Origin of elemental carbon in snow from western Siberia and northwestern European Russia during winter–spring 2014, 2015 and 2016

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Abstract. Short-lived climate forcers have been proven important both for the climate and human health. In particular, black carbon (BC) is an important climate forcer both as an aerosol and when deposited on snow and ice surface because of its strong light absorption. This paper presents measurements of elemental carbon (EC; a measurement-based definition of BC) in snow collected from western Siberia and northwestern European Russia during 2014, 2015 and 2016. The Russian Arctic is of great interest to the scientific community due to the large uncertainty of emission sources there. We have determined the major contributing sources of BC in snow in western Siberia and northwestern European Russia using a Lagrangian atmospheric transport model. For the first time, we use a recently developed feature that calculates deposition in backward (so-called retroplume) simulations allowing estimation of the specific locations of sources that contribute to the deposited mass.

EC concentrations in snow from western Siberia and northwestern European Russia were highly variable depending on the sampling location. Modelled BC and measured EC were moderately correlated ($R = 0.53–0.83$) and a systematic region-specific model underestimation was found. The model underestimated observations by 42% (RMSE = 49 ng g$^{-1}$) in 2014, 48% (RMSE = 37 ng g$^{-1}$) in 2015 and 27% (RMSE = 43 ng g$^{-1}$) in 2016. For EC sampled in northwestern European Russia the underestimation by the model was smaller (fractional bias, FB > −100%). In this region, the major sources were transportation activities and domestic combustion in Finland. When sampling shifted to western Siberia, the model underestimation was more significant (FB < −100%). There, the sources included emissions from gas flaring as a major contributor to snow BC. The accuracy of the model calculations was also evaluated using two independent datasets of BC measurements in snow covering the entire Arctic. The model underestimated BC concentrations in snow especially for samples collected in springtime.
1 Introduction

Black carbon (BC) is the strongest light-absorbing component of atmospheric aerosol and is formed by the incomplete combustion of fossil fuels, biofuels and biomass (Bond et al., 2013). It is emitted directly into the atmosphere in the form of fine particles. BC is a major component of “soot”, a complex light-absorbing mixture that also contains organic carbon (OC) (Bond et al., 2004). Combustion sources emitting BC include open biomass burning (BB; forest, savanna, agricultural burning), residential biofuel combustion, diesel engines for transportation or industrial use, industrial processes and power generation, or residential coal combustion (Liu et al., 2011; Wang et al., 2011).

BC is important on a global perspective because of its impacts on human health and on climate. As a component of fine particulate matter (PM$_{2.5}$), it is associated with negative health impacts, including premature mortality (Lelieveld et al., 2015; Turner et al., 2005). It absorbs solar radiation, has a significant impact on cloud formation and, when deposited on ice and snow, it accelerates ice melting (Hansen and Nazarenko, 2004). BC has a lifetime that can be as long as 9–16 days (Bond et al., 2013). After its emission, BC can travel over long distances (Forster et al., 2001; Stohl et al., 2006) and reach remote areas such as the Arctic. Arctic land areas are covered by snow in winter and spring, while the Arctic Ocean is partly covered by ice. Sea ice has a much higher albedo ($\approx 0.5–0.7$) compared to the surrounding ocean ($\approx 0.06$); thus, presence of sea ice reduces the heat uptake of the ocean. Snow has an even higher albedo than sea ice and can reflect as much as 90% of the incoming solar radiation (Brandt et al., 2005; Singh and Haritashya, 2011). BC deposited on ice lowers its albedo, increases heat uptake by sea ice, accelerates its melting, and therefore decreases surface albedo both directly and indirectly.

Hegg et al. (2009) reported that snow in the Arctic often contains BC at concentrations between 1 and 30 ng g$^{-1}$, which can cause a snow albedo reduction of 1–3% in fresh snow and another 3–9% as snow ages and BC becomes more concentrated near the surface (Clarke and Noone, 1985). This capacity of snow to reflect solar radiation insulates the sea ice, maintains cold temperatures and delays ice melt in summer. After the snow begins to melt and because shallow melt ponds have an albedo of approximately 0.2 to 0.4, the surface albedo drops to about 0.75 or even lower (0.15) as melt ponds grow and deepen (Singh and Haritashya, 2011). These changes have been found to be important for the global energy balance (Flanner et al., 2007; Hansen and Nazarenko, 2004) and, if enhanced by BC, contribute to climate warming (Warren and Wiscombe, 1980).

Although BC in Arctic snow and ice has been found to be important for the Earth’s climate (Flanner et al., 2007; Sand et al., 2015), its large-scale temporal and spatial distributions and exact origin are still poorly quantified (AMAP, 2015). Efforts to determine the concentrations of BC in snow across the Arctic were made by Clarke and Noone (1985), Doherty et al. (2010, 2013), Forssström et al. (2013), Ingvander et al. (2013), and McConnell et al. (2007). This paper presents measurements of elemental carbon (EC) concentrations in snow samples collected in spring 2014, 2015 and 2016 on the Kindo peninsula (White Sea, Karelia), around Arkhangelsk in northwestern European Russia, and in western Siberia. In the latter area, gas flaring emissions are very important. Flaring emissions are highly uncertain because both activity data and emission factors are largely lacking. According to the Global Gas Flaring Reduction Partnership (GGFR) (http://www.worldbank.org/en/programs/gasflaringreduction), nearly 50 billion m$^3$ of gas is flared in Russia annually. Russian flaring emissions in the Nenets and Komi regions and in Khanty-Mansiysk are the major sources in western Siberia and northwestern European Russia. It has been reported that gas flaring in Russia contributes about 42% to the annual average BC surface concentrations in the Arctic (Stohl et al., 2013).

The use of the terms EC and BC has been the topic of several scientific papers (for example, Andreae and Gelencsér, 2006; Bond et al., 2013; Petzold et al., 2013). Petzold et al. (2013) defined BC as a substance with five properties (see Table 1 in Petzold et al., 2013), for which no single measurement instrument exists that is sensitive to all of them at the same time. Consequently, BC cannot uniquely be measured, although some of its properties, such as the absorption coefficient $\sigma_{ap}$ and the EC concentration, both commonly measured in atmospheric monitoring networks across the world, can be measured. Hence, the term BC should be used qualitatively.

In the present study, EC concentrations on ice from three campaigns measured with thermal–optical analysis (TOA) (see Sect. 2.2) are compared to simulation results from the Lagrangian particle dispersion model (LPDM) FLEXPART. The model is used here for the first time to quantify the sources contributing to BC in snow in Russia, adopting a special feature that was developed recently.

2 Methodology

2.1 Collection and storage of snow samples

Fresh snow samples were collected along a north–south transect between Tomsk and the Yamal coast in February–March 2014 (23 samples, Table S1 in the Supplement), while in March 2015 sample collection took place on the Kindo peninsula and near the port of Arkhangelsk in the White Sea (11 samples, Table S1). Finally, in February–May 2016 samples were collected on the Kindo peninsula, in Arkhangelsk, and between Tomsk and Yamal (20 samples, Table S1). These areas have been reported to receive pollution from both urban and gas flaring sources (Stohl et al., 2013). For example, the gas flaring sources located in Yamal and Khanty-
Mansiysk (Russia) are in the main pathway along which sub-Arctic air masses travel to the Arctic (Stohl et al., 2006). All sampling points were located more than 500 m away from roads to minimize the direct influence from local traffic emissions. Information about sample collection such as the location of sampling, the amount of snow collected and the depth at which snow was sampled is reported in Table S1, and the sample locations are plotted in Fig. 1.

Sampling was performed using a metal-free technique using pre-cleaned plastic shovels and single-use vinyl gloves. Samples were stored in polyethylene bags that had been thoroughly washed with 1 M HCl and rinsed with abundant deionized ultrapure water in the laboratory prior to their use. After returning the samples to the laboratory, the snow was allowed to melt at ambient temperature (18–20°C) and immediately filtered through quartz 47 mm fibre filters (2500QAT-UP Pall for samples collected in 2014 and QMA Whatman for samples collected in 2015 and 2016). The filters were dried at 60–70°C, wrapped in aluminum foil and stored in a refrigerator. Quartz fibre filter collection efficiency of BC in liquid samples can be less than 100% (Hadley et al., 2010; Ogren et al., 1983). To what extent this has affected the levels reported in the present study is unknown. Thus, the results presented should be regarded as conservative estimates based on the assumption that some BC might have been lost during filtration.

2.2 Elemental carbon measurements with thermal–optical analysis (TOA)

EC content of the filters was measured at NILU’s laboratories with TOA, using the Sunset laboratory OC–EC instrument operated according to the EUSAAR-2 protocol (Cavalli et al., 2010). A 1.5 cm² punch was cut from the filtered snow samples for the analysis. Transmission was used for OC charring correction. Performance of the OC–EC instruments is regularly inter-compared as part of the joint European Monitoring and Evaluation Programme (EMEP) Aerosols, Clouds, and Trace gases Research InfraStructure Network (ACTRIS) quality assurance and quality control effort (Cavalli et al., 2016).

2.3 Measurements of carbonate (CO$_3^{2−}$)–carbon with thermal–optical analysis (TOA) following thermal-oxidative pretreatment

The content of carbonate (CO$_3^{2−}$)–carbon on the filters was measured with TOA, following thermal-oxidative pretreatment based on the approach described by Jankowski et al. (2008). A punch of 1.5 cm² from each filter was heated at 450°C for 2 h in ambient air to remove OC and EC but not CO$_3^{2−}$–carbon. The filter punch was subjected to TOA immediately (30 s) after thermal-oxidative pretreatment. The split time (between OC and EC) obtained for each filter punch used to determine the filter samples’ content of EC (Sect. 2.2) was also used to apportion CO$_3^{2−}$–carbon to OC and/or EC. The influence of CO$_3^{2−}$–carbon evolving as EC was accounted for using the following equation:

$$EC_{CO_3^{2−}} = EC_{corr} - EC_{CO_3^{2−}},$$

where EC$_{corr}^{CO_3^{2−}}$ is EC corrected for CO$_3^{2−}$–carbon that evolved as EC during TOA, EC is elemental carbon and EC$_{CO_3^{2−}}$ is CO$_3^{2−}$–carbon that evolved as EC during TOA. Applying this correction, EC values were 5–22% lower (see Supplement).

2.4 Emissions and modelling of black carbon

The concentrations of BC in snow were simulated with the LPDM FLEXPART version 10 (Stohl et al., 1998, 2005). The model was driven with operational meteorological wind fields retrieved from the European Centre for Medium-Range Weather Forecasts (ECMWF) of 3 h (for the years 2014 and
in FLEXPART represents an aerosol population with a log-normal size distribution (see Stohl et al., 2005). Assumed aerodynamic mean diameter and logarithmic standard deviation are used by FLEXPART’s dry deposition scheme, which is based on the resistance analogy (Slinn, 1982), and they are consistent with those used in other transport models (see Evangeliou et al., 2016; Shiraiwa et al., 2008). Below-cloud scavenging was determined based on the precipitation rate taken from ECMWF. The in-cloud scavenging was based on cloud liquid water and ice content, precipitation rate, and cloud depth from ECMWF (Grythe et al., 2017). The FLEXPART user manual (available from http://www.flexpart.eu) provides more information. All modelling results for this sampling campaign can be viewed interactively at the URL http://niflheim.nilu.no/NikolaosPY/SnowBC_141516.py.

3 Results

In this section the main results of EC concentrations in snow are presented, in contrast to simulated BC concentrations with FLEXPART. The statistical dependence of the datasets is assessed using the Pearson product-moment correlation coefficient. For further validation, the fractional bias (FB) of each individual sample was calculated together with the mean fractional bias (MFB) for observed and modelled concentrations as follows:

\[
FB = \frac{C_m - C_o}{(C_m + C_o)/2} \times 100\% \quad \text{and}
\]

\[
MFB = \frac{1}{N} \sum_{i=1}^{N} \frac{C_m - C_o}{(C_m + C_o)/2} \times 100\%,
\]

where \(C_m\) and \(C_o\) are the modelled BC and measured EC concentrations and \(N\) is the total number of observations for each year. FB is a useful model performance indicator because it is symmetric and gives equal weight to underestimations and overestimations (it takes values between -200 and 200 %). It is used here to show the locations where modelled BC concentrations in snow over- or underestimate observations. Finally, for the same reasons, the RMSE, which is frequently used to measure differences between values predicted by a model and the values actually observed, was also computed (see Figs. S1–S3 in the Supplement).

3.1 Elemental carbon concentrations measured in snow

The spatial distribution of EC measured in snow samples from northwestern European Russia and western Siberia is shown in Fig. 1c for each of the campaigns (2014, 2015 and 2016) and is also summarized in Table S2. There was large spatial variability in the distribution of EC in snow in 2014 ranging from 3 to 219 ng g\(^{-1}\), with a median (± interquartile range) of 23 ± 49 ng g\(^{-1}\). The highest EC concentrations in 2014 were observed in western Siberia near Tomsk (147 to 219 ng g\(^{-1}\)). FLEXPART emission sensitivities for
these samples showed that the air was coming from the north and the east (see in http://nifheim.nilu.no/NikolaosPY/SnowBC_141516.py). This explains the high concentrations of EC, as most of the anthropogenic BC sources are located in these regions. In the rest of the snow samples for 2014, EC concentrations between 4 and 170 ng g$^{-1}$ were observed. High concentrations were observed near the Ob River coinciding with air masses arriving mainly from Europe. During the 2015 field campaign, EC concentrations were the highest near Arkhangelsk (175 ng g$^{-1}$), for which FLEXPART showed that the air was coming from nearby areas (http://nifheim.nilu.no/NikolaosPY/SnowBC_141516.py). Therefore, it is likely that the samples were affected by direct emissions from the city or the port of Arkhangelsk. During the same campaign, snow samples collected on the Kindo peninsula (on the White Sea coast) showed high variability in EC concentrations (range: 46–152 ng g$^{-1}$, median = 70 ± 34 ng g$^{-1}$). According to FLEXPART emission sensitivities, air masses were transported to Kindo peninsula from central and southern Europe driven by an anticyclone over Scandinavia (http://nifheim.nilu.no/NikolaosPY/SnowBC_141516.py). Finally, for the snow samples collected outside Arkhangelsk, on the Kindo peninsula, and close to the Yamal Peninsula in western Siberia in 2016, EC concentrations ranged between 7 and 161 ng g$^{-1}$ (median = 40 ± 47 ng g$^{-1}$). Outside Arkhangelsk, EC concentrations varied widely from 31 to 161 ng g$^{-1}$ with a median concentration in this region of 61 ± 43 ng g$^{-1}$. This is far below the 175 ng g$^{-1}$ observed in 2015, although there was only one sample collected in that year. On the Kindo peninsula, EC was relatively constant in 2016, ranging between 25 and 35 ng g$^{-1}$ (median = 28 ± 4 ng g$^{-1}$), which is more than 60 % lower compared with the 2015 values (median = 70 ± 34 ng g$^{-1}$). Finally, between Tomsk and Yamal, EC concentration was highly variable (7–119 ng g$^{-1}$) due to the different EC sources affecting snow (median = 50 ± 34 ng g$^{-1}$). For instance, it is expected that gas flaring affects snow close to Yamal, while snow collected in the south (Tomsk) is likely influenced by sources in Europe or local urban emissions. Nevertheless, the highest concentrations (> 100 ng g$^{-1}$) were observed north of 68° N, on the Yamal Peninsula.

We compared the measured EC concentrations in the snow samples with those calculated by FLEXPART. For this, the emission sensitivities were multiplied with the total emission fluxes from ECLIPSE (Sect. 2.4). A scatter plot of modelled and measured snow concentrations is presented in Fig. 1b. The results show a good correlation between modelled BC and measured EC concentrations for the 2015 and 2016 campaigns ($R_{2015} = 0.83$ and $R_{2016} = 0.68$, $p$ value < 0.05), but weaker correlation for 2014 ($R_{2014} = 0.53$, $p$ value < 0.05). The FB for individual samples is shown in Fig. S1. The MFB of the model for the 2014 snow measurements was −42 %, which shows that the model underestimated observations. In total, the model underestimated concentrations by 30–168 % for 17 out of 23 samples, whereas for the rest (six samples) FB values ranged between 20 and 148 % (median MFB: −56 ± 72 %) (Fig. S1). In 2015, the model underestimated observations by 48 % (median MFB: −56 ± 29 %) for 11 out of 12 samples (FB between −101 and −7 %, while one value was found to be 12 %). For 2016, FB values of the simulated concentrations of BC in snow showed another set of underestimation (median: −13 ± 60 %) between 0.3 and 198 % for 12 out of 19 samples. For the remaining seven samples, the model predicted higher concentrations compared with observations (10 to 75 %) (Fig. S1). RMSE values were estimated to be quite high, between 37 and 49 ng g$^{-1}$, due to the large variation in the observed EC concentrations.

The levels of EC in snow presented here are relatively high compared to previously reported concentrations in the Arctic. Apart from Aamaas et al. (2011), who measured maximum EC concentration in snow close to the airport of Svalbard of more than 1000 ng g$^{-1}$, most of the reported levels of EC in the relevant literature are close to our findings. For instance, Ruppel et al. (2014) found that EC concentrations have been increasing up to 103 ng g$^{-1}$ since 1970 in Svalbard. McConnell et al. (2007) reported that the BC concentrations measured at the D4 ice-core site in Greenland were 10 ng g$^{-1}$, at maximum, which most likely originated from BB in the conifer-rich boreal forest of the eastern and northern United States and Canada. Forsström et al. (2013) reported concentrations as high as 88 ng g$^{-1}$ in Scandinavia and lower ones at higher latitudes (11–14 ng g$^{-1}$ in Svalbard, 7–42 ng g$^{-1}$ in the Fram Strait and 9 ng g$^{-1}$ in Barrow). Svensson et al. (2013) collected snow samples from Tyresta National Park and Pallas-Yllästunturi National Park in Sweden. Tyresta is a relatively polluted site located circa 25 km from the city centre of Stockholm with a population of about 2 million people. Pallas-Yllästunturi National Park is located in Arctic Finland and a clean site with no major city influencing the local and regional air. The concentration of EC in Pallas-Yllästunturi was between 0 and 140 ng g$^{-1}$, while in Tyresta the BC concentrations were up to more than 7 times higher (53–810 ng g$^{-1}$). Furthermore, Doherty et al. (2010) in the most complete dataset for the Arctic snow and ice BC reported highly variable concentrations (up to 800 ng g$^{-1}$) for 5 consecutive years (2005–2009). Finally, in the most recent dataset for snow BC, Macdonald et al. (2017) reported BC concentrations ranging from 0.3 to 15 ng g$^{-1}$ for the samples collected near the Alert observatory (see Sect. 4.1).

3.2 Sources and origin of BC

We further analysed the model output in order to calculate relevant contributions from various BC source types to BC concentrations in snow (for method description, see Sect. 2.4). ECLIPSE emissions include waste burning (WST), industrial combustion and processing (IND), surface transportation (TRA), power plants, energy conversion, and extraction (ENE), residential and commercial combus-
Figure 2. Contribution from the various emission categories considered in the ECLIPSE and GFED inventories to simulated BC concentrations in snow in (a) 2014, (b) 2015 and (c) 2016 in western Siberia and northwestern European Russia. BB stands for biomass burning, WST for waste burning, IND for industrial combustion and processing, TRA for surface transportation, ENE for emissions from energy conversion and extraction, DOM for residential and commercial combustion, and FLR for gas flaring. Bars show the relative source contribution (0–100 %, right axis) and are sorted, from left to right, from the northernmost to the southernmost measurement location (coordinates are reported on the bottom as longitude/latitude). Measured EC concentrations in snow are reported with open circles, whereas modelled BC is shown with open rectangles (left axis).

In 2014, TRA contributed about 18 %, on average, to the simulated BC in snow, DOM 28 % and FLR 44 %, whereas ENE and IND were less significant. Maxima of TRA, DOM and FLR contributions were observed at a latitude of about 65° N, where measured EC and modelled BC were similar. An example of the contribution from the aforementioned dominant sources to snow BC concentrations for the highest measured EC concentration in snow is shown in Fig. 3. The transport sector includes emissions from all land-based transport of goods, animals and persons. It is more significant in southern Russia and close to the borders with Kazakhstan and Mongolia, where a large number of major Russian cities (e.g. Moscow, Kazan, Samara, Yekaterinburg, Tomsk, Novosibirsk, Krasnoyarsk) are located and connected with each other by federal highways. Residential and commercial combustion includes emissions from combustion in households and public and commercial buildings. Therefore, it is expected to be high for areas that consist of large population centres (Fig. 3). FLR emissions were found to contribute the most in this example, with a total concentration from this sector of 19.7 ng g⁻¹ (compared with 12.6 and 16.5 ng g⁻¹ in TRA and DOM, respectively) (Fig. 3).

On the Kindo peninsula and in Arkhangelsk, where snow sampling took place in 2015, the main contributions to snow BC were from DOM (47 %), TRA (30 %), BB (7 %) and FLR (6 %) (see Fig. 2). Similar to EC measurements in snow, simulated BC was also higher than in 2014, as the sampling sites were located closer to strong sources in Europe (Kindo) and close to a populated area (Arkhangelsk) with a strong regional impact. The highest concentration of EC was observed on the Kindo peninsula (33.13° E–66.53° N). Figure 4 shows the spatial distribution of emissions that contributed to simulated snow BC at the sampling point where the highest BC concentration was observed. In this case, TRA and DOM emissions from Europe mostly affected snow on the Kindo peninsula whereas FLR emissions were very low due to the long distance from the sampling point. Emissions from an unusual late winter–early spring episode of BB at the borders of Belarus, Ukraine and Russia also affected BC concentrations in snow in northwestern European Russia (Fig. 4). The importance of episodic BB releases in Russia, the miscalculation of satellite-retrieved BB emissions and their impact on Arctic concentrations in early spring has been explained.
by Evangeliou et al. (2016) and Hao et al. (2016). BB emissions, originating mostly from eastern Europe, contributed about 19.4 ng g\(^{-1}\) to the snow concentration at the receptor point (Fig. 4). TRA and DOM emissions were the dominant sources for this sampling point, contributing 33.6 and 47.2 ng g\(^{-1}\), respectively (Fig. 4).

Finally, in 2016, when samples were collected at the Kindo Peninsula, in Arkhangelsk and in Yamal, DOM, FLR and TRA contributed, on average, 31, 29 and 27 %, respectively (see Fig. 2c). Similar to the measured EC concentrations in snow, simulated concentrations of BC in 2016 were lower than those in 2015, on average. The highest measured EC concentration was observed in the Khanty-Mansiysk region (72.94°E–65.36°N), which mirrors the simulated BC concentration at the same point very well. The much higher contribution from TRA at this sampling point (38.6 ng g\(^{-1}\)) (Fig. 5b) is attributed to emissions from southern Russia (e.g. Tomsk), where all the main cities in Russia are located. Another large fraction of TRA emissions comes from central and eastern Europe (see also in http://niflheim.nilu.no/NikolaosPY/SnowBC_141516.py). Similar to TRA, emissions from DOM were mostly transported to Khanty-Mansiysk from central and eastern Europe, as well as from Turkey, contributing 36.6 ng g\(^{-1}\) (Fig. 5). As previously mentioned, the sampling point where the highest EC concentration was measured is located inside the largest gas flaring region of Russia. In addition, the corresponding emission sensitivity maps showed that the air was coming from the south passing directly through this high-emission region, making FLR emissions the highest contributing source (88.8 ng g\(^{-1}\)) (Fig. 5).

4 Discussion

4.1 Cross validation of modelled BC concentrations with public datasets

In this section, we present an effort to further validate our model calculations of BC concentrations in snow. For this purpose, BC concentrations in snow that were adopted from Doherty et al. (2010) were compared with modelled BC concentrations in snow that were simulated with FLEXPART as described in Sect. 2.4. Samples were collected in Alaska, Canada, Greenland, Svalbard, Norway, Russia and the Arctic Ocean during 2005–2009, on tundra, glaciers, ice caps, sea ice, frozen lakes and in boreal forests. Snow was collected mostly in spring, when the combination of snow cover and exposure to sunlight was at a maximum and before the snow had started to melt. Samples of melting snow collected in the summer of 2008 from Greenland and from Tromsø, Nor-
Figure 4. (a) FLEXPART emission sensitivity: (b) contribution from transportation (TRA), (c) residential and commercial combustion (DOM), and (d) gas flaring (FLR) to the maximum measured concentration of snow EC recorded in northwestern European Russia (Kindo peninsula and Arkhangelsk region) during the campaign of 2015.

way, were removed from the study, as we have no knowledge about the depth of the melt layer and effects of the percolation of meltwater through the snowpack. All samples were collected away from local sources of pollution. In many locations (Canadian Arctic, Russia, Greenland, Tromsø and Ny-Ålesund) samples were gathered at different depths throughout the snowpack, giving information on the seasonal evolution of BC concentrations as the snow accumulated (and/or sublimated) throughout the winter. In these cases only the surface BC was taken into account. The snow was melted and filtered, and the filters were analysed in a specially designed spectrophotometer system to infer the concentration of BC (for more information see Doherty et al., 2010). In contrast to our findings for the origin of snow BC in the Russian Arctic, a source apportionment analysis performed on the 2008 and 2009 measurements (Hegg et al., 2010) from this dataset showed that the dominant source of BC in the Arctic snowpack was BB. Specifically in eastern Siberia biomass burning of crops and grasslands contributed more snow BC in high latitudes than boreal forest fires, in contrast to the Canadian Arctic.

A comparison of modelled (FLEXPART) and measured BC concentrations (Doherty et al., 2010) in snow is depicted in Fig. S2. The model captures snow BC concentrations relatively well in most of the Arctic regions except for the Canadian Arctic, where the modelled concentrations of snow in 2007 were significantly higher. Samples from the same region in other years showed moderate agreement with modelled values. Similar to our finding for the new Russian measurements, the model underestimated deposition by 51%. The RMSE was estimated to be 52 ng g$^{-1}$, which is acceptable considering that the variation in snow concentrations in the dataset ranged from 0.3 to 783 ng g$^{-1}$. The highest measured concentrations of snow BC were observed in Russia, where the model showed a good spatial agreement. For instance, the highest values were obtained in western Siberia, close to the gas flaring regions of the Nenets and Komi oblasts, as well as in southeastern and northeastern Russia, where air masses were arriving from high-emitting sources in southeastern Asia. Lower biases in modelled BC concentrations were observed in northern Siberia with the exception of a few samples at the coasts of the Kara Sea and northeastern Siberia. Furthermore, biased BC concentrations were also observed in Greenland and northern Canada. In western Siberia, BC in snow presented in Doherty et al. (2010) between 2005 and 2009 was 80 ± 63 ng g$^{-1}$ on average, which is very close to the average value of measured EC obtained from the sampling 2014–2016 campaigns (50 ± 46 ng g$^{-1}$).

From the total number of samples presented in Doherty et al. (2010) that were used here for validation, only six
were collected on the Yamal Peninsula similar to some of the data presented in the current paper. The rest were collected in the Nenets and Komi regions and in eastern Russia and cannot be directly compared with snow EC measurements from the 2014–2016 campaigns. BC concentrations on the Yamal Peninsula in 2007 ranged from 4.1 to 17.6 ng g\(^{-1}\) (median ± interquartile: 10.3 ± 4.9 ng g\(^{-1}\)). In the same region, we report EC concentrations to be more than double, varying between 6.6 and 55 ng g\(^{-1}\) (median ± interquartile: 27.8 ± 25.5 ng g\(^{-1}\)), whereas there were two samples that showed EC concentrations of more than 100 ng g\(^{-1}\). As mentioned in Sect. 2.1, the sampling of snow for the EC analysis took place more than 500 m away from roads to minimize influence from traffic emissions, while a similar statement is also found in the Doherty et al. (2010) data. It is not clear whether the observed discrepancy arises as a measurement artefact (even though every effort has been taken in both papers to follow a robust protocol) or from real spatio-temporal variation.

Modelled BC concentrations simulated with FLEXPART were also compared with snow BC concentrations from samples collected at the Global Atmosphere Watch observatory at Alert, Nunavut, from 14 September 2014 to 1 June 2015 and they are available in Macdonald et al. (2017). BC concentrations in FLEXPART were simulated as in all previous analyses described in this paper (see Sect. 2.4). Time series of simulated and measured BC are depicted in Fig. S3 for the whole sampling period. As before, a correlation coefficient \((R)\) of 0.63 indicates that our model captures the temporal variation in the measured BC in snow. The RMSE was estimated to be almost 63 ng g\(^{-1}\), a relatively high value. The MFB of 47 % indicates a strong overestimation of snow concentrations, although in many samples the opposite was also observed (Fig. S3). This is in contrast to the previous datasets discussed, for which the model underestimated measurements.

Further analysis was carried out to adequately understand the origin of the aforementioned overestimations in the Canadian Arctic in both datasets (Doherty et al., 2010; Macdonald et al., 2017), as they are shown to be rather systematic. For this reason, we have calculated the average footprint emission sensitivities and the average BC contribution from the major sources in ECLIPSE for the 2007 snow samples in the Canadian Arctic and for Alert samples. We have chosen these samples because they were 3 or more times higher than the

Figure 5. (a) FLEXPART emission sensitivity and (b) contribution from transportation (TRA), (c) residential and commercial combustion (DOM), and (d) gas flaring (FLR) to the maximum measured concentration of snow EC recorded on the Kindo peninsula, in Arkhangelsk and on the Yamal Peninsula (northwestern European Russia, western Siberia) during the campaign of 2016.
observations and in this way we can locate the observed overestimations predicted with FLEXPART (Fig. 6).

Regarding the model overestimation for the 2007 samples, the average footprint emission sensitivity showed that the air was coming from continental regions of Canada with a smaller contribution from Scandinavia (Fig. 6). The highest emission sources for these samples were TRA and DOM, which contributed almost 80% to the snow concentrations, whereas forest fires were less important at the time of sampling. Two hot spots were identified, one along the border of Canada with the US and another, of smaller intensity, in southeastern Asia. A similar emission sensitivity was obtained for the same area of the Canadian Arctic in 2009, only slightly shifted to the north; simulated concentrations were in very good agreement with observations (Fig. S2). This shows that the model overestimation for the 2007 samples is likely attributed to an overestimation of TRA and DOM sources in North America in ECLIPSE for 2007. For the Alert samples, for which the model strongly overestimated BC, the major sources were TRA and FLR, which contributed 55%, and BB, which contributed about 7 ng g\(^{-1}\) (22%) on average (Fig. 6). Anthropogenic BC arriving from Europe and Russia has been previously shown to be important for Alert air pollutant concentrations (Sharma et al., 2013). The model overestimation of BC in snow samples at Alert needs further investigation. It is likely that it originates from anthropogenic emissions in northwestern America or in Europe because forest fires in Canada and Russia, although important for Alert (e.g. Qi et al., 2017), were not significant in the present comparison.

4.2 Model deviation from snow EC measurements and region-specific contribution of sources

It has been shown that measured concentrations of EC in snow in northwestern European Russia and western Siberia were underestimated in FLEXPART (Fig. 2). This was confirmed by the calculated fractional bias (see Sect. 3.2), the spatial distribution of which is shown in Fig. S1. To examine whether this underestimation was due to missing emission sources or errors in modelled transport and deposition, we have calculated the average footprint emission sensitivity for those sampling points, for which FLEXPART strongly (FB < −100%) and slightly (−100% < FB < 0%) underes-
Figure 7. (a) Footprint emission sensitivity from FLEXPART averaged for the sampling points where the model underestimated observations significantly (FB < −100 %) and (b) less significantly (−100 % < FB < 0 %). Black squares show the locations of active fires detected by MODIS (Moderate Resolution Imaging Spectroradiometer) (Giglio et al., 2003). Brown dots show the location of gas flaring sites from the Global Gas Flaring Reduction Partnership (GGFR) (http://www.worldbank.org/en/programs/gasflaringreduction).

estimated the observed values. The average footprint emission sensitivities are shown in Fig. 7 together with the locations of active fires in the last 2 months before the sample collection. The fire data were adopted from MODIS (Moderate Resolution Imaging Spectroradiometer) (Giglio et al., 2003) and the gas flaring facilities from the Global Gas Flaring Reduction Partnership (GGFR) (http://www.worldbank.org/en/programs/gasflaringreduction).

When the model strongly underestimated the measured EC (FB < −100 %), the average footprint emission sensitivity showed the highest values over the Yamal Peninsula and the agglomeration of many gas flares in Khanty-Mansiysk (Fig. 7b). This might confirm the findings of Huang et al. (2014) that gas flaring emissions in the ECLIPSE inventory, while very high, are still underestimated. According to a related study by Huang and Fu (2016), Russia contributes 57 % to the global BC emissions from gas flaring. Underestimation of modelled atmospheric concentrations compared to observations from the Barents and Kara seas was recently also reported by Popovicheva et al. (2017), although the underestimation was relatively small.

When FLEXPART showed a moderate underestimation of EC concentrations in snow (−100 % < FB < 0 %), the emission sensitivity was high near Arkhangelsk and over Scandinavia (Fig. 7). BC emissions in Scandinavia are considered relatively low in most inventories and contribute no more than 6.5 % to the global emissions in ACCMIP (Aerosol Chemistry Climate Model Intercomparison Project) (Lamarque et al., 2013), 6.2 % in EDGARv4.2 (Emission Database for Global Atmospheric Research) (Olivier et al., 2005), 2.1 % in MACCity (Monitoring Atmospheric Composition and Climate and megacity Zoom for the Environment) (Hollingsworth et al., 2008; Stein et al., 2012) and 3.3 % in ECLIPSE (Klimont et al., 2017). The highest emission sensitivity was found over northwestern Russia (Fig. 7), a region that includes Murmansk. Pollution levels in Murmansk could be high due to emissions from local industry, mining, heating and transport (Law and Stohl, 2007). Another potential source region was the Nenets–Komi area and western Kazakhstan, where a few other flaring facilities are located (Fig. 7).

Figure 7 shows that the underestimation of observed EC concentrations in snow strongly depends on the region where samples are collected. In western Siberia, the underestimation was larger than in northwestern European Russia. For this reason, we have computed the average region-specific emission sensitivities and the average region-specific contribution from the major polluting sources identified in the ECLIPSE dataset. We distinguish between three regions: northwestern European Russia, western Siberia (north of 62° N) and western Siberia (south of 62° N) (Figs. S4–S6).

For the samples collected in northwestern European Russia (Fig. S4), an average contribution of 21.6 ng g$^{-1}$ from all sources was estimated to have originated mainly from TRA (7.7 ng g$^{-1}$) and DOM (10.4 ng g$^{-1}$) sources in Finland. The contribution from BB and FLR emissions was insignificant (8 and 6 %, respectively), whereas the rest of the ECLIPSE sources were negligible (IND, ENE, WST). For the samples collected at high latitudes in western Siberia, the average contribution from all sources was more than 4 times higher (86 ng g$^{-1}$) than those observed in northwestern European Russia (Fig. S5). FLR emissions accounted for 40 % of the total contribution, which reflects the proximity of the sampling site to the main flaring facilities of Russia. The average contribution from TRA activities in Europe and southeastern Russia to the northern part of western Siberia was 24 %. DOM emissions in eastern Europe also contributed another 28 %. Finally, for the samples that were collected in the southern part of western Siberia an average contribution of 47.4 ng g$^{-1}$ was estimated from all sources included in
ECLIPSE (Fig. S6). The highest contributing categories were TRA and DOM, whereas FLR appeared to contribute less, although the sampling site is close to the Khanty-Mansiysk flaring region. This is attributed to the prevailing winds that forced flaring emissions to a northernmost direction opposite to the location of the sampling stations (see Fig. S6).

Overall, the region-specific analysis of the sources contributing to modelled BC in snow showed that the DOM, FLR and/or TRA sources might explain the model underestimation in the high Arctic. However, in the most recent assessment of BC of the higher Arctic (Popovicheva et al., 2017; Winiger et al., 2017), it was shown that ECLIPSE captures levels of BC quite well, whereas FLR emissions might have a smaller impact in the central Siberian Arctic (Tiksi) than previously estimated. Surprisingly, the average contribution from BB in lower latitudes was extremely low in all western Siberia (Figs. S5 and S6), despite the fact that sampling took place in springtime, when BB becomes important. Evangeliou et al. (2016) reported that using a different dataset, which is based on the same approach as GFED but includes updated emission factors for Eurasia, surface concentrations of BC in the Arctic stations can be substantially higher. This shows the need for further investigation of BC sources in the Russian Arctic.

5 Conclusions

We have analysed snow samples collected in western Siberia and northwestern European Russia in 2014, 2015 and 2016 with respect to EC. This region is of major interest due to its large uncertainty in BC emissions and because it is located in the main transport route of BC to the Arctic. An effort to constrain the sources that contribute to measured concentration of BC in snow was made using the LPDM FLEXPART (version 10).

The observed EC levels in snow varied widely within and between regions (3–219 ng g$^{-1}$ for 2014, 46–175 ng g$^{-1}$ in 2015 and 7–161 ng g$^{-1}$ in 2016) and are in the upper range of previously reported concentrations of EC and BC in snow in the Arctic region. However, the observed levels presented here appear typical for western Siberia, which is subject to high domestic Russian emissions as well as to transport from distant European ones.

The snow BC concentrations predicted by the model are in fair agreement with EC observations over western Siberia and northwestern European Russia ($R = 0.5$–0.8). However, the calculated negative MFB values ($-48\%$ to $-27\%$) showed that the model systematically underestimated observations in Russia. This underestimation strongly depended on the region where the samples were collected. In northwestern European Russia, the main contributing sources were TRA and DOM mainly from adjacent regions in Finland. TRA and DOM contributed twice as much to snow BC sampled at low latitudes of western Siberia ($< 60^\circ$ N) as compared to samples collected over regions above $60^\circ$ N; the majority of these emissions originated from highly populated centres in central Europe. Finally, in higher latitudes of western Siberia ($> 60^\circ$ N), snow BC concentrations were further increased mainly due to FLR emissions from facilities located close to the snow sampling points.

The modelled BC concentrations in snow were further investigated using two independent public measurement datasets that include samples from all over the Arctic for the period from 2005 to 2009 and from Alert in 2014 and 2015. The model captured levels of BC fairly well despite the large variation in measured concentrations. An exception was observed in North America in spring 2007 and at the Alert observatory in late winter–early spring 2015. In both cases, the major sources were along the Canadian border with the US and in western Europe. Considering the fact that similar deviations were not observed in samples collected in the area during other years, it is likely that some of the prevailing sources of BC in this region show strong temporal variability in their emissions, and this is not taken into account in the ECLIPSE inventory. Previously reported average measurements of BC concentrations in snow in western Siberia and northwestern European Russia were $80 \pm 43$ ng g$^{-1}$, which is about $30\%$ higher than the EC measurements presented here ($50 \pm 46$ ng g$^{-1}$).

Data availability. All data used for the present publication can be obtained from the corresponding author upon request.

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Author contributions. NE designed and performed the modelling experiments and wrote the paper. VPS organized and performed the sampling of EC; KEY performed all the TOA of the snow samples. SE modified the FLEXPART model for the calculation of footprint emission sensitivities for deposited mass. ES wrote an algorithm that computes the starting date of the FLEXPART releases based on the water equivalent volume from ECMWF. OSP, VOK, VBK, AAL, DPS and SNV assisted the sampling campaigns in western Siberia and northwestern European Russia during 2014–2016. RLT and AS supervised the study and wrote parts of the paper.

Competing interests. The authors declare that they have no conflict of interest.

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