- 1 Reply to the editor and reviewer
- 2 We are very grateful for the comments of editor and reviewer. All responses or changes have been
- 3 shown below and marked in blue.
- 4 Thank you very much
- 5 Kind regards,
- 6 Fangping Yan
- 7 (on behalf of the co-authors)

- 9 In general: The English needs still improvement and should be carefully proofread by a native
- 10 speaker.
- Response: The manuscript was improved by a native speaker (website: www.GeoEditing.co.uk)
- and major modifications were shown in the new version.
- 13 L22: You do not show that this glacier is a "typical" glacier. Write "of a medium sized valley
- 14 glacier" or similar.
- 15 Response: "Typical glacier" was changed to "medium sized valley glacier".
- 16 L26ff: The comparisons of your results with other studies seem to be a bit arbitrary to me as
- 17 already mentioned in my access review. It is specifically unclear why you specifically highlight
- 18 Mendenhall Glacier.
- 19 Response: We deleted the comparisons, and this part has been changed to: "DOC concentrations in
- snowpits, surface snow and surface ice (superimposed ice) were 332  $\pm$  132  $\mu$ g L<sup>-1</sup>, 229  $\pm$  104  $\mu$ g
- 21  $L^{-1}$  and 426  $\pm$  270  $\mu$ g  $L^{-1}$ , respectively. The average discharge-weighted DOC of proglacial
- streamwater was  $238 \pm 96 \mu g L^{-1}$ ".
- 23 L32f: Not the best woring.
- 24 Response: This sentence was changed to "Moreover, there was a significant relationship between
- 25 DOC and Ca<sup>2+</sup>, therefore, mineral dust transported from adjacent arid regions likely made
- 26 important contributions to DOC of the glacierized regions, although contributions from
- autochthonous carbon and autochthonous/heterotrophic microbial activity cannot be ruled out."
- 28 L43f: Be aware that not all glaciers contribute to sea-level rise. Especially, the meltwater from
- 29 glaciers draining into endorheric basins on the TP contribute rather to the rise of the lake levels
- 30 (e.g. Neckel et al. 2014).

- 31 Response: We changed this expression to "by contributing to sea levels (Jocob et al., 2012) and
- endorheric basins (Neckel et al., 2014)."
- 33 L51f: Avoid citing too many studies in a row. Be more specific regarding the cited studies.
- 34 Mention e.g. the geographical region.
- 35 Response: This part was changed to: "there are studies on DOC concentrations, ages and
- compositions of glaciers in Alaska (Stubbins et al., 2012; Hood et al., 2009), DOC bioavailability
- of glaciers on the Tibetan Plateau and the Greenland Ice Sheet (Spencer et al., 2014; Lawson et al.,
- 38 2014) and DOC storage and release of the whole glacier regions around the world (Hood et al.,
- 39 2015)."
- 40 L53: "found to be diverse". A bit vague. Be more precise.
- 41 Response: The expression was changed to "The sources of glacier DOC are diverse and include
- 42 autochthonous or *in situ* biological activities (Anesio et al., 2009), allochthonous carbon derived
- 43 from overridden soils and vegetation in subglacial systems (Bhatia et al., 2010); terrestrial inputs
- 44 (DOC deposition from vascular plants and dust) (Singer et al., 2012) and anthropogenic sources
- 45 (Stubbins et al., 2012)."
- 46 L75-78: These statements are too general. The glaciers are not only retreating because of increases
- 47 of temperatures and carbon deposition. Precipitation, the timing of the monsoon, radiation play
- 48 also an important role. Especially regarding glacier changes, there are also valid studies outside
- 49 the ITP group which should be considered in addition or instead.
- 50 Response: Thanks for the suggestions. We reduced the references from ITP group and added
- references from others. Therefore, this sentence was changed to: "The TP has the largest number
- 52 of glaciers at moderate elevations. Most of the glaciers on the TP are experiencing intensive retreat
- because of multiple reasons such as climatic conditions (Kehrwald et al., 2008; Bolch et al., 2012;
- Yao et al., 2012; Kang et al., 2015) and anthropogenic carbonaceous particle deposition (Xu et al.,
- 55 2009; Lau et al., 2010; Nair et al., 2013; Kaspari et al., 2014)."
- L86-92: This is a description of the methodology and should be merged with the methods section.
- 57 Response: Adjusted accordingly.
- 58 L123f: Repetition of "Meanwhile.
- Response: The first "meanwhile" was deleted.
- 60 L126: Do not write "These results will be presented in another article". Why don't you present

- here? Write rather "is subject to future work" or similar.
- Response: Adjusted accordingly.
- 63 L. 189: This sentence should be moved to the description of the study area.
- Response: Adjusted accordingly.
- 65 L190: Write first the results of the measurements. The high concentration support then the
- 66 statement that the high dust concentration is probably due to deserted surroundings and the
- frequent dust storms.
- Response: The statement: "DOC concentrations were probably influenced by desert sourced
- 69 mineral dust deposition from adjacent arid regions and the frequent dust storms" was moved to the
- 70 end of the measurement results.
- 71 L200: "Alaskan glaciers". This is the place to be more specific not in the abstract.
- 72 Response: The name of the glacier was added here.
- 73 L216: "glacierised"
- 74 Response: Adjusted accordingly.
- 75 L218ff: Here and elsewhere: Mention first your results and put them then into the context of other
- 76 studies.
- 77 Response: Thanks a lot for the suggestions. Modifications were made for section 3.1.3, 3.2 and
- 78 3.3.3. The modifications were marked in the new version.
- 79 L324-329: Sentence too long and hard to understand. Cut in pieces and improve wording.
- 80 Response: Changed to small pieces, and the statement was modified to: "Based on an average
- 81 proglacial streamwater DOC concentration of 193 μg L<sup>-1</sup> (Table S2) and annual glacial meltwater
- 82 runoff of 66-68.2 km<sup>3</sup> in China (Xie et al., 2006), it was calculated that DOC flux in proglacial
- streamwater of the entire TP glacier was around 12.7-13.2 Gg C ( $Gg = 10^9$  g). This estimate is
- higher than that of DOC deposition (5.6 Gg C) across the glacial region of the TP (Li et al., 2016),
- 85 and agree well with the negative glaciers water balance of the TP. Therefore, the TP glaciers can
- 86 be considered as a carbon source under present environmental conditions."
- 87 332: You have never shown that this glacier is "typical". Rewrite.
- 88 Response: "Typical" was changed to "medium sized valley".
- 89 L342: "for the first time". Unclear what you have done for the first time from this sentence.
- 90 Response: "for the first time" was deleted from the sentence.

- 91 Conclusions:
- 92 The conclusions are slightly too long and reads partly like a discussion. Draw the main
- conclusions of your own study. Avoid referencing to other studies, if not absolutely necessary.
- 94 Response: Thanks a lot for the suggestion. The conclusion part was cut and references were
- 95 removed except Ming et al., 2013, which was used to calculate the radiative forcing of snowpit
- 96 DOC. In addition, many parts belonged to discussion were deleted.
- 97 Reviewers comments:
- 98 Line 337 on: "indicating that mineral dust transported from adjacent arid regions made important
- 99 contributions to DOC of the studied glacierized regions except autochthonous or in situ biological
- activities". This sentence is unclear, and needs rewriting, I'd suggest as follows or similar:
- 101 "indicating that mineral dust transported from adjacent arid regions likely made important
- 102 contributions to DOC of the studied glacierized regions, although contributions from
- autochthonous and heterotrophic microbial activity cannot be ruled out"
- 104 Response: Adjusted accordingly.
- Line 32-34 Similarly, the related text in the abstract is rather clunky as follows and could be
- rewritten similar to above, although the meaning is clear "Based on this finding and the significant
- 107 relationship between DOC and Ca<sup>2+</sup>, the main source of DOC might be desert mineral dust.
- 108 Meanwhile, autotrophic or 34 heterotrophic biological activities and autochthonous carbon could
- also contribute to glacier DOC".
- 110 Response: Adjusted accordingly.
- 111 Line 223 unclear "-4.8-11.4°C, rewrite as -4.8 to 11.4
- 112 Response: Changed.
- Line 238 should be 'supraglacial', rather than 'supraglacier'
- 114 Response: Changed.
- 116 Reference:

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# characteristics of dissolved organic carbon on a typical 182 medium sized valley glacier, northern Tibetan Plateau 183 F. Yan<sup>1,4,5</sup>, S. Kang<sup>1,3</sup>, C. Li<sup>2,3</sup>, Y. Zhang<sup>1</sup>, X. Qin<sup>1</sup>, Y. Li<sup>2,4</sup>, X. Zhang<sup>1,4</sup>, Z. Hu<sup>1,4</sup>, P. Chen<sup>2</sup>, X. Li<sup>1</sup>, B. 184 185 Qu<sup>5</sup>, M. Sillanpää<sup>5,6</sup> 186 <sup>1</sup>Qilian Station for Glaciology and Ecological Environment, State Key Laboratory of Cryospheric 187 Sciences, Northwest Institute of Eco-Environment and Resources, Chinese Academy of Sciences, 188 Lanzhou 730000, China 189 <sup>2</sup>Key Laboratory of Tibetan Environment Changes and Land Surface Processes, Institute of Tibetan 190 Plateau Research, Chinese Academy of Sciences, Beijing 100101, China 191 <sup>3</sup>CAS Center for Excellence in Tibetan Plateau Earth Sciences, Chinese Academy of Sciences, Beijing 192 100101, China 193 <sup>4</sup>University of Chinese Academy of Sciences, Beijing 100049, China 194 <sup>5</sup>Laboratory of Green Chemistry, Lappeenranta University of Technology, Mikkeli, Sammonkatu 12, 195 FIN-50130, Finland 196 <sup>6</sup>Department of Civil and Environmental Engineering, Florida International University, Miami, FL 197 33174, USA 198 Correspondence to: S. Kang (shichang.kang@lzb.ac.cn), C. Li (lichaoliu@itpcas.ac.cn) 199 Abstract. Light-absorbing dissolved organic carbon (DOC) constitutes a major part of the organic 200 carbon in glacierized regions, and has important influences on the carbon cycle and radiative forcing of 201 glaciers. However, few DOC data are currently available from the glacierized regions of the Tibetan 202 Plateau (TP). In this study, DOC characteristics of a typical glacier medium sized valley glacier 203 (Laohugou Glacier No. 12 (LHG Glacier)) on the northern TP were investigated. Generally, DOC 204 concentrations on LHG Glacier were comparable to those in other regions around the world. DOC 205 concentrations in snowpits, surface snow and surface ice (superimposed ice) were 332 ± 132 µg L<sup>-1</sup>, $229 \pm 104 \mu g L^{-1}$ and $426 \pm 270 \mu g L^{-1}$ , respectively, which were slightly higher than those of the 206 207 Greenland ice sheet. DOC concentration of surface ice (superimposed ice) was 426 ± 270 µg L<sup>+</sup>, 208 comparable to that of the Antarctic ice sheet. The average discharge-weighted DOC of proglacial

Concentration, sources and light absorption

streamwater was 238 ± 96 μg L<sup>-1</sup>, which is lower than that of Mendenhall glacier, Alaska. and the annual DOC flux released from this glacier was estimated to be 6,949 kg C yr<sup>-1</sup>, of which 46.2 % of DOC was bioavailable and could be decomposed into CO<sub>2</sub> within one month of its release. The mass absorption cross section (MAC) of DOC at 365 nm was 1.4 ± 0.4 m<sup>2</sup> g<sup>-1</sup> in snow and 1.3 ± 0.7 m<sup>2</sup> g<sup>-1</sup> in ice, similar to the values for dust transported from adjacent deserts. Moreover, there was aBased on this finding and the significant relationship between DOC and Ca<sup>2+</sup>, therefore, mineral dust transported from adjacent arid regions likely made important contributions to DOC of the glacierized regions, although contributions from autochthonous carbon and autochthonous/heterotrophic microbial activity cannot be ruled outthe main source of DOC might be desert mineral dust. Meanwhile, autotrophic or heterotrophic biological activities and autochthonous carbon could also contribute to glacier DOC. The radiative forcing of snowpit DOC was calculated to be 0.43 W m<sup>-2</sup>, demonstrating that DOC in snow need to be taken into consideration in implying the necessity of accounting DOC of snow for accelerating melt of glaciers on the TP.

Key words: dissolved organic carbon, light absorption, LHG Glacier, the Tibetan Plateau

#### 1 Introduction

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Ice sheets and mountain glaciers cover 11 % of the land surface of the Earth and store approximately 6 Pg (1 Pg = 1015 g) of organic carbon, the majority of which (77 %) is in the form of dissolved organic carbon (DOC) (Hood et al., 2015). The annual global DOC release through glacial runoff is around  $1.04 \pm 0.18$  Tg C (1 Tg =  $10^{12}$  g) (Hood et al., 2015). Therefore, glaciers not only play an important role in the hydrological cycle by contributing to sea-level rise level (Jacob et al., 2012) and endorheric basins (Neckel et al., 2014), but also potentially influence the global carbon cycle (Anesio and Laybourn-Parry, 2012; Hood et al., 2015) in the context of accelerated glacial ice loss rates. In addition, a large portion of glacier-derived DOC has proven to be highly bioavailable, influencing the balance of downstream ecosystems (Hood et al., 2009; Singer et al., 2012; Spencer et al., 2014). Although DOC storage in ice sheets is much larger than that of mountain glaciers, the annual mountain glacier-derived DOC dominates the global DOC release (Hood et al., 2015). Currently, there are studies on DOC concentrations, ages and compositions of glaciers in Alaska (Stubbins et al., 2012; Hood et al., 2009), DOC bioavailability of glaciers on the Tibetan Plateau and Greenland Ice Sheet (Spencer et al., 2014; Lawson et al., 2014) and DOC storage and export of the whole glacier regions around the world (Hood et al., 2015)many studies on the concentration, age, composition, storage and release of DOC have been conducted around the world (Hood et al., 2009; Stubbins et al., 2012; May et al., 2013; Bhatia et al., 2013; Lawson et al., 2014; Fellman et al., 2015; Hood et al., 2015). The sources of glacier DOC are diverse and include autochthonous or in situ biological activities (Anesio et al., 2009), allochthonous carbon derived from overridden soils and vegetation (Bhatia et al., 2010); terrestrial inputs (DOC deposition from vascular plants and dust) (Singer et al., 2012) and anthropogenic sources (Stubbins et al., 2012). Research on glacier microbial activity suggests that globally cryoconite holes alone can potentially fix about 64 Gg C per year (Anesio et al., 2009). The sources of glacier derived DOC were found to be diverse (Bhatia et al., 2010; Stubbins et al., 2012; Singer et al., 2012; Spencer et al., 2014), with Meanwhile, there are large variations in glacier DOC concentrations and ages (Singer et al., 2012; Hood et al., 2015; Antony et al., 2011). For example, a study on the Greenland ice sheet showed that the concentration of exported DOC exhibited slight temporal variations during the melting period, with subtly higher values in early May than in late May and July (Bhatia et al., 2013). Additionally, the concentration of total organic carbon in snow across the East Antarctic Ice Sheets exhibited remarkable spatial variations due to the marine source of organic

carbon (Antony et al., 2011). Studies of both radiocarbon isotopic compositions and biodegradable DOC (BDOC) have proposed that ancient organic carbon from glaciers is much easier for microbes to utilize in glacier-fed rivers and oceans, implying that large amounts of this DOC will return to the atmosphere quickly as CO<sub>2</sub> and participate in the global carbon cycle, thereby producing a positive feedback in the global warming process (Hood et al., 2009; Singer et al., 2012; Spencer et al., 2014).

In addition to black carbon (BC), another DOC fraction known as water-soluble brown carbon has also been considered as a warming component in the climate system (Andreae and Gelencsér, 2006; Chen and Bond, 2010). This type of DOC exhibits strong light-absorbing properties in the ultraviolet wavelengths (Andreae and Gelencsér, 2006; Chen and Bond, 2010; Cheng et al., 2011). The radiative forcing caused by water-soluble organic carbon (the same as DOC) relative to BC in aerosols was estimated to account for 2-10 % in a typical pollution area of North China (Kirillova et al., 2014a) and approximately 1 % in a typical pollution area of North China (Kirillova et al., 2013) and at a remote island in the Indian Ocean (Bosch et al., 2014), respectively. Unfortunately, to date, few direct evaluations have been conducted in the glacierized regions around the world, including the Tibetan Plateau (TP), where DOC accounts for a large part of the carbonaceous matter (Legrand et al., 2013; May et al., 2013) and potentially contributes significantly to the radiative forcing—in the glacierized region.

The TP has the largest number of glaciers at moderate elevations. Most of the glaciers on the TP are experiencing intensive retreat because of multiple reasons such as climatic conditions (Kehrwald et al., 2008; Bolch et al., 2012; Yao et al., 2012; Kang et al., 2015) and anthropogenic carbonaceous particle deposition (Xu et al., 2009; Lau et al., 2010; Nair et al., 2013; Kaspari et al., 2014). However, to date, no study has quantitatively evaluated the light absorption characteristics of DOC in the glacierized regions on the TP, despite some investigations of concentrations, bioavailability and sources of DOC (Spencer et al., 2014; Yan et al., 2015). The primary results of these studies have shown that DOC concentrations in snowpits at sites on the northern TP are higher than those on the southern TP (Yan et al., 2015). In addition, a large fraction of the ancient DOC in the glaciers on the southern TP has high bioavailability characteristics highly bioavailable (Spencer et al., 2014). However, knowledge of DOC in TP glaciers remains lacking due to the large area and diverse environments of the TP in contrast to the relatively limited samples and studies. Therefore, this study is numerous snow (n=67), ice (n=42) and proglacial streamwater samples (n=201) were collected from a typical glacier in

the northeastern TP (Laohugou glacier No. 12) based on the preliminary research of snowpit samples (Yan et al., 2015) (Table 1, Fig. 1). The concentrations of DOC and major ions (Ca<sup>2+</sup>, Mg<sup>2+</sup>, Na<sup>+</sup>, K<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, Cl<sup>-</sup>, NO<sub>2</sub><sup>-</sup> and SO<sub>4</sub><sup>2-</sup>) and DOC light absorbance were measured to comprehensively investigate the sources, light absorption properties and carbon dynamics in this glacierized region. The results provide a basis for the study of DOC across the TP and other regions in the future.

#### 2 Methodology

## 2.1 Study area and sampling site

Laohugou Glacier No. 12 (LHG Glacier) (39°05′-40′N, 96°07′-97°04′E 4260-5481 m) is the largest mountain glacier (9.85 km, 20.4 km²) in the Qilian Mountains and is located on the northeastern edge of the TP (Du et al., 2008; Dong et al., 2014a). It divides into two part of western and eastern branches at the elevation of 4560 m a.s.l (Dong et al., 2014a). The glacier is surrounded by extensive large arid and semi-arid regions (sandy deserts and the Gobi desertDesert) and is frequently influenced by strong dust storms (Dong et al., 2014b) (Fig. 1) and covers an area of approximately 53.6 % of the entire LHG glacier basin (Du et al., 2008; Li et al., 2012).

LHG Glacier has typical continental and arid climate characteristics (Li et al., 2012; Zhang et al., 2012b). Precipitation occurs mainly from May to September accounting accounts for over 70 % of the annual total annual amount (Zhang et al., 2012b). The monthly mean air temperature in the ablation zone of the glacier ranges from -18.4°C in December to 3.4°C in July (Li et al., 2012). Like other glaciers on the TP, LHG Glacier has been experiencing significant thinning and shrinkage at an accelerated rate since the mid-1990s (Du et al., 2008; Zhang et al., 2012b).

### 2.2 Sample collection

Two snowpits were dug in 2014 and 2015 at almost the same site-location in the accumulation zone of LHG Glacier. In total, 15 and 23 snow samples were collected from these pits in 2014 and 2015, respectively, at a vertical resolution of 5 cm—for each snowpit. Moreover, 29 surface snow and 42 surface ice samples were collected along the eastern tributary from the terminus to the accumulation zone at an approximate elevation interval of 50 or 100 m—from the terminus to the accumulation zone, and 201 proglacial streamwater samples were collected at the gauge station during the melting period (Fig. 1, Table 1). The concentrations of glacier DOC have been observed to be very low and are prone to contamination, often causing an overestimation of DOC concentrations (Legrand et al., 2013). Therefore, before sample collection, polycarbonate bottles were firstly washed three times by ultrapure

water, then soaked with 1 M HCl for 24 h (Spencer et al., 2009), rinsed three times using ultrapure water, then finally soaked in ultrapure water for over 24 h. during Throughout the sampling periodthe whole sampling procedure, snow samples were collected directly into 125-mL pre-cleaned bottles, surface ice (0-3 cm and 3-5 cm) samples were collected using an ice axe directly into polycarbonate bottles after crushing, while proglacial streamwater samples were filtered immediately after collection before putting being transferred into bottles. All ice and snow sample were filtered as soon as possible after they were melted. To prevent contamination, sampling personnel were careful to avoid touching any other surfaces whilst carrying out the sample collection. During all the processes, the person who takes charge of sample collection should not touch any other things to avoid contamination. Meanwhile, At least one blank was made for every sampling process to confirm that the contamination was low (Table S1). Meanwhile, another batch of samples was also collected for BC analysis following the protocol discussed in detail in our earlier study (Qu et al., 2014); these results will be presented in another articleare subjected to future work. In order to evaluate DOC discharge from the entire TP, DOC concentrations in proglacial streamwater samples from a further five glaciers were also measured during of other five glaciers in monsoon and non-monsoon seasons were measured, respectively (Fig. 1, Table S2).

All the collected samples were kept frozen and in the dark during storage in the field, transportation and in the laboratory until analysis. In addition, four dust fall samples from Dunhuang, a desert location (39°53'-41°35'N, 92°13'-93°30'E) and potential source region for dust deposited on LHG Glacier, were collected to compare the light absorption characteristics of dust-sourced DOC to those of the snowpit and ice samples. Mineral and elemental compositions of desert sands of west China are have been homogenized by aeolian activity (Hattori et al., 2003), so that the dust samples collected in this study are representative of desert sourced dust in west China.

### 2.3 Laboratory analyses

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### 2.3.1 Concentration measurements of DOC and major ions

DOC concentrations were determined using a TOC-5000A analyzer (Shimadzu Corp, Kyoto, Japan) after the collected samples were filteredfollowing filtration through a PTFE membrane filter with 0.45- $\mu$ m pore size (Macherey–Nagel) (Yan et al., 2015). The detection limit of the analyzer was 15  $\mu$ g L<sup>-1</sup>, and the average DOC concentration of the blanks was 32  $\pm$  7  $\mu$ g L<sup>-1</sup>, demonstrating that

contamination can be ignored during the pre-treatment and analysis processing of these samples (Table S1). The major cations (Ca<sup>2+</sup>, Mg<sup>2+</sup>, Na<sup>+</sup>, K<sup>+</sup> and NH<sub>4</sub>) and major anions (Cl<sup>-</sup>, NO<sub>3</sub> and SO<sub>4</sub><sup>2-</sup>) were measured using a Dionex-6000 Ion Chromatograph and a Dionex-3000 Ion Chromatograph (Dionex, USA), respectively. The detection limit was 1  $\mu$ g L<sup>-1</sup>, and the standard deviation was less than 5 % (Li et al., 2007; Li et al., 2010). The average ion concentrations of the blanks were very low and could be ignored (Na<sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup>, F<sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, Cl<sup>-</sup>, NO<sub>3</sub><sup>3</sup> < 1  $\mu$ g L<sup>-1</sup>; NH<sub>4</sub><sup>+</sup> = 1.4  $\mu$ g L<sup>-1</sup>; Ca<sup>2+</sup> = 1.2  $\mu$ g L<sup>-1</sup>).

#### 2.3.2 Light absorption measurements

The light absorption spectra of DOC samples were measured using an ultraviolet-visible absorption spectrophotometer (SpectraMax M5, USA), scanning wavelengths from 200-800 nm at a precision of 5 nm. The mass absorption cross section (MAC) was calculated based on the Lambert-Beer Law (Bosch et al., 2014; Kirillova et al., 2014a; Kirillova et al., 2014b):

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$$MAC_{DOC} = \frac{-\ln|\frac{I}{I_0}|}{c.L} = \frac{A}{c.L} \times \ln(10)$$
 (1)

where  $I_0$  and I are the light intensities of the transmitted light and incident light, respectively; A is the absorbance derived directly from the spectrophotometer; C is the concentration of DOC; and L is the absorbing path length (1 cm).

In order to investigate the wavelength dependence of DOC light absorption characteristics, the Absorption Ångström Exponent (AAE) was fitted by the following equation (Kirillova et al., 2014a; Kirillova et al., 2014b):

$$\frac{A(\lambda_1)}{A(\lambda_2)} = \left(\frac{\lambda_2}{\lambda_1}\right)^{AAE} \tag{2}$$

AAE values were fitted for wavelengths of 330 to 400 nm; within this wavelength range, light absorption by other inorganic compounds (such as nitrate) can be avoided (such as nitrate) (Cheng et al., 2011). The radiative forcing caused by BC has been widely studied (Kaspari et al., 2014; Qu et al., 2014; Ming et al., 2013). Therefore, in this study, using a simplistic model (the following algorithm); in this study, the amount of solar radiation absorbed by DOC compared to BC was estimated:

$$f = \frac{\int_{300}^{2500} I_0(\lambda) \cdot \left\{ 1 - e^{-(MAC_{365} \left( \frac{365}{\lambda} \right)^{AAE_{DOC}} \cdot C_{DOC} \cdot h_{ABL})} \right\} d\lambda}{\int_{300}^{2500} I_0(\lambda) \cdot \left\{ 1 - e^{-(MAC_{550} \left( \frac{550}{\lambda} \right)^{AAE_{BC}} \cdot C_{EC} \cdot h_{ABL})} \right\} d\lambda}$$
(3)

where  $\lambda$  is the wavelength;  $I_0(\lambda)$  is the clear sky solar emission spectrum determined using the Air Mass 1 Global Horizontal (AM1GH) irradiance model (Levinson et al., 2010); MAC<sub>365</sub> and MAC<sub>550</sub> are the mass absorption cross section of DOC at 365 nm and mass absorption cross section of BC at 550 nm, respectively;  $h_{ABL}$  is the vertical height of the atmospheric boundary layer; and AAE<sub>DOC</sub> and AAE<sub>BC</sub> are the Absorption Ångström Exponents (AAEs) of DOC and BC. In this simplistic model, following a previous study, we used MAC<sub>550</sub> = 7.5  $\pm$  1.2 m<sup>2</sup> g<sup>-1</sup> (following Bond and Bergstrom, 2006), AAE for BC was set as 1, and  $h_{ABL}$  was set to 1000 m, which has little influence on the integration from the wavelengths of 300-2500 nm (Bosch et al., 2014; Kirillova et al., 2014a; Kirillova et al., 2014b). It is obvious that the value of f is closely connected to the relative concentrations of DOC and BC.

#### 2.3.3 In situ DOC bioavailability experiment

The bioavailability experiment was conducted <u>during fieldwork</u> from August 17th to 31st, 2015, at the glacier terminus <u>during fieldwork</u>. In brief, surface ice samples were collected in pre-combusted (550°C, 6 h) aluminum basins and melted in the field. The melted samples were filtered through pre-combusted glass fiber filters (GF/F 0.7 µm) into 12 pre-cleaned 125-mL polycarbonate bottles and wrapped with three layers of aluminum foil to avoid solar irradiation. Two samples were refrigerated immediately after filtering to obtain initial DOC concentrations; the others were placed outside at the terminus of the glacier, and 2 samples were refrigerated every 3 days to obtain corresponding DOC values. The BDOC was calculated based on the discrepancies between the initial and treated samples.

#### 3 Results and discussion

## 3.1 DOC concentrations and bioavailability

### 3.1.1 Snowpits

LHG glacier is surrounded by arid and semi-arid regions and frequently influenced by strong dust storms (Dong et al., 2014b) (Fig. 1). Therefore, heavy desert sourced mineral dust deposition contributes to high DOC concentrations on LHG glacier. The average DOC concentration in the snowpit samples was  $332 \pm 132 \,\mu g \, L^{-1}$  (Fig. 2), with values ranging from  $124 \,\mu g \, L^{-1}$  to  $581 \pm \mu g \, L^{-1}$  (Fig. 3). The highest values occurred in the dirty layers (Fig. 3), similar to the pattern observed at the Greenland summit (Hagler et al., 2007) and in glaciers on the southern TP (Xu et al., 2013), indicating that DOC concentrations in the study area were probably influenced by desert sourced mineral dust

deposition from adjacent arid regions and the frequent dust storms. Spatially, our results were higher than those of Xiaodongkemadi Glacier on Mountain Tanggula (TGL) in the central TP and the East Rongbu Glacier on Mount Everest (EV) on the southern TP (Fig. 1) (Yan et al., 2015); however, they follow a similar pattern to that of but similar to the mercury distribution on the TP (Zhang et al., 2012a). In addition, the DOC concentrations on LHG Glacier were also higher than those of Mendenhall Glacier, Alaska\_n glaciers (Stubbins et al., 2012) and the Greenland summit (Table 2) (Hagler et al., 2007).

#### 3.1.2 Surface snow and ice

The average DOC concentration in LHG glacier surface snow was significantly lower than that in surface ice because more impurities are present in the latter (Fig. 2). Like those of the snowpits, DOC concentrations in the glacier surface ice (Fig. 2) were higher than those on the southern TP (Nyainqentanglha Glacier) (Spencer et al., 2014) and subsurface ice (0.5 m beneath the glacier surface) in a European Alpine glacier (Singer et al., 2012) (Table 2) but comparable to that in the surface ice of the Antarctic Ice Sheet (Hood et al., 2015). However, the DOC concentrations in surface snow (Table 2) were higher than those in the Greenland Ice Sheet (Hagler et al., 2007), mainly due to the heavy dust load on LHG Glacier. No significant relationship was found between DOC concentration and elevation for either the surface snow or ice (Fig. S1), suggesting there is no link between altitude and DOC at this glacierne "altitude effect" on DOC in this glacier. Similar weak correlation with altitude at LHG was also found with mercury This finding is similar to the mercury distribution pattern in surface snow of this glacier (Huang et al., 2014). Therefore, the distributions of DOC concentrations in the glacier surface snow and ice were influenced by other complex factors, such as different slopes (Hood and Scott, 2008) and cryoconite holes. Furthermore, DOC concentrations in snow and ice at this glacier were within the range of previously reported values for glacierized regions outside the TP.

#### 3.1.3 DOC bioavailability

Previous studies conducted under controlled conditions (stable temperature) have shown that glacier derived DOC is more bioavailable than terrestrial derived DOC (Hood et al., 2009; Fellman et al., 2010; Spencer et al., 2014). Our The in situ bioavailability experiment results showed that the amount of DOC being consumed decreased exponentially over time ( $R^2 = 0.98$ ) (Fig. 4), with approximately 26.7 % (from 417 µg L<sup>-1</sup> to 306 µg L<sup>-1</sup>) degraded within 15 days during the experiment

(average temperature:  $3.8 \pm 3.7$  °C; range: -4.8 to 11.4 °C). The BDOC would have reached 46.3 % if the experiment duration was extended to 28 days, according to the equation derived from the 15-day experiment (Fig. 4). Despite different incubation conditions, this result agrees well with the reports of BDOC from a glacier on the southern TP (28-day dark incubation at 20 °C, 46-69 % BDOC) (Spencer et al., 2014) and from European Alpine glaciers (50-day dark incubation at 4 °C, 59±20 % BDOC) (Singer et al., 2012). Therefore, the previous results obtained in the laboratory closely reflect *in situ* situation and can be used to estimate the bioavailability of glacier-derived DOC.

### 3.2 Sources of snowpit DOC

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The sources of glacier DOC are diverse and include autochthonous or in situ biological activities (Anesio et al., 2009; Bellas et al., 2013), allochthonous carbon derived from overridden soils and vegetation in subglacial systems (Bhatia et al., 2010); terrestrial inputs (DOC deposition from vascular plants and dust) (Singer et al., 2012) and anthropogenic sources (fossil fuel and biomass combustion) (Stubbins et al., 2012; Spencer et al., 2014). Research on glacier microbial activity suggests that globally only cryoconite holes can potentially fix about 64 Gg C per year (Anesio et al., 2009). Moreover, viral induced mortality at the cost of heterotrophic bacterial community plays a dominant role in carbon cycle and other nutrients transformation in supraglacier ecosystems (Bellas et al., 2013). In this study, major ions were adopted as indicators to investigate the potential sources of snowpit DOC, because the sources of major ions in snowpit samples from Tibetan glaciers have previously been investigated in detail (Kang et al., 2002; Kang et al., 2008; Wu et al., 2011; Yan et al., 2015). Moreover, the DOC profiles in two snowpits varied with the dust content; specifically, DOC concentration of dust layers was much higher than that of clean layers. Furthermore, it was found that DOC and Ca2+ (a typical indicator of mineral dust (Yao, 2004)) were significantly correlated ( $R^2 = 0.84$ , Fig. S2), suggesting that the major source of DOC was desert sourced mineral dust, similarly to the previous investigations of sources of snowpits at this glacier (Yan et al., 2015). Combined geochemical and backward trajectories analysis at LHG Glacier further supports the interpretation that dust particles on the glacier were mainly derived from the deserts to the west and north of the study area (Dong et al., 2014a; Dong et al., 2014b). Despite this potential source, local anthropogenic pollutants (biomass, fossil fuel combustion and other activities) (Yan et al., 2015) and biological activities on glacier surface (Anesio et al., 2009) may also contribute to the glacier DOC.

#### 3.3 Light absorption characteristics of DOC

#### 3.3.1 AAE

The Absorption Ångström Exponent (AAE) is generally used to characterize the spectral dependence of the light absorption of DOC, thereby providing important input data for radiative forcing calculations. The fitted  $AAE_{330-400}$  values ranged from 1.2 to 15.2 (5.0  $\pm$  5.9) for snow samples and from 0.3 to 8.4 (3.4  $\pm$  2.7) for ice samples (Fig. S4). The relatively low  $AAE_{330-400}$  values for ice indicated that the DOC had experienced strong photobleaching caused by long-term exposure to solar irradiation. Previous studies have found that the AAE values of brown carbon in aged aerosols (Zhao et al., 2015) and secondary organic aerosols (SOAs) (Lambe et al., 2013) were much lower compared to that of thethan their respective primary values. Therefore, the wide divergence in AAE values might suggest different chemical compositions of DOC due to multiple factors, such as different sources and photobleaching processes. Regardless, the average AAE value of the snow samples was comparable to that of atmospheric aerosols in urban areas in South Asia (New Delhi, India) (Kirillova et al., 2014b) (Table 2). In general, the  $AAE_{330-400}$  values had a negative relationship with  $MAC_{365}$ , especially in the ice samples (Fig. S4), suggesting that the more strongly absorbing DOC might contribute to lower AAE values, which was also found in otheras has been observed in previous aerosol studies (Chen and Bond, 2010; Bosch et al., 2014; Kirillova et al., 2014b).

### 3.3.2 MAC<sub>365</sub>

The mass absorption cross section at 365 nm (MAC<sub>365</sub>) for DOC is another input parameter for the radiative forcing calculation. The light absorption ability at 365 nm was selected to avoid interferences of non-organic compounds (such as nitrate) and for consistency with previous investigations (Hecobian et al., 2010; Cheng et al., 2011). The MAC<sub>365</sub> was  $1.4 \pm 0.4$  m<sup>2</sup> g<sup>-1</sup> in snow and  $1.3 \pm 0.7$  m<sup>2</sup> g<sup>-1</sup> in glacier ice (Fig. S4), both of which were higher than those of water soluble organic carbon in an outflow in northern China (Kirillova et al., 2014a) and on a receptor island in the Indian Ocean (Bosch et al., 2014). Meanwhile, the values were comparable to DOC concentrations in typical urban aerosols associated with biomass combustion in winter in Beijing, China (Cheng et al., 2011) and in New Delhi, India (Kirillova et al., 2014b) (Table 3). The MAC values for DOC from different sources vary widely. Typically, the MAC<sub>365</sub> of DOC derived from biomass combustion can be as high asreach 5 m<sup>2</sup> g<sup>-1</sup> (Kirchstetter, 2004) (Table 3). Correspondingly, the values for SOAs can be as low as 0.001-0.088 m<sup>2</sup> g<sup>-1</sup> (Lambe et al., 2013). Due to the remote location of LHG Glacier, it was considered that the snowpit

DOC should comprise SOAs with low MAC<sub>365</sub> values; however, the high MAC<sub>365</sub> value of the snowpit DOC indicated that DOC may not be entirely derived from SOAs. Hence, it was proposed that mineral dust-sourced DOC caused the high MAC<sub>365</sub> values in the snowpit samples. For instance, the light absorption characteristics of DOC from both snowpit and ice samples showed similar patterns to those of water soluble organic carbon in dust from the adjacent deserts, further indicating that LHG glacier DOC was transported via desert sourced mineral dust and shared similar light absorption characteristics (Fig. 5). Moreover, the difference in light absorption characteristics (especially for wavelengths larger than 400 nm) between snow/ice samples and aerosols in Beijing, China, also indicated their different sources (Fig. 5). Light absorbance was significantly correlated with DOC concentrations in both snow and ice samples (Fig. S3), indicating that DOC was one of the absorption factors. Nevertheless, the MAC<sub>365</sub> values of surface ice (0-3 cm) were lower than those of subsurface layers (3-5 cm), despite their higher DOC concentrations (Fig. 6), reflecting stronger DOC photobleaching in the surface ice due to the direct exposure to solar irradiation.

### 3.3.3 Radiative forcing of DOC relative to BC

Our results showed that the radiative forcing by DOC relative to that of BC ranged from 2.1 % to 30.4 % (9.5  $\pm$  8.4 %) for snowpit samples and from 0.01 % to 0.5 % (0.1  $\pm$  0.1%) for surface ice samples (Fig. S4). The high radiative forcing ratio of snowpit samples was caused by its higher DOC/BC (0.65) than that of surface ice (0.012) (Fig. S5), and the low ratio of DOC/BC in surface ice was caused by enrichment of BC in surface glacier ice during the intensive ablation period (Xu et al., 2009). Snowpit samples can be considered as broadly representative of to be fresh snow; thus, it is concluded that radiative forcing by DOC is a non-ignorable-trivial contributor in addition to BC in reducing the albedo of a glacier when the glacier is covered by fresh snow. The snowpit samples can directly reflect the wet and dry deposition of atmospheric carbonaceous matter in glacierized regions, and the contribution of radiative forcing of snowpit DOC samples is comparable to that of water soluble organic carbon relative to BC in the atmosphere aerosols to some extent The radiative forcing contributed by water soluble organic carbon relative to BC in aerosols has been proposed to be as high as 2 10 % (Kirillova et al., 2014a), but lower than that of aerosols at the top of the atmosphere for the faster decrease of BC concentrations than brown carbon in the high-altitude atmosphere (. Furthermore, it was estimated that brown carbon accounts for a higher ratio of 20 % of the direct radiative forcing of aerosols at the top of the atmosphere because BC concentrations decrease faster than brown carbon in the high altitude atmosphere (Liu et al., 2014).

### 3.4 DOC export during the melt season

The two-year average discharge-weighted DOC concentration was  $238 \pm 96 \,\mu g \, L^{-1}$  during the melting period, comparable with that of the proglacial streamwater of Mount Nyainqentanglha glacier on the southern TP (Spencer et al., 2014). Seasonally, high DOC concentrations appeared during the low discharge periods (May to July and September to October) (Fig. 7), suggesting that DOC concentrations were slightly enriched to some extent. However, there were no clear diurnal variations in DOC concentrations with discharge, suggesting that the discharge from different parts of the glacier was well mixed at the glacier terminus (Fig. S6).

The seasonal variations in DOC flux were similar to those of the discharge (Fig. 7), indicating that discharge (rather than DOC concentrations) played a dominant role in the DOC mass flux. Hence, the majority of the glacier DOC export occurred during the summer melting season. Over the whole melting season, the annual flux of DOC from LHG Glacier was 192 kg km<sup>-2</sup> yr<sup>-1</sup>, with peak DOC fluxes occurring from mid-late July to late August (70 % of the annual flux). When combined with the value of BDOC determined above, at least 3211 kg C yr<sup>-1</sup> was ready to be decomposed and returned to the atmosphere as CO<sub>2</sub> within one month of its release, producing positive feedback in the global warming process.

When it comes toconsidering the entire TP, it is obvious that proglacial streamwater DOC concentrations (Table S2) showed similar spatial variation to those of snowpit DOC (Li et al., 2016), with high and low value appeared being observed on theat northern and southern TP, respectively, reflecting strong association between good succession of proglacial streamwater DOC concentration and that of snowpit samples. Based on an average proglacial streamwater DOC concentration of 193 µg L¹ (Table S2) and annual glacial meltwater runoff of 66-68.2 km³ in China (Xie et al., 2006), it was calculated that DOC flux in proglacial streamwater of the entire TP glacier was around 12.7-13.2 Gg C (Gg = 109 g) based on an average proglacial streamwater DOC concentration of 193 µg L¹ (Table S2) and annual glacial meltwater runoff of 66-68.2 km³ in China of 66-68.2 km³ (Xie et al., 2006). This estimate; which is higher than that of DOC deposition (5.6 Gg C) at across the glacial region of the TP (Li et al., 2016), and agree well with the negative glaciers water balance of the glaciers of the TP. Therefore, the TP glaciers can be considered as a carbon source under present environmental conditions.

#### 4 Conclusions and implications

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The concentrations and light absorption characteristics of DOC on a typical medium sized valley glacier on the northern TP were reported in this study. The mean DOC concentrations of snowpit samples, fresh snow, surface ice and proglacial streamwater were  $332 \pm 132 \,\mu g \,L^{-1}$ ,  $229 \pm 104 \,\mu g \,L^{-1}$ ,  $426 \pm 270 \mu g L^{-1}$  and  $238 \pm 96 \mu g L^{-1}$ , respectively. These values were slightly higher than or comparable with those of other regions (e.g., European Alps and Alaska) (Singer et al., 2012; Fellman et al., 2015). DOC in snowpit samples was significantly correlated with Ca<sup>2+</sup>, a typical cation in mineral dust, indicating that mineral dust transported from adjacent arid regions likely made important contributions to DOC of the studied glacierized regions, indicating that mineral dust transported from adjacent arid regions made important contributions to DOC of the studied glacierized regions except autochthonous or in situ biological activities. In addition, the light absorption profiles of the snowpit DOC was similar to that of dust from potential source deserts. Based on the previously published radiative forcing data for black carbon in snowpit of LHG (Ming et al., 2013), it was estimated that the radiative forcing caused by snowpit DOC was 0.43 W m<sup>-2</sup>, accounting for around 10 % of the radiative forcing caused by BC. Therefore, in addition to BC, DOC is also an important absorber of agent in terms of absorbing solar radiation in glacierized regions, especially when the glacier is covered by fresh snow. It has also been proven that water-insoluble organic carbon has stronger light absorption ability (Chen and Bond, 2010). Therefore, the total contribution of OC to light absorption in glacierized regions should be higher, which requires further study in the future. Wet deposition is the most effective way of removing carbonaceous matter from the atmosphere (Vignati et al., 2010), and the removal ratio of OC in remote areas is almost the same as that of BC after long range transport from source regions (Garrett et al., 2011). Because snowpit samples directly reflect the wet and dry deposition of carbonaceous matter, it is assumed that the contribution of radiative forcing for water soluble organic carbon relative to BC in the atmosphere in glacierized regions should be close to that of the snowpit samples in this study. Because pProglacial streamwater represented a well-mixed, integrated contribution from different parts of the glacier, so no clear diurnal variations in DOC concentrations were identified. from different parts of the glacier is well mixed, no clear diurnal variations in DOC concentrations have been found. Combined with discharge and the corresponding DOC concentration, it was calculated that approximate 192.0 kg km<sup>-2</sup> yr<sup>-1</sup> of DOC was released from LHG Glacier. It was also estimated that

- approximate 46.3 % of the DOC could be decomposed within 28 days; thus, 3,211 kg C yr<sup>-1</sup> would
- return to the atmosphere as CO<sub>2</sub>, producing providing the potential for positive feedback in the
- 576 warming process. Although the flux of DOC from the studied glacier is small, when it comes to the
- 577 entire TP DOC flux from glaciers of the total TP was around 12.7 13.2 Gg C.

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 Table 1. Sampling information for snow, ice and proglacial streamwater in this study.

Sample type	Sampling time	Resolution*	Sampling site	Number (n)	Index
Snowpit	30th July, 2014	5 cm	4989 m	15	DOC, absorbance, ions
Snowpit	25th August, 2015	5 cm	5050 m	23	DOC, absorbance, ions
Surface fresh snow	4th August, 2014	100 m	4450-4900 m	18	DOC
Surface ice	6th August, 2014	100 m	4350-4900 m	20	DOC
Surface snow	16th July, 2015	50 m	4350-4850 m	11	DOC
Surface ice	15th August, 2015	50 m	4350-4850 m	11	DOC
Surface ice	25th August, 2015	50 m	4350-4600 m	6	DOC, absorbance
Subsurface ice	25th August, 2015	50 m	4350-4600 m	5	DOC, absorbance
Proglacial streamwater	29th-30th July, 2014	2h (day),4h (night)	4210 m	17	DOC
Proglacial streamwater	20th May-9th	Every day	4210 m	184	DOC
	October, 2015				

<sup>755 \*\*</sup> Vertical resolution (snowpit) or horizontal distance (surface snow and ice).

**Table 2**. Comparison of DOC concentrations in snow, ice and proglacial streamwater from the glacier in this study and glaciers in other regions.

Sites	DOC concentration (µg L <sup>-1</sup> )	Sample types	References
Laohugou Glacier (LHG)	$332 \pm 132$	Snowpit	This study
Tanggula Glacier (TGL)	$217 \pm 143$	Snowpit	Yan et al. ( 2015)
Mount Everest (EV)	$153 \pm 561$	$153 \pm 561$ Snowpit	
Mendenhall Glacier, Alaska	190	snowpit	Stubbins et al. (2012)
Greenland Ice Sheet	401 - 57	Snowpit	Hagler et al. (2007)
Laohugou Glacier (LHG)	$229\pm104$	Surface snow	This study
Greenland Ice Sheet	111	Surface snow	Hagler et al. (2007)
Juneau Icefield, Southeast Alaska	100 - 300	Fresh snow/snowpits	Fellman et al. (2015)
Laohugou Glacier (LHG)	$426\pm270$	Surface ice	This study
Mount Nyainqentanglha Glacier	212	Glacier ice	Spencer et al. (2014)
Antarctic Ice Sheet	$460\pm120$	Surface ice	Hood et al. (2015)
Alpine glacier	$138 \pm 96$	Subsurface ice	Singer et al. (2012)
Laohugou Glacier (LHG)	$238 \pm 96$	Proglacial streamwater	This study
Mount Nyainqentanglha Glacier	262	Proglacial streamwater	Spencer et al. (2014)
Mendenhall Glacier, Alaska	$380\pm20$	Proglacial streamwater	Stubbins et al. (2012)

Table 3. Mass absorption cross section (MAC) and Absorption Ångström Exponent (AAE $_{330\text{-}400}$ ) of ice and snow from LHG Glacier and aerosols from other regions.

Site/Source	MAC (m <sup>2</sup> g <sup>-1</sup> )	AAE330-400	λ (MAC)	References
LHG Glacier	$1.4 \pm 0.4 \text{ (snow)}$	$5.0 \pm 5.9 \text{ (snow)}$	265	This study
LHG Glacier	$1.3 \pm 0.7$ (ice)	$3.4 \pm 2.7$ (ice)	365	This study
Biomass smoke	5.0	4.8	350	Kirchstetteret al. (2004)
Secondary organic aerosols	0.001 - 0.088	5.2 - 8.8	405	Lambe et al. (2013)
Wood smoke	0.13 - 1.1	8.6 - 17.8	400	Chen and Bond (2010)
HULIS, Arctic snow	$2.6 \pm 1.1$	6.1 <sup>**</sup>	250	Voisin et al. (2012)
Beijing, China (winter)	$1.79 \pm 0.24$	7.5	365	Cheng et al. (2011)
Beijing, China (summer)	$0.71 \pm 0.20$	7.1	365	Cheng et al. (2011)

 $^{\ast\prime}$  The wavelength range for AAE in this study is 300 - 550 nm.

767 Figure 1. Location map of LHG Glacier No. 12. 768 Figure 2. Average DOC concentrations of ice, snow and proglacial streamwater for LHG Glacier. 769 Figure 3. Variation in DOC concentrations in profiles of studied snowpits. The gray rectangles are dirty layers. 770 Figure 4. Exponential deceases in DOC concentrations during the biodegradation experiment. Note: The blue point 771 is calculated using equations derived from the experimental data (black point). 772 Figure 5. Absorption spectra for DOC in snow and ice of LHG Glacier and the dust from surrounding areas. 773 Figure 6. Comparison of DOC concentrations (A) and  $MAC_{365}(B)$  between surface and subsurface ice. 774 Figure 7. The discharge, DOC concentrations and fluxes exported from LHG Glacier. Note: The concentrations 775 with error bars are used for days with more than one sample.

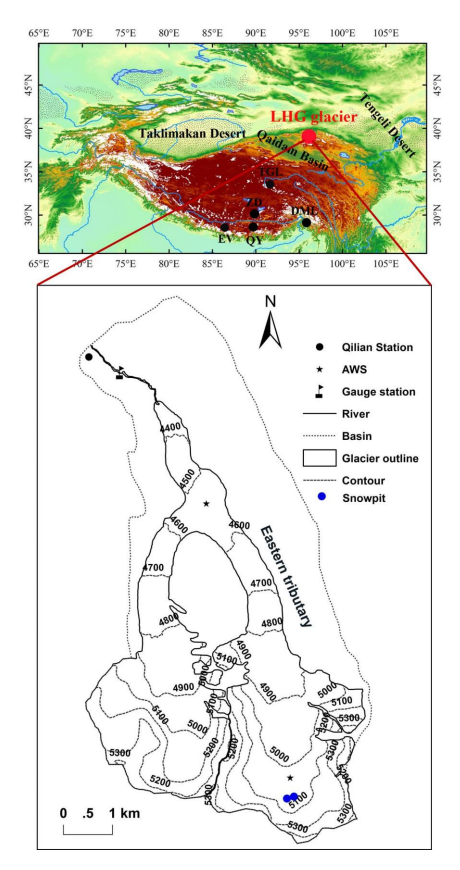


Figure 1. Location map of Laohugou glacier No. 12.

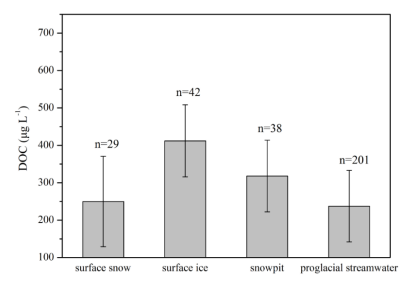
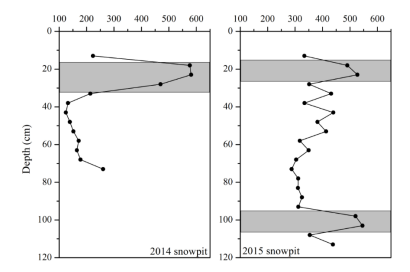


Figure 2. Average DOC concentrations of ice, snow and proglacial streamwater for LHG Glacier.



**Figure 3**. Variation in DOC concentrations in profiles of studied snowpits. The gray rectangles are dirty layers.

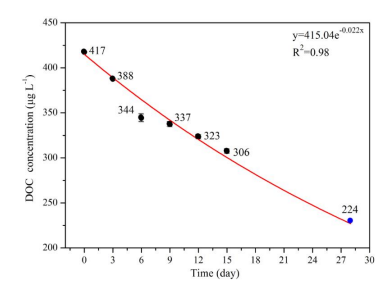


 Figure 4. Exponential deceases in DOC concentrations during the biodegradation experiment. Note: The blue point is calculated using equations derived from the experimental data (black point). Mean values  $\pm$  standard deviations of duplicate treated samples are presented.

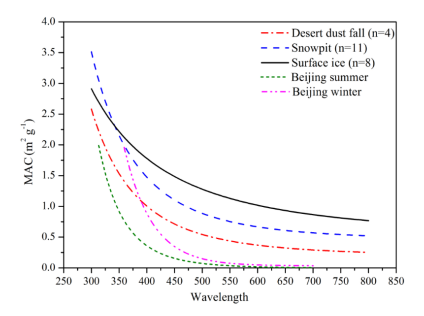
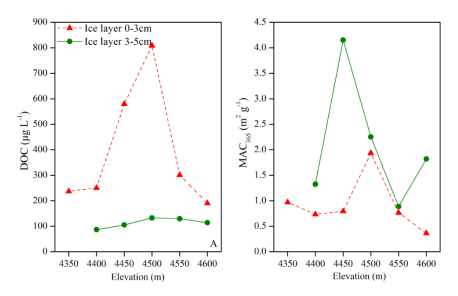
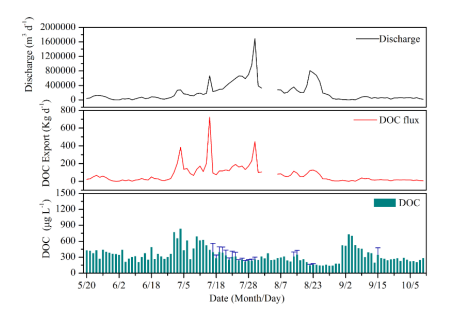


Figure 5. Absorption spectra for DOC in snow and ice of LHG Glacier and the dust from surrounding areas.



 $\textbf{Figure 6}. \ Comparison \ of \ DOC \ concentrations \ (A) \ and \ MAC_{365} \ (B) \ between \ surface \ and \ subsurface \ ice.$ 



**Figure 7**. The discharge, DOC concentrations and fluxes exported from LHG Glacier. Note: The concentrations with error bars are used for days with more than one sample.