Pedogenic and microbial interrelation in initial soils under semiarid climate on James Ross Island, Antarctic Peninsula region

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Abstract. James Ross Island (JRI) offers the exceptional opportunity to study microbial-driven pedogenesis without the influence of vascular plants or faunal activities (e.g., penguin rookeries). In this study, two soil profiles from JRI (one at Santa Martha Cove – SMC, and another at Brandy Bay – BB) were investigated, in order to gain information about the initial state of soil formation and its interplay with prokaryotic activity, by combining pedological, geochemical and microbiological methods. The soil profiles are similar with respect to topographic position and parent material but are spatially separated by an orographic barrier and therefore represent windward and leeward locations towards the mainly southwesterly winds. These different positions result in differences in electric conductivity of the soils caused by additional input of bases by sea spray at the windward site and opposing trends in the depth functions of soil pH and electric conductivity. Both soils are classified as Cryosols, dominated by bacterial taxa such as Actinobacteria, Proteobacteria, Acidobacteria, Gemmatimonadetes and Chloroflexi. A shift in the dominant taxa was observed below 20 cm in both soils as well as an increased abundance of multiple operational taxonomic units (OTUs) related to potential chemolithoautotrophic Acidiferrobacteraceae. This shift is coupled by a change in microstructure. While single/pellicular grain microstructure (SMC) and platy microstructure (BB) are dominant above 20 cm, lenticular microstructure is dominant below 20 cm in both soils. The change in microstructure is caused by frequent freeze–thaw cycles and a relative high water content, and it goes along with a development of the pore spacing and is accompanied by a change in nutrient content. Multivariate statistics revealed the influence of soil parameters such as chloride, sulfate, calcium and organic carbon contents, grain size distribution and pedogenic oxide ratios on the overall microbial community structure and explained 49.9 % of its variation. The correlation of the pedogenic oxide ratios with the compositional distribution of microorganisms as well as the relative abundance certain microorganisms such as potentially chemolithotrophic Acidiferrobacteraceae-related OTUs could hint at an interplay between soil-forming processes and microorganisms.

1 Introduction

In extreme environments, like Antarctica, local climatic conditions such as low temperatures, precipitation or irradiance are important and often limiting factors for soil formation. Even though soils in Antarctica are often poorly developed, they can be highly diverse (Michel et al., 2014; Simas et al., 2008; Bockheim et al., 2015). Therefore, soil
scientific investigations became a relevant topic in Antarctic research, proving that there are actually soils in Antarctica (Jensen, 1916) and identifying soil-forming processes (Ugolini, 1964). Antarctic soil research is mostly located in Victoria Land, continental Antarctica, especially in the McMurdo Dry Valleys (Michel et al., 2014; Ugolini and Bockheim, 2008), in the South Shetland Islands, maritime Antarctica (Simas et al., 2015) and the western Antarctic Peninsula region (APR) (Haus et al., 2015; Hrbáček et al., 2017b; Schaefer et al., 2017; Souza et al., 2014; Pereira et al., 2017).

Soils on continental Antarctica are often saline with thick salt horizons (Souza et al., 2014). Due to environmental stressors such as very low temperatures, low water availability, frequent freeze–thaw cycles and limited organic nutrient contents, soils from continental Antarctica show limiting conditions for higher organisms (Cary et al., 2010). However, diverse microbial communities thrive in a variety of Antarctic habitats, such as permafrost soils (Cowan et al., 2014).

Soils in maritime Antarctica and western APR differ from soils in continental Antarctica according to their stage of development (Balks et al., 2013; Blune et al., 2004; Parnikoza et al., 2017). They show extensive cryoturbation processes with occasional salt crusts at the soil surface (Balks et al., 2013; Bockheim, 1997). Local conditions determine nutrient availability in Antarctic soils (Prietzel et al., 2019). Ca, Mg, K and P contents are generally high in igneous and volcanic rocks, whereas P and N contents are highest in ornithogenic soils. Ornithogenic soils are well known in Antarctica. The World Reference Base for Soil Resources (IUSS Working Group WRB, 2014) defines ornithogenic material (from Greek ornithos, bird, and genesis, origin) as material which is characterized by penguin deposits mainly consisting of guano and often containing a high content of gravel transported by birds (compare with Ugolini, 1972). Soils from the eastern part of the APR (also called Weddell Sea sector) are different, since they are associated with a dry climatic transitional zone between the wet, warmer maritime Antarctica and colder, arid continental Antarctica. Mean temperatures are below 0°C and liquid water supply is sufficient to allow soil-forming processes (Souza et al., 2014). Souza et al. (2014) also showed that cryoturbation is less pronounced in the eastern APR than in the South Shetland Islands. The base saturation (>50%) and electric conductivity (EC) are generally high, whereas the amount of total organic carbon (TOC) is substantially low. Regarding cryoturbation, active layer depth, chemical weathering and soil organic C content, soils from the eastern APR are comparable to soils from inland areas of the Ross Sea region (Balks et al., 2013), though they are formed on different parent material (Daher et al., 2018). In comparison, the eastern APR with its semiarid soils remains one of the least studied areas in Antarctica (Souza et al., 2014; Daher et al., 2018).

Since microorganisms in Antarctica show a broad diversity as revealed by recent molecular phylogenetic and metagenomic methods (Cowan et al., 2014) and contribute to the weathering of minerals in soils (Uroz et al., 2009), they are pivotal to understanding initial soil formation. The bacterial phyla Proteobacteria, Acidobacteria, Actinobacteria, Bacteroidetes, Firmicutes and Gemmatimonadetes, commonly found in temperate soils, also dominate the microbial communities observed in Antarctic habitats (e.g., Bajerski and Wagner, 2013; Cary et al., 2010; Pearce et al., 2012; Chong et al., 2012). The microbial community structure is influenced by local soil chemical parameters, especially pH (e.g., Chong et al., 2010; Siciliano et al., 2014), but also by soil physical parameters such as grain size distribution and soil moisture (Ganzert et al., 2011). Chong et al. (2015) proposed, however, that historical contingency and dispersal limitations could have a stronger influence on differences in community distributions at a regional scale (>1000 km).

At the microscale, microbial activity such as photosynthesis and nitrogen fixation has a distinct influence on soil chemical parameters, e.g., the increase in carbon and nitrogen contents in oligotrophic soils (Ganzert et al., 2011; Cowan et al., 2011; Niederberger et al., 2015). In return, these changes in soil characteristics affect microbial community composition. Conflicting results illustrate the lack of understanding of drivers of soil microbial diversity in high-latitude soils (Souza et al., 2014). Since most of the nonlichenized Antarctic fungi are known to be decomposers and their abundance and distribution is limited by plant-derived nutrients, and bioavailable Carbon (Arenz and Blanchette, 2011), the focus of this study lies on the prokaryotic interplay with soil characteristics and soil formation.

Micromorphological studies in maritime Antarctica and the western APR described sulfurization and phosphatization in ornithogenic soils and mineral transformation on volcanic rocks (Pereira et al., 2013; Schaefer et al., 2008) and paleosols (Kirshner and Anderson, 2011; Spinola et al., 2017). Even though micromorphology offers the opportunity to study constituents of soil and their mutual relations in space and time and to identify soil-forming processes in an undisturbed state (Stoops, 2003), so far no micromorphological study has been published about soil-forming processes in the eastern APR that are influenced either by sulfates or by birds.

Our study sites are located on James Ross Island (JRI) in the eastern APR (Fig. 1) and therefore offer a unique setting to study soil formation and microbial communities in a transitional Antarctic landscape between the wet maritime and dry, colder continental Antarctica. We selected two different soils, representing coastal soils and inland soils of James Ross Island, developed on similar substrate and at similar topographic positions, but differing in local climate conditions and nutrient contents due to their relative position towards the mainly SW winds. The western study site (Brandy Bay – BB) is located in a windward position and is highly influenced by sea spray, while the eastern study site (Santa Martha Cove – SMC), located behind a mountain range, is located in a leeward position (Prietzel et al., 2019). This setting enables...
an investigation of interdependencies particularly between prokaryotic life and soil properties, since the selected soils are not influenced by vascular plants, sulfides, and penguin rookeries. With this, the main goal of our study is to identify major soil and microbiological properties in an extreme environment by combining pedochemical and micromorphological methods with microbial community studies based on high throughput sequence analyses. Thus, we will gain a better general understanding of (i) the main soil-forming processes and (ii) the drivers of soil microbial diversity community structure in the eastern APR. This addresses also the question of whether the variance of pedogenic and microbial properties is larger between depth increments within one profile (e.g., with different distances to the permafrost table) or between different soil profiles, i.e., due to different local environmental conditions.

2 Material and methods

2.1 Regional setting of James Ross Island, maritime Antarctica

James Ross Island is situated east of the Antarctic Peninsula and is the largest island in the western Weddell Sea sector (Hjort et al., 1997). The study area is located on the Ulu Peninsula in the northern part of JRI (Fig. 1). It represents one of the largest ice-free areas of the APR (Nedbalová et al., 2013; Hrbáček et al., 2017b) with the beginning of its deglaciation 12.9 ± 1.2 ka (Nývlt et al., 2014). More than 300 km² of the JRI lowlands are currently ice-free, except for a few glaciers (Engel et al., 2012).

The climate on JRI is semiarid polar continental (Martin and Peel, 1978). The precipitation, mostly snow, ranges between 200 and 500 mm of water equivalent per year with the major share during winter (Davies et al., 2013; Zvěřina et al., 2014). The thickness of the snow cover does not exceed 30 cm but varies due to strong winds (Hrbáček et al., 2017b, 2016a). The annual air temperature ranges between +10 and −30°C on the Ulu Peninsula (Hrbáček et al., 2016a; Láska et al., 2011). The year 2015 marked the warmest summer ever measured on the Ulu Peninsula, having a mean seasonal summer temperature (MSST) of 0.0°C and a maximum air temperature of 13.3°C (Hrbáček et al., 2017a), even though the mean annual air temperature (MAAT) decreased slightly from −6.8°C in 2011 to −7°C in 2015 (Hrbáček et al., 2016b; Láska et al., 2012).

The two study sites are located at Brandy Bay near the western coast and at Santa Martha Cove at the eastern coast of the Ulu Peninsula. Both sites are located at similar topographic positions (small plateaus) and elevation (80 m a.s.l.) with no visible vegetation (Figs. 2 and 3).

BB is located windward towards the mainly southwesterly winds (Hrbáček et al., 2016c; Nývlt et al., 2016), whereas SMC is located leeward, shielded by the Lachman Crags from the stronger winds. This results in less precipitation in the eastern part of JRI (Davies et al., 2013). Therefore, BB can be considered to be a characteristic wind-exposed coastal site with high influence of sea spray, whereas SMC represents a characteristic soil of an inland site with less influence of sea spray.

The substrate of both study sites is basically composed of coarse-grained Cretaceous sandstones and siltstones of the Alpha Member of the Santa Martha Formation (Hrbáček et al., 2017b). The land surface is generally covered by a debris layer of gravels and large clasts mixed with loose sandy regolith, mostly derived from James Ross Island Volcanic Group basalts, which were deposited as debris flows containing mainly basalt and hyaloclastite breccia and palagonite (Davies et al., 2013; Hrbáček et al., 2017b; Salzmann et al., 2011). No nesting birds are found on JRI.

The continuous permafrost on James Ross Island shows an active layer thickness ranging between 40 and 107 cm related to the topographic position on the Ulu Peninsula (Bockheim et al., 2013; Borzotta and Trombotto, 2004).

2.2 Soil sampling

During the austral summer period in 2016 soil samples from BB and SMC (Figs. 4 and 5) were taken. Both profiles were dug until a layer of coarse gravel was found. Bulk samples of both profiles were taken in depth increments (0–5, 5–10, 10–20, 20–50, > 50 cm) and were placed into sterile plastic bags, which were frozen immediately. Continuous cooling at −20°C was ensured by a transfer with the RV Polarstern to Germany. For micromorphological analyses, undisturbed and oriented samples were taken in modified Kubiena boxes (10 cm × 6 cm × 5 cm). Samples for micromorphology were taken at depths of 0–10, 10–20, 20–40, 40–60 and 80–90 cm at SMC. BB samples represent the depths of 10–20, 20–30 and 40–50 cm. Soils were described according to the Food and Agriculture Organization of the United Nations (FAO) (2006) and classified according to the World Reference Base for Soil Resources (WRB; IUSS Working Group WRB, 2015).

2.3 Soil physical and chemical analysis

2.3.1 Grain size distribution

The samples were saturated (100 mL of deionized water) and sonicated (800 J mL⁻¹). Coarse-medium sand (> 200 µm), fine sand (63–200 µm) and coarse silt (20–63 µm) were obtained by wet sieving. The smaller fractions, including medium silt (6.3–20 µm), fine silt (2–6.3 µm) and clay (< 2 µm), were separated by sedimentation. Fractions > 20 µm were dried at 45°C and weighed afterwards. The fractions < 20 µm were freeze-dried before weighing. The different procedures were chosen due to practical reasons: freeze-drying allows submitting the finer fractions to further anal-
yses (particularly carbon and nitrogen content) immediately, while the coarser fractions need milling anyway.

2.3.2 pH, EC, C&N contents, major elements and pedogenic oxides

The pH value was obtained using a pH meter (ph197i, WTW, Germany). Electrical conductivity was measured with a conductivity meter (LE703, Mettler Toledo, USA). Values of pH and electric conductivity were measured from bulk samples < 2 mm in deionized water with a sample-to-water ratio of 1:2.

Carbon (C) and nitrogen (N) contents of the bulk soils were analyzed by dry combustion (Elementar CNS vario Max cube). A total of 300 to 500 mg per sample was analyzed in duplicate. In order to distinguish between the total organic carbon (TOC) content and the total inorganic carbon (TIC), TIC was removed by acid fumigation according to Ramnarine et al. (2011). A total of 5100 mg of the milled bulk soil samples was moistened with 20 to 40 µL of deionized water and put into a desiccator together with 100 mL of 37 % HCl. Afterwards, the samples were dried at 40 °C. Finally, the samples were measured again by dry combustion (EuroVector EuroEA3000 Elemental Analyzer) to obtain the TOC content. TIC content was calculated as follows: TIC = C_{tot} − TOC.

Major elements were analyzed with a wavelength dispersive XRF device (AXS S4 Pioneer, Bruker, USA). Prior to preparation, the samples (ratio of lithium metaborate to soil 1:5) were ground with an agate mill for 12 min. Major elements were used for the calculation of weathering indices.

Pedogenic iron oxides (Fe_{d}) were determined by dithionite–citrate–hydrogen carbonate extraction (Holmgren, 1967). Poorly to noncrystallized Fe oxides (Fe_{o}) were
A 90 cm deep soil profile was taken; (b) soil profile Santa Martha Cove (SMC). It is not covered with vegetation. A 60 cm soil profile was taken; (c) study site Brandy Bay (BB) is close to snowfield. It is not covered with vegetation. A 60 cm soil profile was taken; (d) soil profile Brandy Bay (BB). Scale of the tape measure is in centimeters.

Figure 2. Study sites and soil profiles on James Ross Island; (a) Santa Martha Cove (SMC). It is not covered with vegetation. A 90 cm deep soil profile was taken; (b) soil profile Santa Martha Cove (SMC). Scale of the tape measure is in centimeters; (c) study site Brandy Bay (BB) is close to snowfield. It is not covered with vegetation. A 60 cm soil profile was taken; (d) soil profile Brandy Bay (BB). Scale of the tape measure is in centimeters.

determined by acid ammonium extraction (Schwertmann, 1964). The extractions were analyzed at a wavelength of 238.204 nm by an inductively coupled plasma optical emission spectrometer (Vista Pro CCD Simultaneous ICP-OES, Varian, USA).

2.3.3 Ion chromatography

The initial water content in the investigated soil material was too low to extract sufficient amounts of pore water for ion chromatography. Hence, the soil samples were leached, according to Blume et al. (2011). Five grams of soil material were suspended in 25 mL deionized water, shaken for 90 min and centrifuged at 9000 rpm to separate the soil material from the soil solution and sterile filtered through a 0.22 µm polyethersulfone (PES) filter (Sartorius AG, Germany).

The ion concentrations in leached water samples were analyzed by using two ion chromatography (IC) systems (SYKAM Chromatographie Vertriebs GmbH, Germany). For cations, the IC system consisted of a 4.6 mm x 200 mm Repro sil CAT column (Dr. Maisch HPLC GmbH, Germany), an S5300 sample injector and an S3115 conductivity detector (both SYKAM Chromatographie Vertriebs GmbH, Germany); 175 mg L\(^{-1}\) 18-Crown-6 and 120 µL methanesulfonic acid served as the eluent with a set flow rate of 1.2 mL min\(^{-1}\). The injection volume was 50 µL. The column oven temperature was set at 30 °C. The Cation Multi-Element IC standard (Carl Roth GmbH + Co. KG, Germany) containing NH\(^+_4\), Ca\(^{2+}\), K\(^+\), Li\(^+\), Mg\(^{2+}\) and Na\(^+\) was measured before every replication series. For anions, the IC system consisted of a SeQuant SAMS anion IC suppressor (Merck KGaA, Germany), an S5200 sample injector, a 3.0 mm x 150 mm SykroGel A 01 column and an S3115 conductivity detector (all SYKAM Chromatographie Vertriebs GmbH, Germany). A total of 6 mM Na\(_2\)CO\(_3\) with 90 µM sodium thiocyanate served as the eluent with a set flow rate of 1 mL min\(^{-1}\) and a column oven temperature of 50 °C. The injection volume was 50 µL. The multi-element anion standard containing F\(^-\), Cl\(^-\), Br\(^-\), NO\(_3\)^-, NO\(_2\)^-, PO\(_4\)^3- and SO\(_4\)^2- was measured before every replication series. The standards and samples were measured in triplicates.

2.3.4 Weathering indices and pedogenic oxide ratios

The KN index A (SiO\(_2\) + CaO + K\(_2\)O + Na\(_2\)O) / (Al\(_2\)O\(_3\) + SiO\(_2\) + CaO + K\(_2\)O + Na\(_2\)O) was calculated according to Kronberg and Nesbitt (1981). The index is based on the relative enrichment of the Al and Si oxide phase and the leaching of Na, K and Ca. It ranges between 0 (prevailing chemical weathering) and 1 (prevailing physical weathering). To get more precise information on the ongoing chemical weathering, the chemical index of alteration (CIA) [(Al\(_2\)O\(_3\) / (Al\(_2\)O\(_3\) + Na\(_2\)O + CaO + Fe\(^{2+}\) + K\(_2\)O)) × 100], in which Ca\(^{2+}\) represents the amount of silicate-bound CaO, was calculated according to Nesbitt and Young (1982). The CIA is frequently used as a quantitative measure of feldspar breakdown, assuming that feldspar represents the most abundant and reactive mineral. Higher values indicate increasing weathering intensity. Additionally, the degree of iron release (Fe\(_d\) / Fe\(_s\)) was calculated according to Blume and Schwertmann (1969), which gives information on the iron release from primary Fe-bearing mineral weathering: a longer or more intensive weathering process is indicated by a higher ratio (Baumann et al., 2014; Mirabella and Carnicelli, 1992).

2.4 Micromorphology

Samples for thin section preparation were air-dried and afterwards embedded with a mixture of resin (Viscosooss N55 S, Vosschemie, Germany), stabilized styrene (Merck KGaA, Germany) and hardener (MEKP 505 F, V osschemie, Germany), stabilized styrene (Merck KGaA, Germany), and then one half was ground with the grinding machine (MPS-RC Vacuum, G&N GmbH, Germany) and mounted onto a glass carrier. Then the mounted samples was sawed into slices of about 150 µm thickness. Finally, these slices were ground to a thickness of 25 µm. The preparation followed the instructions given by Kühn et al. (2017). Afterwards, they were analyzed by using a polarizing microscope (ZEISS Axio Imager.A2m, Software AxioVision 4.7.2, Carl Zeiss, Germany) containing NH\(^+_4\), Ca\(^{2+}\), K\(^+\), Li\(^+\), Mg\(^{2+}\) and Na\(^+\).
Figure 3. Images of micromorphological features found at Brandy Bay (BB) and Santa Martha Cove (SMC). Pictures were taking using plane-polarized light (ppl) and crossed polarizers (xpl). (a) BB III: highly separated lenticular platy microstructure, platy aggregates are indicated by green dotted lines, lenticular ms (microstructure) is indicated by black dotted lines, 2.5×, ppl; (b) SMC IV: moderately separated lenticular platy microstructure, indicated by black dotted lines, 2.5×, ppl; (c) SMC I: coarse monic microstructure, 2.5×, ppl; (d) BB II: chitonic c/f-related distribution (the distribution of coarse and fine material within the thin section) and thin link cappings (li) on quartz grains, 20×, ppl; (e) BB III: weathered rock fragment covered by silty capping (cap) and also showing a thick pendent (pen) consisting of silty material and mineral grains, 10×, ppl; (f) SMC I: strongly weathered sandstone fragment with former boundaries, indicated by red dotted line, still visible by capping (cap), 5×, ppl; (g) SMC I: weathered volcanic rock fragment with distinct pellicular alteration pattern, 5×, ppl; (h) BB II: weathered and broken volcanic rock fragment with internal volcanic glass and covered by a thin clay capping (cap), (110–120 µm), 2.5×, ppl; (i) SMC I: weathered volcanic rock fragment with feldspar phenocrysts, covered by a dusty clay-silt capping (80–100 µm) (cap), 2.5×, ppl; (j) SMC I: usage of crossed polarizers makes it possible to tell external coating (cap) from altered internal material, border indicated by grey dotted line, 2.5×, xpl.
2.5 Microbial community analysis

2.5.1 Nucleic acid extraction

For each soil sample (maximum amount of 0.5 g per sample), triplicates of total genomic DNA were extracted using the FastDNA® Spin Kit for Soil (MO BIO Laboratories Inc., USA). The extracted DNA was stored at −20°C and used as a template for the enumeration of target genes by quantitative PCR (qPCR) and next-generation sequencing (Illumina HiSeq).

2.5.2 Quantification of bacterial 16S rRNA gene copy numbers

qPCR was used to quantify total bacterial abundances. All qPCR assays were performed in triplicates on a CFX96 real-time thermal cycler (Bio-Rad Laboratories Inc., CA, USA) and contained 10 µL SensiFAST SYBR Mix (BioLine GmbH, Germany), 5.92 µL PCR water, 0.04 µL of forward and reverse primer (100 µM), and 4 µL template. The quantification of the bacterial 16S rRNA gene was based on the primers 341F (5′-CCTACGGGAGGCAGCAG-3′) and 534R (5′-ATTACCGCGGTCTGCTGG-3′) according to Muyzer et al. (1993). After an initial denaturing phase of 3 min at 95°C, the cycler included 35 cycles of 3 s at 95°C, 20 s at 60°C and 60 s at 72°C plus the plate read. All cycling programs included a melting curve from 60 to 95°C with 0.5°C steps per plate read. The analysis of quantification data was performed with the CFX Manager™ Software (Bio-Rad Laboratories Inc., CA, USA).

2.5.3 Illumina HiSeq sequencing

Unique combinations of tagged 515F (5′-GTGCCAGCMGCCGCGGTAA-3′) and 806R (5′-GACTACHVGGGTWTCTAAT-3′) (Caporaso et al., 2010) primers were assigned to each sample (Tables S1, S2 in the Supplement). For each sample, two PCR reactions were prepared and the PCR product was pooled after PCR to reduce PCR variability. The PCR was performed on a T100™ Thermal Cycler (Bio-Rad Laboratories Inc., CA, USA) in 25 µL reactions, containing 0.125 µL OptiTaq DNA Polymerase and 2.5 µL 10× Pol Buffer B (Roboklon GmbH, Germany), 1 µL MgCl₂ (25 mM), 1 µL dNTP Mix (5 mM), 16.625 µL PCR water, each 0.625 µL of forward and reverse primer (20 µM), and 2.5 µL genomic DNA. The following cycler program was used: initial denaturing step for 3 min at 95°C followed by 10 cycles of 1 min at 94°C, 1 min at 53°C (−0.2°C/cycle) and 1 min at 72°C, followed by 20 cycles of 1 min at 94°C, 1 min at 50°C and 1 min at 72°C, followed by a final extension step for 10 min at 72°C. All barcoded samples were pooled into a single sequencing library by adding an equal amount of DNA (60 ng DNA per sample). Subsequently, a purification of the PCR product pool was achieved by using the Agencourt AMPure XP

Zeiss Microscopy GmbH, Germany) and described following the terminology of Stoops (2003).
– PCR purification (Beckman Coulter, Inc., CA, USA). The Illumina HiSeq sequencing was performed by GATC Biotech AG, Germany.

### 2.5.4 Bioinformatics and statistical analysis

Sequencing was performed on an Illumina HiSeq (2 × 300 bp). Dual-indexed reads were demultiplexed using CutAdapt (options: e0.1; trim-n; Martin, 2011). Barcode base pairs were required to have a Phred quality score of Q25 and no mismatches were allowed. Read pairs were merged using PEAR (options: Q25; p10⁻⁴; o20; Zhang et al., 2013). The orientation of all sequences were standardized by a custom script using the information from demultiplexing. Sequences containing low-quality base pairs were trimmed and filtered using Trimmomatic (quality score of at least Q25 for trailing and leading base pairs, sliding window length of five base pairs, minimum sequence length of 200; Bolger et al., 2014). QIIME (version 1.9.1) (Caporaso et al., 2010) was employed for the clustering of operational taxonomic units (97 % sequence similarity) and their taxonomic assignments. Singletons, OTUs assigned to chloroplasts and mitochondria, and rare OTUs (relative abundance of < 0.1 % within each sample) were removed. Sample triplicates were merged by the mean value of their relative abundance before visualization of the sequencing data and before analysis of correlating environmental factors. For the processing and visualization of the obtained OTU table, R and PAST3 (Hammer et al., 2001) were used. The hierarchical clustering of the samples using the average linkage method was based on the Bray–Curtis dissimilarity. CANOCO5 ( Šmilauer and Lepš, 2014) was used for the canonical correlation analysis (CCA). If the Bonferroni-corrected \( p \) value was < 0.05, a given environmental parameter was included.

### 3 Results

#### 3.1 Field properties and soil classification

Both soils derived from coarse-grained marine sand- and siltstones, which were covered with volcanic clasts. There was a higher contribution of volcanic material in BB than in SMC. The amount of coarse material > 2 mm was larger at the profile BB. Deflation processes led to a residual enrichment of larger grains and pebbles at the soil surface of both profiles. The permafrost table was not reached in both soil profiles, but ground ice was visible at a depth of 85 cm at SMC. Neither SMC nor BB showed any ornithogenic influence. Both sites were unvegetated by cryptogamic or vascular plants. The C horizon was the only distinct soil horizon occurring at SMC, whereas BB shows two changes within horizontal structures by abrupt textural change below 10 and 20 cm. The textural change below 20 cm goes along with a change in textural class: SCL (sand: 52.5 %, silt: 21.9 % and clay: 25.6 %) to CL (sand: 44 %, silt: 27.2 % and clay: 28.8 %). Different from macroscopic features of the soil profiles, both soils showed evidences of a downward transport and accumulation of particles and nutrients, e.g., soluble products most likely originating from sea spray (Table 1). Accumulation starts at a depth of 50 cm at SMC and below 20 cm at BB. Soil color did not change through the profiles. SMC was brown to yellowish brown and BB was brownish yellow.

Both soils were classified as Cryosols (eutric, loamic) according to the WRB (IUSS Working Group WRB, 2015).

#### 3.2 Grain size distribution and soil chemistry

SMC had higher sand contents (mean value 61.7 %, Table 1), while BB was characterized by lower sand contents (mean value 47.4 %) and higher silt and clay contents (mean values 25.3 % and 27.2 % respectively). The grain size distribution varied only slightly with depth and similar clay and silt contents were demonstrated for both soils.

The pH was slightly to moderately alkaline in both profiles and highly alkaline only in the upper 5 cm of BB. The pH values followed opposing trends with depth, increasing in SMC from 7.7 to 8.1 and decreasing in BB from 8.6 to 7.4. The EC ranged between 50 and 60 µS cm⁻¹ in SMC and was substantially higher in BB with a minimum of 350–450 µS cm⁻¹ within 5–50 cm and its highest values around 900 µS cm⁻¹ between 0–5 cm and from 50 cm downwards. According to the EC values, SMC and the middle part of BB can be considered to be salt-free, whereas the salt content in the upper and lowermost part of BB was low (Food and Agriculture Organization of the United Nations (FAO), 2006).

The total inorganic carbon (TIC) content was low in both soils ranging between 0.1 and 0.3 mg g⁻¹ in SMC and between 0.7 and 2.0 mg g⁻¹ in BB. The TOC content ranges from 0.8 to 0.9 mg g⁻¹ for SMC and from 1.4 to 2.6 mg g⁻¹ for BB and increased there slightly with depth. The N content was around 0.4 mg g⁻¹ across both soil profiles. The C/N ratio was generally low with values below 7.5 in both soils; it decreased with depth in SMC (2.6–2.1) and increased with depth in BB (4.0–7.4).

Ion concentrations (Table 1) were parallel to the depth function of the conductivity in both soils; e.g., higher EC and ion concentration characterized BB. Cl⁻ concentrations decreased with depth in SMC from 20.5 to 3.5 µmol g⁻¹ soil as well as in BB from 4522 to 231 µmol g⁻¹ soil. The highest SO₄²⁻ concentrations were observed in the shallow (SMC: 9.6 µmol g⁻¹ soil; BB: 621 µmol g⁻¹ soil) and deepest (SMC: 15.3 µmol g⁻¹ soil; BB: 451 µmol g⁻¹ soil) samples. K⁺, Mg⁺ and Ca⁺ concentrations followed the same trend as SO₄²⁻. Br⁻, NO₃⁻, NO₂⁻, PO₄³⁻, Li⁺ and NH₄⁺ concentrations were below the detection limit.
3.3 Weathering indices and pedogenic oxide ratios

Weathering indices were calculated according to the major element contents (Table 3). The KN index A was at 0.91–0.92 in SMC and only slightly lower at 0.89–0.90 in BB (Table 2). The CIA varied between 53.9 and 54.8 in SMC and between 56.9 and 58.8 in BB. Both indices indicated weak chemical weathering with a slightly higher weathering intensity in BB.

The Fe<sub>d</sub>/Fe<sub>t</sub> ratio showed a decreasing trend from 0.18 to 0.11 with depth in SMC, indicating a decreasing intensity of pedogenic processes with depth. No particular trend was found in BB, but the Fe<sub>d</sub>/Fe<sub>t</sub> ratio is – similar to the CIA – generally higher around 0.20 except for 0.16 in the upper 5 cm.

3.4 Micromorphology

Formation of platy and lenticular aggregates due to repeated freezing and thawing processes was detected. Neither platy and lenticular platy structures nor the results of translocation (eluviation) processes were observed during fieldwork but could be confirmed later using micromorphology.

SMC had a weak to moderately developed pedality and a weak to moderate degree of separation (Table 3). Both pedality and degree of separation are well developed at a depth of 50–60 cm and were the least developed close to the surface and at the bottom of the profile. In contrast, BB had a well-developed pedality and a moderate to high degree of separation with its maximum development close to the bottom of the profile.

Lenticular and subangular blocky microstructures were present in both profiles, whereas lenticular microstructure was dominant in SMC and subangular blocky microstructure was dominant in BB. Lenticular-shaped aggregates were first observed at a depth of 10 cm in profile BB and at 30 cm in SMC (Fig. 3a and b).

Translocation features, like cappings consisting of clay and silt particles welded together with sand-sized quartz grains, were present in the upper part of both profiles. Link cappings occurred in the lower part of both profiles, with lesser and smaller cappings in BB (Fig. 3d). Link cappings were very rare and occurred only where coarse rock fragments were located close to each other. Dusty silt and clay pendants occurred only in the lower part of BB (20–50 cm) (Fig. 3e). The sphericity of mineral grains was smooth in both profiles. The minerals were slightly better rounded in BB (subangular to round) than in SMC (subangular to subrounded).

Weathering processes were identified by pellicular and dotted alteration patterns on rock fragments (mostly in sandstone fragments) in both profiles with a higher number of fragments with dotted alteration patterns than with pellicular alteration patterns. The quantity and intensity of dotted alteration patterns decreased with depth. Larger rock frag-
3.5 Microbial abundance and community structure

The enumeration of the 16S rRNA gene revealed a similar trend for both soil profiles (Fig. 4). The highest abundances with $6.6 \times 10^5$ copies g$^{-1}$ soil (BB) and $1.7 \times 10^6$ copies g$^{-1}$ soil (SMC) were detected in the uppermost depth increment of both soil profiles. Both soils showed a decrease in bacterial abundances with depth. The lowest bacterial abundances in SMC were detected below 50 cm depth with $3.7 \times 10^5$ copies g$^{-1}$ soil, as well as in BB at 20–50 cm depth with $1.7 \times 10^5$ copies g$^{-1}$ soil.

In total, 19,732,536 reads were obtained after merging the forward and reverse reads, demultiplexing, filtering, and deleting chimeric and singleton sequences. Additionally, reads of chloroplast-associated OTUs (36,573), mitochondria-associated OTUs (1,117) and rare OTUs (OTUs with a relative abundance of $< 0.1\%$ in every sample; 4,287,382) were filtered, resulting in 15,407,464 reads (Table S4). The number of reads per sample ranged from 54,122 to 916,583 with a mean value of 513,582. A total of 687 OTUs were clustered. After taxonomic classification, 258 putative taxa were obtained. Shannon's $H$ index was used to estimate and compare the alpha diversity of the different depth increment interval of the soils (Table S5). Both soils showed a similar Shannon $H$ index, which ranged from 3.7 to 4.7 not following any specific trend.

Bacteria dominated the microbial community in both soil profiles (Fig. 4). Higher abundances of Thaumarchaeota (7.2%–12.9%) were found in the upper 10 cm of the soil profile from SMC (Table S4). On a phylum level, the soil profile of SMC was dominated by Proteobacteria (23.4%–57.9%) and Actinobacteria (17.7%–41.3%) but showed also relative high abundances of Acidobacteria (3.9%–14.1%). The microbial community in BB was also mainly composed of Proteobacteria (28.2%–30.8%), followed by Actinobacteria (27.6%–46.6%), Gemmatimonadetes (3.9%–24.7%) and Chloroflexi (5.3%–10.9%). Bacteroidetes were highly abundant (10.5%) in the top 5 cm of BB. Regarding potential phototrophic organisms in the investigated soils, the amount of chloroplast-related reads was relatively low (< 0.2%) in each sample, except for SMC > 50 cm (0.03%–1.30%) and BB 0–5 cm (0.87%–1.01%). Cyanobacteria-related OTUs were rare and only showed low relative abundances in SMC 5–10 cm (0.06%), SMC 10–20 cm (0.62%) and SMC > 50 cm (0.04%).

The distribution of dominant OTUs was reflected by a cluster analysis based on the Bray–Curtis dissimilarity of the investigated depth increments. Samples were clustered according to their origin and depth. On a first level, samples grouped according to depth in upper (0–20 cm) and deeper (20–80 cm) samples, and within these groups they clustered according to location (BB vs. SMC). An ex-
Table 3. Micromorphological features of two soil profiles from Santa Martha Cove (SMC) and Brandt Bay (BB) from James Ross Island, Antarctica. The micromorphological property is shown by the presence (cross) or absence (no cross). (×): occasionally occurring.

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<th>Depth (cm)</th>
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<th>Translocation Features</th>
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**Aggregation.** hp: highly developed pedality; mp: moderately developed pedality; wp: weakly developed pedality; ds: degree of separation; h: highly separated; m: moderately separated; w: weakly separated.

**Groundmass.** Hypersthene (h); Orthopyroxene (o); Clinopyroxene (c); Labradorite (l); Quartz (q); Destiba (d); Albite (a); Celsian (c). Sphene (s); Glaucophane (g). Fissure (f); Single grain mesostructure (sg); Pellicular grain mesostructure (pg). Lenticular mesostructure (ml); Platy mesostructure (mp). Silt (s); Clay (c). Silt content was measured at 10 × magnification. 

**Pedofeatures.** Nodules – t: typic; a: aggregate. hp: hypocoatings; ro: redoximorphic hypocoatings. Coatings – li: link cappings; cap: cappings; pen: pendent. Infillings – ld: loose discontinuous. Degree of roundness and sphericity results separated by “/”. Microstructures separated by “( )”: one ms shows partly features of another ms. Microstructures separated by “×”: two different microstructures were found. Microstructures separated by “-”: no microstructure was found. Microstructures separated by “<1”: microstructure was found only in one sample. Microstructures separated by “<2”: microstructure was found in the remaining samples. Microstructures separated by “<3”: microstructure was not found in the remaining samples. Microstructures separated by “+”: microstructure was not found in the remaining samples. Microstructures separated by “-”: microstructure was not found in the remaining samples. Microstructures separated by “?”: microstructure was not found in the remaining samples. Microstructures separated by “<”:<: microstructure was not found in the remaining samples. Microstructures separated by “>”:<: microstructure was not found in the remaining samples. Microstructures separated by “<”:<: microstructure was not found in the remaining samples.

**Voids Micros.** li cap pen db spv xpv pl vu RS. Microsstructures separated by “×”: two different microstructures were found. Microstructures separated by “( )”: one ms shows partly features of another ms.

**Translocation features.** X: xanthate; Ch: chito-gefuric; O: oee; S: see; Chi: chi; Ce: close enaulic. Color – gb: greyish brown; db: dark brown. b-fabric – u: undifferentiated; gs: granostriated.

**Redomorphomorph features.** V: vughs; M: micros; G: networks. Degree of roundness and sphericity results separated by “/”. Two different microstructures were found. Microstructures separated by “( )”: one ms shows partly features of another ms.

The relationship of OTU distribution and environmental parameters was examined by applying a CCA (Fig. 6). Contents of chloride (18.5 %), calcium (11.8 %), sulfate (5.9 %), and TOC (6 %) and the Fe\textsubscript{Fe}/Fe\textsubscript{O} ratio (12.5 %) formed the optimal subset to explain variations in community structure of the investigated soil profiles (p < 0.05). The adjusted explained compositional variation was 49.9 %. A strong correlation between the unique community of BB 0–5 cm and the saline conditions was observed, mainly caused by high sulfate and chloride concentrations. The remaining samples were arranged according to sample site and depth as already observed in the cluster analysis above.

**4 Discussion**

The interaction of biotic and abiotic processes remains one of the fundamental questions in ecosystem research and further the initial development of soils under harsh environmental conditions, such as Antarctica. So far, only a few studies exist for polar environments that integrate pedogenic and microbiological research (e.g., Aislabie et al., 2008; Cowan et al., 2014; Ganzert et al., 2011; Bajerski and Wagner, 2013). James Ross Island offers an exceptional opportunity to improve our understanding of the interrelations between soil formation and microbiological properties in the absence of plants. The present interdisciplinary study gives profound insights into the state of soil formation and microbial community structure in initial soils in the transition zone between maritime and continental Antarctica.

James Ross Islands is located in the transition zone between warmer and wetter maritime Antarctica and cold and dry continental Antarctica (Souza et al., 2014). In this area, we studied two representative soils 16 km apart, with different exposures to the dominant southwesterly winds. The leeward position of SMC displays formation conditions of a typical inland soil, while BB in its windward position represents coastal soils. As indicated by EC values, BB is influenced by sea spray, while SMC, sheltered behind the Lachman Crags, does not show strong input of soluble salts from sea spray.
parameters. If the Bonferroni-corrected dissimilarities of the OTU data and its associated environmental parameters. If the Bonferroni-corrected \( p \) value was below 0.05, a given environmental parameter was included in the visualization. The amounts of chloride, sulfate, silt, Ca and TOC contents, and the \( \text{Fe}_{\text{d}}/\text{Fe}_{\text{t}} \) ratio explained 49.9 % of the microbial community composition.

The examined soils on JRI were characterized by low TOC (0.9–2.6 mg g\(^{-1}\)) and low total nitrogen contents (approx. 0.4 mg g\(^{-1}\)), which is common for Antarctic soil environments (e.g., Cannone et al., 2008), and relative high pH values (7.4–8.6). The moderately to highly alkaline pH in both soils cannot be explained by the occurrence of CaCO\(_3\), because the soils have a negligible amount with of TIC (\( \leq 2 \) mg g\(^{-1}\)). Low C contents do not only show the missing influence of penguins, but also indicate a relative juvenility of the soils: this indicates that no cations have been leached from the topsoil, and therefore the pH remains neutral to basic (Wilhelm et al., 2016). In addition, the content of basalt clasts in the parent material results in increased soil pH values (Simas et al., 2008; Moura et al., 2012). The opposing trends in the depth function of the pH values are caused by the input of soluble salts from sea spray: wind can transport soluble salts from the sea, causing an additional input of bases simultaneously increasing the pH at BB, while SMC is not affected (Benassai et al., 2005; Russell et al., 2010; Hara et al., 2004; Udisti et al., 2012). Since the substrate was not colonized by plants, lichens or endolithic prokaryotes, and the taxonomic data revealed low abundances of phototrophic organisms, the alkalinization of the substrate by the release of hydroxyl ions in the course of photosynthesis has a minor effect on soil pH. On the other hand, the neutral to basic pH does not significantly affect the soil microbial community structure, which is in accordance with observations in soils from Livingston Island (South Shetland Islands archipelago, maritime Antarctica) by Ganzert et al. (2011). They explained it by the occurrence of a specific soil microbial community, which thrives under low C and N conditions and is not dependent on nutrient input. Therefore, pH is mainly driven by the parent material composition combined with the input of soluble salts in these young soils on JRI.

The additional input of airborne cations by sea spray led to higher sodium and calcium contents and a rejuvenation of the affected depth increments of the soil profile, which can be seen in the lower CIA values in 0–5 cm soil depth of both soils compared to the lower part of the profiles. Ions, for instance sulfate, accumulate close to the permafrost table, which acts as a barrier and therefore explains increasing contents of sulfate with depth. The high amount of sulfate near the surface is most likely caused by sea spray and precipitation, because they are known to carry high amounts of sulfate in coastal areas (Blume et al., 2010).

Chemical weathering, as indicated by the KN index A (Kronberg and Nesbitt, 1981), is only of minor importance, whereas physical weathering is prevalent. The CIA and pedogenic oxide ratios (POR) confirmed the low degree of soil formation. Pedogenic oxides with specific degrees of crystallization relate to intensity and/or duration of pedogenic processes (Baumann et al., 2014; Blume and Schwertmann, 1969; Mirabella and Carnicelli, 1992). The results show that both CIA and both POR are slightly higher at BB compared to SMC. The KN index A and the CIA showed a weak chemical weathering of these mineral soils (Michel et al., 2014). Both indices indicated a more intensive chemical weathering at BB and, thus, indicate a slightly stronger pedogenesis at BB than at SMC. This finding could be explained by the sea- and windward position of BB, which results in an increased water availability and a slightly more leveled microclimate. Since both soils are located in similar topographic positions and derived from similar parent material, CIA and POR results allow the interpretation that soils influenced by coastal conditions tend to be more weathered. Besides physical and chemical weathering, microorganisms play an important role in mineral dissolution and oxidation. Adapted microorganisms colonize minerals and are, depending on nutritional requirements, nutrient availability and mineral type, potential contributors to the weathering of minerals (Uroz et al., 2009). Taxonomical groups, which are usually connected to microbial weathering, are present in the soils, such as *Mussilia, Bacillus* (Ma et al., 2011) and *Polaromonas* (Frey et al., 2010). Interestingly, the relative abundances of these taxa changed according to the degree of weathering. This could indicate a possible interrelation between the occurrence of these potential weathering-related organisms and the degree of weathering of Antarctic soils.
Evaluating weathering using the CIA, it must be noted that the value for BB is most likely underestimated. Ion chromatography results show that Na content is much higher at BB. The high amount of Na is most likely caused by sea spray, which is known to carry high amounts of Na (Udisti et al., 2012). Since the calculation of the CIA takes Na into account (Nesbitt and Young, 1982), the CIA values would be significantly higher if the additional input of sea salts could be excluded. It is very likely that the actual difference in state of weathering between SMC and BB would be much higher.

In conclusion, chemical weathering, even without influence of guano deposits, is of higher importance for the current state of soil formation than the ongoing cryoturbation.

In case of the pedogenic oxide ratios, 12.5% of the total compositional variation could be explained, which indicates a correlation between the microbial community structure and weathering at a very initial stage of soil formation. The pedogenic oxide ratios correlate with the compositional distribution of microorganisms in the investigated soils and with the relative abundances of one Acidiferrobacteraceae-related OTU. Microorganisms of this family are described as autotrophic sulfur and iron oxidizers, which have the capacity to use ferrous iron, thiosulfate, tetrathionate, sulfide and elemental sulfur as electron donors and oxygen or ferric iron as terminal electron acceptor (Hallberg et al., 2011). The reactive iron could potentially be used as terminal electron acceptor in the course of microbial iron cycling (Canfield, 1989). Organic matter, a potential substrate for heterotrophic microbial processes, sorbs on mineral surfaces (Kaiser and Guggenberger, 2000) and could be released in the course of microbial oxidation and reduction of reactive iron phases. In addition to the autotrophic processes, the release of sorbed, organic matter from mineral surfaces could be an additional way to increase the pool of biologically available carbon. The availability of such a mechanism potentially has an influence on the microbial community structure and abundances in oligotrophic environments.

Translocation features are common features in permafrost-affected soils. They often occur together with platy rectangular or lenticular aggregates, caused by reoccurring freeze–thaw cycles (Van Vliet-Lanoë, 1985). Platy blocks and lenses dominated the microstructure in the areas between 20 and 50 cm of both profiles. They were absent near the surface of both profiles and at the bottom of the profile SMC. These microstructures are known to occur in the transition zone between permanently frozen and unfrozen soils (Shur et al., 2005; Van Vliet-Lanoë et al., 2004). Here, the alternating temperature and soil moisture conditions additionally affect the microbial community structure. The frequency of freeze-and-thaw cycles tends to be steady in the middle part of a permafrost-affected soil, whereas weather shifts influence the surface, causing several freeze-and-thaw events per day, which do not result in typical microstructure formation due to insufficient water supply (Van Vliet-Lanoë, 1985). Aggregate formation by reoccurring freeze-and-thaw cycles results in a change in pore shape and size (Van Vliet-Lanoë et al., 2004). Especially during the summer season, intensive insolation causes high evaporation, resulting in dry soil surfaces. Changes in pore space affect microbial habitats, due to larger pores and a more sufficient water supply. This has a severe influence on matter fluxes and soil–environmental conditions, which is reflected in a changing species distribution and, more specifically, the occurrence of different clusters of highly abundant organisms in both soils. Multivariate statistics were performed for soil depth increments considered to be independent. However, when processes are discussed that link between soil horizons, e.g., water and solute flow through the profiles, we account for the limited number of two soil profiles with great care. We could not detect any environmental factors that increase or decrease the correlation between the chosen depth increments. Nevertheless, freeze-and-thaw cycles definitely also occur in the upper part of the profile, as indicated by the well-sorted areas (Van Vliet-Lanoë, 1985), which were described as single grain microstructure. Near the permafrost table aggregates are often formed by frost desiccation and are hence poorly compacted, which makes them unstable upon moistening, which occurs during thawing events and explains the missing platy microstructure at SMC near the bottom of the profile (Van Vliet-Lanoë, 2010). The fact that lenticular-shaped aggregates occur also in the lower part of the profile indicates that the permafrost table is located underneath the layer of coarse gravel at BB.

Although the investigated soils were poorly developed, an abundant and diverse prokaryotic community could be observed. Microbial abundances in both soils showed a decreasing trend with depth. Values of up to \(10^9\) gene copies g\(^{-1}\) soil in the uppermost depth increments are comparable to observed microbial abundances from other cold environments, such as alpine glacial forelands (Sigler et al., 2002), permafrost-affected soils from arctic regions (Liebner et al., 2008) and Antarctic glacier forefields (Bajerski and Wagner, 2013).

Both soils were characterized by a highly diverse community dominated by Proteobacteria, Actinobacteria, Gemmatimonadetes, Acidobacteria and Chloroflexi, which is in accordance with the observations in other continental and maritime Antarctic habitats (e.g., Yergeau et al., 2007; Cary et al., 2010; Ganzert et al., 2011; Bajerski and Wagner, 2013). Substantial differences in geochemical parameters such as conductivity and the change in the community structure on a phylum level were evident as well as the occurrence of depth-dependent clusters (0–20 cm; > 20 cm) of dominant OTUs (Fig. 8). Whereas the upper 20 cm of the soils was dominated by Gammaproteobacteria and Gailliales, the deeper part of the soils showed increased abundances of OTUs related to Acidiferrobacteraceae and Gemmatimonadaceae. This distinct shift correlates with the occurrence of the microstructure related to freezing and thawing and could be related to its changes in the pore space and the availability of oxygen, wa-
ter and nutrients. For instance, Gemmatimonadaceae were a common observation in the soils and showed increased abundances in deeper parts of BB. These organisms have a cosmopolitan distribution in terrestrial environments and depend on the soil moisture condition of the respective soil and soil depth (DeBruyn et al., 2011; Bajerski and Wagner, 2013). Only a few isolates have been described for this phylum (e.g., Zeng et al., 2015) and their exact functions in soil ecosystems remain uncertain. The change in relative abundance of these taxa with depth could be coupled to the changing availability of water, which depends on the microstructure. For example, the amount and size of microaggregates have been shown to be important regarding prokaryotic colonization, leading to genetically distinct communities as well as cell densities in different size classes of aggregates (Ranjard et al., 2000). Thus, in addition to environmental parameters, which shape the overall prokaryotic community, the microstructure of the initial soils could have a substantial influence on species distribution.

Higher abundances of Bacteroidetes- and especially Flavobacteriaceae-related OTUs were observed in the uppermost area of soil from BB, while only showing minor abundances in the deeper soil areas. This area differed from the remaining soil in two regards, namely very high chloride concentrations and a relative high content of coarse sandy material and could select for adapted psychro- and halotolerant Bacteroidetes-related organisms, such as Flavobacteriaceae (e.g., Bajerski et al., 2013). Members of the Flavobacteriaceae family detected in this area, for instance Gillisia sp., were isolated from Antarctic habitats before and were shown to be at least moderately tolerant to saline conditions (Bowman and Nichols, 2005). Putative halotolerant or halophilic Flavobacteriaceae in this area could have a need for high chloride contents. Chloride can be accumulated inside the cell to osmotically balance the cytoplasm with the surrounding habitat (Oren, 2002; Müller and Oren, 2003). Furthermore, the detected Bacteroidetes-related organisms could prefer the coarser, sandy microstructure from this depth increment. The preference of microbial groups for certain grain-size-dependent environments, for instance the sand-sized fraction being preferred by Bacteroidetes, was shown, e.g., in Typic Hapludalfs from central Denmark (Hemkenmeyer et al., 2018).

Both investigated soils were poor in soil organic C as well as N. Organisms with the ability to use oxygenic photosynthesis to fixate CO₂, such as cyanobacteria, were nearly absent in the investigated soils. Low abundances of Cyanobacteria are a common observation for Antarctic soil habitats (Ji et al., 2016). Due to the lack of phototrophic organisms and organic carbon, inorganic compounds and metabolic pathways utilizing those may have a more pronounced role in sustaining the microbial ecosystem at this initial stage of the soils. Several of the most abundant taxa observed in BB and SMC were putative chemosyntrophs involved in nitrogen, iron and sulfur cycling, such as potential ammonia-oxidizing Thaumarchaeota or sulfur/iron-oxidizing Acidifero bacteraceae. Microorganisms can be seen as the primary pioneers of nutrient-poor environments such as Antarctic soils and were shown to have the genetic potential to fix C and N (Cowan et al., 2011; Niederberger et al., 2015), thus increasing C and N contents of these oligotrophic soils. The chemosyntrophic Thaumarchaeota oxidize ammonia aerobically to nitrite (Brochier-Armanet et al., 2008; Vajrala et al., 2013) and were observed in many studies located in Antarctica (Magalhães et al., 2014; Ayton et al., 2010). However, ion chromatography showed that amounts of ammonia as well as nitrite and nitrate were negligible in both soils. Ammonia originating from necromass and products in the course of nitrification could be metabolized directly by the present community, so no accumulation of the different intermediates containing nitrogen takes place. These organisms are reported to have the genetic potential to use the hydroxypropionate/hydroxybutyrate pathway for CO₂ fixation, which is highly efficient and could provide an ecological advantage in oligotrophic environments (Könneke et al., 2014). Further, a part of the community could use atmospheric compounds as an energy source. Atmospheric H₂, CO and CO₂ are scavenged and used as an energy source by microorganisms, especially organisms associated with the phyla Actinobacteria, Chloroflexi, Acidobacteria, Planctomycetes, Verrucomicrobia, and Proteobacteria (Greening et al., 2015; Ji et al., 2017). Operational taxonomic units related to the phylum Actinobacteria and the associated orders Acidimicrobiales and Solirubrobacteriales were highly abundant in the investigated soils. Microorganisms in Antarctic soils, especially bacteria related to the phyla Actinobacteria, AD3 and WPS-2, were shown to generate biomass by consuming H₂, CO₂ and CO from the atmosphere (Ji et al., 2017). The gene for chemosynthetic CO₂ fixation, rbcL1E, was found in multiple orders, including Pseudonocardiales, Acidimicrobiales and Solirubrobacteriales. Similar functional capabilities could be present and active in the investigated soils. Our results show that, in this initial stage of soil development, chemolithoautotrophic lifestyles play an important role for the generation of biomass and initial accumulation of soil organic carbon and nitrogen.

5 Conclusion

The presented soil and microbiological study on initial soils in the semiarid environment of Antarctica shows the current state of soil formation indicated by main soil and microbiological properties and their interplay. The results allow us to draw the following conclusions.

Despite similarities in topographic position and substrates, both profiles have distinct differences in chemistry (content of salts indicated by EC, opposing trends in pH and states of weathering, indicated by WI and POR) and microbiology (depth functions of microbial abundances and diver-
sity, e.g., Proteobacteria, Gemmatimonadetes and Thaumarchaeota abundances), which are caused by the different local environmental conditions at each site.

The EC values of the soils and the depth function of the pH values clearly showed different conditions for soil formation at the two sites caused by the more exposed location of BB towards the mainly southwesterly winds, resulting in a more intense weathering and higher input of salt by sea spray.

Taking weathering and aggregation as indicators of soil formation, we conclude that coastal conditions – in contrast to inland conditions – favor the formation of soils in maritime Antarctica.

Despite different local environmental conditions at each site, the microbial communities differ more distinctly between the depth increments in one profile than between the two profiles. Therefore, we conclude that in this initial stage of soil formation factors such as weathering and microstructure formation, as well as the resulting parameters (e.g., water availability and matter fluxes), are more important drivers of soil microbial community composition than chemical parameters such as EC and pH.

Assuming that prokaryotic life is highly affected by changes in soil structure and vice versa, further investigations in this field should include analyses of (micro-)aggregates.

**Data availability.** Demultiplexed raw sequencing data were submitted to the European Nucleotide Archive (http://www.ebi.ac.uk/ena, last access: 17 June 2018) under accession number PRJEB29853.

**Supplement.** The supplement related to this article is available online at: https://doi.org/10.5194/bg-16-2481-2019-supplement.

**Author contributions.** The project was initiated and designed by DW, PK, TS and CWM. LAM and CWM carried out fieldwork during the PROANTAR field trip led by CEGRS in 2016. LAM, PK, IP and FH performed analyses and interpretation. LAM and PK prepared this paper with contributions from all co-authors. LAM and PK contributed equally.

**Competing interests.** The authors declare that they have no conflict of interest.

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