High Arctic biocrusts: characterization of the exopolysaccharidic matrix

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Received: 17 June 2019 / Revised: 9 September 2020 / Accepted: 10 September 2020 / Published online: 23 September 2020
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Abstract
Biocrusts can be found in a wide array of habitats, where they provide important ecosystem services. These microbial associations are particularly important in High Arctic environments, where biocrust colonize the newly exposed barren soil after glacier retreat and significantly contribute to soil stabilization and nutrient cycling. Starting from incipient, structurally simple biolayers, they develop in complexity, increasing from the glacier terminus. Starting from a simple community structure, mainly constituted by cyanobacteria, heterotrophic bacteria and fungi immersed in a self-secreted extracellular polymeric matrix (cyanobacterial crusts), they later may recruit mosses and lichens (moss crusts and lichen crusts, respectively). The extracellular polymeric matrix protects the biocrust community from abiotic constraints, notably drought and freezing stress, from external physical harming factors, and from predation. The physicochemical characteristics of the extracellular matrix are related to several of its properties, such as its soil-stabilizing effect and water retention. We analysed the chemical (monosaccharidic composition) and macromolecular (molecular weight distribution) properties of the extracellular polymeric matrix of biocrusts with different morphologies collected in northwestern Spitsbergen, Norway. The uronic acid content and molecular weight (MW) distribution of the extracellular polysaccharidic matrices (EPMs) appeared in accordance with the developmental stages of the biocrusts. The MW distribution also showed significant differences between the samples, possibly reflecting differences in microbial enzymatic activities leading to the degradation of high-MW polymers into smaller compounds. The MW distribution profiles presented some important differences, reflecting differences in environmental conditions and, probably, the seasonal variance in microbial community composition that is known to characterize the environment examined in the present study.

Keywords Biocrusts · Arctic environment · Extracellular polysaccharidic matrix (EPM) · Extracellular polymeric substances (EPS)

Introduction
In arid and semiarid environments, microorganisms can be found in association with biocrusts colonizing the topsoil. Biocrusts are complex associations of cyanobacteria, heterotrophic bacteria, microalgae, fungi and lichens in varying proportions that have a primary ecological role (Yoshitake et al. 2010; Williams et al. 2017). They are common in High Arctic tundra, where they develop on newly exposed soil after deglaciation and exert beneficial effects on vegetation establishment and succession (Breen and Lévesque 2008). Biocrusts contribute to soil water retention, soil stabilization and nutrient cycling in thin and undeveloped soil owing to their photosynthetic and nitrogen-fixation potentials (Yoshitake et al. 2010). They originate from pioneer organisms, recognized mainly as cyanobacteria (Pushkareva et al. ...)
to clarify the relationships between the monosaccharidic investigations conducted in sand microcosms attempted to contribute positively to biocrust establishment. Recent relations between EPS physicochemical characteristics (2012; Li et al. 2013). Some studies strongly suggest correlations between EPS physicochemical characteristics and the capability to aggregate loose soil and, in general, to contribute positively to biocrust establishment. Recent investigations conducted in sand microcosms attempted to clarify the relationships between the monosaccharidic composition of EPSs, their MW distribution and the development and physical stability of biocrusts (Mugnai et al. 2018a, b; Mugnai et al. 2020). For example, some EPS properties, such as a hydrophobic or a hydrophilic nature, are determined by the presence of specific types of sugar constituents and functional groups (Pereira et al. 2009; Colica et al. 2015). However, these types of correlative studies are still hampered by the limited information currently available on the physicochemical characteristics of biocrust EPMs, so it is currently difficult to extrapolate general concepts. A limited number of biocrust typologies from a limited number of habitats were analysed for EPM features. Particularly, analyses were performed on induced biocrusts at different stages of development from semiarid soils in Inner Mongolia, China, and natural biocrusts from North American deserts (Rossi et al. 2012a, b; Colica et al. 2014, 2015; Chen et al. 2014). However, to the best our knowledge, there are no data concerning the characteristics of the EPM of biocrusts from High Arctic environments.

In this work, we investigated, for the first time, the chemical and macromolecular features of the EPM of biocrusts with different morphological traits collected on the northwestern coast of Spitsbergen, Svalbard Archipelago. We hypothesized that the morphological differences of the biocrusts also reflected differences in the physicochemical features of the EPM. Additionally, we compared the detected EPM characteristics with those of biocrusts from hot arid and semiarid environments, with the aim of understanding whether different environmental and climatic conditions significantly affect EPM features.

Materials and methods

Study sites and sample collection

The study area is located in the surroundings of the research town of Ny-Ålesund on the northwest coast of Spitsbergen Island (78.9° N, 11.9° E). The environment is classified as Arctic tundra and characterized either as a lime-demanding, moderate snowbed environment colonized by Saxifraga oppositifolia and Cetraria delisei or as an acidophilous moderated snowbed environment. In addition, wetter sites are dominated by mosses and Deschampsia alpina (Krzyzowska 1989). The Svalbard archipelago is characterized by relatively mild climatic conditions. The mean temperature of Ny-Ålesund is 8 °C in summer and −14 °C in winter, although temperatures as low as −35 °C were also registered. The average annual precipitation is 471 mm, falling prevalently (70%) between October and May, mainly as snow (Mann et al. 1986; Williams et al. 2017). The soil parameters are detailed in Online Resource 1.
Sampling and biocrust morphology

Sampling took place from 27th to 31st August 2010 in several locations around Ny-Ålesund (Fig. 1). The sampling strategy was to collect samples with as much morphological diversity as possible and that were representative of the main morphological types of biocrusts surrounding the research station. Sampling sites were significantly far apart from one another (Table 1) to avoid any possible reciprocal interference between them. Additionally, biocrust were collected in

Fig. 1 Maps of Svalbard Archipelago, Norway, and of the sampling sites. Location of Svalbard (a), Topographic map (b) and indication of the sampling sites around the research town of Ny-Ålesund (c). Maps are used courtesy of the Norwegian Polar Institute

| Sample | Sampling location       | Latitude/longitude | Altitude (metres above sea level) | Distances from the nearest sampling sites (kilometres) |
|--------|------------------------|--------------------|-----------------------------------|-------------------------------------------------------|
| ML-1   | Midtre Lovénbreen      | 78° 53' 39.2" N    | 60                                | 0.23 (ML-2)                                           |
|        |                        | 12° 04' 47.8" E    |                                   | 9.44 (BL-1)                                           |
| ML-2   | Midtre Lovénbreen      | 78° 53' 42.2" N    | 58                                | 9.44 (BL-1)                                           |
|        |                        | 12° 05' 15.7" E    |                                   | 10.1 (OS-1)                                           |
| BL-1   | Blomstrandhalvøya, Kongsfjorden | 78°58'41.1" N | 247                               | 11.7 (OS-1)                                           |
|        |                        | 12° 02' 22.6" E    |                                   | 10.5 (BT-1)                                           |
| OS-1   | Ossian Sars, Kongsfjorden | 78° 55' 46.2" N | 121                               | 10.3 (ML-1)                                           |
|        |                        | 12° 31' 24.2" E    |                                   | 0.17 (OS-2)                                           |
|        |                        |                    |                                   | 7.57 (BT-1)                                           |
| OS-2   | Ossian Sars, Kongsfjorden | 78° 55' 50.0" N | 46                                | 10.3 (ML-2)                                           |
|        |                        | 12° 31'44.5" E     |                                   | 11.8 (BL-1)                                           |
|        |                        |                    |                                   | 10.5 (ML-1)                                           |
| BT-1   | Botnbreen, Kongsfjorden | 78° 53' 32.0" N   | 20                                | 3.22 (ML-1)                                           |
|        |                        | 12° 13' 44.6" E    |                                   | 3.06 (ML-2)                                           |
|        |                        |                    |                                   | 7.76 (OS-2)                                           |
spots that were clear of plants and shrubs. Samples were collected by pushing the open side of sterile plastic Petri dishes (92 mm diameter × 16 mm depth) into the soil, subsequently sealing the Petri dishes closed using parafilm, and transporting the dishes to the laboratory.

The morphological traits of the biocrust samples are shown in Fig. 2.

**Chlorophyll a content**

Chlorophyll a was extracted and quantified according to the method reported in Castle et al. (2011). Briefly, dry crust samples were homogenized, weighed, extracted with ethanol at 80 °C, and then cooled at room temperature for 5 min. The absorbance of the supernatants was measured at 665 nm and 750 nm with a UV–Visible spectrophotometer (Cary, Varian Inc.). The value of the absorbance at 750 nm was subtracted from that at 665 nm to account for the residual scattering of the ethanol solution. The corrected absorbance ($A_{665}^c$) was then plotted with the equation:

$$[11.9035 \times (A_{665}^c) \times V] (g \text{ soil}^{-1}) \times L^{-1}$$

where $V$ is the volume of the analysed extract (expressed in litres) and $L$ is the path length (expressed in cm).

**Extraction and quantification of the extracellular polymeric substances**

EPSs were quantified as total extracellular carbohydrates (TECs) and high molecular weight extracellular carbohydrates (HMWECS), the fraction of TECs having a molecular weight (MW) larger than 100 kDa. Dry biocrust samples were homogenized, weighed and treated with Na$_2$EDTA at 0.1 M for 15 min according to Underwood et al. (1995). Afterwards, samples were centrifuged at 3600×g at 8 °C for 20 min, and a fraction of each TEC-containing
supernatant was stored frozen for further analysis. To recover the HMWECs, the remaining supernatants were precipitated in cold ethanol (70% final concentration) at 4 °C for 16 h, a treatment previously reported to be capable of precipitating EPSs with a MW higher than 100 kDa (Brouwer and Stal 2001). HMWEC-containing precipitates were resuspended in deionized water and stored frozen for further analysis. TEC and HMWEC contents were finally determined using a phenol–sulfuric acid assay (Dubois et al. 1956) and expressed as mg per g soil.

Analysis of the monosaccharidic composition of the extracellular polysaccharides

TEC-containing supernatants were confined in dialysis tubes (14,000 Da MW cut-off, Medicell International, London) and dialyzed against distilled water for 24 h, with two changes of water to remove the Na2EDTA that could interfere with the analytical procedures. Furthermore, TECs were dried in a rotary evaporator and then treated with 2 N trifluoroacetic acid (TFA) at 120 °C for 2 h before finally cooling on ice. Before instrumental analysis, samples were ultra-filtered using centrifugal filters (Ultra-4 Amicon, Billerica, MA) at 8200×g for 20 min and then washed three times with deionized water in a rotary evaporator to remove the residual TFA. The monosaccharidic composition was analysed with a Dionex ICS-2500 ion-exchange chromatograph equipped with an ED-50 detector with a gold-working electrode and a CarboxPA1 column with a length of 250 mm and internal diameter of 4.6 mm (Dionex, Sunnyvale, CA). The eluents were deionized water (A), 0.185 M NaOH (B) and 0.488 M sodium acetate (C). Chromatographic conditions were in accordance with those described in Chen et al. (2014). In the first stage (1st to 7th minute), the eluent combination was 84% A, 15% B, and 1% C; in the second stage (7th to 16th minute), the eluent combination was 0% A, 50% B and 50% C; and in the final stage (16th to 30th minute), the eluent combination matched that of the first stage. The flow rate was constant at 1 mL min⁻¹. Sugars in the extracts were identified and quantified according to reference standards. The composition of monosaccharides of the EPSs was expressed as a molar percent, i.e. as moles of the specific sugar divided by the total moles of the sugars present in the EPSs × 100 (De Philippis and Vincenzini 1998).

Molecular weight distribution profile of the extracellular polysaccharidic matrix

Dry TECs were dissolved in deionized water and clarified by ultracentrifugation at 13,000×g to remove coarse particulates. Samples were analysed using a Varian Prostar HPLC chromatograph equipped with two size-exclusion (SEC) columns, Polysep-GFC-P 6000 and 4000 (Phenomenex Inc., USA) connected in series, and a refractive index detector. Samples were analysed with runs of 65 min at a flow rate of 0.4 mL min⁻¹, according to the method of Colica et al. (2015). Apparent MWs of the samples were identified according to standards of dextran at different average MWs (40,000–5000 kDa; 2000 kDa; 485 kDa; 72.6 kDa; 0.34 kDa) purchased from Sigma Aldrich. The standards were first incubated at 40 °C to dryness by evaporation and then solubilized in distilled water before injection into the chromatograph.

Statistical analysis

Biocrust samples were collected in triplicate (biological replicates, N = 3). Each measurement was performed three times on each biological replicate (technical replicates, n = 3). The significance of the data was evaluated using one-way analysis of variance (ANOVA) at 95% significance. To correlate parameters, linear regression analyses were performed, and $R^2$ and p values are reported for each case. Statistical analysis was performed using GraphPad Prism version 5.00 (GraphPad software, USA). Monosaccharidic composition and MW distribution data were organized in matrices and analysed by principal component analysis (PCA) via the PAST 3 software package (https://folk.uio.no/ohammer/past/).

Results

Biocrust classification and chlorophyll a content

According to the thresholds reported in Lan et al. (2013), sample BL-1 is classifiable as a moss crust, since mosses cover at least 75% of the crust surface, while sample BT-1 is classifiable as a semi-moss crust, being colonized by mosses over 20% of the crust surface. Sample ML-1 is technically a lichen crust, as lichens cover at least 20% of the crust surface, and no moss colonization is present. Samples ML-2, OS-1 and OS-2 are classifiable as cyanobacterial crusts.

The highest contents of chlorophyll a were found in samples BL-1, BT-1 and OS-2. The highest values were found in sample BL-1, followed by BT-1 and then OS-2. The remaining samples (ML-1, ML-2 and OS-1) showed lower and non-significantly different contents (Fig. 3).

Extracellular carbohydrate quantification

The TEC and HMWEC contents of biocrusts are reported in Fig. 4. The average TEC content ranged between 1.45 and 3.48 mg g⁻¹ dry soil. Samples ML-1, BL-1 and OS-2 contained larger TEC amounts. Samples BL-1 and OS-2 also showed the highest contents of HMWECs (TECs with a MW > 100 kDa) among the samples. The percentages of...
TECs accounted for by HMWECs ranged from 21% (sample ML-1) to 39% (sample OS-2). No statistically significant correlations were found between extracellular carbohydrates (TECs or HMWECs) and chlorophyll \(a\) content (\(r^2 = 0.105, p = 0.530\) and \(r^2 = 0.115, p = 0.511\), respectively).

Monosaccharidic composition of the extracellular polymeric substances

The polysaccharidic fraction of the EPM was complex in terms of monosaccharidic composition and up to 13 different types of monosaccharides were identified (Table 2). Glucose (Glc), mannose (Man) and galacturonic acid (GalA) were the most representative constituents. Cumulative amounts of Glc and Man ranged from 27.52% (sample ML-1) to 70.02% (sample BT-1) of the total sugar moles. Glc alone was highest in sample BT-1 and lowest in sample ML-1. GalA was present in the largest amount (34.02%) in sample ML-1. The total uronic acid content (i.e. glucuronic acid (GlcA) and GalA cumulative content) was highest in the EPMs of sample ML-1 (44.78%) and sample BL-1 (38.02%) (Table 2).

PCA (Online Resource 2 and 4) showed that Glc and GalA had the highest loadings in the first two principal components, explaining 79% and 16.8% of the total variance, respectively.

Molecular weight distribution of extracellular polymeric substances

The EPMs extracted from the biocrusts showed high polydispersity, but with significant differences among them with regard to the fractions present in larger amounts (Fig. 5). The EPMs of samples ML-1, OS-1 and BT-1 were prominently composed of low-MW fractions (from 72.6 kDa to 72.6–0.34 kDa), while those of samples OS-2 and BL-1 were equally represented by low-MW (from 72.6 kDa to 72.6–0.34 kDa) and high-MW (larger than 2000 kDa) fractions. On the other hand, the EPM from sample ML-2 was prominently composed of high-MW fractions, mainly distributed in the size ranges higher than 485 kDa.

When treated as variables in PCA (Online Resource 3 and 5), the percentages of polymers with a size of 2000 kDa and in the range 72.6–0.34 kDa explained much of the variance in the MW distribution profile. While molecules with a size of 2000 kDa showed high loadings on the first and second components (0.50 and 0.43, respectively), polymers with a size of 72.6–0.34 kDa showed a high loading only on the first component (−0.81).

Discussion

The analysed biocrust samples presented different morphologies and chlorophyll \(a\) contents, a proxy for phototrophic abundance (Colica et al. 2015). The biocrust samples also presented different TEC contents, which is an index of biocrust development in relation to soil stability (Mazor et al. 1996; Belnap et al. 2008) and possibly an indicator of the soil C pool (Mager 2010). The samples were also characterized by different levels of development from cyanobacterial biocrusts to moss and lichen crusts. The level of development is generally associated with differences in community composition and species relative abundances (Hu and Liu 2003; Liu et al. 2017). However, community succession is allowed or restricted according to semi-equilibrium with environmental stresses and the characteristics of a given location (Li et al. 2013). The presence of mosses or lichens, for example, is highly conditioned by moisture, illumination
and possible natural protection from wind erosion (Lan et al. 2014). Consequently, some cyanobacterial crusts may be prevented from shifting to moss or lichen crusts, implying that LOD does not necessarily coincide with crust age.

Samples BL-1 (moss crust), BT-1 (semi-moss crust) and OS-2 (cyanobacterial crust with scarce moss colonization) showed the highest chlorophyll contents. Chlorophyll a content reflected the differences in moss colonization between the three samples, in the order BL-1 > BT-1 > OS-2. Moss crusts are considered the latest and most mature stage of biocrust LOD, characterized by an increased biomass content (Liu et al. 2013), increased nitrogenase activity, and higher EPS productivity compared to those of cyanobacterial biocrusts (Liu et al. 2017). Additionally, moss crusts are characterized by a higher presence of biomass-degrading microorganisms than early-LOD biocrusts, improving their nutrient status (Maier et al. 2018). Accordingly, in these samples, we detected the largest mean relative amounts of TECs and HMWECs. An exception was sample ML-1 (classifiable as a lichen crust), which, although displaying one of the lowest chlorophyll contents, showed a high TEC content, although not a high HMWEC content. In this case, a high TEC content might support improved C fixation rates, a common feature of lichen crusts (Liu et al. 2017).

Our set of samples differed in HMWEC content. This EPS fraction (MW ≥ 100 kDa) is the direct product of microbial synthesis and strongly contributes to biocrust EPM stability (Mugnai et al. 2018a). Several microbial groups synthesize HMWECs, notably cyanobacteria, which are known for producing high-MW polymers (Pereira et al. 2009). Sample OS-2, a cyanobacterial crust also analysed in another study (Felde et al. 2016), was characterized by the dominance of Gloeocapsa sp., a genus known for producing a large, thick EPS envelope. We did not find any statistical correlation between chlorophyll content and HMWEC content. Conversely, a strong positive correlation between chlorophyll and HMWEC contents was found for biocrusts from hot arid environments, thus suggesting that cyanobacteria contribute to their synthesis (Rossi et al. 2012b). However, the lack of correlation may depend on the more rapid community changes characterizing Arctic biocrusts compared to hot dryland biocrusts. According to Schostag et al. (2015), our study sites are characterized by seasonal variation in phototrophic abundance, owing to the decline in the relative abundance of chloroplasts and cyanobacteria starting from mid-summer, with the lowest relative abundances registered in late summer and autumn. Therefore, chlorophyll content may depend on the time of sampling (late August), in which cyanobacterial abundance is the lowest. Nevertheless, since cyanobacterial sheaths and more condensed EPSs remain to cement soil particles even after death or migration of the producing strain (Zhang 2005; Rossi et al. 2018), the secretion events may have preceded the sampling period.

IEC analysis showed that the polysaccharidic fraction of the EPM contained dominant relative percentages of hexoses (glucose and mannose) and, in the case of samples ML-1 and BL-1, uronic acids. In addition, the deoxy sugars fucose and rhamnose were detected in all the samples, in percentages from 4 to 9%. Except for some differences (see below), the EPMs of the samples showed similar monosaccharidic compositions. Such a composition is superimposable to that of cyanobacteria-dominated biocrusts from Chinese hot arid soils (Wang et al. 2009; Colica et al. 2015). The presence

Table 2 Monosaccharidic composition of the EPSs extracted from the six BSC samples, expressed as molar % (i.e. moles of the specific sugar/total moles of the sugars present in the EPSs × 100)

| Sugar               | Abbreviation | ML-1     | ML-2     | OS-1     | OS-2     | BL-1     | BT-1     |
|---------------------|--------------|----------|----------|----------|----------|----------|----------|
| Fucose              | Fuc          | 4.15     | 5.22     | 9.67     | 6.96     | 4.39     | 3.78     |
| Rhamnose            | Rha          | 5.71     | 5.76     | 9.63     | 9.26     | 4.44     | 4.63     |
| Galactosamine       | GalN         | 0.40     | nd       | 0.73     | 0.47     | 1.36     | 1.93     |
| Arabinose           | Ara          | 0.58     | 1.07     | 0.52     | 1.35     | 0.74     | 0.72     |
| Glucosamine         | GlcN         | 0.68     | 0.63     | 1.38     | 0.90     | 1.32     | 0.57     |
| Galactose           | Gal          | 5.92     | 8.56     | 9.08     | 10.00    | 7.06     | 5.48     |
| Glucose             | Glc          | 8.96     | 38.29    | 17.87    | 29.31    | 11.88    | 54.57    |
| Mannose             | Man          | 18.56    | 21.83    | 18.80    | 25.03    | 19.50    | 15.42    |
| Xylose              | Xyl          | 9.67     | 13.54    | 14.99    | 11.80    | 9.28     | 8.66     |
| Fructose            | Fru          | 0.58     | 5.09     | 0.04     | 0.69     | 1.33     | 0.66     |
| Ribose              | Rib          | nd       | nd       | 9.66     | nd       | 0.69     | nd       |
| Galacturonic acid   | GalA         | 34.02    | nd       | 0.87     | 2.87     | 25.12    | 2.24     |
| Glucuronic acid     | GlcA         | 10.76    | nd       | 6.75     | 1.37     | 12.90    | 1.33     |

Values are means of at least three replicates, and SDs never exceeded 5%

nd Not detected
of uronic acids on one side and rhamnose and fucose on the other confers both a hydrophobic and a hydrophilic character to the EPM, depending on the relative abundances, contributing to the establishment of different types of interactions with the surrounding environment (Pereira et al. 2009; Rossi et al. 2012a). Principal component analysis performed on monosaccharide composition showed that variance in the data was explained by the relative contents of glucose and galacturonic acid (Online Resource 2 and 4). Owing to their hydrophilia, uronic acids confer a water-absorptive character to the EPM, maintaining optimal moist conditions for the biocrust community (Pereira et al. 2009). We previously observed that an increase in the relative abundance of uronic acids was associated with biocrust age (Colica et al. 2015). Our current results evidenced a higher uronic acid contents in samples BL-1 and ML-1, which were a moss crust and a lichen crust, respectively, types of biocrusts considered to be of a later stage of development (Belnap et al. 2008). Galacturonic acid was particularly abundant in the EPM of sample ML-1. Conversely, sample ML-2 did not contain uronic acids. The lack of uronic acids is indicative of an incipient biocrust stage (Colica et al. 2015), also supported by the lower levels of chlorophyll and EPSs.

Contrary to the IEC results, SEC revealed a MW distribution differing from that of biocrusts sampled from other climatic zones. A common trait observed among samples from Inner Mongolian Desert biocrusts is the significant presence of fractions with apparent MWs in the range 2000–485 kDa (Chen et al. 2014; Colica et al. 2015). We did not observe common traits in the MW distribution profiles of the EPMs of the present samples. According to PCA, the variance in the data was explained by the relative contents of polymers with an apparent MW of 2000 kDa and those with an apparent MW between 72.6 and 0.34 kDa (Online Resource 3 and 5). The variation in the abundance of polymers belonging to these two size classes is indicative of different microbial activities between the samples, causing variations in EPM synthetic and degradative patterns (Chen et al. 2014). The differences from biocrusts from hot deserts can probably be explained by the more varied and variable environmental conditions characterizing Arctic tundra. Our samples showed three different MW distribution patterns. In the first, characterizing samples ML-1, OS-1 and BT-1, the EPM contained polymers with MWs primarily between 72.6 and 0.34 kDa, corresponding to small polymers or oligomers. This indicates prevalent polymer-degrading activity and an accumulation of small compounds (Chen et al. 2014). The second MW distribution pattern belonged to samples OS-2 and BL-1, in which the EPM was characterized by both high- (≥ 2000 kDa) and low-MW polymers (in the range 72.6–0.34 kDa). In this case, the polymer-degrading activity is coupled with the synthetic activity of high-MW EPSs by cyanobacteria and other producing species. In a third MW distribution pattern, characterizing sample ML-2, the EPM constituted only high-MW polymers, distributed in several size classes from 450 kDa upwards. This is indicative of the low enzymatic activity and prevailing synthetic activity of high-MW polymers. Our results showed that the largest relative contents of low-MW polymers prevailed in the EPMs of mature samples, notably sample ML-1 (50%), a lichen crust; sample OS-1 (41%), a black crust sparsely colonized by lichens; and sample BT-1 (41%), a semi-moss crust also sparsely colonized by lichens. The biocrust level of development is strongly related to microbial enzymatic activity. Miralles et al. (2012) and Liu et al. (2014) demonstrated that cyanobacterial crusts are characterized by the lowest activity of hydrolytic enzymes, which is also positively related to the presence of mosses and lichens. The MW distribution patterns of our samples seem to confirm this assumption. Indeed, sample ML-2, for which we detected an EPM composed of only polymers distributed in the high-MW classes, could be classified as an early-stage biocrust according to sample morphology, lower levels of TECs and HMWECs, and a lack of uronic acids as EPM components.

In conclusion, this study reports for the first time the characteristics of the EPMs of High Arctic biocrusts, specifically providing information on their monosaccharide composition and MW distribution profiles. We underlined several divergent traits compared to those of biocrusts developed in temperate environments. The major detected difference between the samples concern the MW distribution profiles of the EPM, which are the result of the synthetic but also the hydrolytic activity of the microbial communities. Most likely, the more variable environmental conditions at the micro and macro scales compared to those of hot drylands caused biocrusts differing in community structure and composition, which in turn influenced species interactions and hence EPM characteristics. We found that the significant presence of low-MW EPM fractions, possibly an index of hydrolytic enzyme activity, was enhanced in mature biocrusts colonized by mosses or lichens. In later-stage biocrusts, the EPM was also characterized by the marked presence of uronic acids, which are important for all the hydrophilic interactions of the EPSs with the surrounding environment and contribute to ensuring a lower-abiotic-stress microenvironment by maintaining a high level of water in the soil. Furthermore, this work also confirms the possibility, already hinted at in previous studies on biocrusts from hot biomes, of using some parameters to gauge biocrust...
age and maturation. In particular, the relative content of uronic acid in the EPSs and the presence of low-MW EPS fractions, whose content is positively related to enzymatic activity, could be used as possible indices. However, further studies involving samples from more climatic areas are necessary to extrapolate definitive sound correlations.

Acknowledgements This study received funding from the Italian Ministry of University and Research (MIUR) in the framework of the Piano Nazionale di Ricerca sull’Antartide (PNRA 2014–2016), Project “Biological Soil Crusts; microbial biodiversity; metagenomics; metabolomics; microstructure” (WHYCRUST).

Funding Open access funding provided by Università degli Studi di Firenze within the CRUI-CARE Agreement.

Compliance with ethical standards

Conflict of interest The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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