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Co-variation in soil biodiversity and biogeochemistry in northern and southern Victoria Land, Antarctica


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Abstract: Data from six sites in Victoria Land (72–77ºS) investigating co-variation in soil communities (microbial and invertebrate) with biogeochemical properties show the influence of soil properties on habitat suitability varied among local landscapes as well as across climate gradients. Species richness of metazoan invertebrates (Nematoda, Tardigrada and Rotifera) was similar to previous descriptions in this region, though identification of three cryptic nematode species of Eudorylaimus through DNA analysis contributed to the understanding of controls over habitat preferences for individual species. Denaturing Gradient Gel Electrophoresis profiles revealed unexpectedly high diversity of bacteria. Distribution of distinct bacterial communities was associated with specific sites in northern and southern Victoria Land, as was the distribution of nematode and tardigrade species. Variation in soil metazoan communities was related to differences in soil organic matter, while bacterial diversity and community structure were not strongly correlated with any single soil property. There were no apparent correlations between metazoan and bacterial diversity, suggesting that controls over distribution and habitat suitability are different for bacterial and metazoan communities. Our results imply that top-down controls over bacterial diversity mediated by their metazoan consumers are not significant determinants of bacterial community structure and biomass in these ecosystems.

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Key words: habitat suitability, invertebrate diversity, Latitudinal Gradient Project, microbial diversity, nematodes

Introduction

Terrestrial ecosystems along the Antarctic Victoria Land coast, from the McMurdo Dry Valleys in the south (78–76ºS) to Cape Adare (71ºS) in the north, occur across broad scale gradients of biodiversity, climate, and soil properties (Barrett et al. 2006). Insights into the relationships between biodiversity, productivity and ecosystem functioning may be especially discernable in Victoria Land since this region spans a range of ecosystems with no conspicuous source of organic matter and no metazoans, to those with abundant moss, lichen and algal biomass, marine-derived organic matter deposits and complex soil food webs (Rudolph 1963, Freckman & Virginia 1997, Bargagli et al. 1999, Porazinska et al. 2002, Sinclair et al. 2003). This latitudinal gradient of varying soils and biota can be used to test specific hypotheses about the distribution of soil biodiversity and physical and biogeochemical controls over habitat suitability (Howard-Williams et al. 2006).

Little is known about the composition of Antarctic soil microbial communities, and the environmental factors that drive their patterns of diversity. Until quite recently, the low sensitivity of microbial techniques provided only a limited view of microbial communities, but more sophisticated techniques are now revealing more diverse communities than previously recognized (Cowan et al. 2002, Adams et al. 2006, Aislabie et al. 2006, Connell et al. 2006). Victoria Land invertebrate communities are comprised of five phyla of invertebrates: Arthropoda (Collembola, Acari), Tardigrada, Nematoda, Rotifera, and Protozoa (Adams et al. 2006). Diversity within these invertebrate taxa is typically among the lowest on Earth (less than five species). Free-living soil nematodes and tardigrades are the most abundant and widely distributed animals in continental Antarctica (Freckman & Virginia 1997, Petz 1997, Sinclair 2001, Convey & Wynn-Williams 2002, Sohlenius et al. 2004, Convey & McInnes 2005, Adams et al. 2006).

The suitability of Antarctic soil environments as habitats for soil biota occur across a range of biogeochemical properties and climate (Freckman & Virginia 1997, Virginia & Wall 1999). Courtright et al. (2001) formalized a habitat suitability model for metazoan invertebrates by quantifying...
species abundance, life stages and their relation to a suite of soil biogeochemical properties with factor analysis. This model segregates metazoan communities into distinct habitats according to a combination of soil properties, most particularly moisture, organic matter and salinity. More recent work has synthesized these results together with a large number of observations in the Dry Valleys (Freckman & Virginia 1997, Powers et al. 1998, Treonis et al. 1999, Gooseff et al. 2003, Moorhead et al. 2003, Barrett et al. 2004, Parsons et al. 2004, Nkem et al. 2006) in a numerical model based upon logistic regression (Poage et al. unpublished). This model predicts probabilities for the presence of various nematode species and species combinations based upon a small number of chemical parameters (salinity, moisture, carbon, nitrogen, pH) with high statistical significance.

Here we report on work conducted at six sites in northern and southern Victoria Land (Fig. 1), Antarctica in December and January of 2003–2004 (Table I). We investigated the structure (bacterial and metazoan diversity) and functioning (soil respiration) of soil communities and the influence of soil biogeochemical properties (organic matter, inorganic nutrients, physicochemical properties) on habitat suitability. The objective of this study was to investigate controls over the distribution and diversity of soil biota and their activity by quantifying co-variation in soil microbial and metazoan communities with soil biogeochemical properties among local landscapes and across broad climatic gradients in Victoria Land.

Materials and methods

Site descriptions

Taylor Valley (77°S, 162°E, Fig. 1) is the principal site of the McMurdo Dry Valleys Long-Term Ecological Research programme. Arid soils are the most extensive landform of Taylor Valley, occupying > 95% of glacier ice-free surfaces below 1000 m (Burkins et al. 2001). Due to low precipitation (< 10 cm water equivalent yr⁻¹), relative humidity and soil temperatures, availability of liquid water is extremely limited and confined to brief periods during the summer when temperatures exceed freezing (Campbell & Claridge 1982, Doran et al. 2002). Consequently, surface soil moisture content is typically quite low (< 2% by weight), and is an important limitation over biotic community structure (Treonis et al. 1999, Aislabie et al. 2006), and the functioning of soil biota (Treonis et al. 2000, Barrett et al. 2006, Elberling et al. 2006, Gregorich et al. 2006).

Sources of organic matter contributing to Taylor Valley food webs include low rates of C-fixation by photoautotrophic communities inhabiting the surface layers of sandstones and granites, i.e. endolithic communities and lichens (Friedmann et al. 1993, Kappen et al. 1998), microbial mats, lichens and mosses occupying intermittently saturated zones on stream and lake margins (Schwartz et al. 1992, Pannewitz et al. 2003), and ancient organic matter entrained in lacustine and marine sediments deposited during high stands of proglacial lakes or marine incursions (Burkins et al. 2000). Microbial mats (e.g. Phormidium spp., Nostoc spp.) and moss communities (e.g. Byrum spp., Hennediella heimii (Hedw.) Zand.) dominate C cycling and ecosystem processes where they occur (e.g. Schwartz et al. 1992, 1993, Pannewitz et al. 2003, Moorhead et al. 2003), but are confined to areas where

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**Table I.** Location of study sites in Victoria Land Antarctica.

<table>
<thead>
<tr>
<th>Location</th>
<th>Latitude</th>
<th>Longitude</th>
<th>Elevation</th>
<th>Mean annual temperature</th>
<th>Mean DJF temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Taylor Valley</td>
<td>77°S</td>
<td>162°E</td>
<td>50</td>
<td>-20</td>
<td>-5</td>
</tr>
<tr>
<td>Luther Vale</td>
<td>72°22'S</td>
<td>169°53'E</td>
<td>150</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Cape Hallett</td>
<td>72°19'S</td>
<td>170°13'E</td>
<td>10</td>
<td>-15</td>
<td>-1</td>
</tr>
</tbody>
</table>

DJF = December, January and February, i.e. the austral summer. NA = not available.
significant liquid water is often available.

Cape Hallett (72°19’S, 170°13’E) is a small, coastal, ice-free area (72 ha) in northern Victoria Land 100 km south of Cape Adare (Fig. 1), with mean annual temperatures of \(-15.3^\circ\text{C}\) and annual precipitation of 18 cm of water equivalent per year (Dufhorn 1981). About half of the total area of Cape Hallett is occupied by an Adélie penguin (Pygoscelis adeliae (Hombron & Jacquinot)) rookery on the low-lying coastal areas (Seabee Spit) and skua (Catharacta maccormicki Saunders) colonies on the lower portions of an adjacent scree slope (Raytheon Polar Services Company 2003). The Cape Hallett landscape consists of basal scree and moraines colonized by dense moss beds (Byrum spp.), green algae (Praisola spp.) and conspicuous lichen colonies, particularly in the Willett Cove area (Rudolph 1963, Pannewitz et al. 2005). Sources of soil moisture include precipitation, melting of snowfields, and the Bornmann Glacier above the scree slopes. Campbell & Claridge (1987) emphasized that there is higher moisture availability in Cape Hallett soils compared to sites in southern Victoria Land, because in coastal zones of northern Victoria Land snow tends to melt rather than sublimate.

Other ice-free terrestrial environments in northern Victoria Land are found in the vicinity of Cape Hallett in dry cirques and saddles of the Admiralty and Victoria mountain ranges that serve as northern analogues of less-marine influenced terrestrial environments in southern Victoria Land, i.e. the McMurdo Dry Valleys. An unnamed bowl-shaped valley north-east of Luther Peak is such a location (Fig. 1). The soils of this area, hereafter "Luther Vale", occupy approximately 100 ha at 200 m a.s.l. and are underlain by metamorphic rock. Observed sources of water include precipitation and meltwater from snowfields on the north-western side of Luther Peak, which contribute to an ice covered pond.

Sample collection

We identified two sites at each of the three locations across Victoria Land (i.e. Taylor Valley, Cape Hallett, Luther Vale) that differed in conspicuous soil features that influence soil biology, and therefore represented areas of low and high potential productivity, based upon a priori assessment. Transects (20 m) were sampled at randomly determined points at each of these sites. In Taylor Valley, four replicate transects were established at varying distances from two glacial meltwater streams in the Lake Fryxell basin; each transect covered similar topography (i.e. distance from stream, surface soil conditions), and was perpendicular to the axis of stream flow of Von Guerard Stream and Green Creek. Green Creek is among the most diverse and biologically active sites in the Taylor Valley with regard to colonization by microbial mats and mosses, while Von Guerard is characterized by lower density of microbial mats and little to no moss colonization (Alger et al. 1997, McKnight et al. 2004). Likewise, in Luther Vale, four replicate transects were established at sites on the north and south sides of an unnamed meltwater pond; each transect covered similar elevation, slope, and aspect at locations of varying distances from the pond-soil margin. At Cape Hallett, only two transects were sampled at each of two sites (in order to minimize disturbance in the ASPA 106), where transects were established at various distances from recently abandoned penguin mounds (SeaBee Spit), and in moss-dominated soils (Willett Cove).

Five replicate samples were collected from the surface soil (0–10 cm) of each transect along the productivity gradients in the study sites located in Taylor Valley, Cape Hallett and Luther Vale (n = 100). Soil samples were collected using standard aseptic protocols developed in the dry valleys (Freckman & Virginia 1997). Samples were split in the field to provide material for invertebrate and chemical analyses, and for assessment of bacterial diversity. Subsamples for bacterial analysis were collected in duplicate from excavated soil profiles (see below).

Soil CO$_2$ flux

Soil CO$_2$ flux, i.e. soil respiration across the sites to provide a comparative index of ecosystem functioning was measured using a portable gas exchange system (LiCor 6400 Portable Photosynthesis System with 6400-09 soil CO$_2$ flux chamber, LiCor Inc., Lincoln, NE) customized to measure small changes in CO$_2$ (Parsons et al. 2004). Soil temperature (5 cm depth) was measured continually using Hobo temperature loggers (Onset Computer Corp, Bourne, MA). We measured soil CO$_2$ flux over the course of two days at each site, with the exception of the Von Guerard site where equipment malfunction stopped data collection after 24 hr. Here we report midday averages of measurement taken from 12 replicate plots at each site.

Soil chemical analysis

Field moist samples were passed through a #10 sieve and the < 2 mm fraction retained for analysis. Gravimetric soil moisture, conductivity, pH and inorganic N were determined using conventional techniques modified for dry valley soils (Barrett et al. 2002). We determined the elemental concentration and natural abundance of C and N isotopes in acidified soils using an a Delta-Plus XL isotope ratio mass spectrometer interfaced with a Carlo Erba 1500 elemental analyser at Dartmouth College.

Soil metazoans

Nematodes, rotifers and tardigrades were extracted from unsieved soil samples using a sugar-centrifugation technique modified for Antarctic soils (Freckman & Virginia 1997). Nematodes were identified under light microscope.
microscopy based upon morphology and enumerated to species, sex and life stage; densities of tardigrades and rotifers were enumerated. Presence of collembolans and mites were noted, both in the field and in the extractions. The presence of cryptic species was explored via DNA analysis.

DNA was extracted from individual metazoans using DNeasy tissue kits (Qiagen Inc, Valencia, CA). Ribosomal DNA of the internally transcribed spacer 1 region was PCR amplified using the 18S primer designed by Vrain et al. (1992) and the reverse primer of Cherry et al. (1997). Ribosomal DNA of the D2D3 region was amplified and sequenced using the 500-501 primer set of Nadler et al. (2000). The ribosomal small subunit was amplified and sequenced using the 1A 3B primer set of Baldwin et al. (1997). PCR reactions were carried out in 25 μl volumes. Each reaction contained: 2.5 μl 10xPCR buffer, 1.5 μl MgCl2, 1 μl dNTP mixture (10 mM each), 1 μl of 10 pM forward primer, 1 μl of 10 pM reverse primer, 0.25 μl of high fidelity DNA polymerase (Enzypol Ltd, London, Ontario, Canada), 19.55 μl of distilled water and 5 μl of DNA (approximately 10 ng μl-1). All PCR reactions were run in a PTC-100 Thermocycler (MJ Research Inc, Waltham, MA) with the following cycling profile: 1 cycle of 94°C for 7 min followed by 40 cycles of 94°C for 1 min, 50°C for 1 min, 72°C for 1 min. The last cycle was followed by a lengthened primer extension period (72°C for 10 min). PCR products were prepared for sequencing reactions by enzymatic treatment (ExoSap, GE Healthcare, Piscataway, NJ) and sequenced at the Brigham Young University DNA Sequencing Center on Perkin-Elmer/Applied Biosystems automated DNA sequencers. The resulting DNA sequences were used to reconstruct evolutionary relationships, identify independently evolving lineages and inform species delimitation (Wiley 1978, Lynch & Renjifo 1990, Frost & Kluge 1994, Adams 1998, 2001, Lynch 1999, Wiley & Mayden 2000). These lineages also correspond with the MOTU concept (Floyd et al. 2002, Eyualem & Blaxter 2003, Blaxter et al. 2005), in that there are more than three nucleotide differences (in most cases many more) at the SSU locus between “species”.

Shannon-Weiner diversity indices (H') were calculated for nematodes using the equation:

$$H' = -\sum_{i=1}^{s} p_i \ln p_i$$  \hspace{1cm} (1)

where $p_i$ is the proportion of the $i$th species and $s$ is the number of species present in a given soil community. Only nematodes were present in sufficient abundance and diversity necessary to calculate Shannon-Weiner indices.

**Microbial biomass and bacterial diversity**

Microbial biomass was estimated using the chloroform fumigation-extraction (Cheng & Virginia 1993) modified for the low biomass of Antarctic soils. Soils were fumigated with distilled chloroform in a vacuum desiccator cabinet at 65 mbar vacuum for five days. On day 5, pairs of fumigated and unfumigated soils were extracted in 0.5 M K2SO4, filtered through Whatman #42 filters, acidified with 200 μl concentrated HCl and frozen prior to analysis of total organic C and total N on a Schimadzu TOC analyser at the Crary Laboratory at McMurdo Station. Microbial biomass C and N was calculated as the difference between the chloroform-labile C in the pairs of fumigated and unfumigated soils.

Nucleic acids were extracted from soil using a modified cetyltrimethyl-ammonium bromide-polyvinylpyrrolidone-βmercaptoethanol (CTAB) extraction protocol (Coyne et al. 2001) designed for maximum recovery of nucleic acids from low biomass silica-rich soils. Soils (0.7 g) were aseptically measured into 1.5 ml tubes containing 0.5 g each 0.1 mm and 2.5 mm zirconia/silica beads (Biospec Products Inc, Bartlesville, OK). Beads were baked at 250°C for 4.0 ms-1 for 30 sec to lyse the cells. Samples were then centrifuged at 13 200 rpm for 3 min. The supernatant was removed and transferred to a 1.5 ml sterile Eppendorf tube containing 200μl CTAB buffer (100 mM NaH2PO4), and 300 μl SDS lysis buffer (100 mM NaCl, 500 mM Tris pH 8.0, 10% SDS) were added to the tubes. A Fast Prep® FP120 cell disrupter was used at 4.0 ms-1 for 30 sec to lyse the cells. Samples were then centrifuged at 13 200 rpm for 3 min. The supernatant was removed and transferred to a 1.5 ml sterile Eppendorf tube containing 200μl CTAB buffer (100 mM Tris-HCl, 1.4 M NaCl, 20mM EDTA, 2% CTAB, 1% PVP, 0.4% BME). Samples were incubated at 100 rpm and 60°C for 30 min. Subsequent to incubation, 1000 μl chloroform:isoamyl alcohol (24:1) was added to each sample. Samples were mixed by repeated inversion, vortexed for 15 sec and then centrifuged at 13 200 rpm for 5 min. The upper aqueous layer was removed into a sterile 1.5 ml Eppendorf tube.

An additional chloroform extraction was performed by adding 500 μl chloroform:isoamyl alcohol (24:1), vortexing the samples for 10 seconds followed by incubation on a rocking bed for 20 min at room temperature. Samples were then centrifuged at 13 200 rpm for 5 min and the upper aqueous layer was removed into a 1.5 ml sterile Eppendorf tube. DNA was precipitated for at least one hour at -80°C with the addition of ammonium acetate to a final concentration of 2.5 M and 0.54 volume isopropyl alcohol. Samples were then centrifuged at 13 200 rpm for 20 min and the DNA pellet was washed with 1 ml 70% ethanol and air dried in a laminar flow hood for 1 hour. DNA was resuspended in 20 μl sterile LO-TE (3 mM Tris-HCl pH = 8,0.2 mM EDTA) and stored at -80°C until use. The yield and quality of each DNA preparation was evaluated through visualization on a 1% agarose gel and quantified using a NanoDrop ND-1000 spectrophotometer at 260 nm (NanoDrop Technologies, Montchanin, DE).

Bacterial communities were examined using denaturing gradient gel electrophoresis (DGGE, Muyzer et al. 1993) in order to obtain a fingerprint of the microbial populations.
present within the individual soil samples. Although it is subject to biases of the PCR (Suzuki & Giovannoni 1996, Polz & Cavanaugh 1998), DGGE is capable of resolving DNA fragments differing by a single base pair (Myers et al. 1988, Lessa & Applebaum 1993) and detecting amplicons comprising ≥ 1% of the total community (Muyzer et al. 1993, Lee et al. 1996, Murray et al. 1996). For bacterial DGGE analysis, the V3 region (~ 180 bp) of the bacterial 16S rDNA genes present in soil samples was amplified using PCR (Saiki et al. 1988) with primers 338F/GC clamp and 519RC as described by Muyzer et al. (1993). For each sample, 3–25 µl reactions were pooled. Amplifications were performed using 10 ng template DNA, 1x JumpStart PCR Buffer (Sigma, St. Louis, MO), 0.2mM each deoxyribonucleotide triphosphate (dNTP), 2.5 mM MgCl2, Buffer (Sigma, St. Louis, MO), 0.2mM each primer, and 0.5U JumpStart Taq polymerase (Sigma). Three PCA amplifications from each sample, 400 ng of purified PCR product was loaded into lanes on opposite sides of the gel for DGGE analysis, the V3 region (~ 180 bp) of the bacterial 16S rDNA genes present in soil samples was amplified using PCR (Saiki et al. 1988) with primers 338F/GC clamp and 519RC as described by Muyzer et al. (1993). For each sample, 3–25 µl reactions were pooled. Amplifications were performed using 10 ng template DNA, 1x JumpStart PCR Buffer (Sigma, St. Louis, MO), 0.2mM each deoxyribonucleotide triphosphate (dNTP), 2.5 mM MgCl2, Buffer (Sigma, St. Louis, MO), 0.2mM each primer, and 0.5U JumpStart Taq polymerase (Sigma). Three PCA amplifications from each template were performed on a PTC-200 Thermocycler (MJ Research Inc, Waltham, MA) using the following hot start touchdown protocol: 94°C for 2 min, followed by 30 cycles of 94°C for 0.5 min, 65°C (-0.5°C per cycle until 55°C was reached) for 0.5 min, and 72°C for 1 min with a final elongation step consisting of 72°C for 7 min. Each set of amplifications was subsequently pooled for each template and evaluated through visualization on a 1% agarose gel, purified using a QIAquick PCR Purification Kit (Qiagen) and quantified spectrophotometrically as above.

For microbial community DGGE analysis, the PCR products were separated on an 8% acrylamide gel (37.5:1 ratio of arylamide-bisacrylamide) with a 30% to 70% gradient of denaturant (7 M urea and 40% formamide). For each sample, 400 ng of purified PCR product was loaded and run. Additionally, a repeatable standard (300 ng pooled PCR products from three known bacterial isolates) was loaded into lanes on opposite sides of the gel for normalization. All gels were electrophoresed for 5 hr (130V) at 60°C in 1X TAE buffer (40 mM Tris base, 20 mM sodium acetate, 1 mM EDTA) on a DCode Universal Mutation Detection System (Bio-Rad, Hercules, CA). Gels were stained with ethidium bromide (1 µg mL⁻¹) for 30 min, destained in sterile water for 20 min, visualized under UV and digitally photographed using the AlphaImager imaging system (Alpha InnoTech Corp, San Leandro, CA).

### Statistical analysis

Estimates of nematode, rotifer and tardigrade abundance were log(x+1) transformed to satisfy assumptions of normality and uniformity of variance. We performed conventional statistical analyses in SAS 9.1 to test for variation in soil biogeochemical properties, invertebrate abundance and diversity among and within locations in northern and southern Victoria Land (Barrett et al. 2004, Table I). We used principal component analysis (PCA) to characterize the distribution of metazoan and bacteria across ranges of soil biogeochemical properties encountered among the locations sampled in Victoria Land (CANOCO v4.02, Ter Braak 1986). PCA was run on the entire metazoan, bacteria and soil biogeochemical dataset and also on the metazoan-biogeochemistry data independent of the bacterial data to facilitate comparison with earlier work in southern Victoria Land (Courtright et al. 2001, Barrett et al. 2004). Correlation coefficients of nematode species abundances were plotted against the principle components in species-environment bi-plots (Ter Braak 1986, Palmer 1993).

Images of DGGE bands were analysed using Gelcompar II™ (Applied Maths, Kortrijk, Belgium) to allow normalization, grouping and identification of electrophoresis banding patterns within and among samples and gels. Manual band detection was used with an approximate minimal profiling of 12% and a minimal area of 0.6%. Similarities among tracks were calculated using the band-based similarity Dice coefficient with fuzzy logic. Based on these similarity coefficients, dendograms representing hierarchical linkage levels were constructed using the unweighted pair group method with arithmetic averages (UPGMA) clustering algorithm (Sokal & Michener 1958).

### Results

#### Soil properties

Soil biogeochemical and physical properties varied significantly among the locations and sites sampled. Mean

<table>
<thead>
<tr>
<th>Study site</th>
<th>Soil organic C (g kg⁻¹)</th>
<th>Total soil N (mg kg⁻¹)</th>
<th>C:N</th>
<th>Microbial biomass C (mg kg⁻¹)</th>
<th>Ammonium (N₄⁻-N) (mg kg⁻¹)</th>
<th>Nitrate (NO₃⁻-N) (mg kg⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Von Guerard Stream</td>
<td>0.29 ± 0.04</td>
<td>0.08 ± 0.01</td>
<td>3.98 ± 0.40</td>
<td>8.2 ± 10</td>
<td>0.41 ± 0.09</td>
<td>4.4 ± 1.97</td>
</tr>
<tr>
<td>Green Creek</td>
<td>0.35 ± 0.04</td>
<td>0.09 ± 0.01</td>
<td>4.20 ± 0.40</td>
<td>13.1 ± 1.5</td>
<td>0.17 ± 0.06</td>
<td>0.77 ± 0.38</td>
</tr>
<tr>
<td>Luther Vale North</td>
<td>0.53 ± 0.05</td>
<td>0.19 ± 0.01</td>
<td>2.69 ± 0.10</td>
<td>67.3 ± 5.6</td>
<td>0.22 ± 0.04</td>
<td>0.02 ± 0.01</td>
</tr>
<tr>
<td>Luther Vale South</td>
<td>0.69 ± 0.14</td>
<td>0.24 ± 0.02</td>
<td>2.48 ± 0.30</td>
<td>73.0 ± 9.5</td>
<td>0.16 ± 0.01</td>
<td>0.18 ± 0.07</td>
</tr>
<tr>
<td>Willett Cove</td>
<td>17.05 ± 1.80</td>
<td>3.19 ± 0.29</td>
<td>5.18 ± 0.10</td>
<td>204.8 ± 24.0</td>
<td>4.57 ± 0.46</td>
<td>2.94 ± 0.50</td>
</tr>
<tr>
<td>SeaBee Spit</td>
<td>21.90 ± 1.28</td>
<td>9.86 ± 0.68</td>
<td>2.26 ± 0.10</td>
<td>506.1 ± 108.0</td>
<td>718.40 ± 39.00</td>
<td>154.60 ± 54.00</td>
</tr>
</tbody>
</table>

Note: superscripts denote significant differences among means. Sites with the same letter are not significantly different (P < 0.05).
moisture content was highest in soils collected from the sites at Cape Hallett (9.47–25.87% g g\(^{-1}\)) relative to soils collected from sites in Luther Vale (3.91–4.66% g g\(^{-1}\)) and Taylor Valley (2.18–5.36% g g\(^{-1}\), \(F = 109.59, P < 0.0001\)). Moisture content also varied significantly among transects within study sites, particularly in Green Creek (1.11–15.58% g g\(^{-1}\), \(F = 12.07, P < 0.0001\)).

The most alkaline conditions were found in soils collected from Taylor Valley and Luther Vale (pH = 9.5 ± 0.1 and 9.4 ± 0.03, respectively), relative to soils from the sites in Cape Hallett, where pH was more variable and ranged from 5.3–8.4 (\(F = 333.53, P < 0.0001\)). Electrical conductivity of the soil varied among study sites from low values of 47–130 \(\mu\)S cm\(^{-1}\) in southern Luther Vale to 1200 \(\mu\)S cm\(^{-1}\) in marine influenced soils from Seabee Spit (\(F = 17.87, P = 0.0001\)).

Soil organic matter varied significantly among the sites (Table II). Soils from Cape Hallett had organic C and total N concentrations an order of magnitude above soils from Taylor Valley and Luther Vale. Differences in organic matter (C and N) concentrations among the sites were highly significant (\(F = 141.32, P < 0.0001\), \(F = 138.98, P < 0.0001\)). Patterns in microbial biomass C were similar to those of bulk soil organic C (\(F = 18.85, P < 0.0001\)), with the highest levels of microbial biomass found in Cape Hallett soils, followed by Luther Vale and Taylor Valley soils. Nutrient concentrations also varied among the sites (Table II). The highest levels of soil NH\(_4^+\) content were found at Cape Hallett (\(F = 228.69, P < 0.0001\)), particularly in the Seabee Spit site located near the penguin colony. Soil NO\(_3^-\) concentrations were also highest in the Seabee Spit site, followed by the two most disparate sites, Von Guerard and Willett Cove where NO\(_3^-\) concentrations averaged 3–4.4 mg N kg soil\(^{-1}\) (Table II, \(F = 7.08, P < 0.0001\)).

Stable isotopic composition of soil organic matter varied significantly among the study sites (Fig. 2, one-way ANOVA: \(F = 281.65, P < 0.00001\) and \(F = 4.33, P = 0.014\), for \(\delta^{15}N\) and \(13C\)‰, respectively). The greatest differences in stable isotope ratios were for N, with the enriched \(^{15}N/^{14}N\) ratios found in soils collected from Willett Cove and Seabee Spit, and relatively depleted values for soils from Taylor Valley and Luther Vale (Fig. 2). Ranges of \(^{13}C\) content were more similar among soils from all locations; the most notable trends were tight clustering of \(^{13}C\) values for soils from SeaBee Spit and Willett Cove (-24.8 and -18.2 \(\delta^{13}C\)‰, respectively), and broader ranges of \(^{13}C\) contents for Luther Vale and Taylor Valley soils (-15 to -25, and -17 to -32, \(\delta^{13}C\)‰, respectively, Fig. 2).

**Soil respiration**

Rates of midday soil respiration (Table III) in SeaBee Spit were an order of magnitude greater than any of the other sites (2.43 ± 0.96 \(\mu\)mol CO\(_2\) m\(^{-2}\) s\(^{-1}\)). Soil respiration rates observed at the Von Guerard site were the lowest (0.02 ±

**Table III.** Mean mid-day soil respiration and microclimate conditions (± SD) at each site.

<table>
<thead>
<tr>
<th>Study site</th>
<th>Soil respiration ((\mu)mol CO(_2) m(^{-2}) s(^{-1}))</th>
<th>Mid-day soil temperature (5 cm) (°C)</th>
<th>Soil moisture content (% g g(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Von Guerard Stream</td>
<td>0.02 ± 0.015(^a)</td>
<td>10.43</td>
<td>2.18± 0.26</td>
</tr>
<tr>
<td>Green Creek</td>
<td>0.31 ± 0.14(^b)</td>
<td>13.62</td>
<td>5.36± 1.52</td>
</tr>
<tr>
<td>Willett Cove</td>
<td>0.51 ± 0.23(^b)</td>
<td>9.31</td>
<td>25.87± 1.16</td>
</tr>
<tr>
<td>Seabee Spit</td>
<td>2.43 ± 0.96(^a)</td>
<td>9.31</td>
<td>9.47± 1.01</td>
</tr>
</tbody>
</table>

\(^a\)Not included in ANOVA because of missing data.

**Table IV.** Abundance (# kg soil\(^{-1}\)) of soil biota at each site (mean ± SE).

<table>
<thead>
<tr>
<th>Study site</th>
<th>S. lindsayae</th>
<th>Eudorylaimus</th>
<th>Plectus antarcticus</th>
<th>Panagrolaimus davidii</th>
<th>Rotatoria sp.</th>
<th>Heterotardigrada sp.</th>
<th>Hypsibius sp.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Von Guerard Stream</td>
<td>1052 ± 348</td>
<td>25 ± 10</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Green Creek</td>
<td>618 ± 155</td>
<td>322 ± 213</td>
<td>0</td>
<td>0</td>
<td>353 ± 171</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Luther Vale north</td>
<td>336 ± 76</td>
<td>0 ± 4</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>19 ± 19</td>
<td>0</td>
</tr>
<tr>
<td>Luther Vale south</td>
<td>247 ± 104</td>
<td>60 ± 18</td>
<td>0</td>
<td>0</td>
<td>6 ± 4</td>
<td>19 ± 12</td>
<td>0</td>
</tr>
<tr>
<td>Willett Cove</td>
<td>0</td>
<td>0</td>
<td>36 ± 11</td>
<td>0</td>
<td>241 ± 60</td>
<td>788 ± 446</td>
<td>272 ± 171</td>
</tr>
<tr>
<td>Seabee Spit</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>122 ± 40</td>
<td>18 ± 1</td>
<td>0</td>
</tr>
</tbody>
</table>

**Fig. 2.** Stable isotope bi-plot of Victoria Land soils.
0.015 μmol CO2 m⁻² s⁻¹), though these measurements were excluded from the ANOVA because of missing data following equipment malfunction in the field. Linear regression of means suggests that microbial biomass C (MBC) is a significant factor influencing rates of soil respiration (respiration = 0.0045*MBC-0.0078, \( r^2 = 0.96 \), \( P < 0.037 \)). Similar regressions for total soil organic C and soil moisture content were not statistically significant (\( P \) of linear regressions = 0.19 and 0.92, respectively).

**Metazoan diversity**

There were significant differences in soil metazoan community composition and population densities between the study sites (Tables IV & V). Total metazoan richness increased from Taylor Valley in southern Victoria Land, to the northern sites in the vicinity of Cape Hallett, with the greatest diversity found in soils collected from the Willett Cove site at Cape Hallett (Fig. 3a, \( F = 52.57, P < 0.0001 \)). There were similarities in metazoan diversity among intermediate sites, particularly Green Creek and North Luther Vale (Table IV). Collembolans and mites were only observed in northern Victoria Land (Cape Hallett and Luther Vale), though their distribution is known to extend to sites colonized by moss and microbial mat communities previously described in the Dry Valleys (Schwartz et al. 1993, Adams et al. 2006).

Distribution of tardigrades was limited to wet sediments, and were found only in soils collected from the Green Creek and Willett Cove sites. (Table IV). Rotifers (Rotatoria sp.) were ubiquitous across the study sites (though they were present in very low numbers in Luther Vale and SeaBee Spit), with the greatest abundances observed in soil from the Willett Cove site on Cape Hallett where population densities were more than five times greater than other metazoan populations (Table IV). DNA analyses of local tardigrade and rotifer communities confirmed morphological observations that they were dominated by single species. The tardigrade Hypsibius sp. was present at the Green Creek site, while an unknown member of the Heterotardigrada dominated the Willett Cove site (Table IV).

Nematode diversity (\( H' \)) was lowest at the extreme ends of the soil productivity gradient at Von Guerard Stream in Taylor Valley and from the most organic rich soils on SeaBee Spit at Cape Hallett (Fig. 3b, \( F = 9.36, P < 0.0001 \)). DNA sequence profiles of three out of four of the nematode species encountered were congruent with morphological observations (e.g. Andrassy 1998). However, based on DNA sequences of the small and large ribosomal subunits (SSU, LSU), our data indicate that Eudorylaimus is represented by at least three species, which we describe as follows: Eudorylaimus sp.1 was isolated from soils in the

<table>
<thead>
<tr>
<th>Variable</th>
<th>Source</th>
<th>df</th>
<th>F</th>
<th>( P )</th>
<th>Partial ( r^2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td>Site</td>
<td>5</td>
<td>27.71</td>
<td>&lt; 0.0001</td>
<td>0.50</td>
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<tr>
<td>Scottnema lindsayae</td>
<td>Transect</td>
<td>3</td>
<td>5.34</td>
<td>0.0021</td>
<td>0.058</td>
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<tr>
<td>abundance</td>
<td>Site*transect</td>
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<td>3.93</td>
<td>0.0001</td>
<td>0.16</td>
</tr>
<tr>
<td>Total</td>
<td>Site</td>
<td>5</td>
<td>11.46</td>
<td>&lt; 0.0001</td>
<td>0.17</td>
</tr>
<tr>
<td>Eudorylaimus spp.</td>
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<td>3</td>
<td>43.63</td>
<td>&lt; 0.0001</td>
<td>0.39</td>
</tr>
<tr>
<td>abundance</td>
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<td>5.52</td>
<td>0.0001</td>
<td>0.18</td>
</tr>
<tr>
<td>Total</td>
<td>Site</td>
<td>5</td>
<td>24.83</td>
<td>&lt; 0.0001</td>
<td>0.36</td>
</tr>
<tr>
<td>Plectus antarcticus</td>
<td>Transect</td>
<td>3</td>
<td>15.17</td>
<td>&lt; 0.0001</td>
<td>0.13</td>
</tr>
<tr>
<td>abundance</td>
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<td>6.24</td>
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<td>0.20</td>
</tr>
<tr>
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<td>34.66</td>
<td>&lt; 0.0001</td>
<td>0.59</td>
</tr>
<tr>
<td>Panagrolaimus davisii</td>
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<td>2.87</td>
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<td>0.029</td>
</tr>
<tr>
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<td>3.13</td>
<td>0.0015</td>
<td>0.12</td>
</tr>
<tr>
<td>Total</td>
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<td>27.44</td>
<td>&lt; 0.0001</td>
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</tr>
<tr>
<td>Rotifer</td>
<td>Transect</td>
<td>3</td>
<td>2.01</td>
<td>0.12</td>
<td>0.019</td>
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<td>abundance</td>
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<td>4.67</td>
<td>0.0001</td>
<td>0.16</td>
</tr>
<tr>
<td>Total</td>
<td>Site</td>
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<td>8.5</td>
<td>&lt; 0.0001</td>
<td>0.23</td>
</tr>
<tr>
<td>Tardigrade</td>
<td>Transect</td>
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<td>0.48</td>
<td>0.70</td>
<td>0.01</td>
</tr>
<tr>
<td>abundance</td>
<td>Site*transect</td>
<td>11</td>
<td>4.96</td>
<td>&lt; 0.0001</td>
<td>0.32</td>
</tr>
</tbody>
</table>
vicinity of Green Creek and Von Guerard Stream (Taylor Valley), *Eudorylaimus* sp. 2 from Luther Vale, and *Eudorylaimus* sp. 3 from Willett Cove (Fig. 4).

Total nematode abundance was greatest in soils collected from the Green Creek site, followed by soils collected from the Von Guerard Stream and Willett Cove sites (Fig. 3c, F = 6.95, P < 0.003), and was lowest in soils collected from SeaBee Spit (Fig. 2c). Variability encountered among transects within study sites was also significant, though typically accounted for a smaller proportion of variance than that contributed by differences among sites (Table V). Nematodes were found in a high proportion of soils examined among the study sites relative to previous work (Freckman & Virginia 1997, Courtright et al. 2001). Ninety percent of soils from the Taylor Valley transects near V on Guerard Stream and Green Creek, 95–100% of soil from Luther Vale and 70–90% of the samples examined from SeaBee Spit contained nematodes. Soils collected from SeaBee Spit had the highest frequency of instances where no metazoans were present (30%).

Patterns of metazoan distribution were strongly associated with variation in the availability of soil water and organic matter and species showed distinct habitat preferences. For example, the log(abundance+1) of *Scottnema lindsayae* Timm was negatively correlated with soil moisture and organic C content (r = -0.52, P < 0.0001 and r = -0.64, P < 0.0001, respectively); population densities were greatest in soils collected from the Von Guerard transects, followed by Green Creek and the Luther Vale sites (Table IV & V). *S. lindsayae* was not found in soils collected from Cape Hallett (Table IV). In contrast, the log(abundance+1) of *Panagrolaimus davidi* Timm was strongly correlated with both soil moisture and organic C content (r = 0.62, P < 0.0001 and r = 0.79, P < 0.0001, respectively), and this species was observed only in Cape Hallet soils where it was found in the greatest densities in the Willett Cove transects (Table IV & V). *Plectus antarcticus* de Man was observed only in wet soils with low electrical conductivity collected from Green Creek, South Luther Vale, and Willett Cove sites (Table IV). Nematodes of the genus *Eudorylaimus* were found in the greatest number of sites relative to other metazoans, with the highest abundances observed in soils collected from the vicinity of Green Creek (Table IV, F = 11.46, P < 0.0001).

**Bacterial diversity**

We were able to extract high-quality DNA from all samples using the newly developed extraction protocol whereby environmental enzyme inhibitors common in soil extracts using more conventional methods were minimized. Samples containing higher microbial biomass (Table II) provided significantly more DNA (both Dice band-based similarity and densiometric curves, data not shown). As expected, PCR with the bacteria-specific primer set including the forward primer with a G-C clamp resulted in a single 180-bp fragment. These PCR products were purified and quantified prior to DGGE separation to assess qualitative bacterial community composition; replicate DNA extractions from the same soil sample (n = 10)
showed highly homogenous DGGE banding patterns.

DGGE profiles revealed an unexpectedly high degree of diversity both within and among soil samples when compared to previous studies, with an average of 9.90 ± 3.23 distinct bands (range 4–21). DGGE profiles also contained regions of increased staining density possibly consisting of merging minor bands. There was no significant difference in number of bands among sampling sites ($F = 0.7, P > 0.5, df = 3$). However, many soils showed unique bacterial assemblages as well as some bands common to many of the soils.

DGGE profiling and subsequent cluster analysis (both Dice band-based similarity and densiometric curves) revealed significant differences in bacterial community structure with the largest effect being the geographic proximity of the soils sampled. For example, bacterial communities from Luther Vale tended to group together in several distinct clusters, while communities from Taylor Valley at Von Guerard and Green Creek generally clustered together (Fig. 5). Four main bacterial groups were associated with various sites from northern and southern Victoria Land (Fig. 5). Those soils that clustered with soils from different field sites did not group with those same soils based on PCA including the biogeochemical properties of soils. There was also a discernable effect based on the a priori designation of the productivity (low vs high) of soils upon bacterial diversity; soils designated as high productivity environments (within individual sites) were more similar to each other than they were to those designated as low productivity environments.

Co-variation in soil biological and biogeochemical parameters

The PCA plot (Fig. 6) illustrates significant co-variation between soil biota and biogeochemical properties. The first four principal components were selected for the analysis of the total biodiversity-biogeochemistry dataset; they accounted for 67% of the total variation. The PCA results showed that the first two principal components accounted
for a larger portion (24% and 20%, respectively) of the total variability when compared to the subsequent 3rd and 4th components (14% and 9%, respectively). The factor loadings on the first component were large for microbial biomass C and N, and total soil N. The largest loadings on the second component were for SOC, total moisture and chlorophyll \(a\). The third component had large loadings for electrical conductivity (salinity) and both inorganic nitrogen species (\(\text{NH}_4^+\), \(\text{NO}_3^-\)). The fourth component had the largest single loading for nematode abundance. The interrelations among the first three components are evident in the projections of components 1, 2, and 3 (Fig. 6). The positions of the samples in 3-dimensional space are related to both the geographic differences among the various sites and their \textit{a priori} designation as high or low productivity environments.

Four principal components accounted for greater than 96% of the total variance in the metazoan species and biogeochemical data that we considered independent of the bacterial data. The first two principal components accounted for 82% of the variance. The factor loadings of the soil biogeochemical data and the metazoan correlations with the first two principal components are plotted in Fig. 7. These data indicate distinct habitat preference for soil metazoans. For example, tardigrades, rotifers and \textit{Panagrolaimus davidi} are correlated with Principal Component 1, and place within the species-environment bi-plot in an area representing high levels of chlorophyll \(a\), soil moisture, organic matter, and microbial biomass. In contrast, \textit{S. lindsayae} was negatively correlated with Principal Component 1, and was most closely associated with high soil pH and low moisture and organic matter content.

**Discussion**

Regional patterns of soil respiration and biodiversity were strongly associated with variability in biogeochemistry. For example, there was a significant correlation between microbial biomass and maximum rates of soil respiration, while metazoan diversity was highest in wet, organic matter rich soils. Bacterial diversity, as indicated by DGGE band richness, did not vary significantly among the sites considered, though composition of bacterial communities did. These differences in bacterial community structure were mainly associated with geography. DGGE profiles did reveal a relatively high level of bacterial diversity both within and among soil samples consistent with recent descriptions of soils from southern Victoria Land (Cowan \textit{et al.} 2002, Aislabie \textit{et al.} 2006, Connell \textit{et al.} 2006). While DGGE is not very sensitive in detecting rare species, it is a useful index for comparing relative diversity and composition among microbial communities. Our results and others are challenging earlier assumptions and descriptions of limited microbial diversity in Antarctic soils (Cowan \textit{et al.} 2002, Connell \textit{et al.} 2006).

Many of the soils examined hosted unique bacterial assemblages while some bands were nearly ubiquitous across all the soils we examined. The distribution of similar bacterial and metazoan taxa across this range of latitudes suggest that dispersal is not a barrier to species distribution (e.g. Moorhead \textit{et al.} 2003, Nkem \textit{et al.} 2006); rather, local differences in soil biogeochemistry and physicochemical properties of soils together determine habitat suitability. The lack of a clear relationship between patterns of metazoan and bacterial diversity and community composition suggests that the respective habitat suitability for these different taxa are driven by distinct combinations of these biogeochemical and physical properties. This pattern is also evident in temperate soil ecosystems, (e.g. Black \textit{et al.} 2003).

There is great interest in the influence of predator/prey dynamics and invertebrate functional types on bacterial community structure and diversity in all soils, though these relationships have proven very difficult to elucidate (Bakonyi 1989, Coleman & Whitman 2005). To this end, we classified each soil sample based upon its relative abundance of nematodes (Table IV) but there was no visible effect of nematode abundance on bacterial community clustering. Nor was there a significant correlation between DGGE band richness and nematode abundance. These results do not exhibit top-down control over bacterial diversity and biomass mediated by metazoan consumers. Although food webs in the Antarctic are simple relative to nearly all other soils (Wall & Virginia 1999), the emerging view of biotic and/or trophic interactions in these simple communities is not (Hogg \textit{et al.} 2006).
Variation in sources of soil organic matter

Variation in the stable isotopic composition of soil organic C and total N among the sites suggests distinct sources of organic matter, particularly between the low lying coastal sites at Cape Hallett and the higher elevation sites of Taylor Valley and Luther Vale (Fig. 2, e.g. Burkins et al. 2000, Barrett et al. 2006). Differences in 15N content were more significant than variability in 13C, indicating a larger diversity of potential N source pools relative to C, along with a greater range of isotopic signatures of these source pools. Sources of soil N in Victoria Land are: atmospheric with highly depleted 15N values (Michalski et al. 2005), in situ N2 fixation with values near atmospheric (−0 8 15N ‰), or marine cycled N with highly enriched values greater than 20 8 15N ‰ (Burkins et al. 2000). Our data indicate that the largest disparities in N source pools are between soils from Cape Hallett which are influenced by marine N sources, and those from Taylor Valley which have atmospheric to slightly depleted values (Fig. 2). Though it is not possible to distinguish between marine aerosols and sea bird guano as sources to Cape Hallett soils, the large concentrations of organic matter and NH4+ suggest that direct input from sea birds is a much more important source than marine aerosols. The 13C values of some soils are greater than the average values for C3 photosynthesis, and may indicate in situ enrichment of organic matter following microbial degradation. The broader range of 13C content for Taylor Valley and Luther Vale may indicate a larger number of potential contributors to soil organic matter that may include C fixed by marine, lacustrine, endolithic and in situ phototrophs (Burkins et al. 2000, Lawson et al. 2004).

The higher elevation areas in northern Victoria Land do not receive significant marine inputs and consequently the soil organic matter concentrations and isotopic signatures are more similar to the dry valleys than to neighboring Cape Hallett, indicating a terrestrial source of organic matter, i.e. algae, moss and lichens. Soil metazoan communities are also similar to the dry valleys, with the nematode S. lindsayae dominating faunal populations (Freckman & Virginia 1997). These similarities between Luther Vale and the dry valleys suggest that the source of C (algal vs ornithogenic) is a key factor in structuring soil invertebrate communities.

Controls over habitat suitability

Soil biogeochemical variables shared significant variance, which is illustrated by the clustering of environmental factor loadings in the PCA (Figs 6 & 7). For example, wet sites tended to be rich in organic matter and microbial biomass C, and these variables were strongly correlated with Principal Component 1 (Fig. 7). Saline, or alkaline soils were either negatively correlated or orthogonal to Principal Component 1 and these generally favorable site conditions for metazoans (Fig. 7). These underlying biogeochemical properties are important influences over the distribution of soil organisms, though the response of individual species to different soil properties varied. For example, S. lindsayae inhabits a quadrant of the Principal Component bi-plot that is strongly correlated with pH, while Panagrolaimus davidii, Rotatoria and tardigrades were most abundant in wet, organic matter rich soils (Fig. 7). Thus different metazoan taxa manifest distinct habitat preferences.

Abundance and distribution of S. lindsayae was unrelated to a priori designations of productivity across transect or latitudes (Table IV & V), or to observed levels of soil organic matter, nutrients and soil respiration (Table II). Numerous studies have demonstrated that this species is well adapted to the arid, nutrient poor habitats that are ubiquitous throughout Victoria Land (e.g. Freckman & Virginia 1997, Treonis et al. 1999, Courtright et al. 2001). In contrast, Plectus antarcticus and Panagrolaimus davidii were confined to narrowly defined and spatially restricted habitats. Plectus antarcticus is found only in wet to intermittently wet soils and sediments where salinity is low (Treonis et al. 1999), while Panagrolaimus davidii inhabits the wet, nutrient rich habitats of rookeries and adjacent soils (Porazinska 2002), that are evidently inhospitable to most other metazoans (e.g. Sinclair et al. 2003), as indicated by the lower diversity and biomass of organisms reported here (Fig. 3, Table III). Penguin rookeries are likely to be unsuitable habitats for most metazoans because of the elevated levels of organic acids, NH4+ and NO3− associated with ornithogenic soils (Cocks et al. 1998). Differences in nematode distribution may also be related to feeding preferences. For example, Panagrolaimus davidii is a bacterial feeder and may favor ornithogenic soils where culturable bacteria are plentiful relative to sites in southern Victoria Land, while S. lindsayae, a yeast and bacterial feeder, may have an advantage in the dry valleys where fungi are more abundant (Connell et al. 2006, Aislabie unpublished).

In contrast to the other nematodes, the genus Eudorylaimus is found throughout Victoria Land, in dry to wet soils and sediments (2–20% g g−1) of the McMurdo Dry Valleys (Freckman & Virginia 1997, Powers et al. 1998, Treonis et al. 1999, Courtright et al. 2001, Barrett et al. 2004), and saturated soils under algal mats and moss communities in northern Victoria Land (Table IV). The only habitats where Eudorylaimus spp. was not found were in the soils of SeaBee Spit, adjacent to active penguin mounds. This broad environmental tolerance may be related to the genetic variation encountered among the Eudorylaimus individuals. Our data indicate that there are at least three species of this genus occupying the distinct sites where it was present (Fig. 4). Thus greater taxonomic resolution revealed important, though subtle differences in habitat preference among three morphologically indistinguishable
species. It is also significant that these three species occur across three out of four of the sites where distinct bacterial communities were identified through DGGE analysis. These results suggest that greater scrutiny of previously identified taxonomic groups may yield greater ecological insight to controls over their distribution and perhaps the potential for biotic interaction among metazoan and microbial taxa.

Conclusions

Soil biodiversity varied across the latitudinal gradient studied, with the most significant differences encountered among sites with distinct sources and concentrations of soil organic matter and physicochemical properties. There were no apparent differences in bacterial diversity among the sites considered, though composition of bacterial communities was significantly different. There was no correlation between bacterial diversity and invertebrate abundance or diversity suggesting that controls over microbial diversity are not mediated by higher trophic level communities (and vice versa). The limited number of sites, and their uneven distribution across the gradient prevents conclusions about latitudinal and/or climate controls over patterns of soil biodiversity. However, significant heterogeneity in edaphic factors among sites contributed to variability in distribution and abundance of soil biota. This suggests that broad scale trends in soil chemistry, for example, the availability and composition of organic matter or the concentration of salts, are important influences over the composition of soil communities.

Acknowledgements

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References


