

Response to minor comment on “Concentrations and source regions of light absorbing particles in snow/ice in northern Pakistan and their impact on snow albedo” by Chaman Gul et al.

We thank the Co-Editor and reviewers for their comments. Please see responses to each of the comments below, coded as follows:

Reviewer comments in black

Responses in blue

Modified text in revised manuscript in green

The authors have provided a comprehensive response to concerns raised by the reviewers on the first submission of this manuscript. Overall, I think the authors have adequately addressed these concerns, though the following issues should be addressed prior to publication:

1. The paper needs to be carefully and thoroughly edited for grammar and clarity. Reading only the quoted passages of edited text in the response to reviewer’s document, I see numerous instances of incomplete sentences, incorrect grammar, and unclear descriptions.

The present manuscript has been thoroughly edited by a native English speaker editor; this is acknowledged in the Acknowledgements section.

2. One question I initially raised was whether the reported albedo reductions are absolute or relative. The authors explained their calculations and describe them as "relative" changes, but I think they are actually "absolute" changes in the context I was inquiring about. I wasn't completely clear on the distinction, however, so I will clarify here what I meant. Let's say two albedo measurements are 0.70 and 0.80. The absolute difference between these could be reported as 0.10 or 10%. The relative change could be reported as: $(0.80-0.70)/0.70 = 14\%$. Hence when I see "albedo change" reported with a "%", I don't know if it is the absolute change or relative change. I believe the authors are reporting absolute change and if so they should clarify this. Simply reporting the change in absolute albedo units (e.g., +0.10) usually negates this problem and is what I usually advise doing.

Thank you for the clarification. We have used the absolute difference in the manuscript and

clarified this with a sentence as follows (line 370 - 371).

“The percentage change in albedo was calculated in absolute terms as the change between albedo values with a pollutant (BC or dust or both) and a reference albedo value with zero pollutants (zero BC and dust concentration).”

3. With regard to the possibility of dust contamination of BC/OC measurements (an issue raised in my first review), the following addition to the paper is quoted several times: "According to our understanding the analysis method and amount of dust loading on the sample can also alter OC/BC ratios." I am glad to see the authors acknowledge this potential source of contamination, but I suggest they elaborate on this with a couple of additional sentences, for the benefit of readers. Specifically, I suggest explaining why dust can result in over-estimations of BC/OC burdens, citing one or papers to this effect, and if possible mentioning the potential magnitudes of overestimation that may occur.

We have added some sentences in the revised manuscript describing the impact of dust on OC/BC analysis as shown below (lines 303–314).

“The method used for analysis and the amount of dust loading on the sample can also affect the OC/BC ratio, as can the presence of metal oxides and calcium carbonate. High iron oxide concentrations can cause BC to pre-oxidize or drop off the filter, while calcium carbonate can be wrongly identified as BC. Laboratory studies have shown that the presence of metal oxides in aerosol samples can alter the OC/BC ratio either by enhancing OC charring or by lowering the BC oxidation temperature (Wang et al., 2010), while higher fractions of metal oxide can increase BC divergence across the thermal optical protocols (Wu et al., 2016). Dust can lead to a greater decrease in optical reflectance during the 250°C heating stage in the thermal/optical method, and thus an incorrect OC/BC ratio (Wang et al., 2012). Carbon detected by the flame ionization detector (FID) before the optical signal attains the initial value is defined as OC and that detected after is defined as BC; dust on the filter results in the FID division being postponed or inefficient, and thus OC being overestimated and BC underestimated or even negative (Wang et al., 2012). Wang et al. (2012) provides a more detailed discussion of OC/BC ratios derived using the thermal optical method.”

4. The following sentence was added to the revised manuscript (quoted in the response to

reviewers): "Warming role of OC was also not included in the current research, which was low but significant in several regions (Yasunari et al. 2015)." It is not clear to me what this means, so I request that this be clarified. Is it meant that albedo reduction from OC was not quantified here?

Yes, the purpose of the sentence was to show that the albedo reduction from organic carbon was not quantified in the present research. The sentence has been modified as shown below (line 624–627).

“The albedo reduction from OC was not quantified. The contribution of OC to total visible absorption in the top snow layer is relatively small compared to that of BC and dust but has been shown to be significant (~19% of the total solar visible absorption) in several regions including northeastern East Asia, and western Canada (Yasunari et al., 2015).”

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Concentrations and source regions of light absorbing particles in snow/ice in northern Pakistan and their impact on snow albedo

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Abstract. Black carbon (BC), water-insoluble organic carbon (OC), and mineral dust are important particles in snow and ice, which significantly reduce albedo and accelerate melting. Surface snow and ice samples were collected from the Karakoram-Himalayan region of northern Pakistan during 2015 and 2016 in summer (six glaciers), autumn (two glaciers), and winter (six mountain valleys). The average BC concentration overall was 2130 ± 1560 ng g⁻¹ in summer samples, 2883 ± 3439 ng g⁻¹ in autumn samples, and 992 ± 883 ng g⁻¹ in winter samples. The average water insoluble OC concentration overall was 1839 ± 1108 ng g⁻¹ in summer samples, 1423 ± 208 ng g⁻¹ in autumn samples, and 1342 ± 672 ng g⁻¹ in winter samples. The overall concentration of BC, OC, and dust in aged snow samples collected during the summer campaign was higher than the concentration in ice samples. The values are relatively high compared to reports by others for the Himalayas and Tibetan Plateau. This is probably the result of taking more representative samples at lower elevation where deposition is higher and the effects of ageing and enrichment more marked. A reduction in snow albedo of 0.1–8.3% for fresh snow and 0.9–32.5% for aged snow was calculated for selected solar zenith angles during daytime using the Snow, Ice, and Aerosol Radiation (SNICAR) model. Daily mean albedo was reduced by 0.07–12.0%. The calculated radiative forcing ranged from 0.16 to 43.45 Wm⁻² depending on snow type, solar zenith angle, and location. The potential source regions of the deposited pollutants were identified using spatial variance in wind vector maps, emission inventories coupled with backward air trajectories, and simple region tagged chemical transport modelling. Central, South, and West Asia were the major sources of pollutants during the sampling months, with only a small contribution from East Asia. Analysis based on the Weather Research and Forecasting (WRF-STEM) chemical transport model identified a significant contribution (more than 70%) from South Asia at selected sites. Research into the presence and effect of pollutants in the glaciated areas of Pakistan is economically significant because the surface water resources in the country mainly depend on the rivers (the Indus and its tributaries) that flow from this glaciated area.

35 **1 Introduction**

36 Carbon is an essential component of atmospheric aerosols, where it appears in the form of black carbon (BC,
37 also known as elemental carbon, EC), and organic carbon (OC). BC is emitted into the atmosphere from incomplete
38 combustion of carbon-based fuels (mainly fossil fuels and biomass) (Jacobson, 2004), while OC can be directly
39 emitted into or formed in the atmosphere. After deposition on snow and ice surfaces, BC particles significantly
40 reduce the snow albedo (hemispheric reflectance) in the visible part of the electromagnetic spectrum, cause snow
41 albedo feedback (Doherty et al., 2013), enhance solar radiation absorption (Warren and Wiscombe, 1980), and
42 accelerate snow melting (Hansen and Nazarenko, 2004). BC, both in air and deposited on snow, is important in net
43 positive forcing of the climate. Clean snow is one of the most reflective natural surfaces on Earth at the ultraviolet
44 and visible wavelengths, while BC is the most efficient light-absorbing species in the visible spectral range
45 (Horvarth, 1993). One ng g^{-1} of BC has almost the same effect on albedo reduction as 100 ng g^{-1} mineral dust at 500
46 nm wavelength (Warren et al., 1982). However, the exact amount of albedo reduction also depends on the refractive
47 index, grain size, solar zenith angle (SZA), snow density, dust particle size and concentration, particle morphology,
48 surface roughness, snow depth, liquid water content, snow shape, and topography (Wiscombe and Warren 1985).
49 Albedo reduction usually results in amplification of the energy absorbed by dirty snow (Painter et al., 2010). An
50 albedo feedback is triggered and amplified by deposition of impurities on the snow surface which reduces snow
51 albedo thus accelerating melting and further reducing albedo (Doherty et al., 2013; Flanner et al., 2009). Albedo
52 feedback is amplified by the presence of light-absorbing particles (Doherty et al., 2013). Studies conducted in
53 Greenland showed that at visible wavelengths 10 ng g^{-1} coarse-grained BC particles in aged snow and 40 ng g^{-1} BC
54 particles in new snow can reduce snow albedo by around 1 to 3% (Warren and Wiscombe, 1985).

55 Increased BC mass concentration and deposition on the Tibetan glaciers over the last 20 years (Xu et al., 2009a)
56 have played a significant role in rapid glacier melting in the region (Xu et al., 2012; Yao et al., 2012). A high
57 concentration of aerosol has deposited on the snow surface and increased the BC content in snow over the southern
58 edge of the Tibetan Plateau to the north of the Himalayas (Gertler et al., 2016). The southern slope of the Himalayas
59 is relatively even more exposed to BC due to emissions from India and transport through southwesterly and westerly
60 winds (Xu et al., 2009; Yasunari et al., 2010). BC deposited on snow in the Himalayan region induces an increase in
61 net shortwave radiation at the snow surface with an annual mean of about $1 \text{ to } 3 \text{ Wm}^{-2}$, producing an estimated 0.05--
62 0.3°C warming (Ménégoz et al., 2014). Deposition of anthropogenic BC has been observed to contribute
63 significantly to the decrease in snow cover extent over recent decades (Dery et al., 2007), and shortening of the
64 duration of the snow cover season by several days (Ménégoz et al., 2013a). The climate warming efficiency of BC in
65 snow is greater than the warming efficiency of other anthropogenic pollutants, including carbon dioxide (Hansen et
66 al., 2005). Another important characteristic of BC is its higher snowmelt efficiency. The snowmelt efficiency of BC
67 in terms of snow cover fraction and snow water equivalent is larger than that induced by increase in carbon dioxide
68 (Qian et al., 2011). The annual snow albedo reduction effect due to BC outweighs the aerosol dimming effect
69 (reduction in solar radiation reaching the surface) by a factor of about six over the global snow cover (Flanner et al.,

70 2009).

71 At present, South and East Asia are considered to be the two largest BC emission regions in the world and likely
72 to remain so (Menon et al., 2010). BC transported from East Asia can be lifted high and moved towards the
73 northeast during the summer monsoon season (Zhang et al., 2015; Cong et al., 2015; Lüthi et al., 2015), affecting the
74 life of glaciers and snow-covered areas.

75 Research into the glaciers of the extended Himalayan region and Tibetan Plateau has prime importance because
76 these glaciers act as a water storage tower for South and East Asia, and shrinking could affect water resources for up
77 to a billion people (Immerzeel et al., 2010). The glaciated area in northern Pakistan may be more exposed to BC
78 effects than that in other regions because potentially it can receive emissions generated from both South and Central
79 Asia as well as from the Middle East. Meltwater coming from these glaciers flows into the river Indus, which has
80 major economic importance for the people of Pakistan.

81 A number of authors have described the concentration and impacts of light absorbing particles in the Tibetan
82 glaciers (for example Qian et al., 2015; Wang et al., 2015; Zhang et al., 2017; Li et al., 2017; Niu et al., 2017).
83 However, until now, no studies have been published related to the concentration of light absorbing aerosols in the
84 surface snow and ice of northern Pakistan, and although several authors have investigated transport pathways over
85 the Himalayan region (e.g. Babu et al., 2011 for the western trans-Himalayas; Lu et al., 2012 for the Tibetan Plateau
86 and Himalayas) little is known about the potential sources and transport pathways of pollutants affecting the
87 Pakistan area.

88 In this study, we looked at the concentration of light absorbing particles (BC, OC, dust) in snow and ice in
89 northern Pakistan, their impact on snow albedo and radiative forcing, and the likely source regions. Albedo was
90 estimated from the BC and dust concentrations identified in collected samples of snow and ice using the online snow
91 albedo simulation SNICAR model (Flanner et al., 2009). Radiative forcing was calculated from the albedo reduction
92 obtained from the SNICAR model together with the incident short-wave solar radiation obtained from the SBDART
93 (Santa Barbara DISORT Atmospheric Radiative Transfer) model. The frequency distribution of aerosol subtypes
94 (smoke, polluted continental, dust, and others) in the atmosphere over the study area was calculated for the snow and
95 ice sampling periods using CALIPSO (Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observations) satellite
96 data from 2006 to 2014 as a further indication of the types of aerosol contributing to the observed deposition. The
97 potential source regions of pollutants were identified using spatial variance in wind vector maps prepared using
98 MERRA-2 reanalysis data, calculation of back air trajectories using the HYSPLIT-4 (Hybrid Single Particle
99 Lagrangian Integrated Trajectory) model, and a simple region tagged chemical transport model (WRF-STEM). The
100 back air trajectories approach has been used in many studies to identify possible source regions for atmospheric and
101 deposited BC (Zhang et al., 2013). Pollutant source regions identified using the different approaches were compared
102 and the most likely source regions of the pollutants identified.

103 **2 Methodology**

104 **2.1 Study area**

105 The study area was located around 35.40°N 74.38°E in the mountains and adjacent mountain valleys of the
106 Karakoram and Himalayan region in northern Pakistan (Figure 1). Snow and ice samples were collected in summer
107 from six glaciers – Passu, Gulkin, Barpu, Mear, Sachin, and Henarche – and in autumn from Gulkin and Sachin
108 (Figure 1). The Passu and Gulkin glaciers are located very near to the Karakoram highway connecting Pakistan with
109 China, and there are a number of small villages (Passu, Hussaini, Gulmit, and others) close by. The Barpu and Mear
110 glaciers are located very close to each other and around 3 km away from the residential area of the Hopar and Nagar
111 valleys. There is a small city (Astore) near the Sachin glacier and some restaurants near its terminus. Winter snow
112 samples were collected from mountain valleys near the Passu, Barpu, and Sachin glaciers, and three other areas to
113 the west with a number of small villages (Figure 1). The average elevation of the selected glaciers was quite low
114 compared to the elevation of the glaciers studied for BC, OC, and dust on the Tibetan plateau by previous
115 researchers. The mountains around the selected glaciers are mostly dry and rocky. The 10-year records (1999–2008)
116 of the two nearby climate stations at Khunjerab (36.83°N, 75.40°E, 4,730 masl) and Naltar (36.29°N, 74.12°E,
117 2,858 masl) show mean total annual precipitation values of 170 mm and 680 mm, respectively. The daily average
118 temperature during winter and pre-monsoon showed an increasing trend between 1980 and 2014 (Gul et al., 2017).
119 The study area is mostly exposed to the westerlies and emissions from South Asia. Most of the people in the region
120 use wood for cooking and heating.

121 **2.2 Sample collection**

122 A total of 50 surface ice and 49 snow samples were collected from the glaciers in summer 2015 and 2016 (Passu
123 15, Gulkin 31, Barpu 6, Mear 8, Sachin 35, Henarche 4), and 13 in autumn 2016 (Gulkin 7, Sachin 6) at elevations
124 ranging from 2,569 to 3,895 masl (Figure 1). Eighteen snow samples were collected in winter 2015 and 2016 from
125 nearby mountain valleys at elevations of 1,958 to 2,698 masl; the winter sampling region was divided into six sites
126 (S1 to S6) based on geographical location and elevation (Figure 1). Samples were collected using the “clean hands –
127 dirty hands” principle (Fitzgerald, 1999). Ice samples were collected from the surface (5 cm depth) at different
128 points on the glaciers. The elevation difference between collection points on the same glacier ranged from 30 to 100
129 meters.

130 The samples were preserved in ultra clean plastic bags, allowed to melt in a temporary laboratory near the
131 sampling location, and filtered through quartz-filters immediately after melting. An electric vacuum pump was used
132 to accelerate filtration. The melted snow/ice volume of the samples was measured using a graduated cylinder.
133 Sampled filters were carefully packed inside petri-slides marked with a unique code representing the sample.

134 The snow density of winter snow samples was measured using a balance; snow/ice grain sizes were observed
135 with an accuracy of 0.02 mm using a hand lens (25×) (Aoki et al., 2011); and snow shape was estimated using a
136 snow card. In the models, we assumed external mixing of snow and aerosol particles and spherical snow grains.

137 Snow grain size and snow texture were the largest sources of uncertainty in albedo reduction (Section 3.3). Qian et
138 al. (2015) have summarized the sampling methods available for light absorbing particles in snow and ice from
139 different regions including the Arctic, Tibetan Plateau, and mid-latitude.

140 **2.3 Dust, OC, and BC analysis**

141 Before analysis, sampled filters were allowed to dry in an oven for 24 hours and then weighed using a
142 microbalance. The dust mass on the filters was calculated from the mass difference in weight before and after
143 sampling (Kaspari et al., 2014; Li et al., 2017).

144 There are many methods available for analyzing BC and OC. The three methods considered most effective for
145 measuring BC and water insoluble OC concentrations in snow are thermal optical analysis, filter-based analysis, and
146 single particle soot photometer analysis (Ming et al., 2008). The thermal optical (filter-based) analysis method has
147 been used by many researchers (e.g., Li et al., 2017) and was chosen for the study. This is an indirect method for
148 measuring BC and OC on sampled filters; it follows Beer's law and uses stepwise combustion of the particles
149 deposited on quartz filters (Boparai et al., 2008), followed by measurement of light transmission and/or reflectance
150 of the filters. The BC and OC content in the collected samples was measured using a thermal optical DRI carbon
151 analyzer, similar to the IMPROVE protocol (Cao et al., 2003). The temperature threshold applied to separate the two
152 species is described in Wang et al. (2012). A few (<10) filters had higher dust loads; for these the method was
153 slightly modified using a 100% helium atmosphere and temperature plateau (550°C). A very few (<5) samples with
154 very dense dust concentrations were not properly analyzed by the instrument and were excluded from the results.
155 The extremely high dust value of one sample from Passu (15 times the level in the next highest sample) which had
156 low values of other pollutants was excluded as a probable error. In some cases, a single sample was analyzed two or
157 three times to ensure accurate results were obtained.

158 The CALIPSO models also define multiple aerosol sub-types – clean continental, clean marine, dust, polluted
159 continental, polluted dust, smoke, and other – using the 532-nm (1064-nm) extinction-to-backscatter ratio. The
160 frequency of these different aerosol subtypes in the atmosphere over the study region was investigated using
161 CALIPSO data for the same months in which ice and snow samples were collected, i.e. January, May, June, and
162 December, over the period June 2006 to December 2014. The CALIPSO Level 2 lidar vertical feature mask data
163 product describes the vertical and horizontal distribution of clouds and aerosol layers (downloaded from
164 <https://eosweb.larc.nasa.gov/project/calipso/aerosol_profile_table>). The aerosol subtypes were classified in the
165 downloaded data using the observed backscatter strength and depolarization values. The details of the algorithm
166 used for classification are given in Ali H. Omar et al. (2009). The percentage contributions of individual aerosol
167 subtypes were plotted using MATLAB (MathWorks, Inc.).

168 The frequencies of different subtypes were calculated along the specific paths followed by CALIPSO over the
169 study region.

170 **2.4 Albedo simulations and estimation of radiative forcing**

171 Snow albedo was estimated for each of the 18 winter samples and the average calculated for samples at each of
172 the sites (1 to 6). Albedo from two sites – S1 (Sost), which had the highest average concentration of BC and dust, and
173 S6 (Kalam), which had the lowest average concentration of BC and dust – were further explored using the SNICAR
174 model (Flanner et al., 2007). The aim was to quantify the effect of BC, dust, and mass absorption cross-section (MAC)
175 on albedo reduction. Sensitivity model experiments were carried out using various combinations of BC, dust, and
176 MAC values, while other parameters were kept constant (parameters for sites 1 and 6 shown in supplementary
177 materials, Table S1). Snow albedo was simulated for different daylight times, with the SZA set in the range 57.0–88.9°
178 based on the position of the sun in the sky for the sampling date and locations. The daily mean was calculated from
179 the mean of the albedo values simulated for 24 different SZA values (one per hour), and the daytime mean from the
180 mean of the albedo values simulated for 10 SZA values (one per hour during daylight). The mid-latitude winter clear-
181 sky option was selected for surface spectral distribution. The parameters used for sensitivity analysis are shown in
182 Table S1. MAC values of 7.5, 11, and 15 m²/g were selected based on a literature review (Que et al., 2014; Pandolfi
183 et al., 2014). In order to reduce the uncertainty, the dust concentration in the samples was divided into four diameter
184 classes (as per the model requirements): size 1 (0.1–1.0 μm) was taken to be 2%, size 2 (1–2.5 μm) to be 13%, size 3
185 (2.5–5 μm) to be 31%, and size 4 (5–10 μm) to be 54% of total dust mass present in the sample, based on results
186 published by others (Gillette et al., 1974; Mahowald et al., 2014). Radiative forcing (RF) was estimated for the same
187 samples following Eq. (1):

$$188 \quad RF_x = R_{in-short} * \Delta \alpha_x \quad (1)$$

189 where $R_{in-short}$ denotes incident short-wave solar radiation (daily mean), as measured by the SBDART (Santa Barbara
190 DISORT Atmospheric Radiative Transfer) model, and $\Delta \alpha_x$ denotes the daily mean reduction in albedo, as simulated
191 by the SNICAR model.

192 **2.5 Source regions of pollutants**

193 Three methods were used to identify the potential source regions of pollutants found at the study site: wind
194 maps, emissions inventory coupled with back trajectories, and a region-tagged chemical transport modeling analysis.

195 Wind vector maps were prepared using MERRA-2 reanalysis data (available from the National Aeronautics and
196 Space Administration [NASA] <https://gmao.gsfc.nasa.gov/reanalysis/MERRA-2/docs/>). The U and V wind
197 components were combined into a matrix around the study area for each individual month and then plotted against
198 latitude/longitude values to show the spatial variance of monthly wind stress at 850 mb using arrows to indicate the
199 direction and intensity of wind.

200 Air trajectories were calculated backwards from the sampling sites (S1: 36.40°N 74.50°E; S6: 35.46°N 72.54°E)
201 to identify potential source regions for the pollutants using the web version of the Hybrid Single Particle Lagrangian
202 Integrated Trajectory (HYSPLIT-4) model (Draxler and Hess, 1998). The HYSPLIT-4 model has been used by others
203 to compute air mass trajectories to identify possible source regions (Ming et al., 2009; Zhang et al., 2013). Reanalysis

204 meteorological data from the same source as the wind data (<https://www.esrl.noaa.gov/psd/data>) were used as input
205 data in the HYSPLIT model for May, June, and December 2015, and January 2016. HYSPLIT was run in a seven-day
206 backward trajectory mode with trajectories initiating every six hours (0, 6, 12, and 18) on a daily basis from 4 May to
207 19 June 2015 (77 days during summer) and from 1 December 2015 to 31 January 2016 (62 during winter). The
208 HYSPLIT model results were combined with Representative Concentration Pathways (RCPs) emission data for 2010
209 (available from http://sedac.ipcc-data.org/ddc/ar5_scenario_process/RCPs.html; data file <RCPs_anthro_BC_2005-
210 2100_95371.nc>) to identify the source location. This comprises emission pathways starting from an identical base
211 year (2000) for multiple pollutants, including BC and OC; the file description indicates that the inventory includes
212 biomass burning sources. The RCP inventory has the same emissions sectors as the Hemispheric Transport Air
213 Pollution (HTAP) emission inventory used in the modelling approach for identifying source regions (see below),
214 including fuel combustion, industry, agriculture, and livestock, but the HTAP inventory has a higher resolution (0.1 x
215 0.1 degree) than the RCP inventory (0.5 x 0.5 degree). Lamarque et al. (2010) give a more detailed discussion of the
216 inventory and sectors (12) used in the base year calibration of the RCP. Monthly CALIPSO satellite-based extinction
217 data from 2006 to 2014 were used to calculate the vertical profile for aerosol extinction over the study region. The
218 CALIPSO extinction profile was constructed for selected months in 2006 to 2014 – May and June for summer and
219 December and January for winter (Figure S1). The exponential equation $X = (\log(10.46) - \log(Y))/10.29$, where
220 Y is the vertical height of individual trajectories in kilometers and X indicates the extinction against the height of
221 trajectories, was used to calculate the extinction profile for the trajectory heights. The normalized extinction profile
222 was obtained by assuming that surface extinction = 1 (Figure S1).

223 The WRF-STEM model was used as a third approach for identifying the origin (source regions) of air masses
224 carrying pollutants. The WRF-STEM model uses region-tagged carbon monoxide (CO) tracers for many regions in
225 the world to identify geographical areas contributing to observed pollutants (Adhikary et al., 2010). Region tagged
226 CO tracers are used as a standard air quality modeling tool in various regional and global chemical transport models
227 to identify pollution source regions (Chen et al., 2009; Park et al., 2009; Lamarque and Hess, 2003). The WRF-STEM
228 model domain was centered on 50.377° E longitude and 29.917° N latitude, with a model horizontal grid resolution
229 of 45 x 45 km with 200 grids in the east-west direction and 125 north-south. The meteorological variables needed for
230 the chemical transport were derived from the Weather Research and Forecast (WRF) meteorological model (Grell et
231 al., 2005) using FNL data (ds083.2) available from the UCAR website as input data. The main aim of the simulation
232 was to identify the geographic locations contributing to the observed pollutants at the field sites. The HTAP version 2
233 emission inventory, which comprises multiple pollutants including BC and OC, was used for the WRF-STEM
234 modeling (available from <http://edgar.jrc.ec.europa.eu/htap_v2/>). This emission inventory includes major sectors
235 such as energy, industry, transport, and residential, but not large scale open agricultural and open forest burning. The
236 simulations applied in our study used the anthropogenic emissions from the HTAP inventory. Thus the results indicate
237 the amount of pollutants reaching the study area from day-to-day planned and recurring activities in domestic,
238 transport, industrial, and other sectors. The model was run for a month prior to the field campaign dates to allow for

239 model spin up (normal practice for a regional chemical transport model), and then for the months of December, January,
240 and June, to match the field campaign dates.

241 **3. Results and discussion**

242 **3.1 BC, OC and dust concentrations**

243 The minimum, maximum, and average concentrations of BC, OC (water insoluble organic carbon), and dust in
244 the ice and snow samples are given in Table 1. The OC, and BC concentration values were blank corrected by
245 subtracting the average value of the field blanks. Blank concentrations were used to calculate detection limits as
246 mean \pm standard deviation. The average BC concentration overall was 2130 ± 1560 ng g⁻¹ in summer samples, 2883
247 ± 3439 ng g⁻¹ in autumn samples (both from glaciers), and 992 ± 883 ng g⁻¹ in winter samples. The average OC
248 concentration overall was 1839 ± 1108 ng g⁻¹ in summer samples, 1423 ± 208 ng g⁻¹ in autumn samples, and $1342 \pm$
249 672 ng g⁻¹ in winter samples. There was considerable variation in individual samples, with summer values of BC
250 ranging from 82 ng g⁻¹ (Gulkin glacier) to $10,502$ ng g⁻¹ (Henarche glacier), autumn values from 125 ng g⁻¹ (Gulkin
251 glacier) to 6481 ng g⁻¹ (Sachin glacier), and winter samples from 79 ng g⁻¹ (Kalam) to 5957 ng g⁻¹ (Sost).

252 The lowest BC (82 ng g⁻¹) and OC (128 ng g⁻¹) concentrations were observed in summer samples collected from
253 the Gulkin and Sachin glaciers, respectively. The average values of BC and OC were low in all samples from the
254 Passu glacier, even though it lies close to the Karakoram highway which links Pakistan with China. The low
255 concentrations of BC may have been due to the east facing aspect of the glacier shielding it from pollutants
256 transported from west to east. Slope aspect of a glacier is known to be important for snow cover dynamics (Gul et
257 al., 2017); dust concentrations are known to vary with slope aspect due to the effects of wind direction on
258 deposition.

259 The highest average concentration of BC was found in autumn samples from the Sachin glacier, and the highest
260 average concentration of OC in summer samples from the same glacier. The average concentration of BC was much
261 greater in autumn than in summer on the Sachin glacier, but somewhat greater in summer than in autumn on the
262 Gulkin glacier, indicating highly spatiotemporal patterns in the deposition of particles. The marked difference on the
263 Sachin glacier may have reflected the difference in the direction of air, which comes from Iran and Afghanistan in
264 summer and the Bay of Bengal via India in autumn, with the generally lower deposition on the Gulkin glacier more
265 affected by other factors (such as slope aspect of the glacier and status of local emissions near the glacier). There
266 was no clear correlation between the average BC concentration in glacier samples and glacier elevation. However,
267 winter snow samples showed a weak increasing trend in average BC with site elevation (Table 1, Figure S3).

268 Most summer samples were collected from surface ice (Figure S2 a), but a few samples for Gulkin and Sachin
269 were collected from aged snow on the glacier surface (Figure S2 b,c). Dust was visible on the relatively aged snow,
270 and the BC and OC concentrations in these snow samples were much higher than those in ice. The highest average
271 BC values in winter were also observed in aged snow (from Sost) and the lowest in fresh snow (from Kalam) (Table

272 1). Generally, snow samples collected within 24 hours of a snowfall event were considered as a fresh snow.

273 We analyzed the ratios of OC to BC in the different samples as in atmospheric fractions this can be used as an
274 indicator of the emission source, although apportionment is not simple and only indicative. The BC fraction is
275 emitted during combustion of fossil fuels, especially biomass burning in rural areas in winter, and urban emissions
276 from road transport. The OC fraction can be directly emitted to the atmosphere as particulate matter (primary OC)
277 from fossil fuel emissions, biomass burning, or in the form of biological particles or plant debris; it can also be
278 generated in the atmosphere as gases are converted to particles (secondary OC). In general, lower OC/BC ratios are
279 associated with fossil fuel emissions and higher OC/BC ratios with biomass burning. Overall, there was no clear
280 correlation between BC and OC concentrations in our samples. In most cases, the concentration of OC was greater
281 than the concentration of BC, which might indicate a greater contribution from biomass burning in the emissions,
282 but in a few the concentration of BC was greater than that of OC, which might indicate a greater contribution from
283 coal combustion. The lowest OC/BC ratio of 0.041 was observed in a summer sample from Henarche glacier, and
284 the highest ratio of 5 in a winter sample from Kalam. The higher value at Kalam may indicate greater contributions
285 from biomass burning than from fossil fuel combustion in the region. In summer samples, the average concentration
286 of OC was greater than the average concentration of BC in samples from four of the six glaciers, but it was much
287 lower in Barpu and Henarche. In winter, individual snow samples indicated that concentration of OC was greater
288 than BC at low elevation sites and vice versa; the average OC was greater than average BC at all except the highest
289 elevation site (Table 1).

290 However, these results should be considered with care. There are a number of factors that can affect the OC/BC
291 ratio in snow and ice samples apart from the concentrations in the atmosphere. Spatio-temporal variability of the
292 OC/BC ratio may indicate the contribution of different sources, seasonal variation, and frequent change in wind
293 direction. In deposited samples, low OC/BC ratios can result from a reduction in OC (Niu et al., 2017), greater
294 contributions from BC enrichment and OC scavenging, and/or the contribution of different emission sectors
295 (including quantity, combustion conditions, and fuel type). Post deposition processes of scavenging and enrichment,
296 which are influenced by snow melt rate, can cause water soluble OC to be under-represented as meltwater removes
297 OC but not BC, with OC and BC being redistributed primarily by meltwater rather than by sublimation and/or
298 dry/wet deposition. Thus the OC/BC ratio often reflects the impact of dilution of dissolved organic carbon and
299 enrichment of primary organic carbon during snow/ice melting, with differences in OC/BC ratios reflecting
300 differences in the enrichment process. The low OC/BC ratio in the samples from Henarche, the glacier at the lowest
301 elevation, could, for example, be due to preferential washing out of OC particles with meltwater. Overall, there was
302 a higher positive correlation between BC and dust than with OC, suggesting that for BC and dust, particle
303 precipitation and enrichment processes were similar. **The method used for analysis and the amount of dust loading
304 on the sample can also affect the OC/BC ratio, as can the presence of metal oxides and calcium carbonate. High iron
305 oxide concentrations can cause BC to pre-oxidize or drop off the filter, while calcium carbonate can be wrongly
306 identified as BC. Laboratory studies have shown that the presence of metal oxides in aerosol samples can alter the**

307 OC/BC ratio either by enhancing OC charring or by lowering the BC oxidation temperature (Wang et al., 2010),
308 while higher fractions of metal oxide can increase BC divergence across the thermal optical protocols (Wu et al.,
309 2016). Dust can lead to a greater decrease in optical reflectance during the 250°C heating stage in the thermal/optical
310 method, and thus an incorrect OC/BC ratio (Wang et al., 2012). Carbon detected by the flame ionization detector
311 (FID) before the optical signal attains the initial value is defined as OC and that detected after is defined as BC; dust
312 on the filter results in the FID division being postponed or inefficient, and thus OC being overestimated and BC
313 underestimated or even negative (Wang et al., 2012). Wang et al. (2012) provides a more detailed discussion of
314 OC/BC ratios derived using the thermal optical method.

315 A wide range of values has been reported by different authors for BC concentrations in snow and ice samples from
316 different regions (Table S2). The concentrations of BC in our samples were higher than those reported by many authors
317 (Table S2), but were comparable with the results reported by Xu et al. (2012) in the Tien Shan Mountains, Li et al.
318 (2016) in the northeast of the Tibetan plateau, Wang et al. (2016) in northern China, and Zhang et al. (2017) in western
319 Tien Shan, Central Asia. High concentrations indicate high deposition rates on the snow and ice surface, but there are
320 several possible reasons for a wide variation in values apart from differences in deposition rates, including differences
321 in sampling protocols, geographical/sampling location and elevation of sampling site (Qu et al., 2014), and year/season
322 of sampling. The majority of samples were from the ablation zone of the glaciers. Strong melting of surface snow and
323 ice in the ablation zone could lead to BC enrichment and high BC concentrations, as observed by Li et al. (2017) for
324 glaciers on the Southern Tibetan Plateau. The sampling season (May to September in our study) is an important factor
325 because rapid enrichment occurs as snow melts during the melting season. The peak melting period is May to
326 August/September, thus the concentration of BC, OC, and dust in our samples would have been increased as melting
327 progressed due to the enrichment in melting snow and scavenging by the melting water. In most cases, snow and ice
328 samples were collected quite a long time after snow fall, and the concentration of pollutants would also have increased
329 in the surface snow and ice due to dry deposition. It seems likely that the pollutants in surface samples would be
330 affected by sublimation and deposition until the next melt season (Yang et al., 2015). In some of the cases in our study,
331 the average concentration of BC, OC, and/or dust for a particular glacier/site was increased as a result of a single
332 highly concentrated sample, reflecting the wide variation that results from the interplay of many factors.

333 Enrichment is more marked at lower elevations as the temperatures are higher which enhances melting and
334 ageing of surface snow, while deposition also tends to be higher because the pollutant concentrations in the air are
335 higher (Wang et al., 2012; Nair et al., 2013). Previous studies have tended to focus on the accumulation area of
336 glaciers (e.g. ice cores and snow pits) where enrichment influences are less marked, and on high elevation areas,
337 where deposition is expected to be lower, in both cases leading to lower values. In our study, the majority of samples
338 collected in summer and autumn were collected from the ablation area of debris-covered glaciers where enrichment
339 influences are marked due to the relatively high temperature, and this is reflected in the relatively high values of BC,
340 OC, and dust. Li et al. (2017) showed a strong negative relationship between the elevation of glacier sampling
341 locations and the concentration of light absorbing particles. Stronger melt at lower elevations leads to higher

342 pollutant concentrations in the exposed snow. Equally, BC may be enriched in the lower elevation areas of glaciers
343 as a result of the proximity to source areas, as well as by the higher temperatures causing greater melting. Thus the
344 main reason for the high concentrations of BC, OC, and dust in our samples may have been that the samples were
345 taken from relatively low elevation sites. Human activities near the sampling sites in association with the summer
346 pilgrimage season probably also contributed to an increase in pollutant concentrations. Our results do not necessarily
347 indicate that all the glaciers in the Karakoram region are substantially darkened by BC. The ablation zones of debris
348 covered glaciers which are at relatively low elevations and near to pollution sources may be more polluted than
349 other glacier areas.

350 **3.2 Frequency distribution of aerosol sub types in the atmosphere**

351 The analysis of aerosol types using the CALIPSO data identified smoke as the most frequent aerosol type over
352 the study region in both summer and winter, indicating that biomass burning may be the dominant source of
353 emissions. Figure 2 shows the average frequency of different aerosol types in May/June (summer) and
354 December/January (winter) over the period 2006 to 2014 in the form of a box plot. The frequency of different
355 aerosol subtypes in June from 2006 to 2014 is shown in Figure S4; smoke had the highest frequency (39%),
356 followed by dust (21%), polluted dust (12%), and other (20%). This type of aerosol measurement in the atmosphere
357 was useful for our current study because it provides observation based data over the study region, whereas the other
358 approaches used (such as modeling) were based on interpolation not observation. Pollutant deposition depends on
359 the concentration of pollutants in the atmosphere and the results are consistent with the high concentration of BC
360 (from smoke) and dust particles in the glacier and snow surface samples.

361 **3.3 Snow albedo reduction**

362 The albedo of individual winter snow samples was calculated using the SNICAR model and then averaged for
363 each site (S1 to S6). Figure 3a shows the average for each site across the visible and infrared spectrum. Two sites
364 were chosen for further analysis: S1 (Sost) which had the highest average concentration of BC, and S6 (Kalam)
365 which had the lowest average concentration of BC. The albedo was simulated for different MAC values and SZA for
366 samples at the two sites as described in the methods. The values for average albedo of samples from the two sites
367 simulated for MAC values of 7.5, 11, and 15 m²/g and SZA of 57.0–88.9° (daytime) under a clear sky ranged from
368 0.39 (site S1, BC only, midday, MAC 15 m²/g) to 0.85 (site S6, dust only, early evening, MAC 7.5–15 m²/g). The
369 detailed values are shown in Table S3.

370 The percentage change in albedo was calculated in absolute terms as the change between albedo values with a
371 pollutant (BC or dust or both) and a reference albedo value with zero pollutants (zero BC and dust concentration).
372 Table 2 shows the calculated percentage reductions in daily minimum, maximum, and mean broadband snow albedo
373 at different MAC values (7.5, 11, 15 m²/g) resulting from the average BC, dust, and combined BC and dust
374 concentrations found in samples at each of the sites. The reduction was strongly dependent on BC concentration and

375 almost independent of dust concentration, and increased with increasing MAC value. The results suggest that BC
376 was the dominant forcing factor, rather than dust, influencing glacial surface albedo and accelerating glacier melt.
377 BC was found to play an important role in forcing in the northern Tibetan plateau (Li et al., 2016), whereas in the
378 central Tibetan plateau and Himalayas, dust played a more important role (Qu et al., 2014; Kaspari et al., 2014). The
379 MAC value affected the albedo more in the visible range than at 1.2 μm (near infrared) wavelength (Fig 3c,d). The
380 combined concentration of BC and dust, or BC alone, strongly reduced the snow albedo for a given combination of
381 other input parameters. The effect at the low pollutant site (S6) was small: the values for daytime snow albedo at
382 0.975 μm due to BC, or BC plus dust with different MAC and SZA, ranged from 0.70 to 0.83, with a reduction in
383 daily mean albedo of 1.8 to 2.9%, and those for dust alone from 0.79 to 0.85, with a reduction in daily mean albedo
384 of less than 0.1%. The effect at the high pollutant site (S1) was much more marked: BC or BC and dust reduced
385 daytime snow albedo to values ranging from 0.39 to 0.64, a reduction in daily mean albedo of 8.8 to 12.0%, but the
386 effect of dust alone was still low with values of 0.70 to 0.78, again a reduction in daily mean albedo of less than
387 0.1%.

388 Both the snow albedo and the impact of light absorbing particles depend on a range of factors including the SZA,
389 snow depth, snow grain size, and snow density. For example, the snow albedo reduction due to BC is known to be
390 less in the presence of other light absorbing particles as these will absorb some of the available solar radiation
391 (Kaspari et al., 2011). The snow albedo calculated for our samples was strongly dependent on the SZA with albedo
392 increasing with decreasing SZA, especially at near infrared wavelengths (Table S3).

393 The impact of snow ageing was also investigated. The winter samples from S1 (Sost) were aged snow, whereas
394 those from S6 (Kalam) were fresh snow (Table 1, Figure S5 b,c). Not only was dust clearly visible on the surface of
395 the aged snow, the grain size was large and the snow was dense. The aged snow had a much higher concentration of
396 BC and dust, which reduced the albedo, but the extent of reduction is also affected by other factors. Albedo
397 reduction by BC and dust particles is known to be greater for aged snow than for fresh snow (Warren and
398 Wiscombe, 1985). In our samples, the calculated reduction in snow albedo for high MAC values (15) compared to
399 low MAC values (7.5) was greater in aged snow than in fresh snow (Figure 3b). The effective grain size of snow
400 increases with time as water surrounds the grains. Snow with larger grain size absorbs more radiation because the
401 light can penetrate deeper into the snowpack, thus decreasing surface albedo (Flanner et al., 2006). In the melting
402 season, the snowpack becomes optically thin and more particles are concentrated near the surface layer, which
403 further increases the effect on albedo.

404 The estimated reduction in snow albedo by dust and BC (up to 29% of daytime maximum value, Table 2) was
405 higher than that reported by others for High Asia based on farmers' recordings (e.g 1.5 to 4.6% reported by Nair et
406 al., 2013) and in the Himalayas (Ming et al., 2008; Kaspari et al., 2014; Gertler et al., 2016). However, although the
407 values were relatively high, they were at the same level or lower than the estimates for albedo reduction of 28% by
408 BC and 56% by dust in clean ice samples, and of 36% by BC and 29% by dust in aged snow samples, reported by
409 Qu et al. (2014) for surface samples from the Zhadang glacier, China. Simulation results by Ming et al. (2013a)

410 showed BC, dust, and grain growth to reduce broadband albedo by 11%, 28%, and 61%, respectively, in a snowpack
411 in central Tibet. Dust was the most significant contributor to albedo reduction when mixed inside the snow and ice,
412 or when the glacier was covered in bare ice. In our case BC was a more influential factor than dust during a similar
413 study period to that reported by Li et al. (2017), indicating that BC plays a major role in albedo reduction.

414 The possible reasons for the relatively high values for albedo reduction in our samples include the lower
415 elevation of the sampling locations, relatively high concentrations of BC and dust, high MAC values, low snow
416 thickness, underlying ground quality, presence of small and large towns near the sampling sites, and predominance
417 of aged snow samples. Most of the samples collected in winter were from places with a snow depth less than 50 cm
418 (Figure S5a), thus mud, stones, and clay below the snow layer would be expected to increase the absorption of solar
419 radiation and reduce the albedo.

420 The high albedo reduction in the visible range of the electromagnetic spectrum could be due to the relatively
421 high concentration of surface (~1cm) snow impurities. The total amount of deposited particles in the surface layer of
422 aged snow was relatively high, indicating a high deposition rate of atmospheric pollutants.

423 Flanner et al. (2007) reported that BC emission and snow ageing are the two largest sources of uncertainty in
424 albedo estimates. The uncertainties in our estimated albedo reduction include the BC type (uncoated or sulfate coated),
425 the size distribution of dust concentration, the accuracy of snow grain size, snow texture, snow density, and albedo of
426 the underlying ground. Sulfate-coated particles have an absorbing sulfate shell surrounding the carbon; recent studies
427 confirm that coated BC has a larger absorbing power than non-coated BC (Naoe et al., 2009). We used uncoated black
428 carbon concentration in the SNICAR model, but the pollutants at the remote site are presumed to be mainly from long
429 range transport, thus the BC may have gained some coating. The albedo reduction for sulfate-coated black carbon was
430 calculated to be 3–8.5% higher, depending on the MAC and SZA values, than for uncoated black carbon at the low
431 concentration site S6 (Figure S6). The snow grain size (snow aging) and snow texture are also large sources of
432 uncertainty. The effect of snow grain size is generally larger than the uncertainty in light absorbing particles and varies
433 with the snow type (Schmale et al., 2017). The albedo reduction caused by 100 ng g⁻¹ of BC for an effective snow
434 grain radius of 80 μm, 100 μm, or 120 μm was calculated to be 0.017, 0.019, or 0.021, respectively. The snow grain
435 size was measured with a hand lens with an accuracy of 20 μm, thus the associated uncertainty in the albedo results
436 was at least 0.002. The snow grain shape was measured with the help of a snow card, but a spherical shape was
437 assumed for snow grains in the (online) SNICAR albedo simulation model. The albedo of non-spherical grains is
438 higher than the albedo of spherical grains (Dang et al., 2016), and the shape of snow grains and/or ice crystals changes
439 significantly with snow age and meteorological conditions during and after snowfall (LaChapelle 1969). A number of
440 recent studies (e.g., Flanner et al., 2012; Liou et al., 2014; He et al., 2014, 2017) have shown that both snow grain
441 shape and aerosol-snow internal mixing play an important role in snow albedo calculations.

442 **3.4 Radiative forcing (RF)**

443 Radiative forcing (RF) is a measure of the capacity of a forcing agent to affect the energy balance in the

444 atmosphere – the difference between sunlight absorbed by the Earth and energy radiated back to space – thereby
445 contributing to climate change. Changes in albedo contribute directly to radiative forcing: a decrease in albedo
446 means that more radiation will be absorbed and the temperature will rise. In snow and ice, the additional energy
447 absorbed by any pollutants present also increases and accelerates the melting rate.

448 Various authors have described the impact of albedo change in snow and ice on radiative forcing. Zhang et al.
449 (2017) reported that a reduction in albedo by 9% to 64% can increase the instantaneous radiative forcing by as much
450 as 24.05–323.18 Wm⁻². Nair et al. (2013) estimated that in aged snow a BC concentration of 10–200 ng g⁻¹ can increase
451 radiative forcing by 2.6 to 28.1 Wm⁻²; while Yang et al. (2015) reported radiative forcing of 18–21 Wm⁻² for aged
452 snow in samples from the westernmost Tibetan Plateau. The authors used different atmospheric conditions in the
453 forcing estimates: Zhang et al. (2017) used mid-latitude winter with clear sky and a cloudy environment, Nair et al.
454 (2013) mid-latitude winter, and Yang et al. (2015) clear-sky and cloudy conditions .

455 We calculated the radiative forcing in the samples assessed for daytime albedo and daily (24h) mean albedo. The
456 radiative forcing at different daylight times caused by BC deposition varied from 3.93 to 43.44 Wm⁻² (3.93–11.54
457 Wm⁻² at the low BC site and 20.88–43.45 Wm⁻² at the high BC site), and that by dust from 0.16 to 2.08 Wm⁻² (0.16–
458 0.30 Wm⁻² at the low BC site and 1.38–2.08 Wm⁻² at the high BC site) (detailed values given in Table S4), indicating
459 that BC was the dominant factor. The radiative forcing due to combined BC and dust was very similar to that for BC
460 alone. In contrast, studies by others have shown higher forcing by dust than by BC based on the optical properties
461 and size distribution of dust particles (Qu et al., 2014). In our study, the increase in daily mean radiative forcing
462 ranged from 0.1% for dust only at the low pollutant site to 14.9% for BC at the high pollutant site. However, dust
463 forcing varies strongly with dust optical properties, source material, and particle size distribution. The properties for
464 dust are unique in each of the four size bins used in the SNICAR online model. These size bins represent partitions
465 of a lognormal size distribution. We used an estimated size of dust particles and generic dust properties in the model,
466 but some dust particles can have a larger impact on snow albedo than the dust applied here (e.g., Painter et al.,
467 2007).

468 Both radiative forcing and albedo reduction increased with decreasing daytime SZA, indicating higher melting at
469 midday compared to morning and evening. Figure 4 shows the daily mean albedo reduction and corresponding
470 radiative forcing caused by BC for fresh (low BC) and aged (high BC) snow with different MAC values. Snow aging
471 (snow grain size) plays an important role in albedo reduction and radiative forcing. According to Schmale et al. (2017)
472 the effect of snow grain size is generally larger than the uncertainty in light absorbing particles, which varies with
473 snow type. Snow aging reduces snow albedo and accelerates snow melt, but the impact of snow aging on BC in snow
474 and the induced forcing is complex and includes spatial and seasonal variation (Qian et al., 2014).

475 An increase in MAC value from 7.5 to 15 led to an increase in radiative forcing by 1.48 Wm⁻² in fresh snow and
476 4.04 Wm⁻² in aged snow. This suggests that when the surface of snow, ice, and glaciers experience strong melting,
477 enrichment with BC and dust could cause more forcing. Previous studies of ice cores and snow pits probably
478 underestimated the albedo reduction and radiative forcing in glacier regions as samples were taken from high

479 elevation areas where there is less ageing and melting and thus lower surface enrichment of BC and dust than at
480 lower elevation. Our results are higher than those reported in other studies on the northern slope of the Himalayas
481 (Ming et al., 2012), western Tibetan Plateau (Yang et al., 2015b), and Tien Shan mountains (Ming et al., 2016).
482 However, they are comparable to values for radiative forcing reported more recently by others, for example for the
483 Muji glacier (Yang et al., 2015), Zhadang glacier (Qu et al., 2014), in high Asia (Flanner et al., 2007; Nair et al.,
484 2013), and in the Arctic (Wang et al., 2011; Flanner, 2013). The results suggest that enrichment of black carbon (in
485 our case) and mineral dust (other authors) can lead to increased absorption of solar radiation, exerting a stronger
486 effect on climate and accelerating glacier melt.

487 **3.5 Potential source regions**

488 **3.5.1 Wind vector maps**

489 Figure 5 shows the spatial variance of wind vector maps (U and V) at 850 hPa in May, June, January, and
490 December prepared using MERRA-2 reanalysis data for the year 2015/2016. The wind blows primarily from west to
491 east but there were variations over the year. Central Asia contributed some part of the air in May and June. In May,
492 the prevailing air masses were from Syria, Turkey, Turkmenistan, Iraq, Azerbaijan, northwest Iran, Afghanistan,
493 Nepal, southwest China, and southern Pakistan; the trend was similar in June with but with a smaller contribution
494 from Nepal and southwest China. In December and January (winter), the western trade winds were stronger than the
495 easterlies and the wind blew from Azerbaijan and northwest Iran, reaching the study site via Syria, Iraq,
496 Turkmenistan, and Afghanistan.

497 **3.5.2 Coupled emissions inventory with back air trajectory**

498 Trajectory analysis using the HYSPLIT model showed that in May and June 2015 air parcels reached the study
499 site along three different pathways: one from north Asia (Russia) via Central Asia (Kazakhstan), one from western
500 Asia (Cyprus and Syria) via Central and Southern Asia (Georgia), and one via India, which was more local (Figure
501 6). The trajectories in summer had distinct pathways, while those in winter were dispersed in all directions, partially
502 covering West, East, and South Asia, and completely covering Central Asia. Figure 6 shows the product of
503 extinction and emission calculated along the pathways of trajectories calculated using the vertical profile for aerosol
504 extinction over the study region obtained from the monthly CALIPSO satellite-based extinction data. Scattering and
505 absorption decreased exponentially with increasing elevation (Figure S1) but was still visible at elevations above 5
506 km in summer.

507 The RCP emission data combined with back trajectories and extinction data showed that the hotspot regions of
508 pollution that affected the study sites during winter were mainly to the southwest rather than very distant (Figure
509 6b). Iran, Turkmenistan, Azerbaijan, Georgia, the eastern part of Turkey, and the southwestern part of Russia all
510 showed comparatively high pollutant emissions in winter which moved towards northern Pakistan. The western part
511 of Kazakhstan, Uzbekistan, and northeastern Turkey emitted particularly high concentrations of pollutants.

512 Combination of the back-trajectory results and surface-wind direction analysis indicated that during the sampling
513 months, aerosols were significantly influenced by the long-range transport of pollutants coming from Central and
514 South Asia, with a small contribution from West and East Asia. This differs somewhat from previous reports which
515 suggested that the Tibetan Plateau and Himalayan region are mainly affected by pollutants from East and South Asia
516 (Zhang et al., 2015). An increasing trend has been reported for black carbon emissions in Central and South Asia
517 over the past 150 years (Bond et al., 2007), and a significant increase has been found in black carbon concentrations
518 in glacier snow in west China in the last 20 years, especially during the summer and monsoon seasons (Ming et al.,
519 2008). In South Asia, the largest source of atmospheric black carbon is emission from biomass and biofuels used for
520 cooking and heating (dung, crop residues, wood) (Venkataraman et al., 2005).

521 The results indicate that only a low level of pollutants (minor contribution) reached the study area from
522 northwest China. BC particles emitted from distant low latitude source regions such as tropical Africa barely reach
523 the Tibetan Plateau and Himalayan regions because their emissions are removed along the transport pathways during
524 the summer monsoon season (Zhang et al., 2015).

525 **3.5.3 Chemical transport modelling**

526 The contribution of pollutants from potential source regions was also investigated using the WRF-STEM model
527 with tagged carbon monoxide tracers and source regions of East Asia, South Asia, Central Asia, the Middle East,
528 Europe, the Russian Federation, and West Asia. (The individual countries in the regions are listed in Table S5).

529 Figure 7 shows the results of the model simulations for summer (1 June to 4 July 2015) and winter (15
530 December 2015 to 17 January 2016) at two glacier sites (Sachin and Shangla) where the model terrain elevation was
531 close to the observation terrain elevation. The model simulations showed Pakistan to be the major contributor of
532 pollutants in summer (77% at Shangla and 43% at Sachin) followed by the South Asian countries; and the south
533 Asian countries in winter (47% at Shangla and 71% at Sachin) followed by Pakistan, which is in line with the
534 findings by Lu et al. (2012) that South Asia contributed 67% black carbon in the Himalayas. There were minor
535 contributions of 2–7% of pollutants from Afghanistan, Iran, Central Asia, and the Middle East, and extremely small
536 amounts from East Asia, Europe, Africa, West Asia, and China. The contribution from Iran, the Middle East, and
537 Europe was greater in winter than in summer, while the contribution from Central Asia and China was greater in
538 summer than in winter. The proportion of daily contributions fluctuated considerably: with higher contributions from
539 Iran, the Middle East, and Europe on individual days in winter, ranging for example from 2–30% for the Middle
540 East.

541 The concentration of hydrophobic BC (BC1), hydrophilic BC (BC2), and total black carbon ($BC = BC1 + BC2$)
542 given by the model for the Sachin glacier grid point in the summer and winter seasons is shown in the supplemental
543 material (Figure S7). In the model, freshly emitted BC particles are hydrophobic and gradually acquire a
544 hygroscopic coating over time. A time series analysis of BC1 and BC2 concentrations shows the influence of both
545 freshly emitted BC and aged BC reaching the observation location. The highest concentration of BC1 was observed

546 on 20th December 2015 and the second highest on 25th June 2015, indicating an influence from freshly emitted air
547 masses in both the summer and winter months. Future studies (BC tracer) will evaluate the details of the different
548 source regions of the BC reaching the glaciers compared to region-tagged CO tracers.

549 **3.5.4 Comparison of the different approaches used to identify potential source regions**

550 The high BC concentration in the atmosphere over the study region was attributed to long-range transport from
551 urban source regions. Potential source regions of the pollutants deposited on glaciers and snow were identified using
552 wind vector mapping with MERRA-2 reanalyzed data, calculation of back air trajectories using the HYSPLIT-4
553 model, and chemical transport pathways using the WRF-STEM tagged chemical transport model. The back
554 trajectory results indicated that the majority of pollutants in summer were from Central and South Asia, and in
555 winter from Iran, Pakistan, Iraq, Turkmenistan, Azerbaijan, Georgia, Jordan, Syria, Tunisia, Ukraine, Libya and
556 Egypt. The WRF-STEM model indicated that most anthropogenic pollutants were from Pakistan and South Asia
557 during both summer and winter. However, both approaches showed a reasonable contribution from Central Asian
558 countries and limited contribution from East Asian countries in summer. The wind vector maps also indicated that
559 the study site was mostly affected by westerly winds. All three approaches showed a reasonable contribution from
560 neighboring countries such as Afghanistan, Pakistan, Iran, and India in specific months. Overall, the results indicate
561 that South, Central, and West Asia were the major sources of the pollutants detected at the sampling sites.

562 There was some mismatching in source regions among the three approaches. The WRF-STEM model and wind
563 vector maps both identified a small contribution from East Asia, but this was not identified in the back trajectories
564 approach. Similarly, the wind vector maps and back air trajectories showed a dominant contribution from the west,
565 while the WRF-STEM model showed a major contribution from Pakistan and South Asia. The differences in the
566 results obtained by the different methods may be due in part to the complex topography of the region and the
567 different altitudes used by the methods; the coarse resolution of the WRF-STEM model; and differences in the
568 emission source inventories and meteorological parameters used by the WRF-STEM and HYSPLIT-4 models. The
569 limitations of using back trajectories to identify source region is discussed further in a paper by Jaffe et al. (1999).

570 Furthermore, the atmospheric BC concentration over the Himalayas has significant temporal variations
571 associated with synoptic and meso scale changes in the advection pattern (Babu et al., 2011) which can affect
572 pollutant transport and deposition. The large uncertainty among different emission inventories can also affect the
573 results, especially in the Himalayan region.

574 **4 Summary and conclusion**

575 Black carbon (BC) and organic carbon (OC) concentrations were measured using thermal optical analysis of
576 snow and ice surface samples collected from glacier and mountain valleys in northern Pakistan in summer, autumn,
577 and winter. The samples contained high concentrations of BC, OC, and dust in low elevation glaciers and surface
578 snow in mountain valleys. The samples from Sost contained the highest average concentration of BC in mountain

579 valley snow (winter) and those from Kalam the lowest, probably due to the impact of snow age and increased
580 concentration of black carbon and dust (the Sost samples were aged snow and Kalam samples fresh snow). The
581 average concentration of BC in surface samples from the Sachin glacier was higher in autumn than in summer; the
582 BC values in summer snow samples collected from the Sachin and Gulkin glaciers (aged snow from the glacier
583 surface) were much higher than those in ice. The average BC concentration in summer samples collected from
584 glaciers was $2130 \pm 1560 \text{ ng g}^{-1}$ and that in autumn samples $2883 \pm 3439 \text{ ng g}^{-1}$. The average concentration of OC
585 was $1839 \pm 1108 \text{ ng g}^{-1}$ in summer samples, $1423 \pm 208 \text{ ng g}^{-1}$ in autumn samples, and $1342 \pm 672 \text{ ng g}^{-1}$ in winter
586 samples, with the highest variability in summer samples. The individual lowest BC (82 ng g^{-1}) and OC (129 ng g^{-1})
587 concentrations were observed in summer samples collected from the Gulkin and Sachin glaciers, respectively. Dust
588 and other pollutants were clearly visible on aged snow and ice surfaces; the results indicate considerable enrichment
589 during ageing. The pollutant concentrations in our samples were relatively higher than those reported by others in
590 earlier studies, which tended to focus on the accumulation area of glaciers (e.g. ice cores and snow pits), where
591 enrichment influences are less marked and measured values are likely to be lower, and high elevation areas, where
592 deposition of pollutants is expected to be lower. It is likely that pollutant concentrations were underestimated in
593 these earlier studies, particularly when there was strong surface melting.

594 Snow albedo was calculated for winter samples using the SNICAR model with various combinations of BC and
595 dust concentrations, three values for MAC, and a range of values for SZA ($57\text{--}88.89^\circ$ during daytime), with other
596 parameters kept constant. BC was the major component responsible for albedo reduction, dust had little effect. The
597 reduction by BC ranged from 2.8 to 32.5% during daytime, which is quite high, with albedo reduced to below 0.6.
598 The reduction was greater for higher concentrations of BC and greater MAC. The reduction in 24 h average albedo
599 ranged from $<0.07\text{--}2.9\%$ for fresh snow samples and $<0.05\text{--}12.0\%$ for aged snow. Changes in albedo contribute
600 directly to radiative forcing: a decrease in albedo means that more radiation will be absorbed and the temperature
601 will rise. The radiative forcing by BC was also higher than that caused by dust, indicating that BC was the dominant
602 factor. The daytime albedo values in winter snow samples ranged from 0.39 to 0.82 with BC alone or BC plus dust,
603 and from 0.70 to 0.85 with dust alone; the corresponding radiative forcing was $3.93\text{--}43.44 \text{ Wm}^{-2}$ for BC alone, 4.01--
604 43.45 Wm^{-2} for BC and dust, and $0.16\text{--}2.08 \text{ Wm}^{-2}$ with dust alone. The radiative forcing calculated from the daily
605 mean albedo reduction ranged from 0.1% for dust only at the low pollutant site to 14.9% for BC at the high pollutant
606 site. The potential source regions of the pollutants deposited on glaciers and snow were identified using spatial
607 variance in wind vector maps, emission inventories coupled with back air trajectories, and region tagged chemical
608 transport modelling. The wind vector maps identified Central Asian and South Asian countries (such as Azerbaijan,
609 Turkmenistan, Pakistan, Afghanistan, Syria, Iraq, Turkey) as more important. The trajectory analysis coupled with
610 emission inventories showed that air parcels reached northern Pakistan along three pathways, one from north Asia
611 (Russia) via Central Asia (Kazakhstan), one from western Asia (Cyprus and Syria) via Central and southern Asia
612 (Georgia), and one via India. Combination of the back-trajectory results and surface-wind direction analysis
613 indicated that aerosols were significantly influenced by the long-range transport of pollutants from Central and

614 South Asia. The region-tagged chemical transport model indicated that Pakistan and South Asia were the main
615 contributors of pollutants. Analysis based on the WRF-STEM model identified a significant contribution from
616 Pakistan (up to 77%) and South Asia (up to 71%) at selected sites. Overall, the results indicate that Central, South,
617 and West Asia were the major sources of the pollutants detected at the sampling sites, with only a small contribution
618 from East Asia.

619 The overall uncertainty of the BC, and OC concentrations was estimated taking into account the analytical
620 precision of concentration measurements and the mass contribution from field blanks. The uncertainty in the BC and
621 OC mass concentrations was calculated from the standard deviation of the field blanks, the experimentally
622 determined analytical uncertainty, and the projected uncertainty associated with filter extraction. The major source
623 of uncertainty was the effect of dust on the OC/BC measurements.

624 The albedo reduction from OC was not quantified. The contribution of OC to total visible absorption in the top
625 snow layer is relatively small compared to that of BC and dust but has been shown to be significant (~19% of the
626 total solar visible absorption) in several regions including northeastern East Asia, and western Canada (Yasunari et
627 al., 2015). Snow grain size (snow aging) and snow texture were probably the main sources of uncertainty in the
628 albedo reduction/radiative forcing calculations. The measured grain size was generally different from the effective
629 optical grain size used in the SNICAR modeling, and although snow grain shape was measured, the results were not
630 used in the online SNICAR albedo simulation model, which assumes a spherical shape for snow grains. This could
631 slightly affect the results, because the albedo of non-spherical grain is higher than the albedo of spherical grains
632 (Dang et al., 2016).

633 The possible uncertainties on the modeling side relate to the use of CO as a tracer for light absorbing particles to
634 identify the source region. Uncertainties are also attributed to errors in the emission inventories, simulated
635 meteorology, and removal processes built into the model. The physics and chemistry of removal of BC and -CO
636 differ, especially in the wet season. However, we analyzed the model during pre-monsoon and relatively dry periods
637 when there should be a relatively good correlation in the transport of CO and BC. The global emission inventories
638 used are unable to capture emissions at a local scale, and the contribution of local sources may also be
639 underestimated by coarse-resolution models. High resolution models and emission inventories at a local scale are
640 required to capture local emissions.

641 Better constrained measurements will be required to obtain more robust results. High resolution satellite imagery,
642 high resolution models, and continuous monitoring will help to reduce the present uncertainty.

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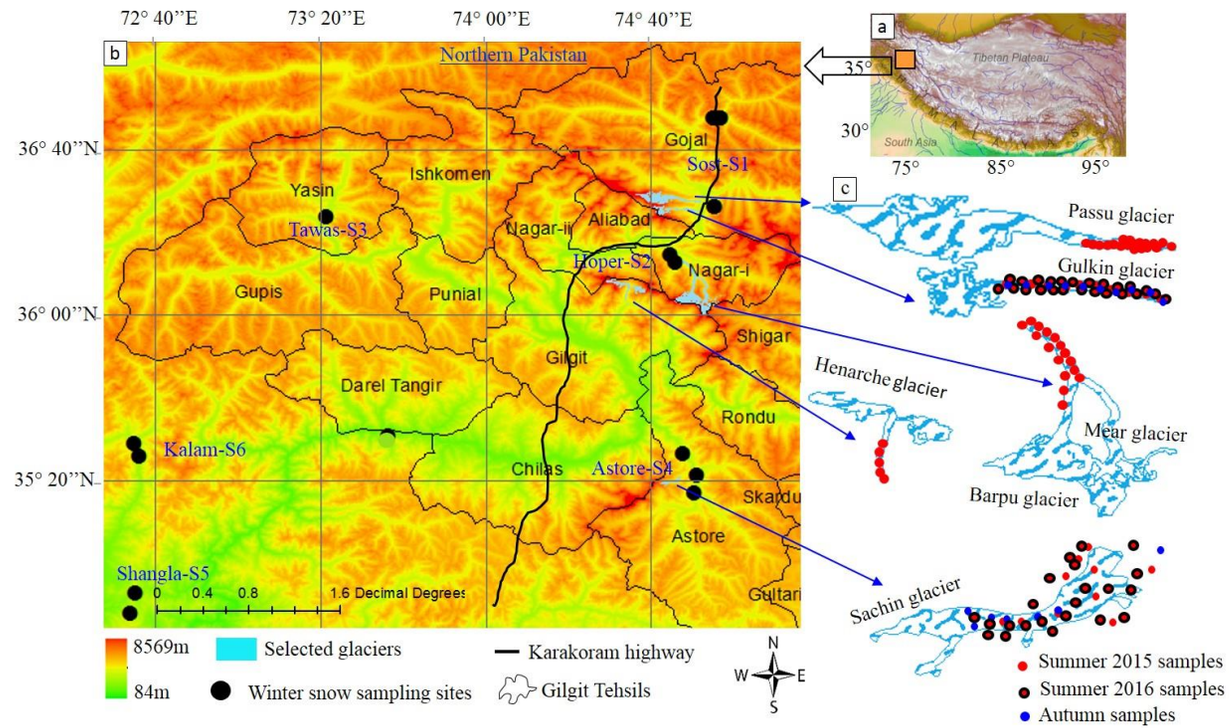
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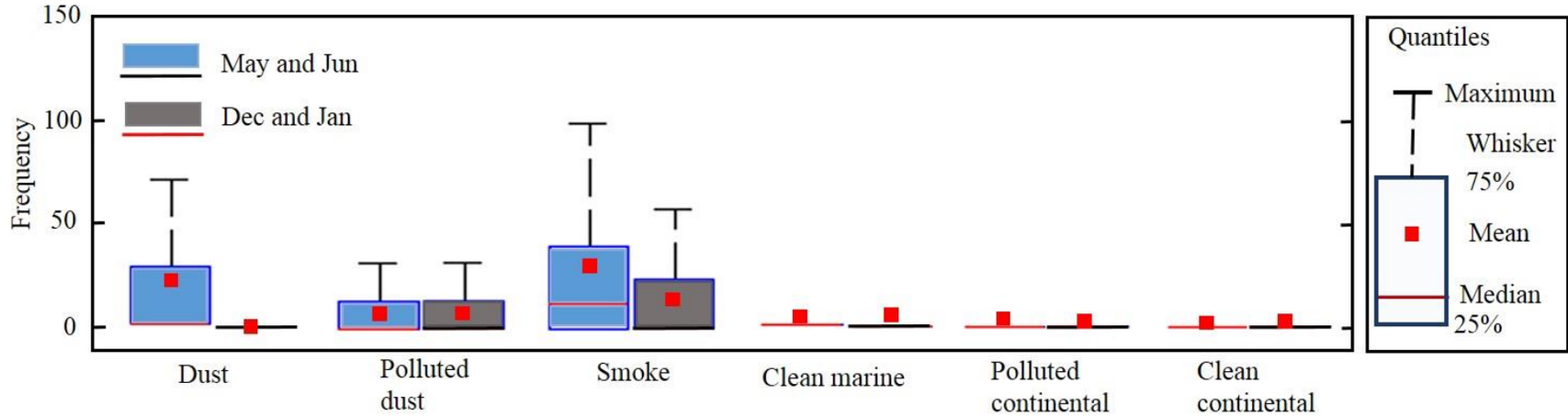


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884 **Figure 1. The study area and sampling sites: (a) Himalayan mountain range and Tibetan Plateau, (b) winter sampling sites (solid black circles), (c) glaciers selected for**
885 **summer and autumn sampling**

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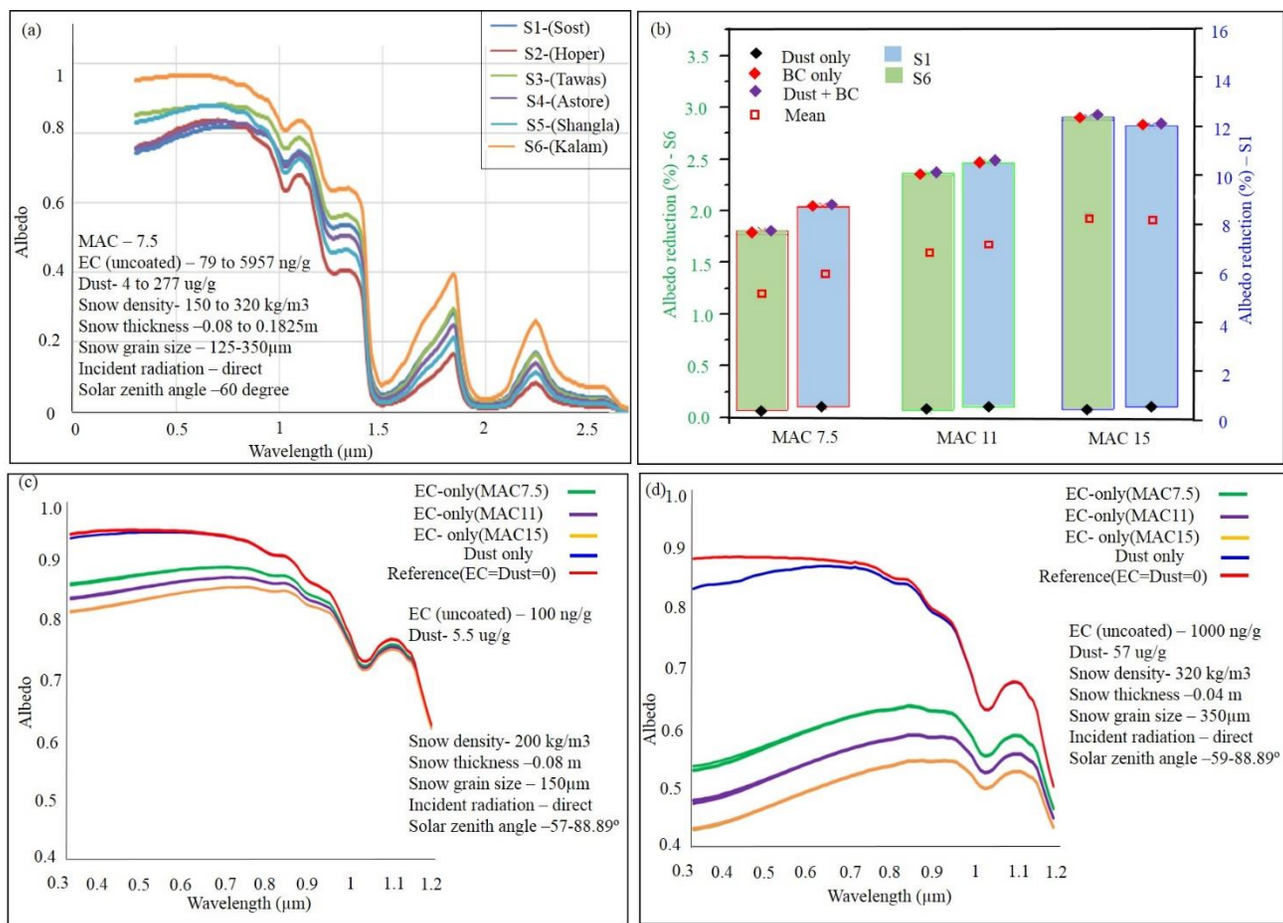
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889 **Figure 2. Frequency distribution of aerosol subtypes in the atmosphere over the study region calculated from CALIPSO data, average for the study months in 2006 to**
 890 **2014**

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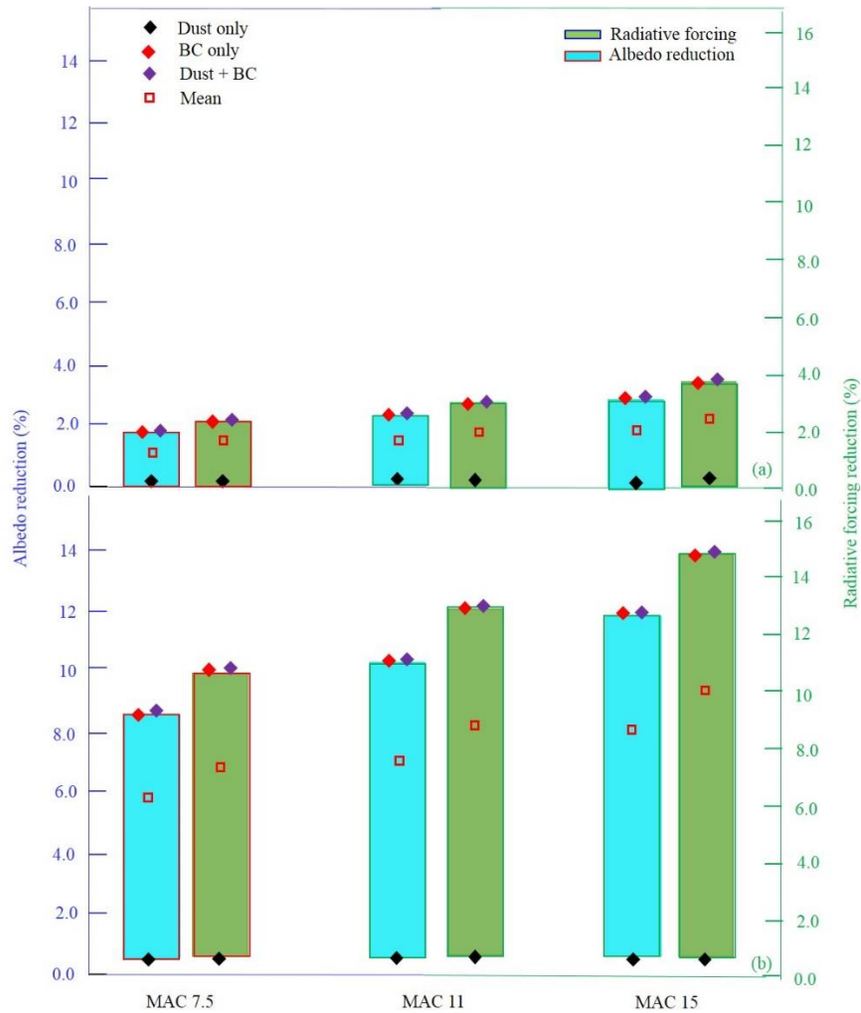
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Figure 3. Spectral variation in albedo for winter sampling sites and selected mass absorption cross-section (MAC) values, (a) average albedo of samples at each of the sites (b) daily mean albedo reduction of fresh snow (site S6) and aged snow (site S1) snow, (note different scales of y axis) (c) albedo of fresh snow site S6, (d) albedo of aged snow site S1

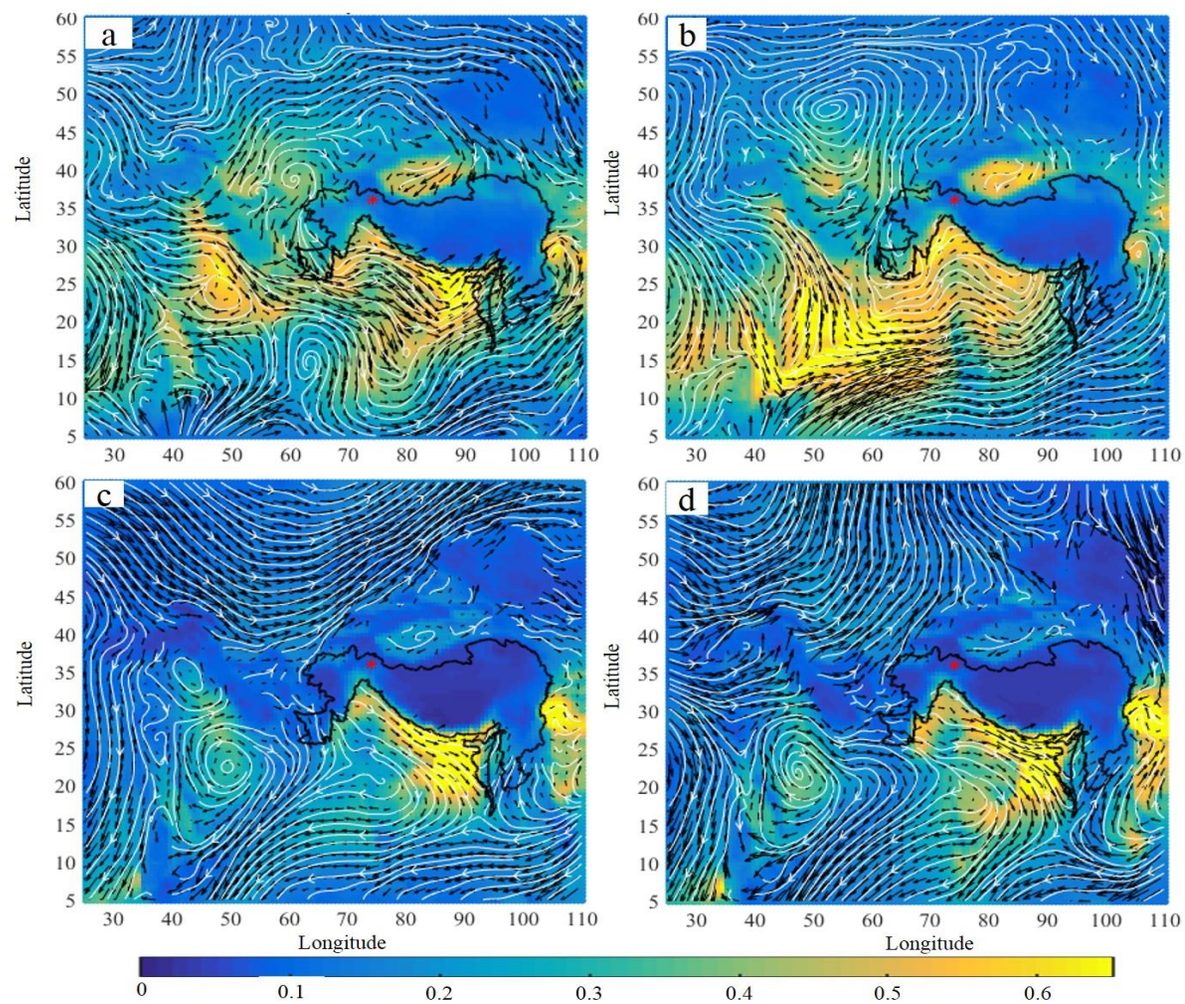
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899 **Figure 4. Daily mean radiative forcing reduction and albedo reduction (%) caused by black carbon and dust for different mass absorption cross-section values (MAC) in**
 900 **(a) fresh (low black carbon) and (b) aged (high black carbon) snow samples (note different scales of y axis)**

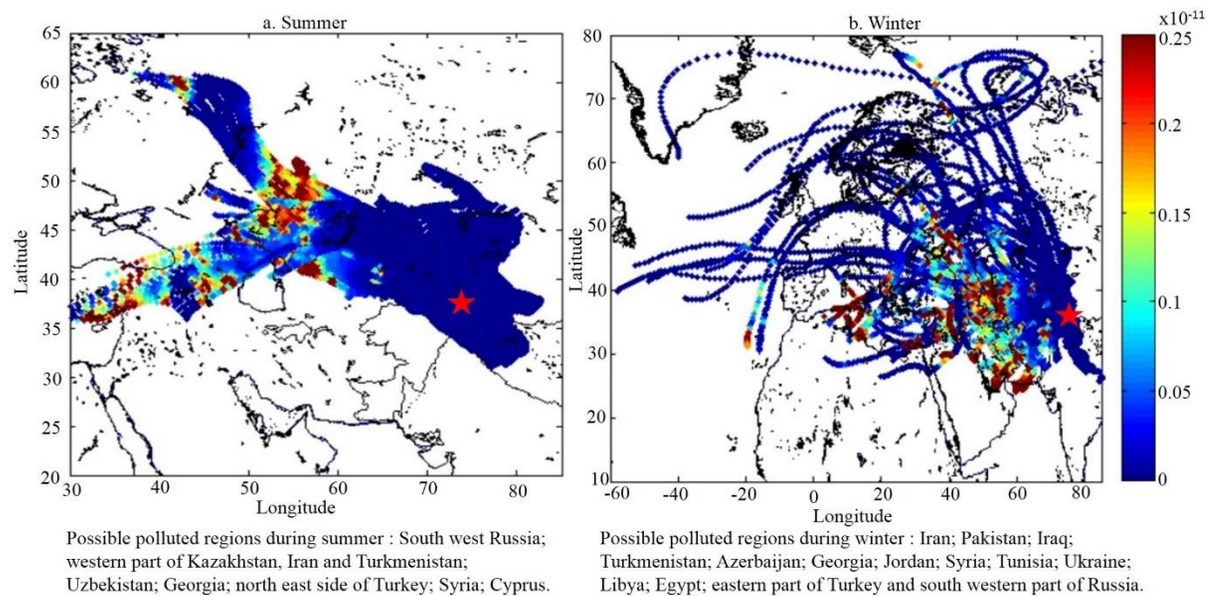


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903 **Figure 5. Monthly average horizontal wind patterns at 850 hPa during a) May, b) June, c) December, and d) January, corresponding to approximately 2,500 masl, from**
 904 **GES DISC. Red star indicates the position of the study area, and white lines indicate streamlines. The background colors show monthly mean aerosol optical depth.**

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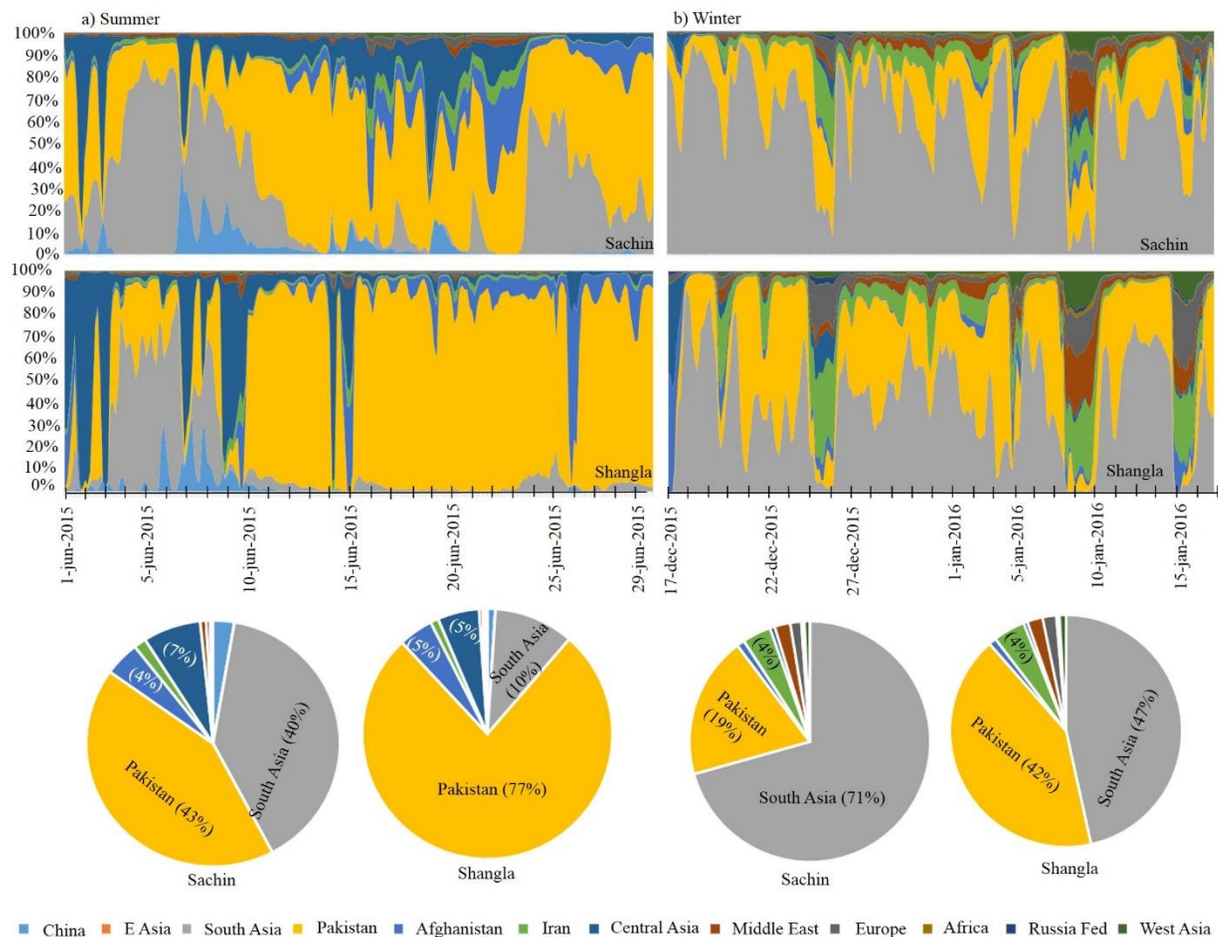
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Figure 6. Source contribution regions of pollutants identified using an emissions inventory (Representative Concentration Pathways) coupled with back trajectories: a) 77 simulated days, b) 63 simulated days. Red star indicates the position of the study area.



911
 912 **Figure 7. Source contribution regions of carbon monoxide for selected sites identified by WRF-STEM during (a) summer and (b) winter**
 913

914 **Table 1. Concentration of black carbon, organic carbon, and dust in summer, autumn, and winter samples in 2015 and 2016**

Glacier/ Site	No.	Elevation (masl) min–max	BC min–max (avg) (ng g ⁻¹)	OC min–max (avg) (ng g ⁻¹)	Dust min–max (avg) (μgg ⁻¹)	Type ^a / snow age in days	OC/BC ^b	Year
Summer (May 2015/ May 2016)								
Barpu	6	2901–3405	877–5994 (2938)	244–1228 (691)	292–5250 (1998)	DCI	0.07–1.38	2015
Gulkin	31	2741–3319	82–5676 (1327)	238–8514 (1594)	31–2039 (648)	DCIS	0.169–3.76	2015/16
Henarche	4	2569–2989	778–10502 (4820)	275–4176 (1628)	225–2723 (993)	Ice	0.04–1.63	2015
Mear	8	2961–3539	222–3656 (1593)	703–6588 (2992)	33–656 (211)	DCI	0.72–4.88	2015
Passu	14	2663–3158	87–734 (346)	132–1810 (741)	28–524 (196)	DCI	1.85–4.80	2015
Sachin	35	3414–3895	257–4127 (1769)	128–7592 (3348)	5.6–2495 (314)	DCIS	0.08–0.53	2015/16
Total	98							
Autumn (October 2016)								
Gulkin	7	2741–3319	125–1028 (451)	266–3574 (1276)	60–767 (253)	DCIS	1.29–3.59	2016
Sachin	6	3414–3895	4342–6481 (5314)	543–3478 (1571)	124–1348 (546)	DCIS	0.11–0.53	2016
Total	13							
Winter (Dec 2015/ Jan 2016)								
S1-Sost	6	2873–3092	482–5957 (2506)	378–2934 (1039)	29–311 (131)	2–17 d	0.25–0.78	2015
S2-Hopar	2	2602–2794	229–1064 (646)	330–1976 (1153)	23–129 (76)	1–15 d	1.4–1.8	2016
S3-Tawas	1	2437	650	1320	16	8–17 d	2.03	2016
S4-Astore	3	2132–2396	450–2640 (1305)	914–3645 (2161)	55–171 (97)	4–7 d	1.38–2.33	2016
S5-Shangla	2	2324–2373	367–1110 (739)	1302–2856 (2079)	13–49 (31)	8–9 d	2.5–3.5	2016
S6-Kalam	4	1933–2101	79–123 (107)	214–558 (347)	4–6 (5)	1 d	2.3–5	2016
Total	18							

915 ^a type = snow or ice type; DCI = debris-covered ice; DCIS = debris-covered ice and aged snow

916 ^b range of OC/BC in individual samples

917

918 **Table 2. Snow albedo reduction (%) by black carbon, dust, and black carbon plus dust at the site with the lowest average pollutant concentration (S6) and the site with the**
 919 **highest average pollutant concentration (S1), under different mass absorption cross-section (MAC) values**

Pollutant	MAC value (m ² /g)	Low concentration site (S6)			High concentration site (S1)		
		Daytime min	Daytime max	Daily mean	Daytime min	Daytime max	Daily mean
Black carbon	7.5	2.8	5.1	1.8	15.6	23.9	9.0
	11	3.7	6.9	2.3	19.2	28.6	10.5
	15	4.6	8.3	2.9	22.3	32.5	12.0
Dust	7.5	0.1	0.2	0.07	0.9	1.6	0.05
	11	0.1	0.2	0.07	0.9	1.6	0.05
	15	0.1	0.2	0.07	0.9	1.6	0.05
Black carbon and dust	7.5	2.9	5.2	1.8	15.7	24.0	8.8
	11	3.8	6.8	2.4	19.2	28.6	10.5
	15	4.6	8.3	2.9	22.3	32.5	12.0

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