Organic carbon fluxes of a glacier surface: a case study of Foxfonna, a small Arctic glacier

Short title: Organic carbon fluxes of an Arctic glacier surface

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Abstract (252 words)

Arctic glaciers are rapidly responding to global warming by releasing organic carbon (OC) to downstream ecosystems. The glacier surface is arguably the most biologically active and biodiverse glacial habitat and therefore the site of important OC transformation and storage, although rates and magnitudes are poorly constrained. In this paper, we present measurements of OC fluxes associated with atmospheric deposition, ice melt, biological growth, fluvial transport and storage (in superimposed ice and cryoconite debris) for a supraglacial catchment on Foxfonna glacier, Svalbard (Norway), across two consecutive years. We found that in general atmospheric OC input (averaging 0.63 ± 0.25 Mg a⁻¹ total organic carbon, i.e. TOC, and 0.40 \pm 0.22 Mg a⁻¹ dissolved organic carbon, i.e. DOC) exceeded fluvial OC export (0.46 \pm 0.04 Mg a⁻¹ TOC and 0.36 \pm 0.03 Mg a⁻¹ DOC). Early in the summer, OC was mobilised in snowmelt but its release was delayed by temporary storage in superimposed ice on the glacier surface. This delayed the export of 28.5% of the TOC in runoff. Biological production in cryoconite deposits was a negligible potential source of OC to runoff, whilst englacial ice melt was far more important on account of the glacier's negative ice mass balance (-0.89 and -0.42 m a⁻¹ in 2011 and 2012, respectively). However, construction of a detailed OC budget using these fluxes shows an excess of inputs over outputs, resulting in a net retention of OC on the glacier surface at a rate that would require c. 3 years to account for the OC stored as cryoconite debris.

Keywords: organic matter, cryoconite, carbon fluxes, cryosphere, DOC, TOC, Svalbard

Introduction

Arctic and mountain glaciers store organic carbon (OC) in dissolved and particulate forms. OC originates from recent and past incorporation into the glacial system, and encompasses both the ancient organic matter from fossil fuels (Stubbins et al., 2012) and the OC currently supplied by active microbial ecosystems (Anesio et al., 2009; Hodson et al., 2008). This OC, especially its dark particulate forms, is most likely contributing to the darkening of glacier surfaces (Cook et al., 2017; Takeuchi et al., 2001; Xu et al., 2012). Climate warming has accelerated the melting of Arctic glaciers (Gardner et al., 2011; Moholdt et al., 2010) and such darkening amplifies this process through the 'bioalbedo' effect (Bøggild et al., 2010; Cook et al., 2017; Takeuchi et al., 2014). Glacier shrinkage leads to greater OC fluxes reaching downstream ecosystems: Hood et al. (2015) estimate such an increase in glacial dissolved OC (DOC) flux for 13%. The main source of this increased DOC release are currently mountain glaciers, which makes their bioavailable OC cycles arguably the most rapidly changing amongst glaciers. OC in those is deemed predominantly allochthonous (Singer et al., 2012; Stibal et al., 2008; Xu et al., 2013). Therefore, the inputs and outputs of OC in such systems depend on the physical processes in the glacier system, and are at least partially climate-driven (by air temperature, precipitation, wind speed and direction changes). However, the link remains poorly defined due to lack of detailed field studies in constrained systems which would incorporate a mass balance scheme to reflect these climate-driven changes.

The work in this field to date comprises several studies which have examined carbon fluxes and storage on ice surfaces (e.g. Chandler et al., 2015; Cook et al., 2010, 2012; Hodson et al., 2010a; Musilova et al., 2017; Stibal et al., 2010, 2012b; Takeuchi, 2002). Further articles report regional or global estimations of OC release

from glaciers (Bhatia et al., 2013; Hood et al., 2015; Liu et al., 2016). Table 1 reports the main findings of such studies, the flux estimates, mainly for OC deposition in snow, removal in runoff and storage within glacial ice. However, due to lack of fieldbased data, none of these studies has incorporated summer precipitation or dust deposition. Furthermore, in order to consider the role of the darkening glacier surfaces in altering glacial OC cycles, the melt release from englacial storage and runoff removal from the glacier surface need to be treated as separate terms, with the supraglacial OC storage as an intermediate stage. Such supraglacial OC storage requires an appreciation of the autochthonous as well as the allochthonous inputs of OC from a range of sources (Stibal et al, 2006) and their likely biological transformation within snow and glacier ice (Stibal et al., 2012a). Therefore, we aimed in this paper to add those usually missing fluxes to the OC cycle study, and to integrate OC inputs, storage and release with a mass balance scheme. By expanding the typical scope of glacial OC fluxes considered, we intended to link the supraglacial carbon cycle to the physical features of the glacial system, with its spatial and temporal variability. As a result, we make a crucial step towards understanding the role played by the supraglacial environment in the overall glacial carbon exchange, and thus also predicting the glacial carbon cycle reaction to climate warming.

These elements were included in a conceptual model for the organic carbon mass balance of a hypothetical glacier (Figure 1). In this paper we quantified these terms for Foxfonna glacier, Svalbard (Norway), using data from two consecutive wholesummer field campaigns.

Materials and methods

Field site

The field site was on the cold-based Foxfonna glacier (Koziol, 2014; Liestøl, 1974), in Nordenskiöld Land, Svalbard (Norway; Fig. 2). A supraglacial catchment covering almost the entire elevation range of the glacier was instrumented (Fig. 3) between June and August in 2011 and 2012. Due to interannual variations in the timing of spring snowmelt, pre-melt snow sampling was carried out in April 2011 and June 2012. The surface area of the catchment varied only slightly between the investigated years, amounting to 1.27 km² in 2011 and 1.31 km² in 2012. Meteorological data (air temperature, relative humidity, wind speed, incident radiation) was measured at the local automatic weather station, however the nearest weather station measuring precipitation was 17.5 km away (WMO station at Svalbard Lufthavn).

The choice of a small, predominantly cold-based glacier is both beneficial and limiting. The data collected here refer to a certain class of glaciers, simpler to understand and representing the likely future for many current glacial systems. However, the OC flux scheme described there does not incorporate subglacial processes, calving and the pronounced autochthonous OC production in the interior of ice-sheets.

Atmospheric deposition (A)

Net atmospheric deposition *A* was calculated as the sum of deposition from "winter" (A^{acc}) and "summer" (A^{abl}) inputs (Equation 1). OC in the late winter snowpack was assumed to represent the sum of OC scavenged from the atmosphere and deposited directly by wet and dry deposition (*sensu* Schlesinger and Bernhardt, 2013) during

the winter snow accumulation period. OC in the late pre-melt snowpack was measured at 10 sites in 2011 and 14 sites in 2012 (Fig. 3). At each site a snow core was collected into a pre-cleaned plastic zip-lock sample bag and slowly melted in the dark. Sample aliquots of 50 mL were sealed in pre-rinsed centrifuge tubes and stored at -20°C for transport to the University of Sheffield (UK). These aliquots were then analysed using a Sievers 900 TOC analyser. At 5 sites the vertical distribution of OC in the snow cover was also studied by extracting samples from 4-6 samples from stratigraphic layers.

During the ablation season, there is frequent washaway of deposited OC, necessitating frequent sampling of atmospheric inputs using dust traps modified from Hall and Upton (1988) and Irvine-Fynn et al. (2012). Trap details are available in Supplementary Information. The atmospheric flux 'A' was then determined as:

$$(1) \quad A = A^{acc} + A^{abl}.$$

Fluvial export (Qc)

Meltwater runoff was sampled every 2–3 days for most of the melt season on the proglacial stream (Site B1 on Fig. 1.). Every other measurement was accompanied by sampling the streams on the glacier snout (B2, B3 & B4 on Fig. 1.). All stream water samples were collected using pre-rinsed centrifuge tubes and frozen at -20°C for transport.

The proglacial stream (B1) water level was measured hourly using a Druck PDCR submersible pressure transducer, enabling discharge to be calculated. It was calibrated with salt dilution discharge gauging on 26 occasions in 2011 and 23 occasions in 2012 (multiple rating curves were used due to channel geometry change during the melt season). The relative standard errors (*RSE*) of the discharge estimates (Hodgkins, 2001) were 13.9% in 2011 and 18.9% in 2012.

Net glacier mass balance contribution (Ic)

The *lc* term represents the loss of englacial OC (see Figure 1) via surface ablation. This was estimated from observations of glacier ice ablation and the spatially distributed OC content of glacier ice in six shallow cores (Fig. 3) collected at the end of summer 2011 using a stainless steel Kovacs ice auger. In each case, a 1-m deep glacier ice section was sampled at 20 cm intervals by collecting ice chips with clean, powder-free gloves. The drill was rinsed with 18 M Ω deionised water between and prior to sampling. The amount of glacier ice ablation was estimated from an annual net glacier mass balance survey conducted at ablation stakes A1–5 (Fig. 3) on 2nd September each year. The glacier-ice samples were processed and stored in the same manner as the snow samples.

Superimposed ice (SI)

In order to measure the maximum transient OC storage in superimposed ice, sampling coincided with its maximum observed thickness at the end of July (just prior to snow cover loss) in both years. Cores were collected from 9 locations in 2012, but just one location in 2011 (E7 in Fig. 3, where 3 cores were taken). Therefore, the 2011 estimate was scaled up using the ratio between deposition measured at this point and the spatial average of deposition in 2012. The samples were stored and processed following the protocol adopted for snow and glacial ice samples.

Biological production (Δbio)

Cryoconite debris on the glacier surface is by far the most biologically active microbial habitat on the glacier surface (Hodson et al., 2008). Net ecosystem production (*NEP*) and respiration (*R*) rates of cryoconite were therefore measured by

means of *in situ* 24 hour incubations of cryoconite following the methodology of Hodson et al. (2010b), using 300 mL sterile Whirlpak® bags as incubation vessels. Biological production (Δbio) was assumed to be equivalent to *NEP*, and the primary productivity (*PP*) was calculated from Equation 2, in order to facilitate comparison with other glacial ecosystem productivity studies:

(2) $\Delta bio = NEP = PP - R.$

Five sets of six incubations were prepared in 2011 and three sets of six incubations in 2012, using three locations at different elevations (A1, A2 and A4 in Fig. 3). Corrections for carbonate dissolution in all incubations were made following major cation determination using Dionex ICS-90 ion chromatography, and by assuming that increases in Ca²⁺ and Mg²⁺ concentrations during the incubations came solely from weathering of carbonates (Hodson et al., 2010b). Thus, the remaining inorganic carbon flux is assumed to equal the TOC flux here.

 Δbio estimates are only presented as TOC fluxes here because the relative importance of DOC and POC (particulate organic carbon or biomass) in the *PP* or *R* fluxes was impossible to determine using the method adopted.

Storage in supraglacial debris (SOC)

Beneath the snow and superimposed ice, cryoconite was assumed to be the dominant supraglacial store of particulate OC, on account of there being no medial moraines. Cryoconite was systematically surveyed by photographing the glacier surface and collecting samples for quantifying OC at 24 locations at 200 m distance intervals (Fig. 3). At each location, \geq 16 pictures were taken using a Sony DSLR-A390 digital camera with a fixed focal length and height above the ice surface (1 m), producing images of *c*. 0.145 m² (± 0.040, 1 SD). The survey was conducted at the

end of 2011 summer season, but was not repeated in 2012 due to early snowfall. Photos were processed using ImageJ software, following Irvine-Fynn et al. (2010), with a manual threshold selection to account for spatial differences in sediment hue. Once the areal extent of the cryoconite was determined, a transfer function was used to estimate the mass (see section *Flux calculations*).

Analytical procedure

Total organic carbon (TOC) and dissolved organic carbon (DOC) analysis were conducted on freshly melted samples using a Sievers 5310 TOC Analyzer. DOC was defined as the 0.7 μ m Whatman GF/F filtrate of the sample. TOC and DOC measurements in the laboratory gave very low and repeatable analytical blanks (0.029 ± 0.009 ppm, 1 SD), while the operational blanks ranged from 0.088 to 0.099 ppm and are given in Table S3 (Supplementary Information). For comparison, the blank-corrected concentration values and their variability are provided in Table S4 (Supplementary Information). POC concentration (C_{POC}) was obtained from equation 3:

 $(3) C_{POC} = C_{TOC} - C_{DOC},$

where C_{TOC} and C_{DOC} are concentrations of the total and dissolved fractions of OC.

In the case of the cryoconite samples, a chemical digestion method was employed after detecting significant mass loss due to clay particle destruction whilst using the Loss On Ignition method. The organic matter content of the cryoconite samples was therefore established gravimetrically following chemical digestion with 1M KOH. From the mass loss, the OC content was estimated using an OC to organic matter mass ratio of 0.254, which was found in cryoconite on Longyearbreen, a glacier located within 20 km radius from Foxfonna (Hodson et al., 2010b).

Flux calculations

Winter deposition (A^{acc}) was estimated from the arithmetic mean loading of the individual snow cores (Equation 4):

4)
$$A^{acc} = \overline{(SWE \times k_S \times OC^{snow})} \times A_c$$
,

where *SWE* = snow water equivalent, OC^{snow} = concentration of TOC or DOC in the snow cores, k_S = correction applied to account for the corer being too short to sample the entire depth of the snowpack, and A_c = catchment area.

Summer deposition (*A^{abl}*) was also calculated as an arithmetic mean using Equation 5:

(5)
$$A^{abl} = \frac{1}{d_A} \sum_n \overline{V \times OC^{DT} \times a \times k_{DT}} \times A_c,$$

where d_A = fraction of total ablation period monitored using dust traps , n = number of sampling intervals (7-10 days), for which A^{abl} was estimated, and V and OC^{DT} are the volume of water and OC concentration measurements collected at each dust trap. The terms *a* and k_{DT} represent the trap area (0.0397 m²) and an efficiency factor as described by Sow et al. (2006).

The calculation of fluvial export (Qc) was achieved using Equation 6:

$$6) \quad Q_{c} = 1/d_{Q} \times Q_{obs} \times \overline{OC},$$

where d_q = fraction of the runoff season monitored for discharge, Q_{obs} = total observed water flux and \overline{OC} = arithmetic mean of DOC or TOC concentration. The arithmetic mean was suitable for the total flux calculation because of the lack of

dependence of \overline{oc} upon Q_{obs} . The OC concentration values were deduced from the streams at the glacier snout (B2, B3 and B4 sites).

The release of OC from glacier ice (Ic) was calculated using Equation 7:

(7)
$$Ic = \sum_{i} (Bn_i \times \overline{OC}^{GI}_i \times A_{Z_i}),$$

where \overline{OC}_{i}^{GI} = mean OC content of the melted depth of glacier ice, Bn_i = total depth

of melt, A_{Z_i} = surface area of each elevation zone. This equation assumes that firn,

the previous year's superimposed ice and glacier ice have the same OC concentration, which is unlikely. However, the very small accumulation area of the glacier (see section *Glacial ice melt (Ic)* of *Results*) means contributions from both firn and "old" superimposed ice were negligible in this study.

The transient OC store in superimposed ice (*SI*) was not included in the annual budget calculations, but was in any case estimated from equation 8:

(8) $SI = \overline{OC}^{SI} \times d_{SI} \times 0.85 \times A_c$,

where \overline{OC}^{SI} = arithmetic mean OC concentration in each superimposed ice core, d_{SI}

= superimposed ice thickness, and 0.85 = assumed density in g cm⁻³.

Estimates of Δbio assumed that *NEP* was constant in time and equal to the average of measured daily values ($\overline{NEP_d}$). The spatial coverage of cryoconite activity was calculated using Equation 9 (after Cook et al., 2012; Hodson et al., 2010a):

(9) $\Delta bio = \overline{NEP_d} \times \sum_{i=1}^n (d \times a \times c \times M_d),$

where d = number of days when a particular fraction of the glacier surface was free from snow; c = mean proportion of area *a* that is covered by cryoconite, and $M_d =$ conversion factor from cryoconite area to dry mass, derived from the complete sampling of ten cryoconite holes of known surface area (0.059 g cm⁻²) (Cook et al., 2010). The term *d* was estimated from oblique photos of the glacier surface taken every 1–4 days (*n* being the number of those periods), while c was deduced from the 2011 photographic survey across the glacier, mentioned in section *Storage in supraglacial debris (SOC)* of *Materials and methods*.

The above terms were then combined according to equation 10 to provide the total carbon mass balance for the glacier (see Hodson et al. (2005) and Fig 1):

(10)
$$A + lc - Qc \pm \Delta bio = \Delta SOC,$$

where $\triangle SOC$ is the change in OC storage on the glacier surface at the end of the ablation season. Since we have accounted for snow and ice, it is most likely to represent a dust layer contributing to cryoconite.

The error estimations for these fluxes are described in *Supplementary Information* and the values are reported together with the results.

Results

Atmospheric deposition (A)

Figure 4 and Table 2 show how the deposition of OC varied in space and time. A^{acc} TOC in 2011 (0.138 ± 0.051 Mg a⁻¹) was much smaller than in 2012 (0.484 ± 0.219 Mg a⁻¹), whilst the opposite was the case for A^{abl} TOC (0.473 ± 0.171 Mg a⁻¹ in 2011 and 0.172 ± 0.048 Mg a⁻¹ in 2012). In the winter of 2012, the distribution of DOC in A^{acc} was uneven, with the middle part of the glacier (500–550 m elevation band) showing the lowest DOC flux. Conversely, in the winter of 2011 the upper elevation bands contained less DOC than the lower part of the glacier (Fig. 4). Otherwise, TOC concentration variability was closely connected to SWE changes (Spearman rank correlation r = 0.482, p<0.018), unlike DOC concentrations (no significant correlation).

Organic carbon transport by meltwater runoff (Qc)

The runoff at the glacier snout in 2011 was generally more stable than in 2012 (Fig. 5), although two slowdowns in runoff progression occurred in August: due to a snowfall event from 10–12 August 2011, and a cooling towards the end of the ablation season, after 27 August 2011. The second half of August 2012 was characterised by a similarly small amount of runoff. The high flows during each year were linked to snow or ice melt in July, sometimes combined with heavy rainfall (between 7 and 14 July 2011 and around 8 August 2012). The importance of melt was emphasised by a significant correlation between discharge and temperature across both years (Spearman rank r = 0.475, p<0.001).

The OC concentrations in runoff (Fig. 5) showed no significant correlation with discharge nor time of the day (for Q, r = -0.219, p>0.05 for DOC and r = 0.272, p>0.05 for TOC; for diurnal cycle, r = -0.192, p>0.05 for DOC and r = -0.108, p>0.05 for TOC; all data merged for 2011 and 2012). There was, however, a progressive increase in both TOC and DOC concentrations with day number across the melt season, especially in 2012 (r = 0.565, p<0.001 for DOC and r = 0.389, p<0.02 for TOC, in pooled data from both seasons). In the Qc fluxes (Table 2), DOC represented the largest fraction of the TOC runoff yield in both years (67.2 ± 6.6 and $89.5 \pm 7.7\%$ in 2011 and 2012, respectively).

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Glacial ice melt (*lc*)

During 2011 and 2012, the net glacier ice mass balance (*Bn*) in the studied catchment was estimated at -0.89 and -0.42 m a⁻¹, respectively. Therefore, glacier ice ablation dominated *lc* for both TOC and DOC (Table 2 and Figure 7) and is likely to do so every year under the present climatic regime. No superimposed ice accumulation was observed at the end of either summer, although there is likely to have been some within the small firn-covered area (<0.3 km²) where crevasses and steep slopes prevented its investigation. However, its limited extent and the steep topography that facilitated water runoff both mean that the quantity of OC stored there was trivial. Therefore, the estimated *lc* fluxes were solely based on *Bn* and the glacial ice OC concentrations, and the 26% difference between the annual TOC fluxes for 2011 and 2012 stemmed mainly from the net mass balance variability. The difference was, however, smaller than the *Bn* difference of 53% due to a high OC concentration in the upper 20 cm layer, which contributed (in 2012) 55% of TOC and 49% of DOC content in the annual *lc* flux.

Transient store of OC in the superimposed ice (SI)

Maximum superimposed ice layer thicknesses during July 2011 and 2012 were 21 (± 2, 1SD) cm and 18 (± 2, 1SD) cm, respectively, indicating a broadly similar quantity of snowmelt refreezing on the glacier surface during the early ablation seasons. This refrozen ice layer was found to represent a flux capturing 0.115 ± 0.042 Mg a⁻¹ TOC and 0.081 ± 0.032 Mg a⁻¹ DOC in 2011, and 0.129 ± 0.048 Mg a⁻¹ TOC and 0.114 ± 0.045 Mg a⁻¹ DOC in 2012, when at maximum thickness. The values for 2011, therefore, show that the temporary storage of OC in SI can be the same order of magnitude as the A^{acc} (Table 2). Furthermore, the storage in *SI* was also spatially variable (Figure 4), although less so than snow inputs to the system.

Biological activity (Δbio)

Table 3 shows that the *NEP* was on average 10.96 (± 5.31 1SD) μ g C g⁻¹ day⁻¹ in 2011 and 23.43 (± 7.17 1SD) μ g C g⁻¹ day⁻¹ in 2012, implying that the ecosystem was therefore net autotrophic. This is in agreement with the results of incubations conducted in 2009 (Hodson, Unpub. Data; locations A2 and A4). Both *R* and *PP* increased with elevation in 2011 and 2012 (data not shown). The final budget contribution of *Δbio* was, however, small: 0.030 ± 0.095 Mg a⁻¹ TOC in 2011 and 0.023 ± 0.071 Mg a⁻¹ TOC in 2012.

Long-term storage on the glacier surface (SOC)

The average cryoconite coverage on the glacier was estimated from the image analysis to be 8.25% (\pm 2.25%, 1 SD), with a debris mass of 56.4 \pm 14.0 Mg according to the mass conversion of 0.059 g cm⁻² and the ablation area. Given the average OC concentration in the debris of 2.03% (\pm 1.20%, 1 SD), the corresponding TOC store in the supraglacial debris was 1.14 \pm 0.23 Mg in the end of summer 2011. The lower part of the glacier had both the greatest proportion of its surface covered with cryoconite and the highest OC mass per unit area gathered in this debris layer (Fig. 6).

Discussion

In this study, the OC deposition, storage and export have been measured in a glacial system over two consecutive years. The combined OC budget calculations produced positive estimates for ΔSOC during both years (Figure 7). Total atmospheric deposition therefore marginally exceeded the fluvial export, and the biological component was small compared to the other fluxes in both years. As a result, ΔSOC was quantitatively similar to *lc* in both years.

Atmospheric inputs

The atmospheric DOC can originate from human activity (local and distant pollutant emissions; see Kozak et al., 2013; Stubbins et al., 2012), vegetation (e.g. plant emissions of volatile organic compounds, or VOCs; Lindwall et al., 2015), and from excretions of the airborne organisms, such as bacteria (Sattler et al., 2001). Due to the sampling of winter deposition in the end of the season, it may also include DOC that comes from post-depositional transformations of POC, e.g. through photodegradation or supraglacial ecosystem activity in decomposing e.g. humic-like compounds. Furthermore, the dry deposition of gaseous organic compounds may also contribute to the final DOC content of snow (see also McNeill et al., 2012). Among POC, bigger microbial cells should be counted (the threshold 0.7 µm particle diameter being higher than cell size of some microbes), alongside wind-blown plant tissue, organic-rich rock and soil dusts, particles related to human activity, and secondary aerosols formed through aggregation of airborne particles.

OC concentrations in the snow were highly variable and subject to the highest error brackets among the OC fluxes, but the means were consistent with other studies (e.g. Xu et al., 2006; Yan et al., 2015). Lafrenière and Sharp (2011) found higher DOC concentrations (280 – 410 ppb) in mature low-latitude snow perhaps indicating the OC content of lower latitude snow might be greater than in the High Arctic due to more abundant vegetation in and around the catchment. The connection between proximity to vegetation and OC content of snow is also consistent with the pattern of mostly allochthonous OC on small glaciers (Stibal et al., 2008) and the observations of macroscopic plant fragments on the surface of Foxfonna. It is likely that windborne transport is also important, given the intensive wind redeposition of snow observed in this site and the interdependence of TOC

concentration and SWE. However, this latter relationship may be partly invoked by the scavenging of OC during snowfall (Kang et al., 2009).

The OC flux from winter snow deposition may be reduced by post-depositional processes, including photo-oxidation during the spring and summer season (Grannas et al., 2004). Photodegradational loss of organic matter from Arctic snowpack of 0.480-4.08 mg C m⁻² d⁻¹ was estimated by Voisin et al. (2012), indicating a potentially important but poorly constrained component of the glacial carbon budget. Furthermore, this process is a potential mechanism of transformation of POC into DOC. Its impact was limited in this study by the late sampling and the consideration of net A^{acc} value.

Rates of atmospheric OC deposition at Foxfonna were highly variable in both space and time. Temporal variation between the two years of monitoring (Table 2 and Figure 7) can be explained by meteorological conditions during the sampling periods. In July 2011, 16 mm of rainfall fell over the catchment in one week, enhancing A^{abl} (Fig. 3). Other unusual meteorological events included winter rainfall and late-spring snowfall in 2012. High precipitation events are usually associated with warm air inflow to Svalbard from southerly latitudes, which may deliver industrial pollutants (e.g. Hodson et al., 2009; Krawczyk et al., 2008; Kühnel et al., 2013). To better predict glacial carbon budgets, a framework for determining the impact of individual meteorological events is required.

Fluvial export

Fluvial export of TOC and DOC from the glacier surface were similar between years (Table 2). The Qc flux for DOC (0.27 g m⁻² yr⁻¹ in both years) was also consistent with Singer et al's (2012) estimate of 0.17 g C m⁻² yr⁻¹ for glaciers in the European Alps. DOC concentrations in runoff reported here are also similar to those

found in other studies (Bhatia et al., 2013; Downes et al., 1986; Hood et al., 2009; Lafrenière and Sharp, 2011) which showed mean annual concentrations of 0.1–4.1 ppm. Our data from Foxfonna (0.35 ppm in 2011 and 0.25 ppm in 2012) are at the lower end of this range, perhaps due to smaller contributions from subglacial and forefield processes compared to other sites (Bhatia et al., 2013; Downes et al., 1986; Hood et al., 2009).

The seasonal increase in TOC and DOC of Foxfonna proglacial runoff may be connected to a shift in hydrological regime. A clear change in runoff amplitudes after 28th July 2012 coincided with the supraglacial catchment losing most of its snow cover, exposing supraglacial stores of OC. Future studies should test this hypothesis because this will impact estimates of OC release from glaciers with consequences for downstream extraglacial ecosystems (Bradley et al., 2016; Fellman et al., 2015; Hood et al., 2015).

Liberation of OC from melting glacier ice and supraglacial biological activity

Mean OC concentrations in Foxfonna glacial ice were 0.933 ppm in the upper 20 cm and 0.410 ppm for the rest of the 1 m core, due to a strong negative down-core gradient. This is higher than reported by Jenk et al. (2009) in a deep Alpine ice core (0.015 – 0.066 ppm). This suggests that the near surface ice might represent a store of OC. Since Foxfonna is a predominantly cold polythermal glacier, the melting isotherm is often very close to the surface. A thin surface layer of ice acts as a shallow perched aquifer through which OC can be transported in flowing meltwater. This is often referred to as the *weathering crust* (Cook et al., 2016; Irvine-Fynn et al., 2011). It is also the most likely environment for the transformation of DOC into POC and the opposite; through dissolution, adsorption, microbial decomposition of organic matter or growth, and photodegradation.

Biological activity in the surface layer of ice may explain some of the additional OC found there, either because OC is leached from cryoconite (Cook et al., 2016) or the weathering crust is a distinct habitat in its own right (Edwards et al., 2014), sometimes referred to as "dirty ice" (Musilova et al., 2017). On Foxfonna, the upper 20 cm of glacial ice is a small store of OC compared to the cryoconite. Even if all the carbon stored in the uppermost glacial ice was produced in situ, the flux related to biological production there would still be an order of magnitude smaller than the glacier-wide supraglacial carbon storage in cryoconite.

The magnitudes of surface OC storage suggest that probably the most important biological OC flux is related to cryoconite. Its *NEP* values on Foxfonna are high compared to those from other glaciers in the area, where a net heterotrophic system (R>PP) has been reported (Stibal et al., 2008; Telling et al., 2010). Stibal et al. (2008) measured *PP* on Werenskiöld glacier, Svalbard, to be *c*. 4.3 µg C g⁻¹ yr⁻¹, whilst Telling et al. (2010) reported *PP* rates of 17.3 µg C g⁻¹ d⁻¹, a mean *R* of 20.1 µg C g⁻¹ d⁻¹ and *NEP* of -1.3 µg C g⁻¹ d⁻¹ for three Svalbard glaciers located near Ny Alesund. Although it is unclear whether cryoconite ecosystems are mostly net autotrophic or net heterotrophic, the finding from this study that the *Δbio* connected to cryoconite is a very small component of the entire OC budget is important. Furthermore, since the *NEP* values found on Foxfonna are an order of magnitude higher than on other Svalbard glaciers, it is likely that the flux is also negligible in other, similar settings.

A second active microbial habitat is surface algal blooms found on snow or ice. In Greenland these were found to promote net autotrophy both on the GrIS and the Mittivakat glacier (Lutz et al., 2014; Yallop et al., 2012), contributing to ice surface darkening. However, intense blooms were not observed on Foxfonna and they are not widespread on Svalbard glaciers. Occasional snow algal blooms, with moderate biomass, do result in red colouration of the snow surface where snow is melting. Therefore, this issue will be treated here as an unquantified error in $\triangle SOC$. Any significant biological activity in the snow would even increase the estimated net retention of OC upon Foxfonna calculated here.

The relation between OC fluxes and its long-term storage

Accumulation and ablation conditions varied between 2011 and 2012, yet ΔSOC was similar. This suggests net retention of OC is typical for this glacier. Cryoconite debris is probably the dominant OC store. This is supported by estimating the quantity of ΔSOC that was particulate (i.e. " ΔSOC^{POC} ") and the magnitude of OC in cryoconite deposits (Equation 11):

(11) $\Delta SOC^{POC} = \Delta SOC^{TOC} - \Delta SOC^{DOC}$,

 ΔSOC^{POC} was 0.310 ± 0.109 Mg a⁻¹ in 2011 and 0.424 ± 0.117 Mg a⁻¹ in 2012. In this case, ΔSOC^{POC} accumulation over *c*. 3 years would account for the cryoconite *SOC* store (1.14 ± 0.23 Mg). Since cryoconite holes are known to persist over timescales of years to decades (Hodson et al., 2010b; Takeuchi et al., 2010) cryoconite is a plausible sink for the excess ΔSOC^{POC} in this study.

Supraglacial processes that disturb and redistribute cryoconite will regulate the residence time of supraglacial OC to some extent. In particular, wash-out or melt-out of cryoconite holes during periods of turbulent heat flux dominated energy balance, rainfall or extreme surface melt and invasion of cryoconite holes by supraglacial rills, streams and crevasses are potential redistribution mechanisms (Cook et al., 2016).

An important part of the glacial OC fluxes is the dissolved organic matter, which comprises 63% of the combined atmospheric flux, 53% of the release from ice melt,

77% of riverine OC and 80% of the transient superimposed ice store. By contrast, it represented between just 54% (2011) and 36% (2012) of ΔSOC , DOC is therefore more likely to be transferred to the glacier forefield by runoff than POC, and thus influence downstream ecosystems (Hood et al., 2009).

Conclusions

This study provides a simultaneous investigation of inputs, throughputs and outputs of organic carbon (OC) upon an Arctic glacier surface. Comparison of the fluxes shows net retention of OC is likely a common occurrence. The physical retention concerns both the allochthonous inputs of atmospheric OC and the OC released from the uppermost layers of glacier ice by surface melt. The estimated rates of storage of particulate OC (POC) were consistent across the two years (2011 and 2012) and represented enough OC to supply the amount of carbon associated with cryoconite debris within *c*. 3 years. This surface carbon may contribute to glacier surface darkening, raising the pressing need to better understand the long-term fate of supraglacial OC storage, and its feedback to melt processes, including the physical redistribution this OC is undergoing.

The processes leading to apparent OC accumulation on the glacier surface need to be balanced by a yet unknown process in the long-term, or else the supraglacial layer of POC would be increasing indefinitely and eventually suppress ablation. Therefore, two potentially important processes involved in the removal of POC from the glacier surface are proposed here: organic-rich debris erosion during extreme melt or rainfall events, and POC transfer to internal storage within the glacier. The former process would drive irregular supplies of OC to the glacier forefield and could supply nutrient to downstream ecosystems. The latter would increase the lag between OC release from melting ice and its removal from the site, which could lead to multiple cycles of the same particle burial and re-emergence on the glacier surface.

The processes leading to OC retention are active also for dissolved organic species, despite these being removed more rapidly from the glacier surface than POC. The highest ratio of dissolved to total organic carbon was found in the refrozen meltwaters (superimposed ice) and supraglacial runoff, and hence the superimposed ice is proposed here as an important temporary store for dissolved OC. Yet, it is unclear how this medium obtains and releases its OC content, for which we recommend further investigation, as we do for the underexplored way of OC transformation which is photodegradation.

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Glacial OC flux type (or concentration which may be used to derive it)		Value	Location	Reference
		Atmospheric depos	sition	
Concentrations in snowpits	DOC	$[\mu g C L^{-1}]$ 519.7 ± 136.3 658.7 ± 78.9 642.3 ± 499.1	Glaciers in China: Tien Shan Glacier No. 1 (TS) Laohugou	(Liu et al., 2016)
Deposition rates	DOC	~19.8 ± 1.0 Gg	Glacier No. 12 (LHG) Total rate for all	(Liu et al., 2016)
		a ⁻¹	Chinese glaciers	
		Meltwater and rui	noff	
Release through melt	DOC	\sim 15.4 ± 6.1 Gg a ⁻¹	Total rate for all Chinese glaciers	(Liu et al., 2016)
Proglacial river runoff concentrations and flux	DOC	0.1.1.1.000	Greenland Antarctic	(Bhatia et al., 2013) (Downes et al.,
		0.1-4.1 ppm	Southern Alaska	1986) (Hood and Berner, 2009)
			Banff National Park, Alberta, Canada	(Lafrenière and Sharp, 2011)
	POC	4.1-13.2 ppm ≈ 0.9 Tg a ⁻¹	Greenland	(Bhatia et al., 2013)
Runoff flux	DOC	0.52 – 0.56 Gg a^{-1} ≈ 0.9 g $m^{-2} a^{-1}$ 0.13–0.17 Tg a^{-1} 0.36–1.52 Tg a^{-1}	Leverett Glacier, Greenland Upscaled to Greenland Ice Sheet	(Lawson et al., 2014)
Runoff flux	DOC	0.34 Gg a ⁻¹ ≈ 0.17 g m ⁻² a ⁻¹	European Alps	(Singer et al., 2012)
Glacier runoff	DOC	1.04 ± 0.18 TgC yr ⁻¹ 0.58 ± 0.07 TgC yr ⁻¹	Worldwide Mountain glaciers	(Hood et al., 2015)
	POC	1.97 Tg yr ⁻¹ 0.70 Tg yr ⁻¹	Worldwide Mountain glaciers	
Biological fluxes				
Supraglacial ecosystem OC fixing	OC	64 Gg a ⁻¹	Global estimate (during the summer season)	(Anesio et al., 2009)
Average glacial	OC	1.3 µg C g⁻¹ d⁻¹	a selection of	(Telling et al.,

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ecosystem			Svalbard	2010)
productivity rates			glaciers	,
Photosynthesis and	OC	10^{1} – 10^{2} Gg C a ⁻¹	Greenland Ice	(Hodson et al.,
respiration fluxes			Sheet	2010a)
		Storage		
Supraglacial storage	OC	$0.213 \text{ g TOC m}^{-2}$	Midtre	Estimate based on
in cryoconite		(or 213 kg TOC	Lovénbreen,	debris mass from
		km⁻²).	Svalbard	Hodson et al.
				(2007) and OC
				content of this
				debris by Telling
				et al. (2012)
Storage in glacier ice	DOC	~3.96 ± 0.87 Tg	Estimated for all	(Liu et al., 2016)
			Chinese glaciers	
Englacial storage	OC	15.4 Pg	Worldwide	(Priscu et al.,
				2008)
Storage in glacier ice	DOC	4.48 ± 2.79 Pg	Global (all	(Hood et al., 2015)
			glaciers and ice	
			sheets)	
		0.07 ± 0.01 Pg	Mountain	
			glaciers	

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Table 2. Organic carbon (OC) fluxes [Mg a^{-1}] of Foxfonna glacier in 2011 and 2012. Note that $\triangle SOC$ is a value calculated from Equation 10 for comparison purposes.

Year	2011		2012	
Flux	TOC	DOC	TOC	DOC
	1		100	DOC
Budget compon	ents			
A A ^{acc}	0.138 (± 0.051)	0.037 (± 0.050)	0.484 (± 0.219)	0.233 (± 0.196)
A ^{abl}	0.473 (± 0.171)	0.377 (± 0.142)	0.172 (± 0.048)	0.145 (± 0.046)
Qc	0.522 (± 0.040)	0.351 (± 0.027)	0.401 (± 0.032)	0.358 (± 0.029)
lc	0.513 (± 0.174)	0.259 (± 0.112)	0.382 (± 0.130)	0.216 (± 0.094)
Δbio	0.030 (± 0.095)	-	0.023 (± 0.071)	-
∆SOC	0.632 (± 0.121)	0.322 (± 0.095)	0.660 (± 0.121)	0.236 (± 0.112)
Temporary storage ^a				
SI	0.099 (± 0.037)	0.070 (± 0.028)	0.114 (± 0.042)	0.101 (± 0.040)

^a This denotes superimposed ice at the maximum of its formation (late July). This term is not a part of the budget sum since ablation in both years exceeded the refrozen layer thickness.

ted

Table 3. Components of biological growth (Equation 4) in µg C g⁻¹ sediment day⁻¹

(± 1 SD).

Ð	
\mathbf{C}	
C	
Y	

Δbio	2011	2012
component		
NEP	10.96 ± 5.31	23.43 ± 7.17
PP	24.14 ± 4.90	36.30 ± 10.09
R	13.19 ± 1.96	12.87 ± 5.98

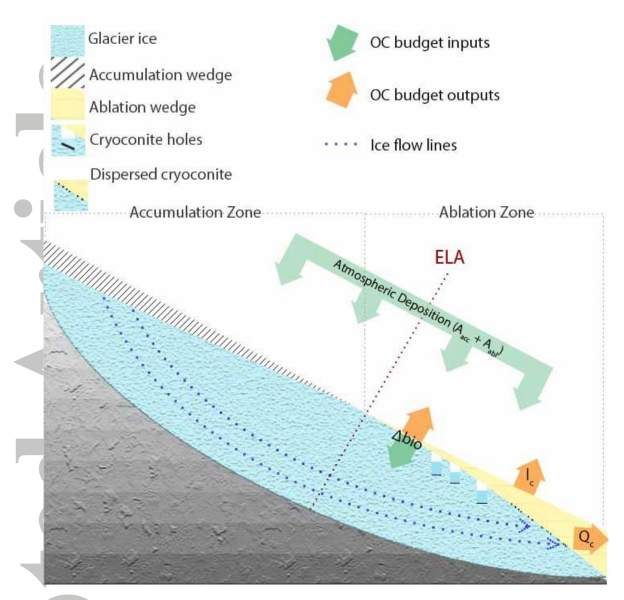


Fig. 1. Conceptual understanding of the important OC fluxes on the glacier surface. Abbreviations as in Equations 1-7.

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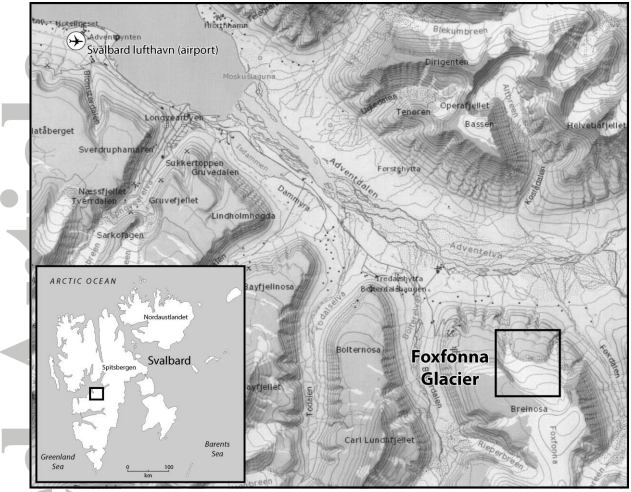


Fig. 2. Location of the study area. Background map source: toposvalbard.npolar.no,

courtesy Norwegian Polar Institute.

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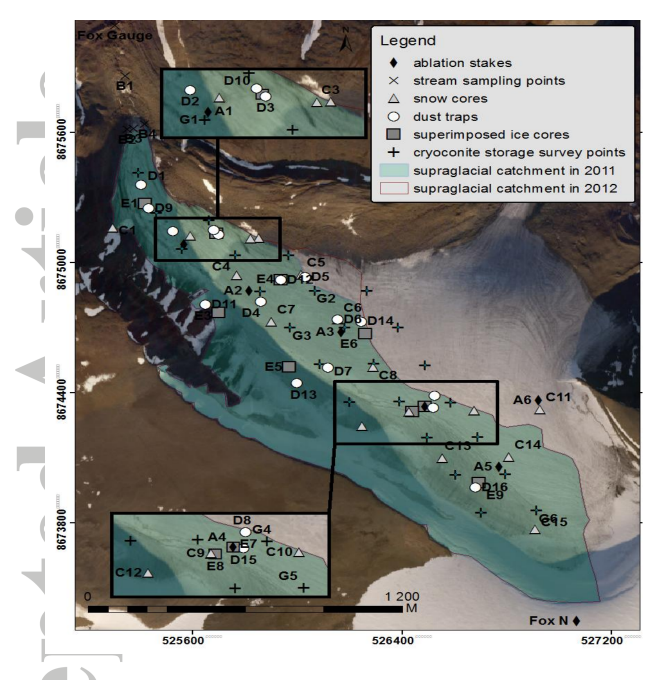


Fig. 3. Sampling points on Foxfonna glacier in 2011 and 2012, showing ablation stakes (A1-6, Fox N), stream sampling locations (B1-4), snow cores (C1-15), dust traps (D1-16), superimposed ice sampling points (E1-9) and glacial ice cores (G1-6). Fox Gauge denotes the location of 2011 discharge monitoring station, and the green catchment area applies to 2011. Background orthophotomap (2006) was made available by Store Norske Spitsbergen Kulkompani AS.

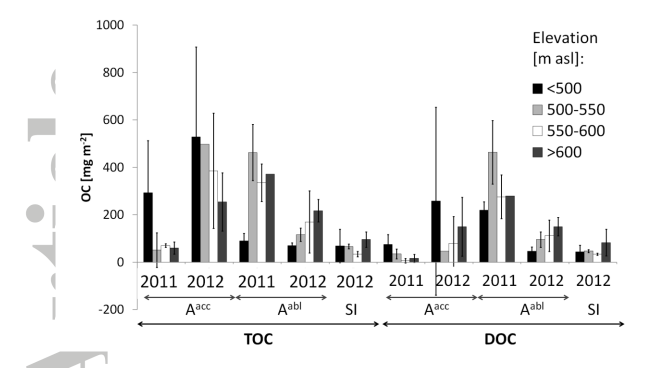


Fig. 4. Variability of atmospheric deposition of OC (per unit area) in the accumulation (A^{acc}) and ablation season (A^{abl}) , and the amount of transient storage as superimposed ice (*SI*): all in 50 m elevation zones on Foxfonna glacier. Where more than one value contributed to the mean, 1 SD error bars are shown.

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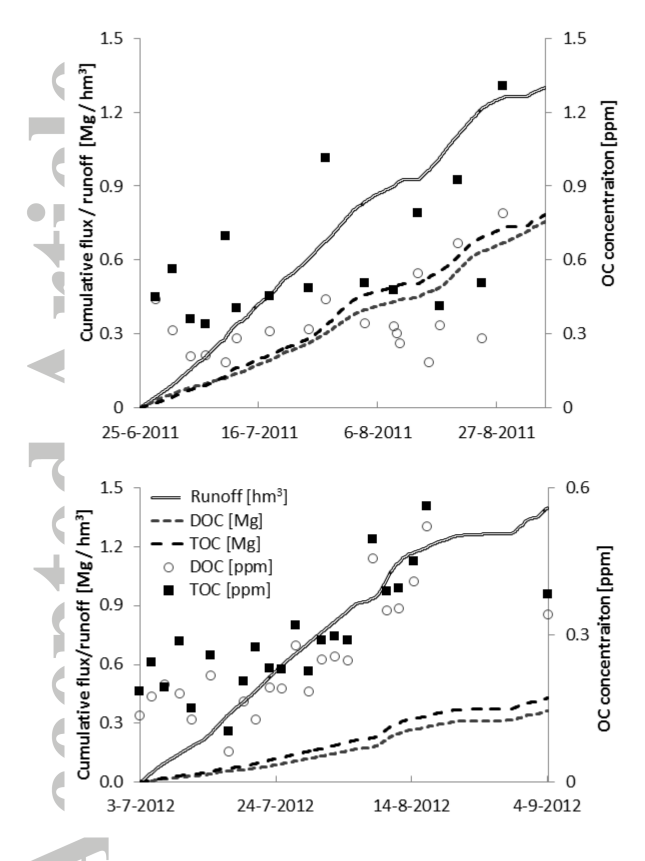


Fig. 5. Cumulative runoff and OC flux in 2011 (top) and 2012 (bottom), complemented with the OC concentrations measured at the glacier snout.

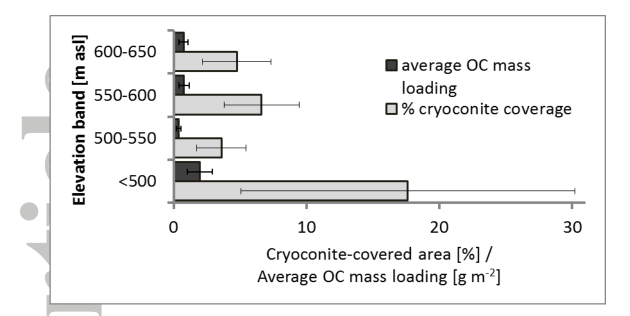


Fig. 6. Cryoconite surface coverage in 50 m elevation bands of Foxfonna glacier, and the supraglacial storage of OC in these zones (as surveyed on 25th August 2011). Error bars as 1 SD (spatial variability).

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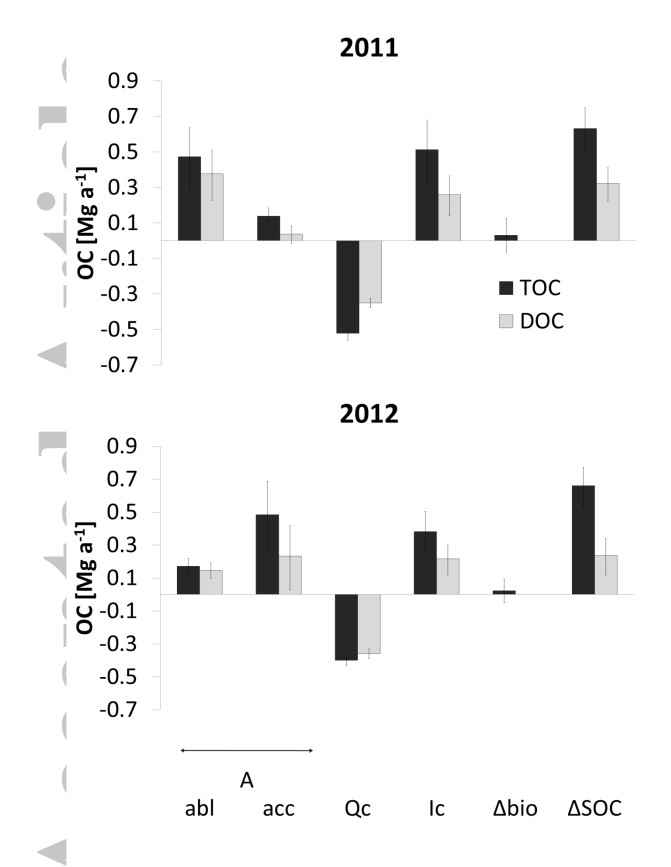


Fig. 7. Organic carbon fluxes of Foxfonna glacier for 2011 and 2012. Abbreviations: see Equations 1 and 2: $abl = A^{abl}$, $acc = A^{acc}$.