimated from monthly average phytoplankton biomass concentrations obtained from the CALIOP satellite sensor data. The correlation coefficients between scattering coefficients measured at Dome C and





- phytoplankton biomass concentrations and a biogenic secondary aerosol emission proxy are high. It may take a month for the biogenic aerosol to be formed and get transported from the sea level to Dome C. The scattering coefficients measured at SPO correlated much better with the SSA emission fluxes in the Southern Ocean than the scattering coefficients measured at Dome C. The difference between the scattering coefficients at these sites
- 5 may either be due to geographical locations, different size ranges measured by the instruments or both.
- 6

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## 7 Data Availability

- 8 The Dome C aerosol physical measurement data will be made openly available for the final publication at the
- 9 EBAS database (http://ebas.nilu.no). The SPO data are there already. The Dome C aerosol chemical composition
- 10 data will be available upon request by writing to RT, BS or MS.
- 11

# 12 Author contributions

- 13 AV contributed to designing the measurements of aerosol physical properties at Dome C, processing and 14 analyzing data and writing most of the manuscript with contributions from all coauthors; HG did all the FLEXPART modeling and processed GFED data and wrote the associated text; JB wrote the codes for data processing; TP 15 16 reviewed and commented the manuscript; MB, CL and AL were responsible for the installation and long-term 17 operation of the aerosol instruments at Dome C; SB, RT, and MS were responsible for the sampling and analyses 18 of the PM1 and PM10 filter samples; VV contributed to designing and administration of the measurements at 19 Dome C; PS and EA were responsible for the SPO aerosol measurements. All authors reviewed and commented 20 the manuscript. 21
- 22 Competing interests
- 23 Tuukka Petäjä is editor of ACP.
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## 1 Tables

- 2 Table 1. Seasonal variation of the fractions of volume concentration and scattering coefficients in the size range
- 3 measured by the DMPS of the respective values calculated from the combined size distributions measured with
- 4 the DMPS and the OPC at Dome C in December 2007 July 2009.

	fV(DMPS),	%	fσ <sub>sp</sub> (DMPS,46	7), %	fσ <sub>sp</sub> (DMPS,53	80), %	fσ <sub>sp</sub> (DMPS,66	60), %
month	average ± std	median	average ± std	median	average ± std	median	average ± std	mediar
JAN	86.8 ± 10.7	91.6	85.2 ± 9.9	88.7	78.5 ± 12.6	82.5	66.1 ± 15.8	69.6
FEB	85.7 ± 14.0	90.6	82.8 ± 14.3	87.2	76.1 ± 15.9	81.1	64.0 ± 17.5	68.6
MAR	79.0 ± 20.9	88.3	77.0 ± 20.2	85.0	70.2 ± 21.6	78.2	58.6 ± 22.2	65.2
APR	72.8 ± 22.7	83.2	73.6 ± 20.4	80.5	66.4 ± 22.0	72.9	55.0 ± 23.0	59.4
MAY	55.4 ± 24.3	53.1	62.5 ± 20.9	63.9	54.5 ± 22.0	54.6	42.9 ± 22.1	40.7
JUN	49.8 ± 18.3	51.2	61.7 ± 14.6	64.5	52.7 ± 14.6	54.9	39.8 ± 13.2	41.1
JUL	49.5 ± 17.9	50.2	62.2 ± 15.1	65.3	53.4 ± 15.2	55.7	40.8 ± 14.2	41.9
AUG	54.4 ± 15.5	56.0	68.3 ± 12.2	69.9	60.0 ± 13.0	60.7	47.4 ± 13.5	46.7
SEP	62.6 ± 14.6	64.4	73.8 ± 9.8	74.6	66.1 ± 11.4	66.4	54.0 ± 13.3	53.4
OCT	64.6 ± 14.5	66.3	74.1 ± 10.5	74.4	66.4 ± 12.2	66.4	54.2 ± 14.0	53.5
NOV	74.7 ± 13.7	79.1	77.4 ± 10.5	80.2	69.4 ± 12.0	72.0	56.0 ± 13.2	57.7
DEC	80.4 ± 14.4	84.1	80.1 ± 10.9	83.4	72.1 ± 12.1	75.4	57.7 ± 12.4	60.5
vear	70.5 ± 21.8	76.7	74.3 ± 17.0	78.4	66.7 ± 18.4	70.3	54.3 ± 19.1	56.2

5 6

7 Table 2. Noise of scattering and absorption coefficients calculated from the particle number size distributions

8 and the PSAP data.  $\sigma_{sp}$  (DMPS, PM<sub>0.8</sub>) and  $\sigma_{sp}$  (DMPS, PM<sub>10</sub>). Noise was estimated as explained in section 2.5.

	$\begin{array}{cccccccccccccccccccccccccccccccccccc$			ו <sup>-1</sup>					
Scattering	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$				p(DMPS,PN	И <sub>10</sub> )			
$\Delta t \setminus \lambda(nm)$	467	530	660	467	530	660			
1 h	5.2×10 <sup>-2</sup>	4.2×10 <sup>-2</sup>	2.7×10 <sup>-2</sup>	7.2×10 <sup>-2</sup>	6.5×10 <sup>-2</sup>	5.3×10 <sup>-2</sup>			
24 h	1.1×10 <sup>-2</sup>	0.85×10 <sup>-2</sup>	0.55×10 <sup>-2</sup>	1.5×10 <sup>-2</sup>	1.3×10 <sup>-2</sup>	1.1×10 <sup>-2</sup>			
$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$				ering correctio	on, Mm <sup>-1</sup>				
	0.0164>	κδσ <sub>sp</sub> (DMP\$	S,PM <sub>0.8</sub> )	0.0164>	κδσ <sub>sp</sub> (DMP\$	S,PM <sub>10</sub> )			
1 h	0.85×10 <sup>-3</sup>	0.68×10 <sup>-3</sup>	0.44×10 <sup>-3</sup>	1.2×10⁻³	1.1×10⁻³	0.87×10 <sup>-3</sup>			
24 h	0.17×10 <sup>-3</sup>	0.14×10 <sup>-3</sup>	0.091×10 <sup>-3</sup>	0.24×10 <sup>-3</sup>	0.22×10 <sup>-3</sup>	0.18×10 <sup>-3</sup>			
Absorption		$\delta\sigma_{\text{ap,nsc}}$		δσ	$\sigma_{ap}(\sigma_{sp}(PM_0$	8))	δα	$\sigma_{ap}(\sigma_{sp}(PM_1$	<sub>0</sub> ))
$\Delta t \setminus \lambda(nm)$	467	530	660	467	530	660	467	530	660
1 h	3.2×10 <sup>-3</sup>	2.5×10 <sup>-3</sup>	2.7×10 <sup>-3</sup>	4.0×10 <sup>-3</sup>	3.2×10 <sup>-3</sup>	3.2×10 <sup>-3</sup>	4.3×10 <sup>-3</sup>	3.6×10 <sup>-3</sup>	3.6×10 <sup>-3</sup>
24 h	0.65×10 <sup>-3</sup>	0.52×10 <sup>-3</sup>	0.56×10 <sup>-3</sup>	0.82×10 <sup>-3</sup>	0.66×10 <sup>-3</sup>	0.65×10 <sup>-3</sup>	0.89×10 <sup>-3</sup>	0.74×10 <sup>-3</sup>	0.74×10 <sup>-3</sup>

9 10

- 11 Table 3. Noise and detection limits of eBC concentration calculated from the noise of the absorption coefficients
- 12 presented in Table 2.

		eBC noise, ng m <sup>-3</sup>	
	$\delta(eBC(\sigma_{ap,nsc}))$	$\delta(eBC(\sigma_{ap}(\sigma_{sp}(DMPS,PM_{0.8}))))$	$\delta(eBC(\sigma_{ap}(\sigma_{sp}(DMPS,PM_{10}))))$
 1 h	0.327	0.415	0.464
 24 h	0.067	0.085	0.100
	$2\delta(eBC(\sigma_{ap,nsc}))$	$2\delta(eBC(\sigma_{ap}(\sigma_{sp}(DMPS,PM_{0.8}))))$	$2\delta(eBC(\sigma_{ap}(\sigma_{sp}(DMPS,PM_{10}))))$
 1 h	0.65	0.83	0.93
24 h	0.13	0.17	0.20





					F	ercentiles		
	λ (nm)	Ν	Ave ± std	5	25	50	75	95
	Scattering co	pefficient, $\sigma_{g}$	<sub>ap</sub> , Mm <sup>-</sup> '					
Filter samples	550	4705	0.01 . 0.01	0.059	0.420	0.04	0.00	0.75
4	550	400	$0.30 \pm 0.23$	0.040	0.110	0.24	0.41	0.70
$\sigma_{so}(DMPS, PM_{0.8}))$	467	21987	0.35 + 0.64	0.034	0 095	0 19	0.40	1.06
Sp(2.1.1 0, 1 110.8/)								0.80
								0.46
	000	2.00.	0110 2 0102	0.010	0.011	0.00	0.10	0.10
$\sigma_{sp}(DMPS, PM_{10}))$	467	21987	$0.48 \pm 0.96$	0.052	0.138	0.26	0.52	1.34
	530	21987	0.41 ± 0.88	0.046	0.120	0.22	0.44	1.12
	660	21987	0.31 ± 0.73	0.034	0.089	0.16	0.31	0.78
	Absorption c	oefficient σ						
$\sigma_{ap}(\sigma_{sp}(DMPS,PM_{10}))$				< 0.001	0.0029	0.0096	0.021	0.051
	530	15815			0.0029	0.0085	0.019	0.044
	660	15657	0.012 ± 0.012	< 0.001	0.0033	0.0077	0.016	0.036
	$\sigma_{ap}$ , calcul	ated using	V2010					
$\sigma_{ap}(\sigma_{sp}(DMPS,PM_{10}))$	467	15778	0.016 ± 0.018	< 0.001	0.0033	0.0099	0.022	0.054
	530	15815	0.012 ± 0.015	< 0.001	0.0023	0.0076	0.017	0.041
	660	15657	0.009 ± 0.011	< 0.001	0.0016	0.0057	0.013	0.031
		In	tensive AOPs					
:	Single-scatte	ering albedo	$\omega_{o} = \sigma_{sp}(DMPS,PM)$	<sub>10</sub> )/(σ <sub>sp</sub> (DMPS	S,PM <sub>10</sub> ) + c	s <sub>ap</sub> )		
	σ	ap calculate	d using B1999					
								0.99
$\omega_{o}(\sigma_{sp}(\text{DMPS},\text{PM}_{10}))$				0.85	0.92	0.96	0.99	1.00
			-					
$\omega_{o}(\sigma_{sp}(DMPS,PM_{10}))$	530	15815	$0.95 \pm 0.05$	0.86	0.93	0.96	0.99	1.00
;	Scattering År	ngström exp	ponent $\alpha_{sp}$ ( $\lambda$ =467/6	60 nm)				
$\alpha_{sp}(\sigma_{sp}(DMPS,PM_{0.8}))$	467/660	21987	2.31 ± 0.28	1.90	2.10	2.25	2.54	2.79
$\alpha_{\text{sp}}(\sigma_{\text{sp}}(\text{DMPS},\text{PM}_{10}))$	467/660	21987	$1.35 \pm 0.40$	0.77	1.05	1.30	1.66	2.04
	Absorption Å	ngström ex	ponent (λ=467/660	nm) $\alpha_{abs}$ for	σ <sub>ap</sub> > 3 δσ <sub>a</sub>	σ		
	σ	ap calculate	d using B1999					
$\alpha_{ap}(\sigma_{ap,nsc})$	467/660	15607	$0.86 \pm 0.34$	0.29	0.67	0.87	1.02	1.37
$\alpha_{ap}(\sigma_{sp}(DMPS,PM_{10}))$				< 0	0.56	0.82	0.99	1.40
	σ	ap calculate	d using V2010					
$\alpha_{ap}(\sigma_{sp}(DMPS,PM_{10}))$	467/660	10270	1.37 ± 0.38	0.72	1.17	1.37	1.59	2.03

1 Table 4. Descriptive statistics of aerosol optical properties at Dome C in 2008 – 2013.

2 3

4

5





1	Table 5. Statistical summary of mass concentrations estimated from particle number size distributions, sums of
2	ion concentrations of $PM_1$ and $PM_{10}$ filter samples and the PSAP data at Dome C in 2008 – 2013. The statistical
3	values of the $PM_1$ and $PM_{10}$ are those of all individual filters, the statistical values calculated from the DMPS and
4	PSAP data are those of running 24h-averaged data, see details in the text. m(DMPS,PM <sub>0.8</sub> ): mass concentration
5	calculated from the particle number size distributions measured with the DMPS assuming particle density of 1.7
6	g cm <sup>-3</sup> ; eBC: equivalent Black Carbon concentration calculated from the absorption coefficients at $\lambda$ =530 nm
7	calculated by using the B1999 algorithm without any scattering corrections and with B1999 and V2010
8	algorithms using $\sigma_{sp} = \sigma_{sp}(DMPS, PM_{10})$ for the scattering corrections and assuming MAC = 7.78 m <sup>2</sup> g <sup>-1</sup> . fPM <sub>10</sub> :
9	scattering-corrected eBC mass fraction calculated from eBC/m(DMPS,PM <sub>10</sub> )×100%; fPM <sub>0.8</sub> : scattering-corrected
10	

 $10 \qquad \text{eBC mass fraction calculated from eBC/m(DMPS, \text{PM}_{0.8}) \times 100\%.}$ 

			Percentiles						
	N	Ave ± std	5	25	50	75	95		
Mass concentrations									
PM₁₀ filters, ng m⁻³	1765	162 ± 161	30	73	126	201	394		
PM₁ filters, ng m⁻³	468	82 ± 64	13	33	66	114	208		
m(DMPS, PM <sub>0.8</sub> ), ng m <sup>-3</sup>	21987	123 ± 161	12	31	70	150	382		
m(DMPS, $PM_{10}$ ), ng m <sup>-3</sup>		171 ± 255	23	54	108	199	483		
eBC from $\sigma_{ap}(\lambda = 530$	) nm) calcula	ited using B1999							
eBC(no $\sigma_{sp}$ correction), ng m <sup>-3</sup>	15815	$2.6 \pm 2.6$	0.36	0.94	1.9	3.2	7.6		
$eBC(\sigma_{sp}(DMPS,PM_{10})))$ , ng m <sup>-3</sup>	15815	1.7 ± 2.0	< 0.2	0.37	1.10	2.4	5.7		
feBC(m(DMPS,PM <sub>0.8</sub> )), %	15815	2.6 ± 2.7	< 0.1	0.53	1.62	4.0	7.8		
feBC(m(DMPS,PM <sub>10</sub> )), %	15815	1.6 ± 1.7	< 0.1	0.35	1.16	2.5	4.7		
eBC from $\sigma_{ap}(\lambda = 530$	) nm) calcula	ated using V2010							
$eBC(\sigma_{sp}(DMPS,PM_{10})))$ , ng m <sup>-3</sup>	15815	1.6 ± 1.9	< 0.2	0.29	0.98	2.2	5.3		
feBC(m(DMPS,PM <sub>0.8</sub> )), %	15815	2.4 ± 2.6	< 0.1	0.41	1.48	3.7	7.3		
feBC(m(DMPS,PM <sub>10</sub> )), %	15815	1.5 ± 1.6	< 0.1	0.28	1.06	2.3	4.5		





## 1 Figures

						<ul> <li>PSAP (26264 H)</li> <li>DMPS (22072 H)</li> <li>PSAP &amp; DMPS &amp; WIND (15815 H)</li> <li>GRIMM OPC (8958 H)</li> <li>PM10 (N=1765)</li> <li>PM1 (N=468)</li> </ul>
2008	2009	2010	2011	2012	2013	

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3 Figure 1. The periods of the PSAP, the DMPS, the Grimm OPC and the PM<sub>1</sub> and PM<sub>10</sub> filter sample data. The

4 number of hours of accepted data and the number of samples are shown in parentheses for the continuous

5 instruments and the filter samplers, respectively.

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Figure 2. Average particle size distributions in summer (DJF) and in winter (JJA) in December 2007 – July 2009 when both the DMPS and the Grimm OPC were operational. Left: average and median (a) number, (c) volume, and (e) scattering size distributions at  $\lambda$  =530 nm; right (b, d, and f): cumulative fractions of the respective parameters in the size range D<sub>p</sub> < 7.5 µm. which corresponds to the aerodynamic particle size range D<sub>a</sub> < 9.8 µm.









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Figure 3. Seasonal cycle of the contribution of the size range measured by the DMPS to a) volume concentration
and b) scattering coefficient at the PSAP wavelengths in December 2007 – July 2009 when both the DMPS and
the Grimm OPC were operational. The circle shows the average, the horizontal line the median, the box the 25<sup>th</sup>
to 75<sup>th</sup> percentile range, and the whiskers the 5<sup>th</sup> to 95<sup>th</sup> percentile range in each month.





8 Figure 4. Wind and absorption coefficient. a) Hourly-averaged non-scattering-corrected absorption coefficients 9  $(\sigma_{ap,nsc}, Eq. (1))$  observed at wind speed WS > 2 m/s in 5° wind direction (WD) sectors. The lines present the 10 percentiles of the cumulative  $\sigma_{ap,nsc}$  distribution in each WD sector. f(WD sector): fraction of wind data from each 11 sector. CS 75<sup>th</sup> perc.: Contamination sector determined from the 75<sup>th</sup> percentiles of the cumulative  $\sigma_{ap,nsc}$ 12 distribution. CS 99<sup>th</sup> perc.: Contamination sector determined from the 99<sup>th</sup> percentiles of the cumulative  $\sigma_{ap,nsc}$ 13 distribution. b) Distribution of WS and WD as a wind rose.







Figure 5. Time series of scattering coefficients calculated from the DMPS ( $\sigma_{sp}$ (DMPS,PM<sub>10</sub>) at  $\lambda$  = 530), PM<sub>1</sub> and PM<sub>10</sub> ( $\lambda$  = 550) filter data measured at Dome C and measured with the nephelometer at the South Pole Station (SPO) ( $\lambda$  = 550) and  $\sigma_{ap,nsc}$ ( $\lambda$  = 530) measured with the PSAP at Dome C. The  $\sigma_{sp}$  from the DMPS and the nephelometer and  $\sigma_{ap,nsc}$  are running 24-hour averages at each hour (± 12 hours), and the  $\sigma_{sp}$  from the PM1 and PM10 filters are those calculated for each filter. The red box within the 2011 time series shows the period presented in more detail in Fig. 6 and the red asterisk symbol (\*) for which the footprint in Fig. 7 was calculated.

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Figure 6. Four-month time series (May-August 2011) of  $\sigma_{sp}$ ,  $\sigma_{ap}$  and  $\omega_o$  at  $\lambda = 530$  nm at Dome C and  $\sigma_{sp}$  at at  $\lambda = 550$  nm at the South Pole Station (SPO): a) upper estimate of  $\sigma_{sp}$  (=  $\sigma_{sp}$ (DMPS,PM<sub>10</sub>)), lower estimate of  $\sigma_{ap}$  (=  $\sigma_{ap}(\sigma_{sp}(DMPS, PM_{10}))$ ) and upper estimate of  $\omega_o$ . b) lower estimate of  $\sigma_{sp}$  (=  $\sigma_{sp}(DMPS, PM_{0.8})$ )), upper estimate of  $\sigma_{ap}$  (=  $\sigma_{ap}(\sigma_{sp}(DMPS, PM_{0.8}))$ ) and lower estimate of  $\omega_o$ . In both a) and b) also the non-scattering-corrected absorption coefficient is shown. All values are running 24-hour averages at each hour (± 12 hours). The numbers 1 - 4 are discussed in the text.

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4 Figure 7. FLEXPART footprint of the overall highest day of scattering in winter 2011, on Jul 28, 2011, indicated by

5 the number (3) in Fig. 6.







Figure 8. Comparison of scattering coefficients calculated from the DMPS vs. scattering coefficients calculated from the PM<sub>1</sub> and PM<sub>10</sub> filter sample data at Dome C, all at  $\lambda = 550$  nm. The scattering coefficients calculated from the DMPS data were averaged for the sampling times of the PM<sub>1</sub> and PM<sub>10</sub> samples and interpolated to  $\lambda =$ 550 nm. a) lower estimate of  $\sigma_{sp}$  (=  $\sigma_{sp}$ (DMPS, PM<sub>0.8</sub>)) vs.  $\sigma_{sp}$ (PM<sub>10</sub>), b) upper estimate of  $\sigma_{sp}$  (=  $\sigma_{sp}$ (DMPS,PM<sub>10</sub>)) vs.  $\sigma_{sp}$ (PM<sub>10</sub>), c)  $\sigma_{sp}$ (DMPS, PM<sub>0.8</sub>)) vs.  $\sigma_{sp}$ (PM<sub>1</sub>), d)  $\sigma_{sp}$ (DMPS,PM<sub>10</sub>)) vs.  $\sigma_{sp}$ (PM<sub>1</sub>). N: number of data points. The red line shows the linear regression line that is forced through zero. The regression equations show the slope ± standard error of the slope, the squared correlation coefficient and the p value of the slope.

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2 Figure 9. Seasonal cycles of scattering and absorption coefficients and single-scattering albedo. a) Scattering 3 coefficient ( $\sigma_{sp}$ ) calculated from the sums of analyzed ion concentrations in PM<sub>1</sub> filters at  $\lambda$  = 550 nm, b)  $\sigma_{sp}$ 4 calculated from the sums of analyzed ion concentrations in  $PM_{10}$  filters, c) the lower estimate of  $\sigma_{sp}$  = 5  $\sigma_{sp}(DMPS,PM_{0.8})$ , d) the upper estimate of  $\sigma_{sp} = \sigma_{sp}(DMPS,PM_{10})$ , e) absorption coefficient  $\sigma_{ap}$  calculated with the 6 algorithms of B1999 and V2010 (Eqs. (17) and (18)) by using the  $\sigma_{sp}$  lower estimate for scattering correction, f) 7  $\sigma_{ap}$  calculated with the two algorithms by using the  $\sigma_{sp}$  upper estimate for scattering correction, g) single-8 scattering albedo  $\omega_o$  calculated by using the  $\sigma_{so}$  lower estimate for both  $\sigma_{so}$  and  $\sigma_{ao}$  and h)  $\omega_o$  calculated by using 9 the  $\sigma_{sp}$  upper estimate for both  $\sigma_{sp}$  and  $\sigma_{ap}$ .







Figure 10. Seasonal cycles of the wavelength dependency of a) scattering and b) absorption. In a) the Ångström exponent  $\alpha_{sp}$  was calculated from the size distributions measured at Dome C ( $\sigma_{sp}$ (DMPS,PM<sub>0.8</sub>) and  $\sigma_{sp}$ (DMPS,PM<sub>10</sub>) for the wavelength range 467 – 660 and measured at the South Pole Station with a nephelometer. The SPO  $\alpha_{sp}$  was calculated for the wavelength range 550 – 700 nm. In b) the absorption Ångström exponent  $\alpha_{ap}$  was calculated for the  $\sigma_{ap}$  without scattering correction, and by using the B1999 and V2010 algorithms with scattering corrected using  $\sigma_{sp} = \sigma_{sp}$ (DMPS,PM<sub>10</sub>), In all of them the data with  $\sigma_{ap} > 3\delta\sigma_{ap}$  were used.

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2 Figure 11. Seasonal cycles of a) aerosol mass concentration calculated from the particle number size distributions 3 m(DMPS,PM<sub>10</sub>) and m(DMPS,PM<sub>0.8</sub>), the sum of ions analyzed from PM<sub>1</sub> and PM<sub>10</sub> filters, b) mass fraction of the 4 sum of nssSO<sub>4</sub><sup>2-</sup> and MSA in PM<sub>1</sub> and PM<sub>10</sub> filters, c) equivalent black carbon (eBC) concentration calculated from 5 the non-scattering-corrected absorption coefficients, and from  $\sigma_{ap}$  corrected with the  $\sigma_{sp}$  upper estimate 6  $(\sigma_{sp}(DMPS, PM_{10})), d)$  mass fraction eBC calculated as the ratio of eBC corrected with the  $\sigma_{sp}$  upper estimate to 7 m(DMPS,PM<sub>0.8</sub>) and m(DMPS,PM<sub>10</sub>). Comparison values: a) monthly average mass concentration calculated from 8 particle volume concentrations at Troll (F2014: Fiebig et al., 2014), average gravimetric PM<sub>10</sub> mass concentration 9 at Dome C (A2011: Annibaldi et al., 2011), c) monthly median eBC concentrations measured at Neumayer (NM) 10 with a MAAP and an Aethalometer (AE) (W2013: Weller et al., 2013), at SPO with an Aethalometer (S2016: 11 Sheridan et al., 2016), and at Syowa with an Aethalometer (H2019: Hara et al., 2019). The small insert in b) shows 12 the scatter plot and linear regression of monthly average  $\alpha_{sp}$  (DMPS,PM10) vs. monthly average mass fraction of 13 the sum of nssSO<sub>4</sub><sup>2-</sup> and MSA in PM<sub>10</sub> filters.







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Figure 12. Monthly average and median  $\omega_0$  vs. eBC mass fraction calculated as the ratio of eBC to a) m(DMPS,PM<sub>0.8</sub>) and b) m(DMPS,PM<sub>10</sub>). The dashed lines represent fittings of  $\omega_0$ (feBC) =  $\omega_0$ (0)exp(-k·feBC) with the data. The continuous lines in b) represent simulations with a core-shell (CS) model for lognormal number size distributions with geometric standard deviation GSD = 1.8 and geometric mean diameter GMD shown in the legend.







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Figure 13. The residence time of 60000 50-day FLEXPART backtrajectories arriving at Dome C in 2008 – a 2013 as a function of latitude and altitude above ground level. The residence time is colorcoded from blue to yellow to show increasing time spent in each grid cell. a) The residence times of the inert tracer backtrajectories. White line: the average altitude of trajectories at each latitude. b) Same as the in (a) but for the BC tracer. Black line: the average altitude of the BC tracer, the white line is for comparison average altitude of the inert tracer. The black triangle shows the latitude of Dome C (75°06'S). The grey text boxes between a) and b) show the approximate latitude range of some major BC emitting regions.







Figure 14. The relative difference RD, Eq. (19), of transport for the two lowest layers (>1000 magl) between the
highest and the lowest 10% of a) eBC concentration and b) single-scattering albedo. Positive values indicate a

5 relative increase in transport, and negative values a decreased transport from a given area by the highest 10%

- 6 of measurements.
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2 Figure 15. Seasonal air mass transport by region calculated as the fraction of time the FLEXPART 50-day

3 trajectories spent over Antarctica, the surrounding oceans and continents.







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Figure 16. Seasonal cycles of fire-related BC emissions from the surrounding continents and eBC concentration
 at Dome C. a) BC emissions and eBC concentrations, b) RTW BC emissions, c) f(t) × BC emissions 1 month earlier,

4 d) f(t) × BC emissions 2 months earlier. The numbers show the squared correlation coefficient of linear regression

5 between eBC concentrations and  $f(t) \times BC$  emissions with the shown time shifts.







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2 Figure 17. Normalized seasonal cycles of a) major natural sources of light-scattering aerosols from the oceans, b) 3 scattering coefficients at Dome C and c) scattering coefficients at SPO.  $\sigma_{sp}(DMPS,PM_{0.8})$  and  $\sigma_{sp}(DMPS,PM_{10})$ : 4 scattering coefficient calculated from particle number size distributions;  $\sigma_{sp}$ (nss-SO<sub>4</sub><sup>2-</sup>,PM<sub>1</sub>): scattering 5 coefficient calculated from non-seasalt sulfate concentrations in PM<sub>1</sub> filter samples;  $\sigma_{so}$  (nephelometer, SPO): 6 scattering coefficient measured with the nephelometer at SPO; Cphyto: phytoplankton biomass concentration in 7 2008 – 2013 calculated as monthly averages from the time series presented by Behrenfeld et al. (2016) using 8 CALIOP satellite data in Zone 1 (45° − 55°S), Zone 2 (55° − 65°S), and Zone 3 (65° − 75°S); I<sub>T</sub>(50°S), I<sub>T</sub>(60°S) and 9 I<sub>T</sub>(70°S): average global radiation at latitudes 50°S, 60°S and 70°S at longitude 0° at 10 − 14 UTC; PAR(Zone 3): 10 photosynthetically active radiation from Behrenfeld et al. (2016); SSA: Sea Spray Aerosol flux modeled according 11 to Grythe et al. (2014).







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3 the PM<sub>1</sub> filter samples  $\sigma_{sp}(nssSO_4^{2-}, PM_1)$  at Dome C vs. normalized monthly averages of C<sub>phyto</sub>PAR in polar latitude

4 zone with and without a time shift.