Iron oxides in the cryoconite on the glaciers over Tibetan Plateau: abundance, speciation and implications

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Abstract. Cryoconite is a mixture of impurities and ice visually represented by dark colors present in the ablation zone of glaciers. As an important constituent of light-absorbing impurities on the glacier surface, iron oxides influence the radiative properties of mineral dust and thus its impact on ice melting processes. In particular, the distinct optical properties between hematite and goethite (the major iron oxide species) highlight the necessity to obtain accurate knowledge about their abundance and geochemical behaviour. Cryoconite samples from five glaciers in different regions of the Tibetan Plateau (TP) and surroundings were studied. The iron abundance in the cryoconite from TP glaciers ranged from 3.40% to 4.90% by mass, in accordance to typical natural background level. Because the light-absorption capacity of mineral dust essentially depends on the presence of iron oxides (i.e. free iron), iron oxides were extracted and determined using diffuse reflectance spectroscopy. The ratios of free to total iron for the five glaciers ranged from 0.31 to 0.70, emphasizing that iron in the form of oxides should be considered rather than total iron in the albedo and radiative modelling. Furthermore, the goethite content in iron oxides (in mass fraction) ranged from 81% to 98%, showing that goethite was the predominant form among the glaciers. Using the abundance and speciation of iron oxides as well as their optical properties, the total light absorption were quantitatively attributed to goethite, hematite, black carbon and organic matters at 450 nm and 600 nm wavelengths. We found that the goethite played a stronger role than BC at shorter wavelength for most glaciers. Such findings were essential to understand the relative significance of anthropogenic/natural effect, and then taking the proper mitigation measures.

1 Introduction

The light-absorbing impurities (LAI:s) in glaciers can significantly reduce the surface albedo of snowpack and absorb more solar energy (Warren and Wiscombe, 1985). LAIs were recognized as a
major contributor to the glacier and ice sheet melting (Qian et al., 2015 and references therein), along with the rising air temperatures (IPCC, 2014). The composition of LAIs on the glacier surface is very complex. Their major constituents include black carbon, brown carbon, mineral dust, as well as biogenic matter (Baccolo et al., 2017; Kaspari et al., 2015; Pu et al., 2017; Takeuchi, 2002). The composition of LAIs is particularly complex when considering mountain glaciers from the Himalayas and Tibetan Plateau (TP) (Fig. 1). In summer, their surfaces are commonly covered by incoherent impurities (granular sediment) with dark color, which was termed as cryoconite (Baccolo et al., 2017; Dong et al., 2016).

Presently, tremendous attention has been paid to the black carbon in glaciers (Kaspari et al., 2011; Wang et al., 2015; Yasunari et al., 2010), because the glaciated areas of the TP are in the vicinity of intensive BC source region, e.g. South Asia (Cong et al., 2015). In contrast, few research has focused on dust in this region despite dust is apparently the predominant constituent of impurities on the glacier surface, especially in its ablation area (Qian et al., 2015). Based on field research at Mera Glacier of Nepal Himalayas, Kaspari et al. (2014) pointed out that when dust loading is high, the snow albedo reduction and subsequent radiative forcing caused by dust will overwhelm the impact related to black carbon.

To quantify the relative contribution of dust and black carbon as well as other substances is challenging (Painter et al., 2010). In the snow albedo simulation models like SNICAR (Flanner and Zender, 2006), dust concentration (micrograms of dust per gram of ice) was employed to represent the loading of dust, without considering the dust composition. Actually, the light-absorption capacity of dust essentially depends on the presence of iron oxides (also commonly termed as “free iron”) (Alfaro et al., 2004; Formenti et al., 2014; Moosmüller et al., 2012; Shi et al., 2012). The most common iron oxide species in mineral dust are hematite ($\text{Fe}_2\text{O}_3$) and goethite ($\text{FeO(OH)}$), which have distinct optical properties in terms of absorption and wavelength dependence (Balsam et al., 2014). At the same time, some parts of iron are incorporated into the crystal lattice of alumino-silicates (defined as structural iron), and they do not contribute to the absorption of solar light (Lafon et al., 2006).

Up to now, the iron abundance and especially its mineral phases in glacial area are not well understood (Hawkings et al., 2018; Shahgedanova et al., 2013). The degree to which iron oxides contribute to solar absorption and reduction of snow albedo remains uncertain. To constrain the uncertainties of estimating the radiative forcing of cryoconite, particularly in the Tibetan Plateau glacier area, we designed this research to address several key issues. Firstly, what is the abundance of iron in the cryoconite of mountain glaciers? How much fraction of the total iron exists as iron oxides with efficient light-absorption capability? What is the relatively proportion of hematite and goethite, considering their distinct optical characteristics? What is their spatial variation in different types of glacier? Furthermore, how do the iron oxides impact the absorbing properties of cryoconite?
2 Field sampling and laboratory measurements

2.1 Field sampling

In order to consider geographic variability, five glaciers in different regions of TP and surroundings were chosen for the cryoconite sampling (Fig. 2). A detailed descriptions of the collection sites is given in Table 1. The Urumqi No. 1 Glacier (hereafter donated as UG, 43°06’N, 86°49’E) presents two branches covering 1.646 km². It is located in eastern Tien Shan. The air circulation regime there is dominated by the westerlies in summer and by the influence of the Siberian High during winter (Wang et al., 2014). Laohugou Glacier (LHG, 39°28’N, 96°32’E), with a length of 10 km and an area of 20 km², is at the northern slope of western Qilian Mountains with typical continental climatic conditions (Dong et al., 2014). That area is surrounded by large sandy deserts in Northwest China, like Taklimakan Desert to the west, Qaidam Basin to the southwest and the Gobi Desert to the north. Xiaodongkemadi Glacier (XDK, 33°03’N, 92°04’E) is located on the northern slope of Tanggula Mountains, at the center of Tibetan Plateau. Previous studies showed that Tanggula Mountains represent the northern boundary of the area influenced by the South Asian monsoon (Tian et al., 2001). Palong #4 Glacier (PL, 29°15’N, 96°56’ E), located in the southeast Tibetan Plateau, is a typical temperate glacier. It is strongly influenced by the South Asian summer monsoon intruding via the Brahmaputra Valley, and it is characterized by high accumulation (2500-3000 mm) and ablation rates on an annual scale (Yang et al., 2015). Baishui No. 1 Glacier (BS, 27°6’N, 100°11’E), with a length of 2.26 km and an area of 1.32 km², is the largest glacier in the Yulong Mountains, at the southeastern edge of Tibetan Plateau. It is characterized by high precipitation, low snow line, and relatively high temperatures (equilibrium line mean annual value -6 °C, summer value 1-5 °C) (Niu et al., 2013).

Cryoconite samples were collected using a stainless-steel scoop on the surfaces of five glaciers described above. Samples were preserved in NALGENE HDPE wide-mouth bottles (500 ml) and kept in frozen until analysis. In the laboratory, the cryoconite samples were freeze-dried into powder. In this work, concentrations and fractions are referred to as dry cryoconite mass. Therefore, our measurements mainly reflect processes occurring in summer on the glacier surface.

2.2 Elemental analysis by ICP-MS

A portion of cryoconite sample (about 20 mg) was dissolved under a laboratory hood using HF+HNO₃ mixture through three steps. In detail, the sample was first transferred into a PTFE high-pressure digestion vessel, and 1 mL HF and 1 ml HNO₃ were added. The digestion vessel was then ultrasonic treated for 20 min, and evaporated to nearly dry on a hot plate. Another 1 mL HF and 1 ml HNO₃ were added, and digested in an oven at 190 °C for 24 h. After cooling, the vessel was opened and evaporated to nearly dry again (on the plate at 150 °C), then followed by a second addition of HNO₃. This procedure was repeated to remove HF completely. Then 1mL HNO₃ and 3mL H₂O were added to the vessel which was placed into the oven for another 24h at 150 °C. After cooling, the final solution was diluted with pure water to about 50mL.
After treatment, each sample was measured by inductively coupled plasma-mass spectrometry (ICP-MS, Thermo X7, Thermo-Elemental Corp.) for Fe and other elements. Indium, Rhodium and Rhenium solution were used as internal standards. The accuracy and precision of trace elements was ascertained based on repeated measurement of the USGS Geochemical Reference Standard (Andesite, AGV-2). The measured and certified values for Fe agree well, with recovery better than 95%. The detailed description of the analytical protocol in our laboratory could be found in previous work (Wu et al., 2009).

2.3 Total organic carbon and black carbon

The contents of organic carbon were determined by a total carbon analyzer (TOC-V, Shimadzu). The accuracy of the TOC analysis was ±5%. The separation and analysis of black carbon in the cryoconite were adopted from the procedures previously developed for sediments (Cong et al., 2013; Han et al., 2011). Specifically, the samples were first freeze-dried, grinded into powder and weighed. Then, HCl (2N) was added, to remove carbonates, silicates, and some kinds of metal oxides. The solution was centrifuged to remove the supernatants. Then mixture (1:2) of HCl (6N) and HF (48%) were added into the residue and reacted further. Finally, the residual solid was diluted with ultrapure water, filtered by quartz fiber filter with even distribution on their surface (QMA grade; Whatman International Ltd, England). The quartz filters were analyzed for BC using a DRI model 2001 carbon analyzer. For quality control, standard reference material (marine sediment, NIST SRM-1941b) was also analyzed (Cong et al., 2013). Our BC values compare well to the value reported by Han et al. (2007) (> 95%, n=5), which indicates the analytical method used is reliable and repeatable.

2.4 Extraction and quantification of iron oxides

Cryoconite samples were treated with Citrate-Bicarbonate-Dithionite (CBD) method three times to completely extract iron oxides (Ji et al., 2002; Lafon et al., 2004; Mehra and Jackson, 1958). Then dissolved Fe$^{3+}$ concentrations in the CBD solution were determined by a UV-2100 spectrophotometer (UNICO Inc., Shanghai) to obtain the iron mass in the form of oxides, Fe(ox), relevant to the light-absorption in the visible light. Uncertainties for the Fe(ox) analysis were less than 5%. The remaining iron, i.e. the structural iron, was calculated by subtracting the Fe(ox) from total iron.

\[
\text{Fe(struc)} = \text{Fe(tot)} - \text{Fe(ox)}
\]

Here, \(\text{Fe(tot)}\) is the total iron concentration achieved from ICP-MS elemental analysis.

2.5 Hematite/goethite measurement using Diffuse Reflectance Spectroscopy

Given the low abundance of Fe in the cryoconite, the speciation of iron oxides can not be achieved by traditional mineralogical analysis methods like X-ray diffraction (XRD). In this study, diffuse reflectance spectroscopy (DRS) was employed to distinguish and quantify hematite and goethite. Measurements were conducted using a Perkin-Elmer lambda 900 spectrophotometer (Perkin-Elmer
Corp., Norwalk, CT) equipped with a diffuse reflectance attachment. Analyses were performed for spectra in the range from 400 to 700 nm with an interval of 2 nm. Detailed procedures have been well described previously (Ji et al., 2002; Lu et al., 2017). Hematite and goethite are the two main coloring agents in mineral dusts and are characterized by distinct colors, red and yellow, respectively. Hematite was chosen to be quantified rather than goethite, because hematite is a more intense (effective) coloring agent than goethite. The limit of detection for Hm can be as low as 0.01% by weight (Balsam et al., 2014; Deaton and Balsam, 1991; Ji et al., 2002; Lu et al., 2017).

A set of calibration samples containing known hematite were measured. Then the percent reflectance in red color band (630-700 nm, redness) was used as an independent variable in a transfer function for calculating hematite, which was established through regression as following:

\[
\text{Hm (wt. %)} = 1 \times 10^{-6} \cdot e^{27.37 \cdot \text{Redness}} \quad (R^2_{\text{adj}} = 0.9301, \text{RMSE} = 0.3018)
\]

Assuming the CBD-extracted Fe (in the form of iron oxides) are only constituted by hematite (Hm) and goethite (Gt), the content of goethite could be calculated using the following equation:

\[
\text{Gt (wt. %)} = 1.59 \times \frac{\text{Fe}_{\text{ox}} - \text{Hm}}{1.43}
\]

The reproducibility standard deviation of reflectance at all wavelengths was less than 0.15% (Lu et al., 2017).

### 2.6 Light absorption of cryoconite

Measurements of light absorption were performed using an ISSW spectrophotometer in Lanzhou University, China. The experimental strategy was mainly based on the method described by Doherty et al. (2010) and Wang et al. (2013). The ISSW measurement system are specially designed to be sensitive to light absorption and to avoid the interference of light scattering (Grenfell et al., 2011). The ISSW spectrophotometer could provide the spectral absorption properties of cryoconite, by weighting the transmitted light (I) for a sample and that for a blank filter (I_0). The relative attenuation (x) described by the natural logarithm of I_0/I:

\[
x = \ln \left( \frac{I_0(\lambda)}{I(\lambda)} \right)
\]

The spectrum of light attenuation was further calibrated by a set of black carbon standards (fullerene soot, Alfa Aesar, Inc., Ward Hill, MA, USA) (Fig. S1). Because the ISSW spectrophotometer is only sensitive to the signal of light absorption but not scattering of LAIs on filters due to its integrating sandwich structure, the light attenuation by samples on filter at specific wavelengths will be converted to equivalent BC mass loading (L_{BC}, µg C cm^{-2}), which allows to calculate absorption optical depth \(\tau(\lambda)\) by cryoconite: \(\tau = L_{BC} \beta_\lambda\). Where \(\beta_\lambda\) is mass absorption coefficient (MAC) of standard black carbon (i.e., fullerene, 6.3 m² g⁻¹ at 550 nm) (Grenfell et al., 2011; Zhou et al., 2017). Then the light absorption capacity of cryoconite was calculated by dividing the absorption optical depth by the mass loading on filter:

\[
\text{MAC} = \frac{\tau}{L}
\]
The absorption Ångström exponent (AAE) describes the wavelength dependence of the light absorption by particles (Ångström, 1929). The value of AAE could be obtained by the formula of: \[ \text{AAE} = -\frac{\ln(\tau_1/\tau_2)}{\ln(\lambda_1/\lambda_2)}, \] where \( \tau_1 \) and \( \tau_2 \) are the light attenuation calculated at given wavelengths \( \lambda_1 \) and \( \lambda_2 \), respectively. The standard deviation between repeated measurement ranged from 0.29% to 7.83% (mean value: 2.92%).

3 Results and discussion

3.1 Organic carbon and black carbon contents

The total organic carbon and black carbon mass fractions of the cryoconite from the five glaciers are presented in Figure 3. The most striking feature was that BS exhibited the highest TOC content (9.70 ± 0.99 % in mass faction), about 4 times higher than other four glaciers. Similarly, the BC in the cryoconite from BS were also significantly higher than other glaciers, i.e. UG, LHG, XDK and PL. For the black carbon concentration, BS glacier also has the highest abundance (1.99 ± 0.28% in mass), indicating the strong anthropogenic (fossil fuel and biomass burning) influence there. For the remaining four glaciers, their black carbon contents were comparable, ranging from 0.06 ± 0.01% (in total mass of dried cryoconite) of XDK to 0.13 ± 0.03% of PL. For comparison, Di Mauro et al. (2017) reported the black carbon values in cryoconites from Morteratsch Glacier (Swiss Alps), with the range of 0.30 - 0.4 % in mass faction.

3.2 Abundance of elemental Fe (total) and free-Fe (iron oxides)

The iron contents found in cryoconite samples from UG, LHG, XDK, PL and BS glaciers averaged 4.62%, 4.28%, 3.40%, 4.18% and 4.90%, by mass, respectively (Figure 4). Our data were similar to the previous reported iron contents in dust particles preserved in ice cores across Tibetan Plateau (Wu et al., 2012), which ranged from 3.38% to 5.41%. The iron in the cryoconite on the TP glaciers represents a natural background level. Lower iron contents were found in the dust layers deposited on snow cover in northern Utah, USA (the Wasatch Range), which varied from 1.73 to 2.85% by mass (Reynolds et al., 2014). Given the scarce information of Fe abundance available in the glacier area, we also briefly summarized the data in mineral dust from various desert regions worldwide for comparison (Table 2). The determined Fe contents in desert aerosols from ZBT (ZhengBeiTai) and Yulin in North China were 5.38% and 7.7% in total dust aerosol mass, respectively, somewhat higher than our values of cryoconite over glaciers. The reported values of Fe content from Sahara and Arabian Peninsula generally varied in the range from 2.0 to 11% by mass (Gao et al., 2001; Gomes and Gillette, 1993; Zhang et al., 2015), depending on the locations and the transport process. Recently, Caponi et al. (2017) reported the mass fraction of Fe in global dust aerosols (PM$_{10.6}$) ranged from 2.8% (Namibia) to 7.3% (Australia). In addition, in this study there was no systematic variation of the Fe concentrations with altitude, which indicated that the cryoconite on each glacier was homogeneous mixture. The comparable levels of iron (3.4% - 4.9%) found in the five glaciers in different parts of TP demonstrated
that there is little regional variability of elemental iron in the cryoconites, which is likely related to the mineralogical composition of parent soils.

Because only iron oxides (free iron) could effectively control the absorbing property of mineral materials, the content of free iron is more concerned in radiative and climate modelling. Table 2 shows the means of free and structural iron in cryoconite (percentage in total mass) from the five glaciers, with their standard deviations. Interesting, the highest value of total Fe and the highest free-to-total ratio were found in the samples from BS. The color of BS samples was darker than others visually, and they also present the highest TOC contents among the five glaciers (Fig. 3).

The ratios of free to total iron for the five glaciers ranged from 0.31 to 0.70 (Table 2). That means substantial Fe are trapped in the crystal lattice (i.e. structural Fe) and has no direct relationship with the light absorption. This finding is generally in agreement with that of Lafon et al. (2004) and Caponi et al. (2017) for desert aerosols (Table 2). Namely, only about half total-iron is under the form of iron oxides. Therefore, our result clearly demonstrates that the total iron is not suitable to be directly used in the albedo and radiative modelling, although this has been a common practice in previous research (Kaspari et al., 2014; Wang et al., 2013). If this point was considered, the contribution of iron-containing minerals to the total light absorption on the glacier surface will decrease almost 50%, namely, the other light-absorbing components like black carbon and brown carbon should account for much larger fraction correspondingly.

### 3.3 The speciation of iron oxides

In previous modelling studies of the dust radiative forcing, hematite was usually assumed to be the major absorbing iron oxides (Sokolik and Toon, 1999). However, in this study goethite was found more abundant than hematite for all the five glaciers (Fig. 4 and Table 2). The goethite in total iron oxide mass ranged from 81% (for XDK) to 98% (for BS), showing that goethite is the predominant form of iron oxides. The ratios of goethite to hematite in our study were even higher than those reported for desert aerosols (Table 2) (Formenti et al., 2014; Lafon et al., 2006; Shen et al., 2006). For example, Lafon et al (2004) reported the iron oxides in the dust from Northwest China with about half of total iron in the form of iron oxides, and the abundance of goethite (73% of the total iron oxide mass) was higher than hematite (27%). Shen et al. (2006) determined comparable goethite/hematite composition data in dust aerosols from North China. i.e. 64% for goethite in total iron oxide mass and 36% for hematite in Dunhuang, 63% and 37% in Yulin, and 68% and 32% in Tongliao, respectively. While in the dust samples collected on the snow from American West (Wasatch Range, Utah), the amounts of goethite and hematite in dust samples were found roughly equal using Mössbauer spectroscopy (Reynolds et al., 2014).

Beside the pedogenic characteristics, the dominance of goethite over hematite may be also ascribed to the glacier surface environment. Goethite formation is favored in moist and cool conditions, while hematite commonly occurs in warm and dry environment (Reynolds et al., 2014). That is also why the ratios of goethite to hematite were frequently used as indicators of paleoclimate (e.g. precipitation and temperature) (Schwertmann, 1971). Taking into account the cold and humid conditions on the glacier surface, mineralogical transformation of hematite to goethite is highly expected to happen, resulting
more goethite. Microorganisms may also play a crucial role in the degradation of rocks and minerals in terrestrial environments through changing the transformation rates, pathways, and even the end products (Fru et al., 2012). This is especially plausible for BS glacier with the highest TOC content (an indicator of high microbial activity) as well as iron oxides, but more research is needed to test this hypothesis.

3.4 Contribution to light absorption by cryoconite components

Optical properties of goethite and hematite, including MAC and AAE parameters, are critical to assess their role in the light absorption. A wide range of MAC values have been reported in the literatures. According to the previous work by Alfaro (2004), the mass absorption coefficient of iron oxides (goethite : hematite, 73% : 27%) in dust from Northwest China desert was measured as 0.56 m² g⁻¹ at 660 nm, and it will increase about 6 times at shorter wavelength (325 nm) (AAE ≈ 3). This value was further employed to evaluate the albedo and radiative forcing effect of dust in snow of Himalayas (Nepal) (Kaspari et al., 2014). Recently, Utry et al. (2015) reported the MAC value of hematite (purity > 95%) as 0.54 m² g⁻¹ at 532 nm, based on the measurements of a multi-wavelength photoacoustic instrument. Wang et al. (2013) choose the MAC of goethite of 0.9 m² g⁻¹ (550 nm) and AAE value of 3 to assess the contribution of mineral dust to the total absorption of LAIs in North China snow. Based on the laboratory experiments by ISSW, we determined Fe-specific absorption coefficient using the goethite (Stream Chemicals, Inc.) and hematite standard (Sigma Aldrich, Inc.) (Fig. 5). The calculated MAC values at 450 nm for goethite and hematite were 1.55 ± 0.08 m² g⁻¹ and 1.12 ± 0.11 m² g⁻¹, respectively. And the MAC values at 600 nm were 0.15 ± 0.01 m² g⁻¹ and 0.55 ± 0.03 m² g⁻¹, respectively. It should be noted that the MACs reported here are the values of isolated iron oxides (i.e., hematite and goethite). They are much larger than the MAC values of dust, which range from 0.013 m² g⁻¹ to 0.055 m² g⁻¹ over the world (Caponi et al., 2017).

Here we assumed that the total light absorption was entirely and exclusively caused by three components, i.e. iron oxides (goethite and hematite), BC and organic matters. Compared to dust and black carbon, the composition and sources of organic matters over the glacier surface are complicated. Organic matter was a mixture of soil humic and humic-like matters, biogenic particles (e.g., algae, fungi and plant debris) and biomass/fossil fuel burning emissions (Wu et al., 2016), which are often termed as brown carbon (BrC). Considering its diverse sources and complex composition, in this work we did not assume the specific optical parameters for organic matters. Instead, the relative contributions to absorption by organic matters were obtained by subtracting the portions by iron oxides and black carbon from the total absorption. The optical properties of the latter two components are much certain than light absorbing organics. The mass absorption efficiency and AAE value of BC were assumed to be 6.3 m² g⁻¹ (550 nm) and 1.1, respectively (Grenfell et al., 2011).

Because the light absorption capability of iron oxides and organic matters vary with wavelength, here we calculated the relative absorption of these three components at 450 nm and 600 nm, respectively. As shown in Figure 6, at 600 nm (the right panel), the organic matters dominated the light absorption. And BC was the second contributor to the light absorption, especially for the BS glacier with the highest BC concentration. However, the contribution of iron oxides increases dramatically at
450 nm, especially for goethite, due to their high light absorption ability at short wavelength. For the glaciers except BS, the absorption by goethite was larger than BC and approximately equal to organic matters. The increased contribution by goethite at the shorter wavelength was due to its large AAE value (Zhou et al., 2017), which indicated stronger light absorption at short wavelength. While the relative contribution to absorption by hematite appeared to be constant between different wavelengths.

In general, goethite plays a stronger role at shorter band, causing higher fraction of light absorption than BC. Although iron oxides are much less absorbing than black carbon per unit mass, the much high mass concentration of mineral dust in the natural environment may result in total absorption be larger than BC.

4. Summary and Conclusions

The degree to which mineral dust, especially iron oxides, affect the solar absorption and decreases of snow albedo remain uncertain. Despite their importance, the content and speciation of iron oxides in the cryoconite over the glacial surface has not been reported previously.

The iron abundances in the cryoconite from TP glaciers ranged from 3.40% to 4.90% by mass, which were comparable to the upper continental crust (UCC) composition (3.5% of Fe) (Taylor and McLennan, 1995) and implied their natural sources. We further separated and determined iron oxides (free Fe) using the Citrate-Bicarbonate-Dithionite method. The ratios of free to total iron for the five glaciers ranged from 0.31 to 0.70. That meant substantial amounts of Fe were trapped in the crystal lattice (i.e. structural Fe) and had no direct influence on the light absorption. Our result clearly demonstrated that the total iron was not suitable to be directly used in the albedo and radiative modelling, although this was a common practice in previous studies. The iron oxides were further quantified into goethite and hematite, the two major species. The goethite content in iron oxides (in mass fraction) ranged from 81% (XDK) to 98% (BS), showing that goethite was the predominant form of iron oxides. Four glaciers (i.e. UG, LHG, XDK and PL) in the main body of TP demonstrated similar patterns regarding the iron oxide abundance and speciation. However, Baihui glacier was unique in these aspects. Baishui glacier is located at the southeastern margin of TP and closely adjacent to the intensive human activities area, and received more organic matter. The microbes in such environments may perturb the natural distribution of iron.

Taking account of both the abundance of iron oxides and their optical properties, the total light absorption were quantitatively attributed to goethite, hematite, BC and organic matters at 450 nm and 600 nm. Organic matters were found to be the most important light absorber at 450 nm and 650 nm wavelengths. We demonstrated that the goethite played a stronger role than BC at shorter wavelength (i.e. 450 nm) for glaciers except Baishui (BS) glacier. Although iron oxides are much less absorbing than black carbon per unit mass, the high mass concentration of mineral dust in the natural environment may result in total absorption larger than BC. In general, this research provided new observations of the iron-oxides in glaciers, and the results are meaningful for understanding their role on mountain glacier surfaces in Himalayas and Tibetan Plateau, a climate sensitive and environmentally fragile region. Currently, due to the limitation of our field sampling, we can not obtain the spatial distribution of cryoconite on the glacier surface (i.e. the percentage of cryoconite cover in a unit area of glacier...
surface). In future research, we will explore quadrat sampling to reveal the exact change of albedo by cryoconite.

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**References**


Ångström, A.: On the atmospheric transmission of sun radiation and on dust in the air, Geografiska Annaler, 11, 156-166, 1929.


Balsam, W., Ji, J., Renock, D., Deaton, B. C., and Williams, E.: Determining hematite content from NUV/Vis/NIR spectra: Limits of detection, American Mineralogist, 99, 2280-2291, 2014.


Han, Y. M., Cao, J. J., Yan, B. Z., Kenna, T. C., Jin, Z. D., Cheng, Y., Chow, J. C., and An, Z. S.: Comparison of Elemental Carbon in Lake Sediments Measured by Three Different Methods and 150-Year Pollution History in Eastern China, Environmental science & technology, 45, 5287-5293, 2011.


IPCC: Climate change 2013: the physical science basis: Working Group I contribution to the Fifth assessment report of the Intergovernmental Panel on Climate Change, Cambridge University Press, 2014.


Shahgedanova, M., Kutuzov, S., White, K. H., and Nosenko, G.: Using the significant dust deposition event on the glaciers of Mt. Elbrus, Caucasus Mountains, Russia on 5 May 2009 to develop a method for dating and "provenancing" of desert dust events recorded in snow pack, Atmos. Chem. Phys., 13, 1797-1808, 2013.


Table 1 Summary of cryoconite samples collected from five glaciers over Tibetan Plateau and surroundings.

<table>
<thead>
<tr>
<th>Glacier</th>
<th>Description</th>
<th>Coordinates</th>
<th>Elevation (m, a.s.l.)</th>
<th>Sampling date</th>
<th>Sample Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urumqi #1 Glacier (UG)</td>
<td>Tienshan Mount.</td>
<td>43°06´N, 86°48´E</td>
<td>3800-4000</td>
<td>Aug. 2014</td>
<td>14</td>
</tr>
<tr>
<td>Xiaodongkemadi (XDK)</td>
<td>Tanggula Mount.</td>
<td>33°03´N, 92°04´E</td>
<td>5400-5600</td>
<td>Aug. 2014</td>
<td>6</td>
</tr>
<tr>
<td>PaLong #4 (PL)</td>
<td>Southeast TP</td>
<td>29°15´N, 96°56´E</td>
<td>4700</td>
<td>Sep. 2015</td>
<td>2</td>
</tr>
<tr>
<td>Baishui #1 Glacier (BS)</td>
<td>YuLong</td>
<td>27°6´N, 100°11´E</td>
<td>4600-4800</td>
<td>Aug. 2014</td>
<td>9</td>
</tr>
</tbody>
</table>
Table 2 The abundances (by mass) of total iron, free iron, hematite, goethite and their ratios determined in cryoconites from TP glaciers, as well as other data available in the literatures.

<table>
<thead>
<tr>
<th>Locations</th>
<th>Description</th>
<th>Total iron (%)</th>
<th>Free iron (%)</th>
<th>Free-total iron ratio</th>
<th>Hematite %</th>
<th>Goethite %</th>
<th>Gt-Hm ratio</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>UG</td>
<td>Cryoconite</td>
<td>4.62 (±0.22)</td>
<td>1.41 (±0.29)</td>
<td>0.31 (±0.07)</td>
<td>0.24 (±0.02)</td>
<td>1.98 (±0.47)</td>
<td>8.26 (±2.27)</td>
<td>This study</td>
</tr>
<tr>
<td>LGH</td>
<td></td>
<td>4.28 (±0.17)</td>
<td>1.50 (±0.30)</td>
<td>0.35 (±0.07)</td>
<td>0.28 (±0.02)</td>
<td>2.08 (±0.48)</td>
<td>7.54 (±1.83)</td>
<td>This study</td>
</tr>
<tr>
<td>XDK</td>
<td></td>
<td>3.40 (±0.18)</td>
<td>1.93 (±0.48)</td>
<td>0.56 (±0.12)</td>
<td>0.56 (±0.08)</td>
<td>2.44 (±0.69)</td>
<td>4.30 (±0.90)</td>
<td>This study</td>
</tr>
<tr>
<td>PL</td>
<td></td>
<td>4.18 (±0.13)</td>
<td>1.53 (±0.28)</td>
<td>0.37 (±0.08)</td>
<td>0.23 (±0.01)</td>
<td>2.18 (±0.44)</td>
<td>9.63 (±1.88)</td>
<td>This study</td>
</tr>
<tr>
<td>BS</td>
<td></td>
<td>4.90 (±0.21)</td>
<td>3.43 (±0.53)</td>
<td>0.70 (±0.10)</td>
<td>0.10 (±0.01)</td>
<td>5.35 (±0.85)</td>
<td>55.2 (±10.4)</td>
<td>This study</td>
</tr>
</tbody>
</table>

| Swiss Alps | Cryoconite | 4.0 (±0.7) | | | | | | Baccolo et al. (2017) |
| Utah, USA  | Dust on snow | 1.73-2.85 | | | | | | ~1<sup>b</sup> Reynolds et al. (2014) |

- Niger, Sahara  Desert Aerosol  6.3 (±0.9)  2.8 (±0.8)  0.44 (±0.11) Lafon et al. (2004)
- Niger, Sahel  Desert Aerosol  7.8 (±0.4)  5.0 (±0.4)  0.65 (±0.04) Lafon et al. (2004)
- Yulin, China  Desert Aerosol  7.7 (±0.3)  3.7 (±0.4)  0.48 (±0.03) Lafon et al. (2004)
- ZBT, China  Desert aerosol  5.38 (±0.2)  3.0 (±0.2)  0.43 (±0.01) ~3.0 Lafon et al. (2006)
- West Africa  Desert Aerosol  0.38-0.72  0.09-0.26<sup>c</sup>  0.21-0.49<sup>c</sup>  0.96-3.1 Formenti et al. (2014)

<sup>a</sup>Values in brackets represent the standard deviations; <sup>b</sup>Mössbauer spectroscopy indicated roughly equal amounts of hematite and goethite, while reflectance spectroscopy showed goethite was dominant iron oxides; <sup>c</sup>Using X-ray absorption (XAS). Note: the total and free iron data in this table were obtained from ICP-MS, so it refers to the elemental Fe. While hematite and goethite were measured as Fe oxides.
Fig. 1 Surface of mountain glacier (upper) in central Tibetan Plateau and dispersed cryoconite (down) on it.
Fig. 2 Topographic map of the Tibetan Plateau and surrounding, with locations of five representative glaciers. Note: the base map was created by Ocean Data View software (Schlitzer, 2017).
Fig. 3 The total organic carbon (blue) and black carbon (grey) in the total mass of cryoconites on the mountain glaciers of Tibetan Plateau and surroundings.
Fig. 4 The free iron and structural iron contents (elemental Fe) measured in the total cryoconite (dried) mass from TP glaciers (Upper), and goethite and hematite (Fe oxides) contents in the total cryoconite mass (Bottom).
Fig. 5 The mass weighted light attenuation by hematite and goethite. Error bars indicate the standard deviation.
Fig. 6 Apportioning of total light absorption (450 nm and 600 nm, respectively) to black carbon, hematite, goethite and organic matters for the cryoconite from five TP glaciers.