FLEXPART v10.1 simulation of source contributions to Arctic black carbon

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Abstract. The Arctic environment is undergoing rapid changes such as faster warming than the global average and exceptional melting of glaciers in Greenland. Black carbon (BC) particles, which are a short-lived climate pollutant, are one cause of Arctic warming and glacier melting. However, the sources of BC particles are still uncertain. We simulated the potential emission sensitivity of atmospheric BC present over the Arctic (north of 66\(^\circ\) N) using the FLEXPART (FLEXible PARTicle) Lagrangian transport model (version 10.1). This version includes a new aerosol wet removal scheme, which better represents particle-scavenging processes than older versions did. Arctic BC at the surface (0–500 m) and high altitudes (4750–5250 m) is sensitive to emissions in high latitude (north of 60\(^\circ\) N) and mid-latitude (30–60\(^\circ\) N) regions, respectively. Geospatial sources of Arctic BC were quantified, with a focus on emissions from anthropogenic activities (including domestic biofuel burning) and open biomass burning (including agricultural burning in the open field) in 2010. We found that anthropogenic sources contributed 82\% and 83\% of annual Arctic BC at the surface and high altitudes, respectively. A large fraction (40\%) of BC in the Arctic at high altitudes comes from anthropogenic emissions in East Asia, which suggests that the rapidly growing economies of developing countries could have a non-negligible effect on the Arctic. To our knowledge, this is the first year-round evaluation of Arctic BC sources that has been performed using the new wet deposition scheme in FLEXPART. The study provides a scientific basis for actions to mitigate the rapidly changing Arctic environment.

1 Introduction

The Arctic region has experienced warming at a rate twice that of the global average in recent decades (Cohen et al., 2014). The Arctic cryosphere has been undergoing unprecedented changes since the mid-1800s (Trusel et al., 2018). Glacier cover in Greenland reached its historically lowest level in summer 2012 (Tilling et al., 2015). Evidence indicates that the emissions and transport of greenhouse gases and aerosols to the Arctic region are contributing to such warming and melting of snow and ice (Keegan et al., 2014; Najafi et al., 2015). Short-lived climate pollutants such as black carbon (BC) particles (e.g., Sand et al., 2016; Yang et al., 2019), sulfate aerosol (Yang et al., 2018), tropospheric ozone, and methane greatly affect the Arctic climate (AMAP, 2015; Quinn et al., 2008).
BC particles are emitted during incomplete combustion of fossil fuels, biofuels, and biomass. BC warms the atmosphere by direct absorption of solar radiation. The deposition of BC on snow and ice surfaces accelerates their melting through decreasing albedo, which contributes to the rapid loss of glaciers. In the Arctic region, ground-based observations have indicated that BC shows clear seasonal variations, with elevated mass concentrations in winter and spring (the so-called Arctic haze) and low values in summer (Law and Stohl, 2007). Such seasonal variations are explained by increased transport from lower latitudes in the cold season and increased wet scavenging in the warm season (Shaw, 1995; Garrett et al., 2011; Shen et al., 2017).

The presence of BC particles in the Arctic is mainly attributed to emissions in high-latitude regions outside the Arctic, such as northern Europe and Russia (Stohl, 2006; Brock et al., 2011). This is partly caused by the polar dome (Stohl, 2006), which is formed because of the presence of constant potential temperature near the surface. The emissions in high-latitude regions are transported to the Arctic region and trapped in the dome, which increases the surface concentration. Recently, Schmale et al. (2018) suggested that local emissions from within the Arctic are another important source, and these are expected to increase in the future.

Although numerous studies have been performed, results regarding regional contributions of BC sources in the Arctic are still inconclusive. For example, ground-based observations and Lagrangian transport model results reported by Winiger et al. (2016) showed that BC in Arctic Scandinavia is predominantly linked to emissions in Europe. Over the whole Arctic region (north of 66° N), Russia contributes 62% to surface BC in terms of the annual mean (Ikeda et al., 2017). Gas flaring in Russia has been identified as a major (42%) source of BC at the Arctic surface (Stohl et al., 2013). Xu et al. (2017) found that anthropogenic emissions from northern Asia contribute 40%–45% of Arctic surface BC in winter and spring. However, the results of some other studies have suggested that Russia, Europe, and southern Asia each contribute 20%–25% of BC to the low-altitude springtime Arctic haze (Koch and Hansen, 2005). Sand et al. (2016) found that the surface temperature in the Arctic is most sensitive to emissions in Arctic countries, and Asian countries contribute greatly to Arctic warming because of the large absolute amount of emissions. With these large disagreements among studies, it is thus necessary to unveil BC sources in the Arctic with high-precision simulations.

Various models have been used to investigate BC sources in the Arctic. Depending on the simulation method, these models are generally categorized as Lagrangian transport models (Hirdman et al., 2010; Liu et al., 2015; Stohl et al., 2013; Stohl, 2006), chemical transport models (CTMs; Ikeda et al., 2017, 2020; Qi et al., 2017; Shindell et al., 2008; Wang et al., 2011; Xu et al., 2017), and global climate models (GCMs; Ma et al., 2013; Koch and Hansen, 2005; Schacht et al., 2019; H. Wang et al., 2014) (Table 1). The treatment of wet-scavenging parameterizations is a key factor affecting the model performance, which determines the uncertainties related to BC particle removal (Kipling et al., 2013; Schacht et al., 2019; Q. Wang et al., 2014). The use of emission inventories is another important factor that affects the simulation results (Dong et al., 2019). The observations of BC that are used for model comparisons may be biased by a factor of 2 depending on the method used (Sinha et al., 2017; Sharma et al., 2017). There are still large uncertainties regarding the sources of BC in the Arctic with respect to emission sectors (anthropogenic sources and open biomass burning) and geospatial contributions (Eckhardt et al., 2015).

The FLEXible PARTicle dispersion model (FLEXPART) had been used to investigate the transport pathways and source contributions of BC in the Arctic (Stohl et al., 1998, 2013; Stohl, 2006). For the FLEXPART model up to version 9, wet removal was treated considering below-cloud and within-cloud scavenging processes (Hertel et al., 1995; McMahon and Denison, 1979), which depend on cloud liquid-water content, precipitation rate, and the depth of the cloud. However, clouds were parameterized based on relative humidity with unrealistic precipitation, clouds frequently extended to the surface, and at times no clouds could be found in grid cells (Grythe et al., 2017). Recently, version 10 of FLEXPART has been developed in which cloud is differentiated into liquid, solid, and mixed phase, and the cloud distribution is more consistent with the precipitation data (Grythe et al., 2017). This improvement in the cloud distribution and phase leads to a more realistic distribution of below-cloud and in-cloud scavenging events. In this study, we quantified region-separated sources of BC in the Arctic in 2010 by using FLEXPART v10.1. We first evaluated the model performance by comparing the results with those based on observations at surface sites. The source contributions of emission sectors and geospatial contributions were evaluated by incorporating the Arctic BC footprint into the emission inventories.

2 Materials and methods

2.1 Transport model

The FLEXPART model (version 10.1) was run in backward mode to simulate BC footprints in the Arctic region. The calculation of wet deposition was improved compared with those in previous versions because in-cloud scavenging and below-cloud scavenging of particles were separately calculated (Grythe et al., 2017). In previous versions of FLEXPART, in the in-cloud scavenging scheme, the aerosol scavenging coefficient depended on the cloud water content, which was calculated according to an empirical relationship with precipitation rate, in which all aerosols had the same nucleation efficiency (Hertel et al., 1995; Stohl et al., 2005). In the new version, the in-cloud scavenging scheme depends on the cloud water phase (liquid, ice, or mixed phase). Aerosols
Table 1. Comparison of BC source contributions in the Arctic surface.

<table>
<thead>
<tr>
<th>Model and versions</th>
<th>Model type</th>
<th>Wet deposition</th>
<th>Grid resolution</th>
<th>Meteorology</th>
<th>Emissions</th>
<th>Domain or sites</th>
<th>Year or season</th>
<th>Major source regions or sectors</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>FLEXPART 6.2</td>
<td>Lagrangian</td>
<td>Stohl et al. (2005)</td>
<td>1° × 1°</td>
<td>ECMWF operational</td>
<td>Unspecified (BC sensitivities were calculated)</td>
<td>Alert, Utqiagvik, Zeppelin</td>
<td>1989–2009</td>
<td>Northern Eurasia</td>
<td>Hirdman et al. (2010)</td>
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<tr>
<td>FLEXPART 6.2</td>
<td>Lagrangian</td>
<td>Stohl et al. (2005)</td>
<td>1° × 1°</td>
<td>ECMWF operational</td>
<td>ECLIPSE4(GAINS), GFED3</td>
<td>Arctic (north of 66° N)</td>
<td>2008–2010</td>
<td>Flaring (42 %), residential (&gt; 20 %)</td>
<td>Stohl et al. (2013)</td>
</tr>
<tr>
<td>FLEXPART 9.2</td>
<td>Lagrangian</td>
<td>Stohl et al. (2005)</td>
<td>1° × 1°</td>
<td>ECMWF operational</td>
<td>ECLIPSE5(GAINS), GFED4.1</td>
<td>Arctic (north of 66.7° N)</td>
<td>2011–2015</td>
<td>Residential and open burning (39 %)</td>
<td>Winiger et al. (2019)</td>
</tr>
<tr>
<td>FLEXPART 10.1</td>
<td>Lagrangian</td>
<td>Grythe et al. (2017)</td>
<td>1° × 1°</td>
<td>ECMWF operational</td>
<td>HTAP2, GFED3, Huang et al. (2015) for Russia flaring</td>
<td>Arctic (north of 66° N)</td>
<td>2010</td>
<td>Flaring (36 %), open burning (18 %), residential (15 %), others (31 %)</td>
<td>Current study</td>
</tr>
<tr>
<td>GEOS-Chem</td>
<td>CTM</td>
<td>Wang et al. (2011)</td>
<td>2° × 2.5°</td>
<td>GEOS-5</td>
<td>Bond et al. (2004), Zhang et al. (2009), GFED3</td>
<td>Alert, Utqiagvik, Zeppelin</td>
<td>April 2008</td>
<td>Asian anthropogenic (35 %–45 %), Siberian biomass burning (46 %–64 %)</td>
<td>Qi et al. (2017)</td>
</tr>
<tr>
<td>GEOS-Chem10.01</td>
<td>CTM</td>
<td>Wang et al. (2011)</td>
<td>2° × 2.5°</td>
<td>GEOS-5</td>
<td>HTAP2, ECLIPSE5, GFED4</td>
<td>Arctic (north of 66.5° N)</td>
<td>2009–2011</td>
<td>Northern Asia anthropogenic (40 %–45 %) in winter–spring</td>
<td>Xu et al. (2017)</td>
</tr>
<tr>
<td>CAM5</td>
<td>GCM</td>
<td>Wang et al. (2013)</td>
<td>1.9° × 2.5°</td>
<td>MERRA</td>
<td>IPCC AR5</td>
<td>Arctic (north of 66.5° N)</td>
<td>1996–2005</td>
<td>Northern Europe in winter, Northern Asia in summer</td>
<td>Wang et al. (2014)</td>
</tr>
</tbody>
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were set as ice nuclei for ice clouds and as cloud condensation nuclei for liquid-water clouds. For mixed-phase clouds, it was assumed that 10% of aerosols are ice nuclei and 90% are cloud condensation nuclei, because BC is much more efficiently removed in liquid-water clouds than in ice clouds (Cozic et al., 2007; Grythe et al., 2017). The below-cloud scavenging scheme can parameterize below-cloud removal as a function of aerosol particle size, precipitation type (snow or rain), and intensity. The biases produced in simulations using the new scheme are therefore smaller than those in the old scheme for wet deposition of aerosols, especially at high latitudes (Grythe et al., 2017).

The Arctic region is defined as areas north of 66° N. The potential BC emission sensitivities at two heights in the Arctic region, i.e., the surface (0–500 m) and high altitudes (4750–5250 m), were simulated. The FLEXPART outputs were set as gridded retention times. We performed tests at 500, 2000, and 5000 m, and chose 500 m as the upper boundary height of the model output. The model was driven with operational analytical data from the European Centre for Medium-Range Weather Forecasts (ECMWF) at a spatial resolution of 1° × 1° with 61 vertical levels. Temporally, ECMWF has a resolution of 3 h, with 6 h analysis and 3 h forecast time steps. The simulation period was set at 60 d backward starting from each month in 2010. The maximum lifetime of BC was set at 20 d, because its suspension time in the upper atmosphere during long-range transport is longer than that at the surface level (Stohl et al., 2013). We implemented the wet deposition scheme in the backward calculations, but it was not represented in the default setting (FLEXPART v10.1, https://www.flexpart.eu/downloads, last access: 10 April 2017).

The chemistry and microphysics could not be resolved by FLEXPART. The model therefore ignores hydrophobic to hydrophilic state changes and size changes of BC, and it assumes that all BC particles are aged hydrophilic particles. This may lead to an overestimation of BC removal and hence force underestimation of simulated BC concentration, especially of fossil fuel combustion sources where BC could be in the hydrophobic state for a few days. A logarithmic size distribution of BC with a mean diameter of 0.16 µm and a standard deviation of 1.96, in accordance with our ship observations in the Arctic, was used (Taketani et al., 2016). The particle density was assumed to be 2000 kg m\(^{-3}\), and 1 million computational particles were randomly generated in the Arctic region for the backward runs.

Four ground-based observations made during the period 2007–2011 were used to validate the model performance. The potential BC emission sensitivity at 0–500 m a.g.l. (above ground level) from a 0.1° grid centered at each site was simulated. Other model parameterizations were consistent with those for the Arctic region, except that 200 000 computational particles were released.
2.2 Emission inventories

We focused on BC sources from anthropogenic emissions and open biomass burning. The Hemispheric Transport of Air Pollution version 2 inventory (HTAP2) for 2010 was used for monthly anthropogenic BC emissions (Janssens-Maenhout et al., 2015), which include sectors from energy, industry, residential, and transport. It is worth noting that the residential sector includes not only combustions of fossil fuels but also biofuels. However, as it has been reported that BC emissions in Russia were underestimated in HTAP2, we used the BC emissions reported by Huang et al. (2015) for Russia, in which the annual BC emissions were 224 Gg yr\(^{-1}\). For open biomass burning, we used the monthly BC emissions from the Global Fire Emissions Database version 3 inventory (GFED3) (van der Werf et al., 2010) for the purposes of intercomparison with other studies, as this version is widely used. The term “open biomass burning” here indicates burning of biomass in the open field as is determined by the remote sensing measurement basis, including forest, agricultural waste, peat fires, grassland and savanna, woodland, deforestation, and degradation, where biofuel burning for residential use is not included. Geospatial distributions of emissions from anthropogenic sources and open biomass burning in January and July are shown in Fig. S1 in the Supplement.

2.3 Calculation of Arctic BC source contributions

The source contributions to Arctic BC were derived by incorporating the gridded retention time into the column emission flux, which was derived from the emission inventories in each grid. Calculations for anthropogenic sources and open biomass burning were performed separately and the sum was used. For anthropogenic sources, the regions were separated into North America and Canada (25–80° N, 50–170° W), Europe (30–80° N, 0–30° E), Russia (53–80° N, 30–180° E), East Asia (35–53° N, 75–150° E and 20–35° N, 100–150° E), and others (the rest) (Fig. 1a). For open biomass burning sources, the regions were separated into Alaska and Canada (50–75° N, 50–170° W), Siberia (50–75° N, 60–180° E), and others (Fig. 1b).

2.4 Observations

BC levels simulated by FLEXPART were compared with those based on surface observations at four sites: Utqiaġvik, USA (formerly Barrow) (71.3° N, 156.6° W; 11 m a.s.l.); Alert, Canada (82.5° N, 62.3° W; 210 m a.s.l.); Zeppelin, Norway (78.9° N, 11.9° E; 478 m a.s.l.); and Tiksi, Russia (71.6° N, 128.9° E; 8 m a.s.l.). Aerosol light absorption was determined by using particle soot absorption photometers (PSAPs) at Utqiaġvik, Alert, and Zeppelin, and an Aethalometer at Tiksi. For PSAP measurements, the equivalent BC values were derived by using a mass absorption efficiency of 10 m\(^2\) g\(^{-1}\). The equivalent BC at Tiksi, which was determined with an Aethalometer, was obtained directly. These measurement data were obtained from the European Monitoring and Evaluation Programme and the World Data Centre for Aerosols database (http://ebas.nilu.no, last access: 1 December 2017) (Tørseth et al., 2012).

It is worth noting that uncertainties could be introduced by using different BC measurement techniques. An evaluation of three methods for measuring BC at Alert, Canada, indicated that an average of the refractory BC determined with a single-particle soot photometer (SP2) and elemental carbon (EC) determined from filter samples give the best estimate of BC mass (Sharma et al., 2017). Xu et al. (2017) reported that the equivalent BC determined with a PSAP was close to the average of the values for refractory BC and EC at Alert. In this study, we consider that the equivalent BC values determined with a PSAP at Utqiaġvik, Alert, and Zeppelin to be the best estimate. There may be uncertainties in the equivalent BC observations performed with an Aethalometer at Tiksi because of co-existing particles such as light-absorbing organic aerosols, scattering particles, and dusts (Kirschstetter et al., 2004; Lack and Langridge, 2013). Interference by the filter and uncertainties in the mass absorption cross section could also contribute to the bias observed in measurements made with an Aethalometer at Tiksi.

3 Results and discussion

3.1 Comparisons of simulations with BC observations at Arctic surface sites

FLEXPART generally reproduced the seasonal variations in BC at four Arctic sites well (Pearson correlation coefficient, \(R = 0.53–0.80\), and root-mean-square error, RMSE = 15.1–56.8 ng m\(^{-3}\)) (Fig. 2). Winter maxima were observed for the four sites, while a secondary elevation was observed for Alert and Tiksi. At Utqiaġvik, the observed high values of BC were unintentionally excluded during data screening in the forest fire season in summer (Stohl et al., 2013); the original observed BC is supposed to be higher as was reflected by the simulation. This seasonality is probably related to relatively stronger transport to the Arctic region in winter, accompanied by lower BC aging and inefficient removal, as simulated by older versions of FLEXPART (Eckhardt et al., 2015; Stohl et al., 2013).

From January to May at Utqiaġvik and Alert, the mean BC values simulated by FLEXPART v10.1 were 32.2 and 31.2 ng m\(^{-3}\), respectively, which were 46 % lower than the observations (59.3 and 58.2 ng m\(^{-3}\), respectively). This is probably related to the inadequate BC emission in the inventory, although seasonal variations in residential heating are included in HTAP2, which would reduce the simulation bias (Xu et al., 2017). Simulations by GEOS-Chem using the same emission inventories also underestimated BC levels at
Figure 1. Regional separation for quantification of BC in the Arctic from (a) anthropogenic and (b) open biomass burning sources.

Figure 2. Observed (filled circles) and modeled (bars) seasonal variations in BC mass concentrations at Arctic sites. Contributions from anthropogenic sources (blue) and open biomass burning (red) in each month are shown. Monthly averages of observed (filled circles) and simulated (bars) BC were conducted for 2007–2011 at Alert, Canada (82.5° N, 62.3° W), and Zeppelin, Norway (78.9° N, 11.9° E); for 2009 at Utqiaġvik, USA (71.3° N, 156.6° W); and for 2010–2014 at Tiksi, Russia (128.9° E, 71.6° N). R and RMSE indicate correlation coefficient and root-mean-square error (ng m$^{-3}$), respectively.

Utqiaġvik and Alert (Ikeda et al., 2017; Xu et al., 2017). The underestimation by FLEXPART could also be partly contributed by the assumption that all particles are hydrophilic, where the BC scavenging could be overestimated. The corresponding uncertainties are larger in winter months, when there are more sources from fossil fuel combustion.

At Zeppelin, the FLEXPART-simulated BC (39.1 ng m$^{-3}$ for annual mean) was 85% higher than the observed value (21.1 ng m$^{-3}$ for annual mean), especially in winter (112%...
higher). It has been reported that riming in mixed-phase clouds occurs frequently at Zeppelin (Qi et al., 2017). During the riming process, BC particles act as ice particles and collide with the relatively numerous water drops, which form frozen cloud droplets, and then snow is precipitated. This results in relatively efficient BC scavenging (Hegg et al., 2011). Such a process could not be dealt with by the model. At Tiksi, FLEXPART underestimated BC (74.4 ng m\(^{-3}\) for annual mean) in comparison with observation (104.2 ng m\(^{-3}\) for annual mean). Other than the hydrophilic BC assumption and underestimated BC emission in the simulation as in the cases for Utqiaġvik and Alert, the observations at Tiksi by an Aethalometer could contain light-absorbing particles other than BC, resulting in higher observed concentrations if compared with those obtained by SP2, EC, or PSAP.

Anthropogenic emissions are the main sources of BC at the four Arctic sites from late autumn to spring, whereas open biomass burning emissions make large contributions in summer. From October to April, anthropogenic emissions accounted for 87%–100% of BC sources at all the observation sites. At Utqiaġvik, open biomass burning accounted for 35%–78% of BC in June–September. There are large interannual variations in both observed and simulated BC (Fig. S2). In June–August 2010, the mean contributions of open biomass burning to BC were 6.3, 2.4, and 8.6 times those from anthropogenic sources at Alert, Zeppelin, and Tiksi, respectively. In this study, we focused on BC in the Arctic region in 2010.

3.2 Potential emission sensitivity of Arctic BC

The potential emission sensitivities (footprint) of Arctic BC showed different patterns with respect to altitude. The Arctic surface is sensitive to emissions at high latitudes (>& 60° N). Air masses stayed for over 60 s in each of the 1° grids from the eastern part of northern Eurasia and the Arctic Ocean before being transported to the Arctic surface in the winter, represented by January (Fig. 3a). In comparison, during the summer, represented by July, BC at the Arctic surface was mainly affected by air masses that originated from the Arctic Ocean and the Norwegian Sea (Fig. 3b). These results imply that local BC emissions within the Arctic regions, although relatively weak compared with those from the mid-latitude regions, could strongly affect Arctic air pollution. Local BC emissions are important in the wintertime because the relatively stable boundary layer does not favor pollution dispersion. Recent increases in anthropogenic emissions in the Arctic region, which have been caused by the petroleum industry and development of the Northern Sea Route, are expected to cause deterioration of air quality in the Arctic. Socioeconomic developments in the Arctic region would increase local BC emissions, and this will be a non-negligible issue in the future (Roiger et al., 2015; Schmale et al., 2018).

BC at high altitudes in the Arctic is more sensitive to mid-latitude (30–60° N) emissions, especially in wintertime. In January, air masses hovered over the Bering Sea and the North Atlantic Ocean before arriving at the Arctic (Fig. 3c). A notable corridor at 30–50° N covering Eurasia and the US was the sensitive region that affected BC at high altitudes in the Arctic in January. These results indicate that mid-latitude emissions, especially those with relatively large strengths from East Asia, eastern America, and Europe, could alter the atmospheric constituents at high altitudes in the Arctic. Central to east Siberia was the most sensitive region for BC at high altitudes in the Arctic in July. These results suggest that pollutants from frequent and extensive wildfires in Siberia in summer are readily transported to high altitudes in the Arctic. Boreal fires are expected to occur more frequently and over larger burning areas under future warming (Veira et al., 2016); therefore, the atmospheric constituents and climate in the Arctic could undergo more rapid changes.

3.3 Seasonal variations and sources of Arctic surface BC

Arctic surface BC showed clear seasonal variations, with a primary peak in winter–spring (December–March, 61.8–82.8 ng m\(^{-3}\)) and a secondary peak in summer (July, 52.7 ng m\(^{-3}\)). BC levels were relatively low in May–June (21.8–23.1 ng m\(^{-3}\)) and September–November (34.1–40.9 ng m\(^{-3}\)) (Fig. 4a). This seasonality agrees with observations and simulations at Alert, Tiksi, and Utqiaġvik if considering the unintentional data exclusion (Stohl et al., 2013), and previous studies targeting the whole Arctic (Ikeda et al., 2017; Xu et al., 2017). Compared with the study reported by Stohl et al. (2013), the current work using the new scheme produced smaller discrepancies between the simulated data and observations. Although the simulation periods (monthly means for 2007–2011 in this study and for 2008–2010 in the old scheme) and the anthropogenic emission inventories (HTAP2 in this study and ECLIPSE4 in the previous study) are different, the new scheme shows potential for better representing BC transport and removal processes in the Arctic.

The annual mean Arctic BC at the surface was estimated to be 48.2 ng m\(^{-3}\). From October to April, anthropogenic sources accounted for 96%–100% of total BC at the Arctic surface. Specifically, anthropogenic emissions from Russia accounted for 61%–76% of total BC in October–May (56% annually), and these were the dominant sources of Arctic BC at the surface. From an isentropic perspective, the meteorological conditions in winter favored the transport of pollutants from northern Eurasia to the lower Arctic, along with diabatic cooling and strong inversions (Klönecki et al., 2003). In comparison, open biomass burning from boreal regions accounted for 56%–85% (75% on average) of Arctic BC at the surface in summer; open biomass burning emissions from North America and Canada accounted for 54% of total Arctic surface BC in June, and those from Siberia
Figure 3. Footprints of Arctic BC shown as retention time(s) of (a) BC at surface (0–500 m) in January 2010, (b) BC at surface in July 2010, (c) BC at high altitudes (4750–5250 m) in January 2010, and (d) BC at high altitudes in July 2010.

Figure 4. Contributions of anthropogenic sources (prefixed “AN_” in the legend) and open biomass burning ("BB_") from each region to (a) seasonal variations in Arctic surface BC, (b) annual mean Arctic surface BC, (c) seasonal variations in Arctic BC at high altitudes, and (d) annual mean of Arctic BC at high altitudes.
accounted for 59\%–61\% in July–August. Wildfires in the boreal forests in summer had a major effect on air quality in the Arctic.

On an annual basis, anthropogenic sources and open biomass burning emissions accounted for 82\% and 18\%, respectively, of total Arctic surface BC. In which, gas flaring and residential burning (including burning of fossil fuels and biofuels) are accounting for 36\% (28\%–57\% in October–March) and 15\% (13\%–25\% in October–March), respectively (Fig. 5a and b). Our results support Stohl et al. (2013) such that residential combustion emissions, especially in winter, are important sources of Arctic BC (Table 1). We estimated a contribution of gas flaring to Arctic surface BC of 17.5 ng m\(^{-3}\) (36\% of total). In comparison, the value was estimated as 11.8 ng m\(^{-3}\) using an average Arctic surface BC of 28 ng m\(^{-3}\) and a fraction from gas flaring of 42\% evaluated by earlier versions of FLEXPART (Stohl et al., 2013; Winiger et al., 2019). The different contribution could be partly attributed to the difference in gas flaring emission inventory. BC emission from gas flaring in Russia by Huang et al. (2015) was used in the current study, where total BC emission from gas flaring in Russia in 2010 was ca. 81.1 kt, which was larger than the estimate of ca. 64.9 kt by the GAINS inventory (Klimont et al., 2017) used by Stohl et al. (2013). Moreover, adopting the ECLIPSev5 inventory as was used by Winiger et al. (2019), we estimated that gas flaring was contributing 14.4 ng m\(^{-3}\) to Arctic surface BC using FLEXPART v10.1, a value 22\% higher than those obtained using FLEXPART v9. This difference could be attributed to the improvement of the wet-scavenging scheme by FLEXPART v10.1.

A recent study based on isotope observations at the Arctic sites and FLEXPART v9.2 simulation suggested that open biomass burning, including open field burning and residential biofuel burning, contributed 39\% to annual BC in 2011–2015 (Winiger et al., 2019) (Table 1). In comparison, we estimated that residential burning and open biomass burning together account for 33\% of total Arctic surface BC. As the residential burning in our study includes burning of both biofuels and fossil fuels, our results indicated that biomass burning has a relatively smaller contribution. Other than the differences in BC removal treatment between different versions of the model, the contribution difference could also be attributed to the different emission inventories and years (2010 versus 2011–2015).

The geospatial contributions of anthropogenic sources and open biomass burning emissions can be further illustrated by taking January and July as examples. In January, high levels of anthropogenic emissions from Russia (contributing 64\% of Arctic surface BC), Europe (18\%), and East Asia (9\%) were identified (Fig. 6a). Specifically, Yamalo-Nenets Autonomous Okrug in Russia, which has the largest reserves of Russia’s natural gas and oil (Filimonova et al., 2018), was the most notable emission hotspot, which suggests gas-flaring sources. The Komi Republic in Russia was also identified as a strong anthropogenic emitter contributing to Arctic surface BC. These gas-flaring industrial regions in Russia (58–69\°N, 68–81\°E) together contributed 33\% and 31\% of Arctic surface BC for January and the annual mean, respectively. Recently, Dong et al. (2019) evaluated BC emission inventories using GEOS-Chem and proposed that using the inventory compiled by Huang et al. (2015) for Russia, in which gas flaring accounted for 36\% of anthropogenic emissions, had no prominent impact on the simulation performance in Russia or the Arctic. They suggested that the use of a new global inventory for BC emissions from natural gas flaring would improve the model performance (Huang and Fu, 2016). These results suggest that inclusion of BC emissions from gas flaring on the global scale is necessary for further BC simulations.

In Europe, a relatively high contribution of anthropogenic emissions to Arctic surface BC in January was made by Poland (50–55\°N, 15–24\°E; contributing 4\% of Arctic surface BC) because of relatively large emission fluxes in the region (Fig. 5a). Anthropogenic emissions from eastern China, especially those north of \sim 33\°N (33–43\°N, 109–126\°E), contributed perceptibly (5\%) to Arctic surface BC.

In July, contributions from anthropogenic sources shrunk to those from Yamalo-Nenets Autonomous Okrug and Komi Republic in Russia, and contributed a lower fraction (3\% of Arctic surface BC) (Fig. 6b). Few open biomass burning sources contributed in January (Fig. 6c), but contributions from open biomass burning to Arctic surface BC in July can be clearly seen, mainly from the far east of Russia, Canada, and Alaska (Fig. 6d). Open biomass burning emissions from Kazakhstan, southwest Russia, southern Siberia, and northeast China also contributed to Arctic surface BC, although at relatively low strengths (Figs. 5d and S1d). However, the contributions from open biomass burning could be higher, as the MODIS burned area, the basis of GFED emission inventories, was underestimated for northern Eurasia by 16\% (Zhu et al., 2017). Evangeliou et al. (2016) estimated a relatively high transport efficiency of BC from open biomass burning emissions to the Arctic, which led to a high contribution, i.e., 60\%, from such sources to BC deposition in the Arctic in 2010. A recent study suggested that open fires that burned in western Greenland in summer (31 July to 21 August 2017) could potentially alter the Arctic air composition and foster glacier melting (Evangeliou et al., 2019). Although the footprint of Arctic surface BC showed a relatively weak sensitivity to areas such as forests and tundra, in the boreal regions pollutants from boreal wildfires could have greater effects on the Arctic air composition in summer under future warming scenarios (Veira et al., 2016).

3.4 Sources of Arctic BC at high altitudes

Arctic BC levels at high altitudes showed the highest levels in spring (March–April, 40.5–53.9 ng m\(^{-3}\)), followed by those in late autumn to early winter (November–January, 36.5–
Figure 5. Sectorial contributions from residential combustion (including fossil fuel and biofuel combustions), gas flaring, open biomass burning and others (energy other than gas flaring, industry, and transport) to (a) seasonal variations in Arctic surface BC, (b) annual mean Arctic surface BC, (c) seasonal variations in Arctic BC at high altitudes, and (d) annual mean of Arctic BC at high altitudes.

Figure 6. Spatial distributions of contributions to Arctic BC at surface for (a) anthropogenic contributions in January 2010, (b) anthropogenic contributions in July 2010, (c) open biomass burning contributions in January 2010, and (d) open biomass burning contributions in July 2010.
40.0 ng m\(^{-3}\)), and summer (July–August, 33.0–39.0 ng m\(^{-3}\)) (Fig. 4c). The annual mean Arctic BC at high altitudes was estimated to be 35.2 ng m\(^{-3}\), which is ca. 73 % of those at the surface. Such a vertical profile is in accordance with those based on aircraft measurements over the Canadian High Arctic (Schulz et al., 2019). Similarly to the case for the surface, anthropogenic sources dominated by the residential, transport, industry, and energy sectors (excluding gas flaring) accounted for 94 %–100 % of Arctic BC at high altitudes in October–May (Figs. 4c and 5c). East Asia accounted for 34 %–65 % of the total BC in October–May (40 % annually).

In comparison, using the Community Atmosphere Model version 5 driven by the NASA Modern Era Retrospective-Analysis for Research and Applications (MERRA) reanalysis data and the IPCC AR5 year 2000 BC emission inventory, H. Wang et al. (2014) found that East Asia accounted for 23 % of BC burden in the Arctic for 1995–2005. In summer, open biomass burning in the boreal regions accounted for 40 %–72 % (57 % on average) of Arctic BC at high altitudes, similar to the source contributions to Arctic surface BC. Specifically, open biomass burning sources from Siberia accounted for 40 %–42 % of Arctic BC at high altitudes in July–August. Annually, anthropogenic sources and open biomass burning accounted for 83 % (in which residential sources accounted for 34 %) and 17 %, respectively, of total Arctic BC at high altitudes (Figs. 4d and 5d).

Further investigations of geospatial contributions to Arctic BC at high altitudes in January and July provided more details regarding BC sources. In January, the main anthropogenic BC source in East Asia covered a wide range in China (Fig. 7a). Not only east and northeast China but also southwest China (Sichuan and Guizhou provinces) were the major anthropogenic sources of Arctic BC at high altitudes.

In July, anthropogenic sources made a relatively weak contribution to Arctic BC at high altitudes. The regions that were sources of open biomass burning contributions to Arctic BC at high altitudes were mainly the far east of Siberia, Kazakhstan, central Canada, and Alaska, i.e., similar to the sources of Arctic surface BC. Unlike Arctic surface BC, for which the dominant source regions are at high latitudes in both winter and summer, Arctic BC at high altitudes mainly originates from mid-latitude regions (Figs. 6 and 7). In terms of transport pathways, air masses could be uplifted at low-to-mid latitudes and transported to the Arctic (Stohl, 2006).

Further investigations are needed to obtain more details of the transport processes.

3.5 Comparison of FLEXPART and GEOS-Chem simulations of BC sources

Data for BC sources simulated with FLEXPART were compared with those obtained with GEOS-Chem (Ikeda et al.,
of BC source contributions indicates improved reliability of evaluated source contributions to Arctic BC.

4 Conclusions

The source contributions to Arctic BC were investigated by using a FLEXPART (version 10.1) transport model that incorporated emission inventories. FLEXPART-simulated BC data agreed well with observations at Arctic sites, i.e., Utqiaġvik, Alert, Zeppelin, and Tiksi. The source regions and source sectors of BC at the surface and high altitudes over a wide region in the Arctic north of 66° N were simulated. BC at the Arctic surface was sensitive to local emissions and those from nearby Nordic countries (> 60° N). These results emphasize the role of anthropogenic emissions such as gas flaring and development of the Northern Sea Route in affecting air quality and climate change in the Arctic. Anthropogenic emissions in the northern regions of Russia were the main source (56 %) of Arctic surface BC annually. In contrast, BC in the Arctic at high altitudes was sensitive to mid-latitude emissions (30–60° N). Although they are geospatially far from the Arctic, anthropogenic emissions in East Asia made a notable (40 %) contribution to BC in the Arctic at high altitudes annually. Open biomass burning emissions, which were mainly from Siberia, Alaska, and Canada, were important in summer, contributing 56 %–85 % of BC at the Arctic surface and 40 %–72 % at Arctic high altitudes. Future increases in wildfires as a result of global warming could therefore increase the air pollution level during the Arctic summer. This study clarifies the source regions and sectors of BC in the Arctic. This information is fundamental for understanding and tackling air pollution and climate change in the region.

Data availability. The data set for simulated footprint and BC source contributions is available upon request to the corresponding author.

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Author contributions. CZ and YK designed the study. CZ, MT, and IP optimized the FLEXPART model. CZ performed FLEXPART model simulations, conducted analyses, and wrote the paper. KI and HT provided data for GEOS-Chem simulations and site observations. All authors made comments that improved the paper.

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