- 1 Multi-year black carbon observations and modeling close to the
- 2 largest gas flaring and wildfire regions (Western Siberian Arctic)
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Abstract

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The influence of aerosols on the Arctic system remains associated with significant uncertainties, particularly concerning black carbon (BC). The polar aerosol station "Island Bely" (IBS), located on Bely Island (Kara Sea) in the Western Siberian Arctic, was established to enhance aerosol monitoring in the Arctic. Continuous in-situ measurements from 2019 to 2022 revealed the long-term effects of light-absorbing carbon. During the cold period, the annual average light absorption coefficient was $0.7 \pm 0.7 \text{ Mm}^{-1}$, decreasing by approximately 2-3 times during the warm period. The interannual mean showed a peak in February (0.9 \pm 0.8 Mm⁻¹), a ten times lower minimum in June, and exhibited high variability in August (0.7 \pm 2.2 Mm⁻¹). The absorption Ångström exponent (AAE) indicated presence of mixed and aged BC. An increase of up to 1.5 at shorter wavelengths from April to September suggests contribution from light absorbing brown carbon (BrC). The annual mean equivalent black carbon (eBC) demonstrated considerable interannual variability, with the lowest in 2020 (24 ± 29 ng m⁻³). Significant difference was observed between Arctic Haze and Siberian wildfire periods, with record-high pollution levels in February 2022 (110 \pm 70 ng m⁻³) and August 2021 (83 \pm 249 ng m⁻³). Overall, anthropogenic BC contributed 83% to the total for the entire study period and gas flaring, domestic combustion, transportation, and industrial emissions dominated. During the cold season, >90% of surface BC was attributed to anthropogenic sources, mainly gas flaring. In contrast, during the warm period, Siberian wildfires contributed to BC concentrations by 48%. Notably, unprecedented smoke was transported from Yakutian wildfires at high altitudes in August 2021, marking the most severe fire season in the region over the past four decades.

1 Introduction

Multiple socio-economic drivers and feedbacks, including air pollution (Arnold et al., 2016) influence the natural and human environment of the Arctic. Over the last few decades, the Arctic warms more than three times faster than the global average (AMAP, 2021). The pronounced rapid changes affect atmospheric transport and aerosol relative source contributions (Heslin-Rees et al., 2020). Drier conditions and warmer temperatures are the main cause of enhanced fire activity. Boreal forest fires become more frequent and severe (Rogers et al., 2020), especially in Central Siberia, and Northern America (Kasischke and Turetsky, 2006; Kharuk and Ponomarev, 2017; Veraverbeke et al., 2017). Widespread smoke plumes, particularly in Siberia, lead to substantial deterioration of air quality increasing fine particulate matter (Silver et al., 2024).

Interactions between aerosol and different cloud types, available solar radiation, sea ice, surface albedo, Arctic and lower latitude removal processes, and atmospheric transport patterns. affect Arctic pollution and its climate impacts (Willis et al., 2018), such as the Arctic haze (namely the persistent Arctic air pollution during late winter and early spring). To understand such phenomena and thus reduce their impact, there is a clear need for comprehensive studies of the climate-relevant aerosol processes that occur in the Arctic. A species of major concern is BC, a short-lived climate forcer (Schmale et al., 2021). BC is emitted from the incomplete combustion of fossil fuel and biomass; it is defined as the portion of carbonaceous aerosols, which absorb strongly in the entire climate relevant wavelength region of the solar spectrum (i.e. IR-VIS-UV). BC contributes to Arctic warming in multiple ways (e.g., Lee et al., 2013), including the darkening effect of BC deposited on snow and ice (Flanner, 2013). AMAP (2015) reports that the Arctic equilibrium temperature response is (+0.4°C) due to forcing from atmospheric BC and (+0.22°C) due to snow BC.

At present, the largest uncertainties when assessing aerosol impact on the climate are attributed to BC (AMAP, 2021). To follow-up on this, BC measurements are taken at various polar regions in the European, Siberian, and Canadian Arctic (Stone et al., 2014; Yttri et al., 2014; Popovicheva et al., 2019a; Winiger et al., 2019; Manousakas et al., 2020; Gilardoni et al., 2023). For instance, Stathopoulos et al. (2021) reported on the long-term impact of light-absorbing carbon in the high Arctic by analysing 15 years of data from the Zeppelin station (Svalbard), while Schmale et al. (2022) studied the status of the Arctic haze peak concentrations at 10 Arctic observatories.

There is a large diversity in magnitude and variability of aerosol optical properties, reflecting differences in sources throughout the Arctic (Schmeisser et al., 2018). BC measurements are based on various instrumentations and methods that increase uncertainty (Sharma et al., 2017; Asmi et al.,

2021; Ohata et al., 2021). The optical properties of BC have been previously evaluated against direct mass measurements techniques (Sharma et al., 2004; Eleftheriadis et al., 2009; Yttri et al., 2024). The conversion of light attenuation to absorbing carbon mass concentration is performed by the mass-specific absorption coefficient (*MAC*) (Petzold et al., 2013) that is highly influenced by the aerosol mixing state and non-BC light-absorbing species such as organic matter and mineral dust (Zanatta et al., 2018) and varies in time and space depending on sources and transformations during transport (Bond et al., 2013; Chen et al., 2023). Therefore, it is crucial to quantify the contribution of non-BC component and aging in order to determine the actual *MAC* value experimentally at each site (Singh et al., 2024).

The AAE defined as the relative fraction of wavelength - dependence of absorption of BC versus other light absorbing constituents, also differs from site to site (Schmeisser et al., 2018). A fraction of organic aerosol, the BrC, increases the aerosol absorbing properties at short UV-VIS wavelengths (Sandradewi et al., 2008; Grange et al., 2020; Helin et al., 2021) and dominates the absorption during wildfire seasons (Bali et al., 2024). BrC originates mainly from biomass burning (BB) and can impose strong warming effect in the Arctic, especially in the summertime (Yue et al., 2022).

Despite its remoteness, the Arctic is one of the main receptors of anthropogenic air pollutant emissions from the Northern Hemisphere (Stohl et al., 2013). BC trends and seasonality at three Arctic sites, Alert (Canadian Arctic), Barrow/Utqiagvik (American Arctic), and Zeppelin, Ny-Ålesund (European Arctic) reveal a negative trend of 40% over 16 years due to the anthropogenic emission reduction (Sharma et al., 2013). The recent increase in fires and their earlier starts, due to the ongoing warming, have made wildfires in the Northern Eurasia a significant source of Arctic BC (Evangeliou et al., 2016). Fossil fuel combustion is the major source of BC in the Arctic troposphere (50–94%) (55–68% at the surface and 58–69% in the snow) and BB dominates at certain altitudes (600–800 hPa) between April to September (Qi and Wang, 2019). This agrees with Matsui et al. (2022) who reported that the largest contribution to Arctic BC is from BB sources in Siberia travelling at high altitudes.

Northern Eurasia, particularly Siberia, is a key source region of pollution in the Arctic. Source quantification (Zhu et al., 2020) shows that surface Arctic BC originates mainly from anthropogenic emissions in Russia (56%). The reason for this is that the largest oil and gas producing facilities of Western Siberia are located along the main pathway of air masses that enter the Arctic and thus have a disproportionally large contribution to the Arctic lower troposphere (Stohl, 2006; Stohl et al., 2013). Eleftheriadis et al. (2009) and Tunved et al. (2013) identified these regions as a key source for the highest measured BC in the European Arctic. The impact of long-range transport from these

regions has been previously reported in Ice Cape Baranova station (Manousakas et al., 2020) and Tiksi (Northeastern Siberia) (Winiger et al., 2017; Popovicheva et al., 2019a). Airborne observations over the coast of the Arctic seas have identified the long-term transport of the industrial pollution (Zenkova et al., 2022). Furthermore, efforts have sought to develop BC emission inventories for the Siberian Arctic, based on activity data from local information, improved gas flaring emissions, and satellite data (Huang et al., 2015; Böttcher et al., 2021; Kostrykin et al., 2021; Vinogradova and Ivanova, 2023). To better quantify the source contribution to the Arctic environment, targeted aerosol measurements close to the flaring facilities are needed. The present operating Eurasian Arctic stations are all too far away to allow assessing how air masses are affected by different source categories (Stohl et al., 2013). However, ship campaigns focusing on BC close to main source regions (e.g., gas flaring facilities of the Western Siberia) have provided a better constraint of how anthropogenic and BB sources influence Arctic polution (Popovicheva et al., 2017b).

Another major source of the Arctic BC is wildfires in the Siberian and Far Eastern regions, which have grown in recent summers (Bondur et al., 2020). Airborne observations of BC in Siberia have confirmed impact forest fires (Paris et al., 2009). Eastern Siberia (Yakutia) has been prone to large wildfires due to a combination of hot summers (> 40°C) and low humidity (Tomshin and Solovyev, 2022). For instance, wildfires in summer 2019 in Eastern Siberia occurred along the trans-Arctic transport pathway resulting in enhanced aerosol load observed in Western Canada (Johnson et al., 2021). BB emissions occurring at midlatitudes reached the European Arctic in 2020 influencing aerosol composition (Gramlich et al., 2024).

Despite the necessity for detailed observations in the Northwestern Siberia, a dense observational network is still absent. Towards this, the polar aerosol station on the Bely Island (Kara Sea, Western Siberia) started to operate in August 2019 (Popovicheva et al., 2022, 2023). The significance of high-quality measurements at the IBS is documented, as the station is located along the main pathway of large-scale emission plumes from industrial regions and Siberian wildfires entering the Arctic (Popovicheva et al., 2022). Further investigation performed at IBS in August 2021 showed impact from a long-range transport event with unprecedented high concentrations of carbonaceous aerosol (Schneider et al., 2024).

In this paper, we show improved light absorption long-term measurements and BC seasonal and inter-annual variability in the Western Siberian Arctic from three and a half years (2019-2022) of observations at IBS. BC was calculated in two ways: as *eBC* by an aethalometer and as elemental carbon (EC) by thermal-optical analysis. We further evaluate the seasonal changes in the observed absorption coefficients. Seasonal difference in intensive optical properties is shown by the

wavelength-dependent AAE, which acts as indication of the BrC impact. Estimated site-specific absorption coefficient (SAC) considered the specific seasonal effects of mixing and aging of aerosols at IBS. We further assess the inter-annual variability of origin, transport and main BC sources using modelling tools coupled with the most recent anthropogenic and BB emission datasets.

2 Methods

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2.1 Polar aerosol Island Bely station, location and meteorology

The aerosol IBS of Moscow State University (73020'7.57"N, 70040'49.05"E) is shown in Figure 1a together with other Polar Arctic observatories. Western Siberia is the world's largest gas flaring region with a leading oil and gas production industry (Figure 1b). It is also an area under intensive exposure by Siberian wildfires (Tomshin and Solovyev, 2022; Voronova et al., 2022). A for 5th August 2021 satellite image of smoke plume was obtained from https://worldview.earthdata.nasa.gov. Fires are shown from the Fire Information for Resource Management System (FIRMS) (https://firms.modaps.eosdis.nasa.gov/map) ten days back in time.

The climate at IBS is characterized by a large annual variability determined by alternating periods of the polar night and midnight sun. Basic meteorological parameters, such as temperature, wind speed and direction were obtained every 3 hours from a meteorological station located 500 m away from the IBS. The cycles of temperature, precipitation, snow coverage, wind speed and relative humidity are shown in **Figure S 1**. Annual temperature varied from -39°C to 23°C (mean: -6±12°C) (**Table S 1**). For further analysis, we have split the annual cycle in two periods based on the prevailing temperatures, November-April ("cold period", -15.9±9.1) and May - October ("warm period", 2.8±5.8). High relative humidity of 87±8% was typical for the study period, with less than 80% observed in winter 2020. Precipitation was maximum in summer (22 mm) with constant snow coverage from October to May. Wind was relatively stable, with a mean speed of 6±3 m s⁻¹, which increased in winter up to 17 m s⁻¹ (**Figure S 1**).

Wind patterns for the cold period in **Figure S** *I* show a prevailing wind direction from south, southwest, and southeast. Winds were predominantly continental, rarely occurring from the ocean; significant emission sources from the continent were downwind. In the warm period, the wind patterns were more spatially homogeneous with northeastern direction. Period from June and September was characterized by a frequent occurrence of oceanic air masses and constant wind speeds.

2.2 Aerosol optical and chemical characterization

The aerosol pavilion takes place approximately half a km to the southeast of the meteorological station. An aerosol sampling system composed from three total suspended particle inlets has been installed approximately 1.5 m above the roof and 4 m above the ground. They are equipped with an electric heating wire to prevent rimming and ice blocking of the system. One inlet was used for the real-time light-absorption measurements with air flow 5 L min⁻¹. Two other inlets low-volume samplers (Derenda, Germany) were used for sampling of total suspended particles (TSP) operating at 2.3 m3 h-1 flow (0°C, 1013.25 hPa).

An Aethalometer model AE33 (Magee Scientific, Aerosol d.o.o.) was used to measure the light attenuation caused by particles deposited on two filter spots at different flow rates (Drinovec et al., 2015) and at seven wavelengths from ultraviolet (370 nm) to infrared (950 nm). The "dual spot" technique is applied for real-time loading effect compensation. Optical absorption of aerosols on the filter is influenced by scattering of light within the filter; the enhancement of optical absorption is described by the factor C that depends on the filter material. The producer recommends an enhancement factor of 1.57 for TFE-coated glass fiber filter. The light-absorbing content of carbonaceous aerosol is reported as eBC concentration by aethalometer (eBC_{AET}) for the given wavelength λ , which is determined for each time interval from the change in the light absorption using the MAC. The aerosol optical absorption coefficient (b_{abs}) is therefore:

$$b_{abs}(\lambda) = eBC_{AET}(\lambda) \times MAC(\lambda)$$
 (1)

where eBC_{AET} at 880 nm is determined using the MAC of 7.7 m² g⁻¹. The aerosol optical absorption coefficient for different wavelengths is determined with their MAC values that are equal to 11.58, and 13.14 m² g⁻¹ at 590, and 520 nm, respectively.

To represent the spectral dependence of the light absorption, the AAE was derived by using a fitted power law relationship:

$$b_{abs}(\lambda) = b_{abs}(\lambda_o) \times \left(\frac{\lambda}{\lambda_o}\right)^{-AAE}$$
 (2)

where $b_{abs}(\lambda_o)$ is the absorption coefficient at the reference wavelength λ_o , AAE is a measure of strength of the spectral variation of aerosol light absorption.

BC absorbs strongly in the NIR-VIS (near-infrared and visible) with only moderate increment towards the shorter wavelengths. Light absorbing organic components related to BrC absorb light at shorter wavelengths more effectively than at 880 nm, which is observed as an increased *AAE* (Sandradewi et al., 2008; Grange et al., 2020; Helin et al., 2021). The total light absorption is assumed to include the contribution of both BC and BrC (Ivančič et al., 2022):

$$b_{abs}(\lambda) = b_{abs/BC}(\lambda) + b_{abs/BrC}(\lambda)$$
 (3)

Using Eq. 1, the BrC absorption becomes: $b_{abs/BrC}(\lambda) = b_{abs}(\lambda) - b_{abs}(\lambda_0) \times \left(\frac{\lambda}{\lambda_0}\right)^{-AAE}$ (4)

Light-absorption measurements were performed for three and a half years, from 10 August 2019 to 31 December 2022, with a time resolution of 1 min. Data were cleaned based on analysis of meteorological parameters by examining whether the wind originated from the direction of the meteorological station where diesel generators operated. In such cases, strong peaks of BC were removed from further analysis. Around 6.4 % of the hourly-average data were cleaned from the dataset due to local pollution impact. To avoid the instrumental noise when calculating the *AAE*, the z-score was used that calculates the ratio of difference between a single raw data value and the data mean to the data standard deviation. Outliers (< -3 and > 3 of observation's z-score) were removed from the dataset.

A thermal EC analysis was conducted for the samples in parallel to AE33. Sampling was performed on 47 mm quartz fiber (Pallflex) filters preheated at 600°C for 5 h. The low concentrations of ambient aerosols necessitate that the sampling times reach up to a week, in order to allow the filter loading to exceed the detection limit for relevant aerosol chemistry analyses. The total number of samples limited by the low detection limit of the thermal-optical instrument were 180.

Organic (OC) and EC were measured by thermo-optical transmittance (TOT) analysis (Lab OC-EC Aerosol Analyzer, Sunset Laboratory, Inc.) using the methodology reported in Popovicheva et al. (2019) and Manousakas et al. (2020). Quartz filter samples were heated first up to 650 °C in He atmosphere and then up to 850 °C in a mixture of 2% O2 in He, using the controlled heating ramps of the EUSAAR_2 thermal protocol. OC evolves in inert atmosphere, while the thermal refractory fraction EC is oxidized in the He-O2 atmosphere. Charring correction due to pyrolytic carbon was applied by monitoring the sample transmittance throughout the heating process. The limit of detection for the EC analysis was 0.05 µg C cm⁻². QA/QC procedures of EN 16909:2017 were also applied during TOT analysis. Laboratory and field blanks were prepared and ran following the same analytical procedures as for the samples.

Both methods have important uncertainties (10 - 80%, Sharma et al., 2017; Ohata et al., 2021). The determination of EC by thermo-optical analysis may be impacted by the presence of carbonate carbon (CC), which is quantified during analysis as OC and/or EC. The contribution of CC in fine aerosol is generally considered negligible but its interference may be significant for coarse aerosol and samples heavily impacted by resuspended soil. The split between EC and OC

may be also affected by the presence of light-absorbing species others than EC, such as light absorbing organic carbon. In addition, the presence of mineral oxides, such as iron oxide, might provide oxygen during analysis and lead to pre-oxidation of EC in inert atmosphere. *eBC* might overestimate BC if there are coexisting components such as BrC (Chakrabarty et al., 2010) and dust (Petzold et al., 2009). In addition, the aethalometer response depends on filter loading and multiple scattering by the filter medium and sampled aerosol particles (Backman et al., 2017).

Validations of *eBC* retrievals were performed against results from thermal-optical analysis of EC according to an approach that has been used previously in Sharma et al. (2004), Eleftheriadis et al. (2009) and Yttri et al. (2014). To convert optical absorption at 880 nm to BC mass, the *SAC* was estimated as:

$$SAC = \frac{b_{abs/BC}}{EC} \tag{5}$$

Data processing was performed using Deming's total least-squares regression to compare measurements from different methods and modelling, estimate the *MAC*, and evaluate correlations among variables (R package "Deming"; (Therneau, 2024)). Deming regression fits a couple of variables considering the independent errors of both. The errors are assumed to be normally distributed; the error ratio is 1, and the regression results are equivalent to the orthogonal regression with the intercept forced through zero.

2.3 Atmospheric dispersion modelling and emission inventories

To investigate the air mass transport and possible origin of BC during the study period (2019 – 2023), the Lagrangian particle dispersion model FLEXPART version 10.4 was used (Pisso et al., 2019) driven by hourly reanalysis meteorological fields (ERA5) from the European Centre for Medium-Range Weather Forecasts (ECMWF) with 137 vertical levels (up to approximately 80 km) and a horizontal resolution of 0.5°×0.5° (Hersbach et al., 2020). In FLEXPART, computational particles were released at heights 0 - 100 m from the receptor (IBS) and tracked backward in time in FLEXPART's "retroplume" mode. Simulations extended over 30 days backward in time, sufficient to include most BC emissions arriving at the station, given a typical BC lifetime of 1 week (Bond et al., 2013). The tracking includes gravitational settling for spherical particles, dry and wet deposition of aerosols (Grythe et al., 2017), turbulence (Cassiani et al., 2015), unresolved mesoscale motions (Stohl et al., 2005), and deep convection (Forster et al., 2007). The FLEXPART output consists of a footprint emission sensitivity that expresses the probability of any emission occurring in each grid-cell to reach the receptor. The footprint can be converted to modelled concentration at the receptor, when coupled with gridded emissions from an emission inventory. Modelled concentrations can be calculated as a function of the time elapsed since the emission has occurred (i.e., "age"), which can

be shown as "age spectrum", while masks of specific regions/continents can give the continental contribution to the simulated concentration (i.e., "continent spectrum").

The source contribution to receptor BC is calculated by combining each gridded emission sector (e.g. gas flaring, transportation, waste management etc...) from an emission inventory with the footprint emission sensitivity (as described in the previous paragraph). Calculations for anthropogenic sources (emission sectors are described below) and open BB were performed separately. This enabled identification of the exact origin of BC and allowed for quantification of its source contribution. Anthropogenic emissions were adopted from the latest version (v6b) of the ECLIPSE (Evaluating the CLimate and Air Quality ImPacts of ShortlivEd Pollutants) dataset, an upgraded version of the previous version (Klimont et al., 2017). The inventory includes emissions from industrial combustion (IND), from the energy production sector (ENE), residential and commercial emissions (DOM), emissions from waste treatment and disposal sector (WST), transportation (TRA), shipping activities (SHP) and gas flaring emissions (FLR). The methodology for obtaining emissions from FLR specifically over the Russian territories has been improved in ECLIPSEv6 (Böttcher et al., 2021). Annual total and monthly anthropogenic emissions are shown in Figure S 2. BB was adopted from the Copernicus Global Fire Assimilated System (CAMS GFAS) (Kaiser et al., 2012) because this product provides an estimation of the injection altitude of the fire emissions that is crucial for accurate simulation of the BB dispersion. Annual total and daily fire emissions from CAMS GFAS are shown in Figure S 3.

3 Results

3.1 Aerosol light-absorption

Light-absorption coefficients at 880 nm, $b_{abs}(880)$ were used to infer *eBC* mass concentrations. $b_{abs}(880)$ were plotted as hourly and monthly means during the entire study period (2019-2022) (**Figure 2**). **Table 1** presents the data statistical summary. The mean \pm sigma (median) value of $b_{abs}(880)$ was 0.5 ± 0.9 (0.27) Mm⁻¹ for the entire study period. In the cold period the annual average mean (median) of $b_{abs}(880)$ was 0.7 ± 0.7 Mm⁻¹ (0.5), during the warm period it was 1.9 (2.5) times less. There is a clear seasonality consistent with the Arctic aerosol light absorption from other studies (Stathopoulos et al., 2021; Schmale et al., 2022; Pulimeno et al., 2024) due to the formation of the polar dome and the slow removal processes in the Arctic in winter (Law and Stohl, 2007). 15 years (2001-2015) record at Zeppelin demonstrated that the long-term seasonality of light absorbing carbon (Stathopoulos et al., 2021) $b_{abs}(880)$ was 0.112 Mm⁻¹

(median) in the cold period and 0.035 Mm⁻¹ in the warm period; both values approximately 5 times less than those observed at IBS.

Monthly means of $b_{abs}(880)$ for each year together with intra-annual means for IBS are shown in **Figure 2**. Specifically, annual average $b_{abs}(880)$ exhibits a significant peak during winter and summer for any year. The examination of the overall changes by the inter-annual mean reveals a gradual increase from November $(0.4\pm0.5~{\rm Mm^{-1}})$ to February $(0.9\pm0.8~{\rm Mm^{-1}})$; the latter represents the maximum light absorption observed at IBS. In February, the monthly mean of $b_{abs}(880)$ ranged from 0.4 to 1.7 Mm⁻¹ reaching the maximum $(1.7~{\rm Mm^{-1}})$ in 2022. Thus, Arctic haze is present at IBS in winter months, from December to February. Starting from March $(0.6\pm0.5~{\rm Mm^{-1}})$, the inter-annual mean decreased down to a minimum in June $(0.1\pm0.2~{\rm Mm^{-1}})$ that was 9 times less than that of February. August had the highest light-absorption (mean: $0.7\pm2.2~{\rm Mm^{-1}})$ within the summer months, ranging from 0.2 to 1.5 Mm⁻¹ and showing a maximum of 1.5 in 2021. September and October demonstrated a similar level of variability with June. At Zeppelin, the maximum of the intra-annual (2001-2015) mean was seen in March - April (0.3 Mm⁻¹) (Stathopoulos et al., 2021), coinciding with the Arctic haze phenomenon in late winter-spring that has been widely observed in the European and Canadian Arctic (Sharma et al., 2004; Schmale et al., 2022).

In order to relate the light absorption in visible spectrum to the variability on other locations (Schmeisser et al., 2018; Pulimeno et al., 2024), we calculate b_{abs} at 520 and 590 mn. The mean (median) value of b_{abs} (520) was 0.9 ± 1.6 (0.5) Mm⁻¹ for the entire study period (**Table 1**). At Ny-Ålesund (Svalbard), the annual mean (median) b_{abs} (530) averaged for 2018 to 2022 was 0.22 (0.13) Mm⁻¹ (Pulimeno et al., 2024), approximately 4 times less. Moreover, the absorption coefficient b_{abs} (550) of 0.18 (0.09) Mm⁻¹ recorded for 2012-2014 again in Svalbard (Schmeisser et al., 2018) was 4 times less compared to annual average light absorption at IBS.

We present multi-annual box-and-whisker plots of b_{abs} at 590 nm in **Figure 3**. The wavelength of 590 nm was chosen as the closest to 550 nm reported for the polar station Tiksi (Schmeisser et al., 2018; Schmale et al., 2022). The monthly medians of b_{abs} (590) in February ranged from 0.3 to 2.3 Mm⁻¹, representing the highest values observed in 2022. The highest extended interquartile range (up to 1 Mm⁻¹) was observed in the cold period. Conversely, the summer months exhibited a minimum of approximately 0.1 Mm⁻¹ for b_{abs} (590) with smaller variation of data characterized by the low interquartile range of 0.4 Mm⁻¹.

The annual cycle of b_{abs} (590) reflects the higher aerosol burden during the haze season and the low concentrations during summer at Alert, Barrow/Utqiagvik, Zeppelin, Gruvebadet, and Tiksi

(Schmale et al., 2022). Seasonality of b_{abs} medians at 550 nm for polar stations (Alert, Barrow/Utqiagvik, Tiksi, Zeppelin) from (Schmeisser et al., 2018) and b_{abs} (590) for IBS are presented in Figure 3. All sites demonstrate similar seasonal variations, albeit a different magnitude of light absorption. In February, the maximum $b_{abs}(590)$ (1.1 Mm⁻¹) was observed at IBS; a higher value has been only observed at Tiksi which is explained by the influence from local sources (Popovicheva et al., 2019a). Other stations show the Arctic haze maximum later (in March or April); a sharp decline of $b_{abs}(590)$ was observed at those months at IBS. Values similar to other Arctic stations were recorded at IBS in June, with an annual minimum of around 0.1 Mm⁻¹. Since July, $b_{abs}(590)$ at IBS was higher than at other stations except Tiksi and peaked at 0.8 Mm⁻¹ in December. The polar station Pallas exhibits the opposite behaviour peaking in spring and summer (Schmeisser et al., 2018). Pallas is located relatively south as compared to the rest of the polar stations and, hence, it is influenced by anthropogenic and biogenic emissions from surrounding boreal forests (Asmi et al., 2011). Aerosol optical properties in the IR and visible solar spectrum at IBS are different from European, Canadian and Western high-latitude polar locations due to different source origins, but light absorption coefficients are higher during the haze period (December-February).

3.2 Black carbon and site-specific mass absorption cross-section

EC collocated with light absorption observations is widely used to infer BC (Grange et al., 2020). **Figure 4**a shows concentrations of EC determined for samples collected in parallel with the aethalometer measurements from 10 August 2019 to 31 December 2022, with eBC_{AET} concentrations averaged over the sampling period. Both weekly EC and eBC_{AET} concentrations show the same seasonal variations with a maximum in winter and minimum in summer. EC concentrations are generally smaller than eBC_{AET} . The annual EC mean concentrations ranged from 6.5 to 16.3 ng C m⁻³. The highest EC (0.2 μ g C m⁻³) was recorded in December 2019 and the highest eBC_{AET} (0.4 μ g m⁻³) in December 2019 and January 2022. EC was higher (0.05 \pm 0.03 μ g C m⁻³) in the cold period and decreased (0.02 \pm 0.03 μ g C m⁻³) in the warm period (**Table 1**). Annual average mean EC during the entire study period was 0.03 \pm 0.03 μ g C m⁻³. For comparison, at Zeppelin and Villum the annual mean EC concentrations were 0.012 \pm 0.04 μ g C m⁻³ (2017-2020) (Yttri et al., 2024) and 0.029 \pm 0.03 μ g C m⁻³ (2011-2013) (Massling et al., 2015), respectively.

Annual mean OC concentrations during the entire study period were estimated as $0.45\pm0.3~\mu g$ C m⁻³. At Zeppelin, annual OC (2017-2020) was 3.5 smaller (0.13 $\pm0.1~\mu g$ C m⁻³) (Yttri et al., 2024). Notably, the multi-year average EC and OC levels at IBS are approximately 3 times higher than at Zeppelin, that correlates well with increased light absorption, as described previously. At IBS, OC was $0.4\pm0.2~\mu g$ C m⁻³ in the cold period and increased to $0.5\pm0.4~\mu g$ C m⁻³ in warm period, opposite

to EC (**Table 1**). The ratio OC/EC shows increased OC and decreased EC in the warm period and an opposite tend in the cold (**Figure 4**b). **Figure 4**c depicts the relationship between eBC_{AET} and EC in cold and warm periods. We note the high R² values for the cold period (0.88) and slightly lower ones for the warm one (0.78). During the warm period, seasonal mean values reveal an overestimation of eBC_{AET} that is more pronounced during the warm period, with a slope equal to 2.3. R² values were lower because many EC values were close to the LOD. Seasonal differences are attributed to pollutant sources altering the chemical composition of aerosol at IBS. A positive correlation was observed between eBC_{AET} /EC and OC/EC indicating that BC at IBS is coated with OC leading to the lens effect (Kanaya et al., 2008) and overestimating eBC.

Similar seasonal variation for *eBC* and EC with highest winter and lower summer concentrations has been observed previously at Villum, with a regression slope of 2 and a R² of 0.64 (Massling et al., 2015). At Alert, the median *SAC* during the Arctic haze season (November to April) was 19.8 m² g⁻¹ (Sharma et al., 2004). However, during the non-Arctic haze period from May to October it was significantly higher 28.8 m² g⁻¹ and much more variable. This is explained by aged, internally mixed, and of anthropogenic origin of winter and spring arctic aerosols while summer aerosols were affected by local sources.

Following the definition in Eq.5, we calculate the SAC from the slope of BC light absorption at 880 nm, $b_{abs/BC}$ (880), and EC concentrations. $SAC_{BC,cold}$ (for the cold period) was estimated to be 15.9 m² g⁻¹ while $SAC_{BC,warm}$ was higher (18.1 m² g⁻¹) (**Figure 5**). SAC values at Alert have been reported to be even higher (Sharma et al., 2004), showing that Western Arctic aerosols differ by composition and aging. Recalculations of BC mass with SAC values for cold and warm periods (eBC), separately, were performed according to Eq.1.

Timeseries of daily and monthly mean eBC concentrations from August 2019 to 31 December 2022 are shown in **Figure 2**. Annual mean and median eBC for the entire period were 28.7 ± 54.1 ngm⁻³ and 12.5 ng m⁻³, respectively (**Table 1**); they exhibit a strong year-by-year variability. We note that the eBC values are approximately half of the eBC_{AET} value. Previous studies have evaluated the optical properties of BC against direct mass measurements techniques and also obtained MAC values depended on the location, different from the recommended by aethalometer (Sharma et al., 2004; Eleftheriadis et al., 2009; Yttri et al., 2024). For example, the relationship between BC_{AET} and EC obtained by the thermal technique at Alert station (Canada) during 3 - year measurements was 0.85 (Sharma et al., 2004). Studies at Villum Research Station (Greenland) showed good agreement between measured EC and eBC_{AET} concentrations (Massling et al., 2015) similar to our study. eBC climatology and the statistics for each month and year of study are

presented in **Figure 2** and **Table S 2**, respectively. The annual mean eBC in 2019, 2021 and 2022 was 33 ± 44 , 33 ± 85 , and 32 ± 48 ng m⁻³, respectively, for the entire study period. Statistically significant difference at the 95% confidence level (p-value <0.05, t-test) was observed for the cold and warm periods with means of 44 ± 47 and 19 ± 57 ng m⁻³, respectively. The smallest mean eBC of 24 ± 29 ng m⁻³ occurred in 2020. The latter is likely attributed to the impact of COVID-19 restriction measures to the emissions of BC (Evangeliou et al., 2020).

The general trend of the maximum in winter and minimum in summer well reproduces the typical *eBC* seasonality reported in polar observatories (Stone et al., 2014; Schmale et al., 2022). **Figure 2** shows monthly mean *eBC* concentrations for half of year 2019 and whole - year periods of 2020, 2021, and 2022 as well as annual averaged monthly mean *eBC* climatology for the entire study period. The highest concentration in the cold period was observed in December 2019 (81±64 ng m⁻³), January 2022 (61 ±49 ng m⁻³), February 2022 (106±67 ng m⁻³), and March 2021 (42±33 ng m⁻³) (**Table S 2**). In warm periods we recorded the highest concentrations in September 2020 (31±48 ng m⁻³), August 2021 (83±249 ng m⁻³), April 2021 (35±26 ng m⁻³), and August 2022 (28±54 ng m⁻³).

3.3 Multi-wavelength absorption Angstrom exponent

As shown by Virkkula (2021), pure BC particles surrounded by non-absorbing coatings can have AAE in the range from <1 to 1.7. Compendium of values from different emissions show AAE variation from 0.2 to 3.0 for transport, power plants, and domestic wood burning (Helin et al., 2021). Primary emissions from residential heating (Cuesta-Mosquera et al., 2024) and BB (Popovicheva et al., 2017a, 2019b) have been associated with high AAE of around 3-4. Due to the mixing with background aerosol, coating and aging processes, a large change in the light absorption has been reported at receptors of long-range transported pollution (Cappa et al., 2016). For highly aged aerosols, AAE has been found lower than 1.0 due to large and internally mixed particles (Popovicheva et al., 2022). Spectral absorption was obtained at IBS in the UV to IR spectral region emphasized by the value of $AAE_{350/950}$ equal to 0.96 for the entire study period (**Figure 6**a). Power law fittings of spectral dependence for both and cold periods show similar values, indicating highly mixed and aged BC.

Multiple studies have addressed the sensitivity of the AAE to the range of wavelengths selected for its calculation (Cuesta-Mosquera et al., 2024); the extent of this sensitivity is higher for aerosols containing a substantial contribution of organic species such as BrC. Events affected by regional fire emissions were evident by the light absorption coefficient $AAE_{370/520}$ in the short wavelength range (Ulevicius et al., 2010). In remote Arctic environments, cases with exceeded $AAE_{467/660}$ have been identified to be influenced by BB (Pulimeno et al., 2024). Impact of

intensive wildfires in North America on aerosol optical properties measured at the European Arctic has been associated with increased daily $AAE_{467/660}$ of up to 1.4 (Markowicz et al., 2016). Strong UV absorption has led to increase of up to 1.8, clearly indicating the importance of non-BC light-absorbing component (Ran et al., 2016).

To apportion the wavelength-dependent light absorption, we used a pair of wavelengths (350 and 950 nm) in the whole spectrum, and in shorter wavelengths (370 and 660 nm, 370 and 520 nm). Timeseries of weekly average $AAE_{370/520}$ showed a similar seasonality but wider variation (0.2-3.1) than (0.5-1.7) for $AAE_{370/950}$ (**Figure 6**b). The mean values increased from 0.97 \pm 0.23 for $AAE_{370/950}$ to 1.17 \pm 0.5 for $AAE_{370/520}$ for the entire study period (**Table 1**). Box-whisker plots and annual averaged means of $AAE_{370/950}$ showed no prominent monthly dependence (**Figure 4**c). However, increased $AAE_{370/950}$ above 1.1 was observed in summer months for several years, in July 2020, June 2021 and from May to September 2022 (**Table S 2**). The shorter the wavelength pair, the higher the annual average AAE above 1.0. The largest values of monthly mean (median) $AAE_{370/520}$ were found for April to September with a maximum in June. Such considerable deviation during warm months implies the importance of BrC light-absorbing components within highly mixed Arctic aerosols at IBS.

Light absorption at 370 nm, $b_{abs}(370)$, was used to estimate the BrC mass concentrations. The mean (median) value of $b_{abs}(370)$ was 2.4 times higher than $b_{abs}(880)$ for the entire study period as well as for cold and warm ones (**Table 1**). Monthly means and box-whisker plot of $b_{abs}(370)$ showed trends similar to $b_{abs}(880)$ (**Figure S 4**). Assuming that the wavelength pair λ and λ_0 in Eq. 3 being 370 and 950 nm, respectively, the absorption coefficient for BrC at 370 nm, $b_{abs/BrC}(370)$, is determined by subtracting BC absorption from the total absorption at the same wavelength using the $AAE_{370/950}$ value for entire period (**Table 1**). Monthly $b_{abs/Bc}(370)$ and $b_{abs/BrC}(370)$ as well as the $b_{abs/BrC}(370)$ percentage contribution to total $b_{abs}(370)$ are shown in **Table S 3** for those years when the contribution of BrC absorption was higher than 1%. We note 13% for August 2021 for the warm period and 5 % for February 2022 and December 2021 for the cold period.

3.4 Modelled concentrations of BC

Figure 7a shows the monthly mean eBC and surface BC ($BC_{FLEXPART}$) concentrations simulated with FLEXPART coupled to ECLIPSEv6-GFAS emissions for the entire study period. FLEXPART model performs well in capturing the seasonality of observed features with both high and low concentrations. Annual mean modelled $BC_{FLEXPART}$) (88.4 ng m⁻³) is 37% higher than eBC_{AET} (64.3 ng m⁻³) and 3 times higher than eBC (29.5 ng m⁻³). Annual and monthly means of

 eBC_{AET} show values closer to $BC_{FLEXPART}$ than eBC. This is a reasonable finding because the global emission datasets could not consider local pollution. Almost all simulated BC concentrations, except in February 2020 and 2021, were found within the standard deviation range of measured eBC_{AET} . A good correlation between measurements and simulations, with a Pearson coefficient of 0.72 and 0.82, a root mean squared error (RMSE) of 15 ng m⁻³ and 0.14 ng m⁻³ and a normalised bias of 0.39 and 0.27 was obtained for the cold and warm period, respectively (**Figure** 7b,c).

FLEXPART does not reproduce seasonal variations of BC everywhere over the Arctic. R² and RMSE varied between 0.53-0.80 and 15.1-56.8 ng m⁻³, respectively, depending on the location (Zhu et al., 2020). At Zeppelin, modelled BC (annual mean of 39.1 ngm⁻³) was reported to be 85% higher than the measured value (21.1 ng m⁻³ for annual mean). At Tiksi, modelled BC was underestimated (74.4 ng m⁻³ for annual mean) by 40% compared with observations (104.2 ngm⁻³ for annual mean) (Zhu et al., 2020). Such good result for IBS is due to its closer location to the biggest emission sources.

Figure 8 shows the vertical distribution of simulated BC as a function of time for 2019-2020 years (vertical cross-section). Consistently high vertical BC profiles up to 2 km were observed in the cold period, except in April 2022. In February 2020, a smoke layer of BC concentrations of up to 100 ng m⁻³ was prominent at up to 4 km. On the contrary, in the warm period the smoke resides near the surface, despite a few events of extremely high vertical BC at altitudes up to 8 km and 10 km, which occurred in July 2020 and August 2021, respectively. Nevertheless, the evidence of atmospheric transport from high altitudes during summer months is evident by the elevated modelled BC (>100 ng m⁻³) at high model layers (e.g., July 2019, June-August 2020, June-July 2021 and May-June 2022). In all these periods, $BC_{FLEXPART}$ (violet line in **Figure 8**) was under 40 ng m⁻³ showing that the emission sources are probably far away, and that long-range transport occurred. The low injection altitude of anthropogenic emissions in winter months cause emitted substances to remain close to the emission sources. BC climatology at IBS indicates that the longrange transported anthropogenic emissions in the cold period reside at altitudes up to 2 km and compose a persistent layer (Figure 8). This is further explained by the rapid (about 4 days, or less) low-level transport of air masses to the Arctic troposphere as described in Stohl (2006). However, this cannot be confirmed without targeted high altitude observations.

4 Discussion

4.1 Long-range transport, age and region contributions

Transport mechanisms from the source regions affect the Arctic BC variability and burden (Chen et al., 2023; Zhou et al., 2012). Transport of aerosols to the Arctic leads to high concentrations of BC in winter and spring (Arctic haze) and low values in summer (Law and Stohl, 2007) when the removal processes in the dry and stable Arctic atmosphere are very slow. Synoptic-scale circulation effects promote the effective transport from lower latitudes, namely diabatic cooling of air masses moving over snow-covered ground, high continental pressure in winter, and the intrusion of warm air from lower latitudes (Gilardoni et al., 2023). Seasonal trends of footprint emission sensitivity demonstrate the transport mechanisms from the source regions to the European Arctic (Platt et al., 2021). BC at Zeppelin is affected by significantly different source regions during the warm and cold seasons, while large-scale circulation patterns that affect the pollutant transport from lower latitudes show the opposite behaviour during these two periods (Stathopoulos et al., 2021).

Figure 9 shows a 3.5-year climatology of the surface footprint emission sensitivities at IBS. From December to February, anthropogenic polluted air mass transportation takes place from Eurasia (territories above 40°N), as illustrated by the elevated footprints there. The extension of the Arctic front towards lower latitudes during the cold period facilitates such transport (Stohl, 2006). The warmer it gets in spring, the narrower the area of emission transport. In the transition from spring to summer, transport patterns and meteorological conditions change, such as that the advection of the particulate pollution to the Arctic boundary layer from lower latitudes becomes limited (Bozem et al., 2019). In JJA (June, July, August) footprint is mostly restricted to coastal regions of Eurasia, Greenland, and North America and does not extend deeply into the continents. This is a consequence of the so-called 'polar dome' that prevents warm continental air masses from entering the Arctic lower troposphere (Stohl, 2006). As a result, anthropogenic pollution becomes less significant, and natural aerosol sources prevail (Moschos et al., 2022b, a). In autumn (September, October, November), footprint is similar to the MAM (March, April, May) one completing the annual cycle.

For the entire study period, the monthly mean contribution to surface BC for all years was from air masses with 1-3 (31%) and 3-6 days (22%) aging (**Table S 4**). The highest BC contribution (34%) and (39%) was observed for the shortest age of 1-3 days in DJF (December, January, February) and MAM, respectively (**Figure 8**). In summer, the highest BC contribution (35%) was replaced by a longer age of 6-9 days.

Footprint emission sensitivities of Arctic air masses also constrain the region contributions. The major source regions contributing BC to IBS are the territory of the Russian Federation (including European part of Russia (EURus), Siberia, Far East), Asia, Europe, Northern America, and Ocean. Due to the geographical proximity, EURus/Siberia/Far East contribution (77%) dominated during the entire study period on a basis of the annual average monthly means (**Table S** 4), with a maximum of 83% in SON (**Figure 9**). Its monthly maximum (88%) was recorded in September 2021, and the minimum (60%) in June 2022. Europe was the second region contributor (11%) followed by Asia. The monthly mean contribution of Northern America was up to 12% in JJA, the largest was observed in July 2022 (62%).

4.2 Anthropogenic and biomass burning sources

The time series of monthly mean and annual average monthly mean source contributions to surface BC at IBS are shown in **Figure 10**a. Anthropogenic sources (DOM, TRA, IND, FLR, All others) contribute 97% of the total for the entire study period (Table S 4). A decrease from winter to July and an increase from August to winter were seen. In the cold period, air masses arrived at IBS through the populated regions of Western Europe, EURus, Siberia, and Asia, crossing the biggest oil and gas extraction regions of Kazakhstan, Volga-Ural, Komi, Nenets, and Western Siberia (Figure 1). Because IBS is located north of the largest oil and gas producing regions of Western Siberia, high FLR contribution of 59% and 32% was observed both in the cold and warm period (Table S 4). Annual mean contributions to modelled surface BC from FLR, DOM, TRA, and IND sectors dominated in January and December (60%, 22%, 12%, and 9%, respectively). All other sources were around 2% at that time. BB played the biggest role between April (8%) and October (17%), with maximum in August (80%).

Figure S 5 shows the percentage sectoral contributions on monthly mean BC concentrations for 2021 and 2022, data for 2019 and 2020 was shown in Figure 10a. February 2021 and December 2021 were the leaders of FLR impact with 67.2% and 67.4%, respectively. During February 2022 of the record high BC pollution level observed at IBS, air masses arrived at IBS through the Western Europe, EURus, and Siberia, passing through the flaring facilities of Kazakhstan, Volga-Ural, Komi, Nenets, and Western Siberia. They caused of 50%, 26%, 15%, 8%, 0.2%, and 3.3% monthly average contribution to surface BC from FLR, DOM, TRA, IND, BB and All other sources, respectively. Footprint emission sensitivities on 3rd February 2022 at 12:00-15:00 when eBC reached 310 ng m-3 (Figure S 5) showed air mass transport to IBS straight through the Western Siberian gas flaring region (Figure 10b).

The contribution of FLR dropped significantly from April to a minimum of 18% in June and rose in September. In the winter months when the overestimation of modelled BC concentrations

was recorded (see section 3.4), the highest FLR impact was seen. DOM showed the biggest contribution (18%) from November to February, exactly during the heating season. The light absorption of BrC was significant mostly in wintertime (Table S 3). The latter indicates significant impact of biomass used for domestic heating, in accordance to wood burning contribution of 61% of the total residential emissions in forest regions (Huang et al., 2015).

According to CAMS GFAS (Figure S 3), significant global fire emissions started from June and lasted until the mid of November in 2020 and 2022; the period of fire emissions was shorter but more intensive from July until September 2021. At IBS, the annual mean BB contribution approached 48% of the total in the warm season (Table S 4). It started increasing from April and approached a maximum of 80% in August, whereas TRA, DOM, IND, and All other sources were minimum. From middle June to September, the average monthly BB contribution was larger than all anthropogenic sources. Notably, from April to September, the high mean BB contribution was related to the excess of $AAE_{370/520}$ over 1.0 (maximum: 1.7 in July) (Figure 6). At that time, the air masses transported to IBS were aged (> 6 days) dominating the age spectrum (57%) (Table S 4).

In 2019, 72,400 km2 were burned in Siberia or 42% of the total burned area that occurred in Russia (Voronova et al., 2020). A significant relationship between the burned areas and associated pyrogenic emissions with atmospheric blocking events was reported (Mokhov et al., 2020). August and September showed 50% and 35%, respectively, monthly mean BB contributions, while October and November lower, 30% and 20%, respectively (Figure S4).

In spring 2020, BB BC concentrations simulated with WRF-Chem model were distributed in areas between 40°N and 60°N in Europe, central Siberia, and East Asia, and indicated intensive seasonal agriculture fires in Europe and Siberia (Chen et al., 2023). Spring fires contributed about 12% BB BC to IBS (April and May). The end of June and beginning of July of 2020 was characterized by high altitude BC (Figure 8) indicating high altitude long-range transport. A high BrC content was also observed in July and September 2020 (Table S 3).

In 2021, the monthly mean spring BB contribution approached a maximum of 36% in May. Yakutia (Eastern Siberia) experienced the worst fire season over the last four decades (Tomshin and Solovyev, 2022). Around 150,000 fires occurred, almost twice as much as the previous year (Voronova et al., 2022). August 2021 received 90% contribution from BB as compared to all the other sources. At that time unprecedented high smoke levels were recorded over Western Siberia (Schneider et al., 2024). Satellite image reveals the strong plume from the area of Yakutian wildfires which brought deep smoke to IBS located around 2000 km far away (Figure 1c). The highest *eBC* level of 1800 ng m-3 on 5th August, exceeded the 75th percentile of the entire period

53 times (Table 1). The measured concentrations were 180 times higher than the Arctic background (Figure S 6). Severe smoke affected the visibility near IBS (Figure 1d). Footprint emission sensitivity on 5th August (from 18:00 to 21:00) at the time when *eBC* peaked (1540 ng m-3) confirms that air masses originated from Yakutia and arrived to IBS from the northeast direction (**Figure 10**b). BC for these wildfires was transported at altitude as high as 10 km (Figure 8). Finally, in summer 2022, wildfires took place in Western Siberia and the EURus (Popovicheva et al., 2023); BB contributions in June, July, August 2022 were around 65%, whereas light absorption of BrC was important in May and August 2022 (Table S 3).

5 Summary and conclusions

We presented four years (2019-2022) of observations at the aerosol station IBS with respect to light-absorption characteristics of Western Siberian polar aerosols and its basic cycles, such as seasonality, annual means, and interannual variability. The annual cycle of multi-wavelength light absorption demonstrates higher levels during the Arctic haze season and lower in summer, similar to other Arctic observations. The light absorption coefficient revealed several unique features:

Higher magnitude (around 4-5 times) in comparison with multi-year observations at high-latitude polar stations in European Arctic (annual mean of $0.7\pm0.7~\mathrm{M~m^{-1}}$ in the cold season and 2 times lower in warm). Wintertime maximum was observed in February ($0.9\pm0.8~\mathrm{M~m^{-1}}$) that coincides with the Arctic haze peak; this is different from the European and Canadian Arctic that is usually observed in early spring. The interannual minimum was observed in June whereas August was highly variable with respect to light-absorption due to the Siberian wildfires. Multi-annual monthly means for $b_{abs}(880)$ in the visible spectrum at IBS were found higher than at European, Canadian and Western high-latitude polar locations, due to that IBS is closer to the main Northern Eurasian source regions.

Wildfires caused increased concentrations, usually in August. Increase of the AAE in the UV spectrum between April and September implies coexistence of highly mixed/aged BC and light-absorbing BrC components. Specifically, monthly BrC contribution to total light absorption was 5% in February 2022 and 13% August 2021 likely due to wildfire impact. BrC light absorption coefficient in the UV spectrum showed similar trends as BC, although it exceeded BC by 2.4 times during both cold and warm periods. AAE was equal to 0.96, indicating highly mixed and aged aerosols. AAE in UV spectrum increase up to 1.17±0.5 implies coexistence of light-absorbing BrC components in BB aerosols, with the biggest impact between April and September.

We calculated SAC for the first time at IBS by combining multi-year optical absorption and EC data. Higher SAC of 18.1 m² g⁻¹ in the warm period than in the cold one (15.9 m² g⁻¹) revealed

influence from non-BC light-absorbing species, such as organic matter and mineral dust; SAC values were lower than those observed in the Canadian Arctic indicating different aerosol composition and aging. Mean eBC in the cold and warm periods were equal to 44±47 and 19±57 ng m⁻³, respectively. Record high eBC was found in February 2022 (106±67 ng m⁻³) and August 2021 (83±249 ng m⁻³) during the years of study.

Observations at the IBS station evaluated the relationship between eBC_{AET} and EC under specific atmospheric conditions. eBC, recalculated using site-specific absorption coefficients, reflects seasonal variations and provides insights into aerosol composition. Annual cycles follow typical Arctic trends, with higher eBC concentrations in winter when air masses primarily originate from Russia, Siberia, the Far East, Europe, and Asia. During this period, black carbon from gas flaring dominates, particularly in January, when air masses pass over oil and gas facilities in Kazakhstan, Volga-Ural, Komi, Nenets, and Western Siberia. In summer, biomass burning (BB) from Siberian wildfires surpasses anthropogenic sources, peaking in August 2021, which saw the worst fire season in four decades, bringing heavy smoke to IBS. February 2022 also recorded extreme BC pollution levels.

Modeling analyses indicate that 77% of BC transport originated from Russia, Siberia, and the Far East, followed by Europe (11%), Asia (7%), and North America (4%). In winter, air masses traveled 1-3 days from Eurasia (north of 40°N) to IBS, whereas in summer, transport took 6-9 days. Low-injection altitude anthropogenic emissions created a persistent BC layer up to 2 km in the cold season, reaching 4 km in February 2020, with record concentrations of 100 ng m⁻³. In contrast, wildfire smoke in summer elevated BC layers to higher altitudes.

Anthropogenic emissions accounted for 83% of BC during the study period, dominated by gas flaring (FLR, 59%), domestic heating (DOM, 18%), traffic (TRA, 10%), and industry (IND, 7%) during Arctic haze period. Gas flaring remained the primary contributor year-round (59% in winter, 32% in summer), given IBS's proximity to major oil and gas regions. Residential heating peaked in winter (18%), aligning with enhanced brown carbon (BrC) absorption from wood burning. In February 2022, modeled BC concentrations reached 310 ng m⁻³ as air masses passed through major flaring regions, though overestimation suggests miscalculated source intensities.

BB contributions peaked at 48% in the warm season, surpassing anthropogenic sources from mid-June to September, with a maximum of 80% in August. Extreme vertical BC events reached 8 km in July 2020 and 10 km in August 2021 due to wildfires. In May 2021, BB contributions reached 36% due to strong agricultural fires in Siberia, while in August 2021, 90% of BC at IBS originated from Yakutia's wildfires, 2000 km away.

The increasing intensity and frequency of wildfires at high latitudes highlight the importance of carbonaceous aerosol measurements. These observations provide critical insights into Arctic

aerosol radiative properties, particularly in the UV-VIS spectrum, where enhanced light absorption 663 contributes to amplified Arctic warming, especially in summer. 664 665 666 667 Data availability. All modelling data from this study are available for download from <a href="https://atmohttps access.nilu.no/BELY2 MSU.py. FLEXPART version 10.4 model can be downloaded from 668 https://www.flexpart.eu/downloads. Black Carbon observations are available upon request from O. 669 B. Popovicheva. 670 671 **Supplement.** The supplement related to this article in available online at. 672 673 Author contributions. OBP supervised the station operation, interpreted data and wrote the 674 manuscript. NE performed all the FLEXPART simulations and analyses, wrote and coordinated the 675 paper. MAC analysed the data, prepared the figures and assisted in the interpretation of the results. 676 ED provided supported AAE calculations and evaluation of data quality. NSK supported the 677 research. All authors contributed to the final version of the manuscript. 678 679 **Competing interests.** The authors declare no competing interests. 680 681 **Acknowledgements.** This research was performed in the frame of the development program of the 682 Interdisciplinary Scientific and Educational School of M. V. Lomonosov Moscow State University 683 "Future Planet and Global Environmental Change". Authors thank Magee Scientific for AE33 684 instrumentation support and Dr. Asta Gregorič for data examination. V.O. Kobelev is 685 686 acknowledged data analyses over all study years. 687 Financial support. The article processing charges for this publication were paid by NILU. 688 Developed methodology of aethalometric measurements was implemented in the frame of the RSF 689 project #19-77-3004Π. Authors thanks to Russian Geographical Society for the data treatment 690 support Institute of Environmental Survely, Planing and Assessment (IESPA) partly supported the 691 692 instrumentation and power supply of IBS. 693 694 References

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Figure 1. (a) The IBS between other polar aerosol stations. (b) A map showing IBS in Western Siberia along with oil and gas fields (adopted from https://skytruth.org/, last access: 7 November 2024). Flares of oil and gas fields are indicated for 2020 as black triangles (https://skytruth.org/, last access: 7 November 2024). (c) Satellite image of strong plume from the area of Yakutian wildfires which brought deep smoke to the Bely Island. (d) View to the pavilion of IBS under clear conditions on 25 July 2021, and during the unprecedented smoke event on 5 August 2021. Maps were created using Open-Source Geographic Information System QGIS (https://qgis.org/en/site, last access: 7 November 2024) with ESRI physical imagery (https://server.arcgisonline.com/ArcGIS/rest/services/World_Physical_Map/MapServer/tile/%7Bz%7D/%7By%7D/%7Bx%7D&zmax=20&zmin=0, last access: 7 November 2024) as the base layer, and for MODIS Reflectance true color imagery (MODIS Science Team) and Satellite imagery from 05 of August 2021 (https://worldview.earthdata.nasa.gov, last access: 7 November 2024) with TERRA MODIS fire anomaly layer. Open-source Natural Earth quick start (NEQS) package was used to add layers of natural and cultural boundaries and polygons from ESRI Shapefile storage.

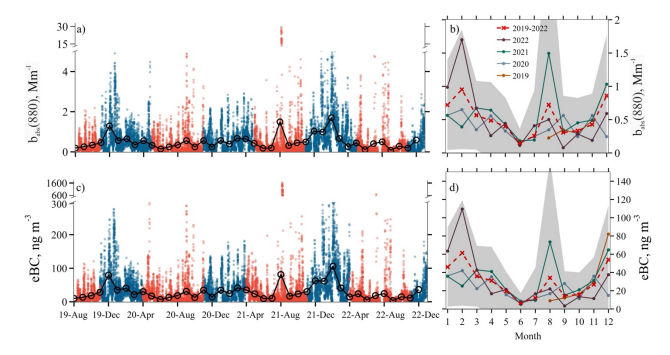


Figure 2. Hourly timeseries and monthly means of (a) $b_{abs}(880)$ and (c) eBC for cold (blue) and warm (red) periods; monthly climatology of (b) $b_{abs}(880)$ and (d) eBC for half year 2019 and 2020, 2021, and 2022. Cross-marks (x) joined by lines show the inter-annual mean; the standard deviation is plotted by shadow area.

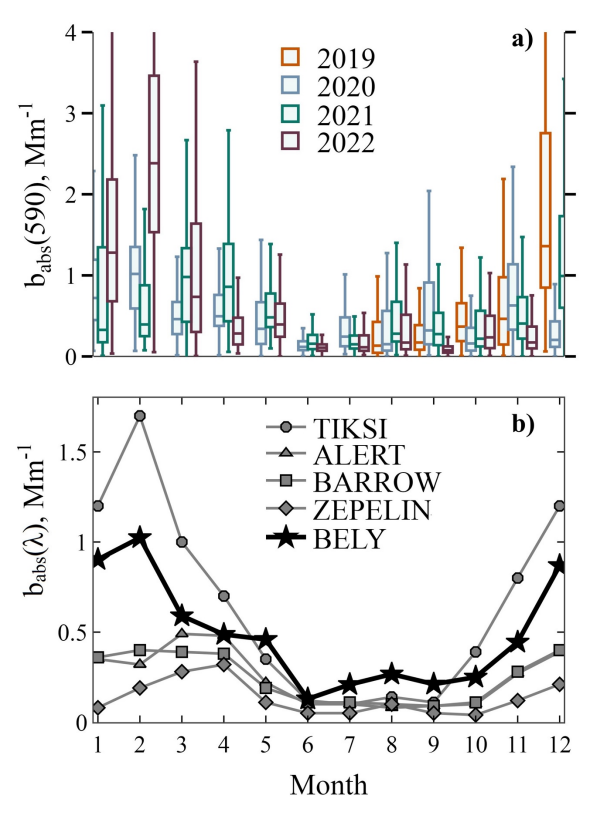


Figure 3. (a) Monthly box-whisker plot for $b_{abs}(590)$ at IBS for half year 2019 and full 2020, 2021, and 2022. The 25th, 50th, and 75th percentiles are shown with boxes, while whiskers extend ± 1.5 times the interquartile range. (b) Seasonality of monthly median of b_{abs} at 550 nm at Tiksi, Alert, Barrow/Utqiagvik, Zepelin for 2012-2014 (Schmeisser et al., 2018), and b_{abs} at 590 nm at IBS for 2019-2022 (this work).

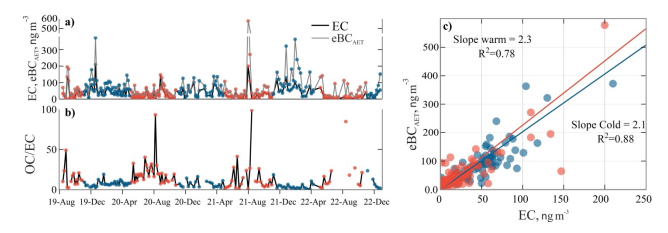


Figure 4. Temporal variation of (a) weekly EC and eBC_{AET} averaged over the whole sampling period and (b) the OC/EC ratio. (c) Scatter plots and orthogonal regressions (solid lines) for measured eBC_{AET} and EC concentrations in cold (blue) and warm (red) period. The figure includes the regression slope, the coefficient of determination (R2).

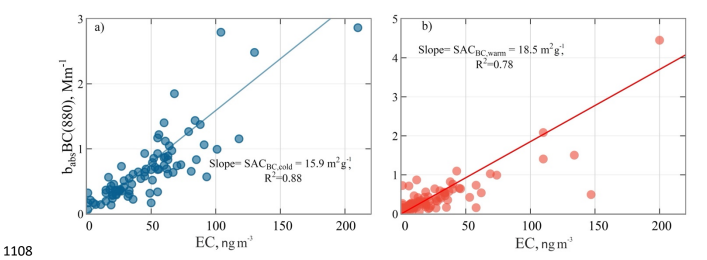


Figure 5. Scatter plots and orthogonal regressions (solid line) for $b_{abs/BC}$ (880) and EC concentrations for the (a) cold (blue) and (b) warm (red) periods. Regression slope defines $SAC_{BC,cold}$ and $SAC_{BC,warm}$.

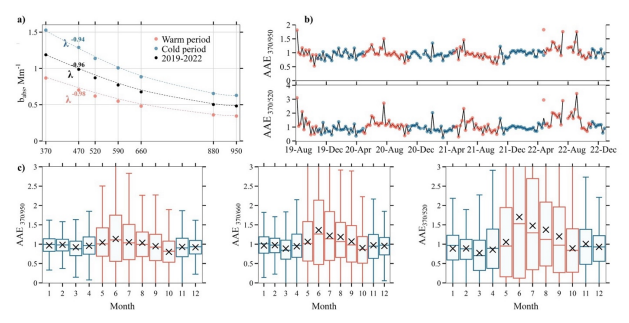


Figure 6. (a) Spectral dependence of light absorption coefficient for 2019-2022, during warm (red) and cold (blue) periods. $AAE_{350/950}$ is the slope of the linear regression in logarithmic scale of a power law regression as described in Eq. 2. (b) Timeseries of $AAE_{370/950}$ and $AAE_{370/520}$. (c) Boxwhisker plots and monthly means of AAE at 370 and 950 nm, 370 and 660 nm, and 370 and 520 nm for the entire period.

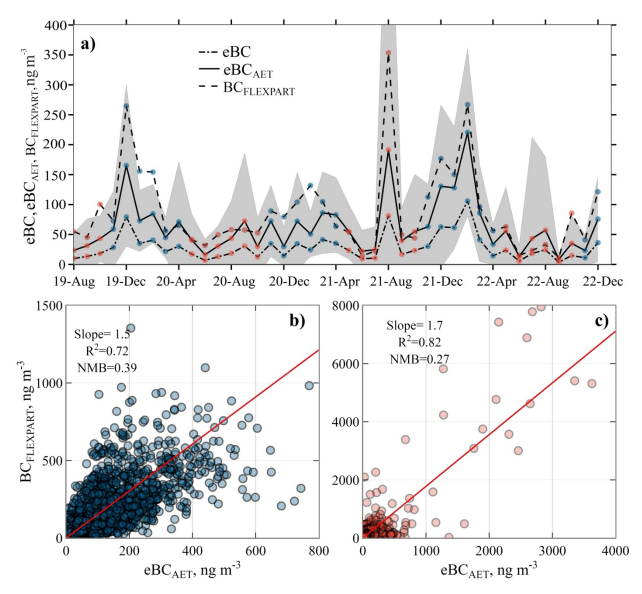


Figure 7. (a) Monthly mean eBC and modelled surface BC concentrations from 10 August 2019 to 31 December 2022. Monthly mean eBC_{AET} (line with crosses) shown with the standard deviation range by shadowed area. Scatter plots and orthogonal regressions (solid lines) for BC_{FLEXPART} calculated over measured eBC_{AET} concentrations for (b) cold and (c) warm period. The figure includes the regression slope, the coefficient of determination (R2).

VERTICAL CROSS-SECTIONS OF MODELLED BC IN IBS

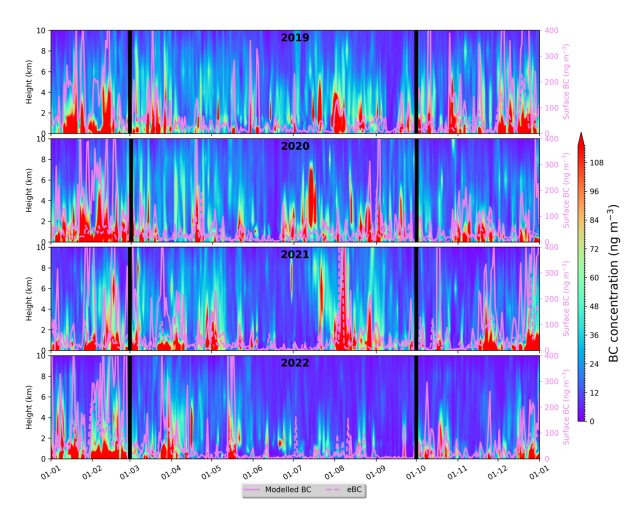


Figure 8. Vertical cross-sections of modelled BC for 2019-2022. Solid and dotted violet lines represent modelled daily surface BC and *eBC*, respectively. Their levels correspond to the right (secondary) axis (also in violet). Boundaries between the cold (November- April) and warm (May-October) are indicated by thick vertical black lines.

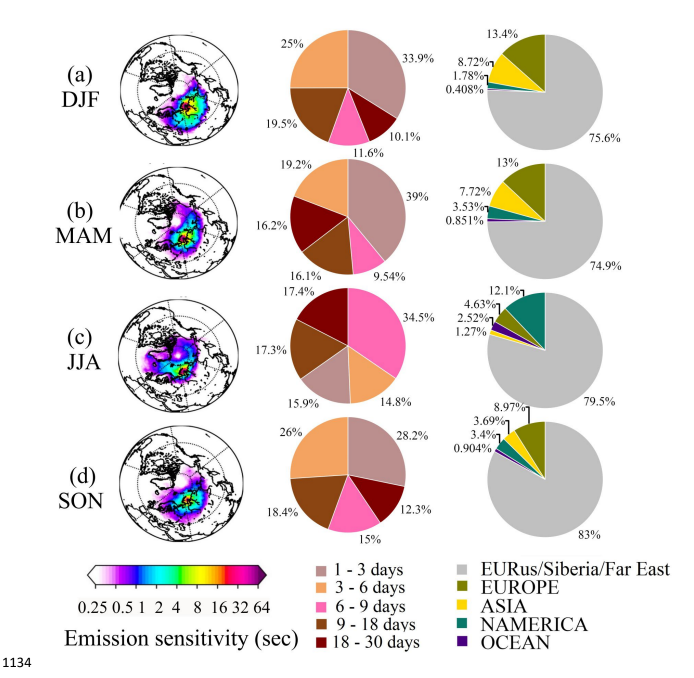
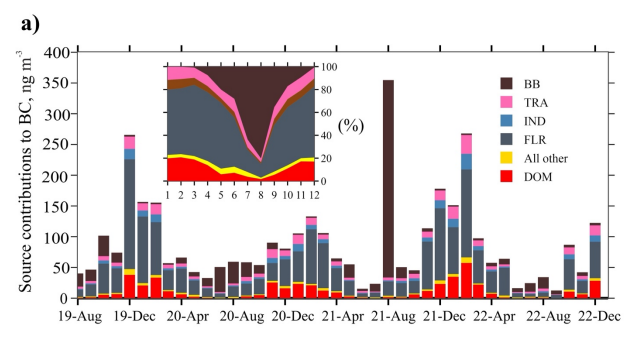


Figure 9. (a-d) Season average footprint emission sensitivity, mean age contribution of emissions from different day-periods back in time and each region contribution to surface concentration of BC.





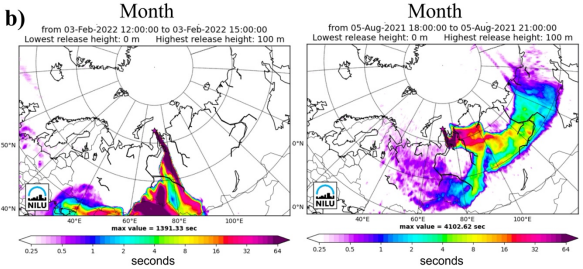


Figure 10. (a) Timeseries of monthly mean contribution from different emission source types to surface BC concentrations for the study period. DOM, BB, TRA, IND, FLR sectors, and All others sources were adopted from ECLIPSEv6 and CAMS GFAS. (b) FES for 3 February 2022 and 5 August 2021 showing the largest probability of emission origin.

TABLES & LEGENDS

Table 1. Statistics of light - absorption coefficients; EC, OC, eBC_{AET} , and eBC mass concentration; absorption Angstrom exponents (AAE) for the study period, cold and warm periods. Mean± standard deviation (1 σ), 1st and 3rd Q quartile (25th and 75th percentiles).

Variable	August 2019 - December 2022				cold (November-April)				warm (May-October)			
	mean±sd	median	1 st Q	$3^{rd} Q$	mean±sd	median	1 st Q	3 rd Q	mean±sd	median	1 st Q	3 rd Q
b _{abs} (880) (Mm ⁻¹)	0.5±0.9	0.3	0.1	0.6	0.7±0.7	0.5	0.22	0.9	0.4±0.9	0.2	0.09	0,4
$b_{abs}(520)$ (Mm ⁻¹)	0.9±1.6	0.4	0.2	1	1.2±1.2	0.8	0.38	1.5	0.6±1.8	0.3	0.1	0,6
$b_{abs}(370)$ (Mm ⁻¹)	1.2±2.4	0.6	0.3	1.4	1.6±1.6	1.1	0.52	2.1	0.9±2.8	0.4	0.2	0,9
EC (ng m ⁻³)	30±30	20	10	50	50±30	40	20	60	20±30	20	10	30
OC (ng C m ⁻³)	459±300	400	300	500	400±200	400	300	500	500±400	400	300	600
* eBC _{AET} (ng m ⁻³)	65±83	36	16	80	84±90	57	25	115	53±158	23	10	45
* <i>eBC</i> (ng m ⁻³)	29±54	13	5	34	44±47	29	12	59	19±57	8.0	4	17
$AAE_{370/950}$	0.96 ± 0.6	0.95	0.7	1.19	0.94 ± 0.4	0.95	0.74	1.1	0.98±0.8	0.95	0.6	1,3
$AAE_{370/520}$	1.0±1.5	0.93	0.4	1.52	0.88±1	0.89	0.49	1.2	1.16±1.9	1.0	0.3	2,0

^{*} *eBC*_{AET} is defined in section 2.2. ** *eBC* is defined in section 3.2.