

Aberystwyth University

Niche partitioning of bacterial communities in biological crusts and soils under grasses, shrubs and trees in the Kalahari

Elliott, David R.; Thomas, Andrew D.; Hoon, Stephen R.; Sen, Robin

Published in: Biodiversity and Conservation DOI:

10.1007/s10531-014-0684-8

Publication date: 2014

Citation for published version (APA):

Elliott, D. R., Thomas, A. D., Hoon, S. R., & Sen, R. (2014). Niche partitioning of bacterial communities in biological crusts and soils under grasses, shrubs and trees in the Kalahari. *Biodiversity and Conservation*, 23(7), 1709-1733. https://doi.org/10.1007/s10531-014-0684-8

General rights

Copyright and moral rights for the publications made accessible in the Aberystwyth Research Portal (the Institutional Repository) are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

• Users may download and print one copy of any publication from the Aberystwyth Research Portal for the purpose of private study or You may not further distribute the material or use it for any profit-making activity or commercial gain

- · You may freely distribute the URL identifying the publication in the Aberystwyth Research Portal

Take down policy

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

tel: +44 1970 62 2400 email: is@aber.ac.uk

\$

•	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684		TYPESET
•	MS Code : BIOC-D-13-00927	CP	V DISK

Biodivers Conserv DOI 10.1007/s10531-014-0684-8

1

3

Niche partitioning of bacterial communities in biological 2 crusts and soils under grasses, shrubs and trees

in the Kalahari 4

5 David R. Elliott · Andrew D. Thomas · Stephen R. Hoon · 6 **Robin Sen**

7 Received: 22 November 2013/Revised: 21 March 2014/Accepted: 24 March 2014

8 © Springer Science+Business Media Dordrecht 2014

9 Abstract The Kalahari of southern Africa is characterised by sparse vegetation inter-10 spersed with microbe-dominated biological soil crusts (BSC) which deliver a range of 11 ecosystem services including soil stabilisation and carbon fixation. We characterised the 12 bacterial communities of BSCs (0-1 cm depth) and the subsurface soil (1-2 cm depth) in 13 an area typical of lightly grazed Kalahari rangelands, composed of grasses, shrubs, and 14 trees. Our data add substantially to the limited amount of existing knowledge concerning 15 BSC microbial community structure, by providing the first bacterial community analyses of both BSCs and subsurface soils of the Kalahari region based on a high throughput 16S 16 17 ribosomal RNA gene sequencing approach. BSC bacterial communities were distinct with 18 respect to vegetation type and soil depth, and varied in relation to soil carbon, nitrogen, and 19 surface temperature. Cyanobacteria were predominant in the grass interspaces at the soil 20 surface (0-1 cm) but rare in subsurface soils (1-2 cm depth) and under the shrubs and 21 trees. Bacteroidetes were significantly more abundant in surface soils of all areas even in 22 the absence of a consolidated crust, whilst subsurface soils yielded more sequences affil-23 iated to Acidobacteria, Actinobacteria, Chloroflexi, and Firmicutes. The common detection of vertical stratification, even in disturbed sites, suggests a strong potential for BSC 24 25 recovery after physical disruption, however severe depletion of *Cyanobacteria* near trees 26 and shrubs may limit the potential for natural BSC regeneration in heavily shrub-27 encroached areas.

Communicated by Guest Editors of S.I. : Biocrust. A1

A2 Electronic supplementary material The online version of this article (doi:10.1007/s10531-014-0684-8) A3 contains supplementary material, which is available to authorized users.

A4 D. R. Elliott (\boxtimes) · S. R. Hoon · R. Sen

A5 School of Science & the Environment, Manchester Metropolitan University, Manchester M1 5GD, UK

A6 e-mail: d.elliott@mmu.ac.uk

A7 A. D. Thomas

Department of Geography and Earth Science, Aberystwyth University, Llandinam Building, Penglais, A8

A9 Aberystwyth SY23 3DB, UK

	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684		□ TYPESET
$\boldsymbol{\boldsymbol{\sim}}$	MS Code : BIOC-D-13-00927	CP	🔽 DISK
			Biodivers Con

28 **Keywords** Biological soil crust · 454 Pyrosequencing · Bacterial community · Valabari cond. Carbon Vacatation

29 Kalahari sand · Carbon · Vegetation

31 Introduction

32 Soils are vital for agricultural productivity, biodiversity and carbon storage (Stringer 2008), 33 properties that are dependent upon the presence and activities of soil microbial communities (Brussard 2012). Most studies on soil microbes have focused on productive agri-34 cultural systems and typically target bulk soil or the plant-associated rhizosphere (e.g. 35 Rousk et al. 2010; Pereira et al. 2012; Fierer et al. 2012a; Phosri et al. 2012). Because 36 37 drylands typically support a patchy vegetation cover, light reaches the soil surface in plant interspaces and facilitates the formation of a complex autotrophic and heterotrophic 38 39 microbial community, which binds soil into a crust. These biological soil crusts (BSC) are 40 a major component of dryland biodiversity (Büdel et al. 2009) and of global carbon and 41 nitrogen cycles (Elbert et al. 2012). However, we have very little information on the microbial content of dryland soils and BSCs, and even less on how the soil microbes affect 42 43 dryland ecosystem function. A key factor is the lack of a detailed characterisation of the 44 microbial composition of BSCs, with the exception of a few important groups such as the 45 Cyanobacteria (e.g. Dojani et al. 2013). Addressing this research gap is essential if future management and conservation of drylands is to be effective in the face of increasing 46 climatic and anthropogenic pressures (Stringer et al. 2012). This is a major area of concern 47 because drylands cover approximately 40 % of the global land surface and support 38 % of 48 49 the human population (Reynolds et al. 2007).

50 Many of the ecosystem functions of BSCs are attributed to Cyanobacteria, which 51 comprise a large fraction of BSC biomass (Gundlapally and Garcia-Pichel 2006). Cya-52 nobacteria can sequester carbon through photosynthesis and fix nitrogen (Elbert et al. 53 2012), enabling BSCs to perform similar ecosystem functions to plants (Bowker et al. 54 2010). BSCs also facilitate numerous ecosystem services of importance to land manage-55 ment, conservation and productivity (Thomas 2012). These include soil stabilisation 56 (Thomas and Dougill 2007), nitrogen fixation (Aranibar et al. 2003), moisture retention 57 (Menon et al. 2011), and modulation of surface runoff (Eldridge and Greene 1994; Belnap 58 2006).

59 Despite the recognition of a significant non-phototrophic component of BSC communities (Garcia-Pichel et al. 2003; Bowker et al. 2010) there have been relatively few studies 60 61 characterising functionally important bacterial heterotrophs in BSCs. Most studies of dryland BSC communities have used a combination of cultivation and molecular 16S 62 63 rRNA gene fingerprinting techniques followed by identification of isolates and molecular 64 types by sequencing. These methods have revealed much about BSC community structure, including relationship to ecosystem functioning (Castillo-Monroy et al. 2011), demon-65 stration of vertical stratification (Garcia-Pichel et al. 2003), similarity of diversity 66 regardless of nearby plants (Nagy et al. 2005), and identification of numerically dominant 67 phyla (e.g. Gundlapally and Garcia-Pichel 2006). However, the methodology employed in 68 69 these studies severely limits microbial identification to no more than a few hundred species 70 or operational taxonomic units (OTUs). This is insufficient to make rigorous community 71 comparisons, particularly in a well-replicated study where more samples must be identi-72 fied, thus yielding a lower number of sequences per sample. Current high-throughput

	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684	🗆 LE	TYPESET
\sim	MS Code : BIOC-D-13-00927	CP	🔽 DISK
Biodivers Conserv			

sequencing technologies such as 454 pyrosequencing overcome this practical limitation

and have recently been used by Steven et al. (2013) to reveal differences in BSC and soil

73

74

81

75 bacterial community structure in relation to parent material in Colorado, USA. In a broader 76 study, Fierer et al. (2012b) went a step further by performing a cross-biome metagenomic 77 survey of soil (0-5 cm depth) microbial communities in tandem with a high-throughput 78 phylogenetic survey, enabling a comparison of functional gene frequencies as well as 79 microbial taxa. This study showed that hot and cold desert communities are quite distinct 80 from forest, grassland and tundra on both a taxonomic and functional level. In this paper we present the first description of bacterial community structure within 82 BSCs and soils of the Kalahari in the south west of Botswana, and assess niche partitioning 83 with respect to depth and nearby vegetation based on high-throughput sequencing of the

84 bacterial 16S rRNA gene. Weakly developed BSCs in the Kalahari Sand soils are found in 85 large areas that are subjected to disturbance by livestock and wildlife activity, which 86 prevents succession into more developed stages (Thomas and Dougill 2006). Grazing also 87 selectively removes palatable grasses and eventually leads to woody shrub encroachment 88 which renders the land useless for continued grazing (Thomas 2012). Thus, both vegetation 89 and co-occurring BSC cover in Kalahari rangelands are strongly affected by human 90 activity and can be influenced by land management decisions relating to factors such as 91 animal stocking density, fencing, and water access.

92 Our hypothesis is that landscape impacts related to grazing pressure such as disturbance 93 and shrub encroachment drives functionally significant soil surface bacterial community 94 changes and thus should be included in land management decision-making. Our specific 95 objectives were to determine whether there are significant differences in bacterial popu-96 lations associated with: (i) soils under tree, shrub and grasses, (ii) subsurface soils and 97 BSCs.

98 Materials and methods

100 Samples were collected from a long-term research site near Tsabong in south west Bots-101 wana (25°56'51"S, 22°25'40"E) at the end of the dry season in November 2011 and during 102 the wet season in March 2012. Soils are formed on Kalahari Sands and are weakly acidic, 103 fine sand-sized Arenosols (FAO 1990), with little or no horizon development. In lightly 104 grazed areas, around 80 % of the surface is covered in a 3-4 mm deep BSC, which has 105 been described in detail elsewhere (Thomas and Dougill 2007; 2012). BSC cover is 106 inversely related to grazing intensity and in frequently grazed areas, cover is typically 107 <10 % of the surface. The organic matter, carbon and nitrogen content of the Kalahari 108 Sand soils is low, reflecting the limited biological productivity and highly oxidising nature 109 of the soils (Thomas et al. 2012). BSCs, however, are enriched in ammonium, total N and 110 organic C compared to the mineral soil (Thomas and Dougill 2007; 2012).

111 Vegetation cover is typical of an open-canopy, fine-leaf savannah, with a mix of 112 perennial (Eragrostis lehmanniana) and annual (Schmidtia kalahariensis) grasses, woody 113 shrubs (Grewia flava and Acacia mellifera) and trees, predominantly Acacia erioloba 114 (Fig. 1). Mean annual precipitation is 331 mm (1996–2013), with a low of 114 mm in 115 2006-07 and a high of 532 mm in 2001-02. Seasonal variations in air temperature are 116 extreme, with summer maxima frequently in excess of 40 °C and winter below 0 °C.

⁹⁹ Study site

~	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684	□ LE	TYPESET
5	MS Code : BIOC-D-13-00927	CP	🖌 DISK
			Biodivers Cons



Fig. 1 Photographs showing examples of the four sampling zones. Letters in square brackets indicate abbreviations used throughout the manuscript. **a** Annual grass [AG] *Schmidtia kalahariensis* interspace. **b** Perennial grass [PG] *Eragrostis lehmanniana* interspace. **c** Shrub [S] *Grewia flava*. **d** Tree [T] *Acacia erioloba*

117 Sample collection and preparation

118 Soils and BSCs were collected from within a fenced 800×500 m paddock where grazing 119 animals had been excluded since the previous year. Sites were selected according to the 120 overlying vegetation type, with three replicate sites under trees (*A. erioloba*), woody shrubs 121 (*G. flava*), perennial (*Eragrostis lehmanniana*) and annual (*Schmidtia kalahariensis*) 122 grasses (Fig. 1).

Soils at all sites, except those under trees, were covered in a BSC, previously classified as type 1 (weakly consolidated with no surface discolouration) and 2 (more consolidated with black or brown speckled surface) by Thomas and Dougill (2006, 2007). Soils underneath trees were not crusted, but unconsolidated and slightly darkened by fragmented litter. There was evidence of severe animal disturbance under all tree canopies, a legacy of cattle seeking shade in the year prior to sampling. Soils under *G. flava* canopies were covered in a dry layer of leaf litter but were also well crusted.

Samples were collected using aseptic techniques from 0 to 1 cm (incorporating the BSC) and 1–2 cm (the soil immediately below the BSC). The same sites were sampled in November 2011 and March 2012, giving a total of 96 samples (48 in each season). Soils were dry at the time of sampling.

🖄 Springer

•••	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684	□ LE	TYPESET
\sim	MS Code : BIOC-D-13-00927	CP	🖌 DISK

134 Physico-chemical analyses

135 Total carbon and nitrogen content of BSC and subsurface soils were determined using a 136 CN element analyser (Leco TruSpec). Soil surface temperature was measured at approximately 2-h intervals during the day at each sampling site for the duration of each field 138 campaign, using an infrared thermometer.

139 Molecular analyses of bacterial community composition

140 DNA was extracted from soil samples within 18 h of sampling using a Powersoil DNA 141 extraction kit (MoBio Inc.). Prior to extraction, samples of approximately 20 g were 142 homogenised by shaking followed by cutting with a scalpel to disaggregate. Extractions 143 were performed according to the manufacturer's instructions except that the soil mass was 144 increased slightly from 0.25 to 0.4 g based on laboratory extraction tests and consultation 145 with the manufacturer. DNA was eluted into 50 µl of buffer (10 mM Tris). Phylogeneti-146 cally informative DNA sequences were obtained from each sample by tag-encoded FLX 147 amplicon pyrosequencing targeting the bacterial 16S rRNA gene (Dowd et al. 2008). This 148 analysis was performed by Research and Testing Laboratory (Lubbock, TX, USA), using a 149 Roche 454 FLX instrument with Titanium reagents.

150 DNA was amplified for pyrosequencing using forward and reverse fusion primers. The 151 forward primer was constructed with the Roche A linker (CCATCTCATCCCTGCGT 152 GTCTCCGACTCAG), an 8-10 bp barcode (see Online Resource 1), and the 341F primer 153 (CCTACGGGAGGCAGCAG) (Muyzer et al. 1993). The reverse fusion primer was 154 constructed with a biotin molecule, the Roche B linker (CCTATCCCCTGTGTGCCTT 155 GGCAGTCTCAG), and the 907R primer (CCGTCAATTCMTTTGAGTTT) (Muyzer 156 et al. 1998). Amplifications were performed in 25 µl reactions with Qiagen HotStar Taq 157 master mix (Qiagen Inc, Valencia, California), 1 µl of each 5 µM primer, and 1 µl of 158 template. Reactions were performed on ABI Veriti thermocyclers (Applied Biosytems, 159 Carlsbad, California) under the following thermal profile: 95 °C for 5 min, then 35 cycles 160 of 94 °C for 30 s, 54 °C for 40 s, 72 °C for 1 min, followed by one cycle of 72 °C for 161 10 min and 4 °C hold. PCR products were visualized with eGels (Life Technologies, 162 Grand Island, New York), pooled equimolar, and size selected before sequencing following 163 manufacturer protocols.

164 Bioinformatics and statistical analyses

165 Sequence data were processed through the QIIME pipeline (Caporaso et al. 2010) and 166 further analyses were performed using R (R Core Team 2012). Denoising, quality filtering, OTU assignment, OTU table generation, and phylogenetic determinations were all per-167 168 formed in QIIME. Sequences shorter than 200 bp or having an average quality score <25169 within a 50 base pair window were discarded. A 97 % sequence similarity was used to 170 define OTUs (approximately species level; Stackebrandt and Goebel 1994) which were 171 assigned using UCLUST (Edgar 2010), and chimeras were removed using chimeraslayer 172 (Haas et al. 2011). After quality control there were on average 1,004 sequences per sample 173 and a total of 2,705 OTUs (further details are provided in Online Resource 1). OTUs were 174 identified through the RDP classifier (Wang et al. 2007) using the Greengenes database 175 release of October 2012 (DeSantis et al. 2006). Identified OTUs were assembled into an 176 OTU table summarising the frequency of observation in each sample. These results formed

177 the basis for the determination and comparison of community structure.

 Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
Article No. : 684		TYPESET
\$ MS Code : BIOC-D-13-00927	CP	🚺 DISK
		Biodivers Cons

178 Thousands of different taxa were detected, so constrained correspondence analysis 179 (CCA) was used to discern community features which specifically relate to vegetation zone 180 or depth. Rare species comprising <0.01 % of the sequences detected in the study were 181 excluded from correspondence analysis because rare species can obscure community 182 patterns and may be differentially detected depending on sample sequencing depth. Cor-183 respondence analysis was based on the Bray-Curtis distance measure and performed using 184 the Phyloseq (McMurdie and Holmes 2013) wrapper to the Vegan package (Oksanen et al. 18 Aqt 2013) for R. Unconstrained correspondence analysis (CA) was performed to visualise the 186 overall community structure (shown in supplementary data Online Resource 2 only) whilst 187 CCA was employed to discern community features which specifically relate to vegetation 188 zone or depth. Permutation tests (n = 1,000) were used to test the significance of measured 189 environmental variables to the ordination.

Differences in community structure were assessed using ADONIS, a permutational
multivariate analysis of variance test in the R package Vegan (Oksanen et al. 2013), to
determine whether communities differ with respect to vegetation zone, depth, or sampling
month. We used the Bray-Curtis distance measure and performed the test at all taxonomic
levels from phylum to species. OTU richness and community diversity were estimated
using the Chao1 and Shannon methods respectively, implemented in the Phyloseq package.
Richness and diversity calculations used the full data set.

197 Kruskal–Wallis tests were used to determine whether each OTU relative abundance 198 differed between vegetation zone, depth, or sampling month. The significance of OTU 199 abundance correlations with continuous variables (e.g. carbon or richness) was assessed 200 using Spearman's test. P-values were corrected to account for multiple comparisons using 201 the false discovery rate method (Benjamini and Hochberg 1995). Significant findings were 202 tested further using post hoc tests to identify the changes responsible. We regarded results 203 with corrected p < 0.05 as being significant.

204 Availability of sequence data

Sequence data and metadata are available on the MG-RAST metagenomics analysis server
 (Meyer et al. 2008) at http://metagenomics.anl.gov/linkin.cgi?project=6691.

- 207 Results
- 208 Soil chemistry

Soil carbon was significantly higher in the soil surface $(0.7 \% \pm 0.1 \text{ SE})$ compared to the subsurface soil $(0.4 \% \pm 0.1 \text{ SE})$, and also differed significantly between vegetation classifications (Fig. 2) but not by month. Total soil carbon and nitrogen were closely correlated and the mean C:N ratio was 9:1. The C:N ratio was significantly higher in BSCs, (ANOVA F = 8.91, df = 1, p = 0.0047), but did not vary significantly with respect to vegetation zones.

215 Bacterial diversity

The Chao1 richness estimate and shannon diversity index (Fig. 3) provide an indication of the total number of species and the microbial diversity (taking account of number of

Article No. : 684	TYPESET
MS Code : BIOC-D-13-00927 🖓 CP	 DISK

Fig. 2 Total carbon and nitrogen in BSC (0–1 cm depth) and soil (1–2 cm depth) samples at each site (n = 6). *Boxes* represent the interquartile range (IQR), and *error bars* extend to the most extreme values within 1.5 * IQR of the *box*. Median values are shown as a *line* within the *box* and *outliers* are shown as *black spots*. Sample coding: *AG* annual grass, *PG* perennial grass, *S* shrub, *T* tree. Note that y axes differ



218 species and evenness) in each niche. Richness and diversity differed significantly with 219 respect to vegetation zone (Kruskal–Wallis Chi squared = 22.91, df = 3, p 220 value = 4.22×10^{-5}) but not depth, although diversity was close to our significance 221 threshold for depth (Kruskal–Wallis Chi squared = 2.93, df = 1, p-value = 0.087). Both 222 diversity (Spearman's rank correlation rho = 0.5, p value = 3×10^{-4}) and richness 223 (rho = 0.5, p value = 3×10^{-4}) were positively correlated with carbon content of the 224 soil. Diversity measures for each sample are provided in Online Resource 1.

225 Bacterial community structure

226 ADONIS showed that the bacterial communities differed by depth and by vegetation zone 227 (p < 0.05, see Online Resource 3 for test statistics) at all taxonomic ranks from phylum to 228 species, but did not differ in relation to sampling month except for at the rank of Family. 229 Interactions were detected (p < 0.05) between vegetation zone and depth at all taxonomic 230 ranks except for Genus (p = 0.051), and also between month and vegetation zones at 231 family level and higher taxa. A total of 28 bacterial phyla were detected, and the top 9 232 shown in Fig. 4 account for 99 % of sequences. Sequences from Actinobacteria and 233 Proteobacteria numerically dominated the samples, together representing 63 % of 234 sequences. Most phyla detection frequencies differed significantly (p < 0.05, see Online 235 Resource 4 for test statistics) with respect to depth or vegetation zone, or both (Table 1). 236 Cyanobacteria and Chloroflexi were more abundant in grass areas, especially the Cya-237 nobacteria which were only rarely detected in tree and shrub areas. Cyanobacteria were 238 also very rare in all subsurface soils (1-2 cm depth), only being found in large number in 239 the BSCs of grass interspaces. Bacteroidetes and Cyanobacteria were significantly asso-240 ciated with BSCs (0-1 cm depth), whilst Acidobacteria, Actinobacteria, Chloroflexi, and 241 Firmicutes were significantly associated with subsurface soil (1-2 cm depth). Phylum 242 composition did not vary significantly in relation to the sampling season for any of the top 243 9 phyla. Data for rarer phyla is also provided in Online Resource 4.

244 Constrained CA was used to generate a visual representation of the microbial com-245 munity structure differences between depths and vegetation zones. Sequences accounting

	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684		□ TYPESET
5	MS Code : BIOC-D-13-00927	CP	DISK
			Biodivers Co



Fig. 3 OTU richness estimation (Chao1) and diversity index (Shannon). Results are *plotted* with respect to **a** sampling site; and **b** sample carbon content. *Boxes* represent the interquartile range (IQR), and *error bars* extend to the most extreme values within 1.5 * IQR of the box (n = 5 or 6). Median values are shown as a *line* within the *box* and *outliers* are shown as *black spots. Error bars* in **b** indicate the *standard error* of the individual Chao1 estimations. Sample coding: AG annual grass, PG perennial grass, S shrub, T tree. Filled symbols = 0–1 cm depth, *hollow symbols* = 1–2 cm depth

246 for <0.01 % of the library were excluded from this analysis, leaving 934 OTUs. The 247 microbial communities were separated by vegetation type (tree/shrub or grasses) on axis 1, 248 and by depth on axis 2 (Fig. 5). Unconstrained CA yielded similar but less well defined 249 patterns (Online Resource 2). Soil carbon (and nitrogen) content and soil temperature were 250 significant to the ordination as determined by permutation tests (n = 1,000). Vectors show 251 that soil carbon and nitrogen increase with axis 1 (direction of tree and shrub samples), 252 whilst soil surface temperature increases in the opposite direction (i.e. in the direction of 253 grass interspace samples).

254 Abundant taxa (OTU level)

The detection frequencies of the most abundant 9 OTUs are shown in Fig. 6. Together these 9 OTUs accounted for 27 % of sequences and all of them had different detection

Article No. : 684	TYPESET
MS Code : BIOC-D-13-00927 CP	🖌 DISK





Fig. 4 Phylum abundance by **a** site and **b** soil carbon content. *Boxes* represent the interquartile range (IQR), and *error bars* extend to the most extreme values within 1.5 * IQR of the *box* (n = 6). Median values are shown as a *line* within the *box* and *outliers* are shown as *black spots*. Sample coding: *AG* annual grass, *PG* perennial grass, *S* shrub, *T* tree. Significance and direction of correlation between phylum abundance and soil carbon is indicated by + or – (determined by Spearman test). Significance codes for positive correlation: +++ < 0.001; ++ < 0.01; + < 0.05. Similar plots for less abundant phyla are included in in Online Resource 4

	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684		TYPESET
\$	MS Code : BIOC-D-13-00927	CP	🖌 DISK
			Biodivers Cons

Table	1	Significance	of	factors	depth	and	vegetation	on	phylum	abundance,	as	determined	by	Kruskal-
Wallis	te	st												

Phylum	Depth	Veg.
Chloroflexi	*	***
Proteobacteria		***
Bacteroidetes	**	***
Acidobacteria	**	**
Firmicutes	**	
Actinobacteria	**	
Gemmatimonadetes		***
Cyanobacteria	**	**
WPS-2		***

Significant effects were further tested by post hoc analyses which are provided in Online Resource 4. Significance codes: *** < 0.001; ** < 0.01; * < 0.05; < 0.1



Fig. 5 Constrained correspondence analysis of the microbial community. Coloured markers indicate individual samples, and dispersion ellipses show the 95 % standard deviation confidence interval for different depths and tree/shrub vs grass interspace classifications. Filled symbols = 0-1 cm depth, hollow symbols = 1-2 cm depth. OTU identification numbers are shown in different colours for the 9 most abundant OTUs belonging to the followng groups: full dataset (*black*), phylum *Cyanobacteria* (*green*), and phylum *Bacteroidetes* (*blue*). Environmental variables with significance p < 0.05, are shown as biplotted vectors (based on permutation tests; n = 1,000)

frequencies (p < 0.05, see Online Resource 4 for test statistics) with respect to depth or vegetation zone (Table 2). Similar plots and tables for the top 9 *Bacteroidetes* and *Cyanobacteria* are shown in Fig. 7 and Table 3, which account for 10 % of sequences. The OTUs of these two phyla were selected because they were detected more often in BSC samples compared to subsurface soil. OTU composition did not vary significantly in

Deringer

•	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684	□ LE	TYPESET
\sim	MS Code : BIOC-D-13-00927	CP	V DISK



Fig. 6 Relative abundance of the top 9 OTUs detected in the study, by **a** site and **b** soil carbon content. *Boxes* represent the interquartile range (IQR), and *error bars* extend to the most extreme values within 1.5 * IQR of the *box* (n = 6). Median values are shown as a *line* within the *box* and *outliers* are shown as *black spots*. Significance and direction of correlation between OTU abundance and soil carbon is indicated by + or - (determined by Spearman test). Significance codes for positive correlation: +++ < 0.001; ++ < 0.01; + < 0.05. Sample coding: *AG* annual grass, *PG* perennial grass, *S* shrub, *T* tree

~	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684		TYPESET
$\boldsymbol{\boldsymbol{\sim}}$	MS Code : BIOC-D-13-00927	CP	🔽 DISK
			Biodivers Con

262

263

264

265

266

267

268

269

270

271

272

relation to the sampling month for any of the top 9 OTUs (Online Resource 4). In addition to the overall top OTUs shown in Fig. 6, we present in Online Resource 4 similar plots and tables for the most abundant OTUs belonging to each phylum. These are included to permit readers interested in particular taxa to easily investigate these in our data, however space and time do not permit detailed presentation of more than a few OTUs here.

The OTUs shown in Figs. 6 and 7 are additionally plotted on the ordination in Fig. 5, illustrating their contributions to the community structure of samples in the ordination. Cyanobacterial OTUs are all clustered near the grass interspace BSC samples (0–1 cm depth) whilst *Bacteroidetes* OTUs are more spread out but tending towards the tree and shrub soil surface samples (0–1 cm depth). The overall most abundant OTUs are spread out on the ordination but with more near the grass area samples.

273 Discussion

274 Phylum level community structure

The distribution of the top 9 bacterial phyla accounting for 99 % of sequences (Fig. 4; Table 1) indicate that *Bacteroidetes* and *Cyanobacteria* are significantly associated with BSCs (Fig. 7; Table 3). The crucial role of *Cyanobacteria* in BSC carbon and nitrogen cycling is already widely recognised, whereas *Bacteroidetes*, although ubiquitous in soil, are not commonly regarded as key BSC community members in the current literature.

280 Using similar methods to this study, Steven et al. (2013) aimed to determine BSC 281 microbial community differences with respect to soil type (sandstone, shale, and gypsum). 282 Their samples were numerically dominated by the same top 6 phyla found in this study: 283 Proteobacteria, Cyanobacteria, Chloroflexi, Bacteroidetes, Actinobacteria, and Acido-284 bacteria. They found Cyanobacteria and Proteobacteria to be associated with BSCs whilst 285 we found Cyanobacteria and Bacteroidetes to be associated with BSCs. The difference 286 may be because Bacteroidetes are particularly dominant in the soil surface under trees and 287 shrubs (e.g. Figure 5) which were not a factor in the work of Steven et al. (2013). In the 288 underlying soil they found enrichment of Chloroflexi, which we also found in addition to 289 Acidobacteria, Actinobacteria, and Firmicutes. Steven et al. (2013) suggested that the 290 Chloroflexi might be involved in anaerobic processes including photoheterotrophy and 291 chemoheterotrophy, which would increase the productivity of BSCs and enable them to 292 continue functioning under a wider range of environmental conditions.

293 Cyanobacteria comprised 8.1 % of our sequences as 38 different OTUs. They were 294 found predominantly in BSCs of the grass areas and are dominated by a single *Phormidium* 295 species (OTU 1912). Other typical cyanobacterial genera found in BSCs include Mi-296 crocoleus, Leptolyngbya, Nostoc, and Scytonema species (Büdel et al. 2009), but these 297 were not identified in our samples. The majority of cyanobacterial sequences obtained 298 could not be classified to genus level however and it is known that molecular identification 299 of Cyanobacteria is problematic (Dojani et al. 2013), so it is possible that these genera 300 were present but not detected or identified. A higher frequency of Cyanobacteria was 301 detected in the perennial grass compared to the annual grass, probably reflecting the more 302 stabilised interspaces of perennial grass being slightly more developed. Due to the dom-303 inance of *Cyanobacteria* in grass interspace BSCs, it is likely that some of the phyla or 304 OTUs found to be more abundant in subsurface soil are not specifically adapted for the soil 305 niche, but may be excluded from the surface by competition.

Table 2 Taxonomic classification of the most abundant 9 OTUs found in the study

	Phylum	CIASS		(I)
54	Proteobacteria	Alphaproteobacteria	Rhodospirillales	Acetobacteraceae			* * *
761	Acidobacteria	Solibacteres	Solibacterales	Solibacteraceae	Candidatus Solibacter		* *
1064	Proteobacteria	Alphaproteobacteria	Rhizobiales	Methylobacteriaceae	Methylobacterium	* * *	
1150	Proteobacteria	Alphaproteobacteria	Rhizobiales	Bradyrhizobiaceae	Balneimonas	*	* * *
1912	Cyanobacteria	Oscillatoriophycideae	Oscillatoriales	Phormidiaceae	Phormidium	* * *	*
2850	Proteobacteria	Alphaproteobacteria	Rhodospirillales	Rhodospirillaceae		* * *	* *
3067	Proteobacteria	Alphaproteobacteria	Rhodospirillales	Acetobacteraceae			*
3302	Actinobacteria	Actinobacteria	Actinomycetales	Kineosporiaceae			***
3323	Actinobacteria	Actinobacteria	Actinomycetales	Pseudonocardiaceae		* * *	

Biodivers Conserv

~	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684	□ LE	TYPESET
\sim	MS Code : BIOC-D-13-00927	CP	V DISK





Fig. 7 Relative abundance of the top 9 OTUs from the phyla **a** Bacteroidetes; **b** Cyanobacteria. *Boxes* represent the interquartile range (IQR), and *error bars* extend to the most extreme values within 1.5 * IQR of the *box* (n = 6). Median values are shown as a *line* within the *box* and *outliers* are shown as *black spots*. Sample coding: *AG* annual grass, *PG* perennial grass, *S* shrub, *T* tree. Similar *plots* for other phyla are included in Online Resource 4

Deringer

Author Proof

Veg.

*

Depth ** Flavisolibacter Flavisolibacter Segetibacter Segetibacter Genus Table 3 Taxonomic classification of the most abundant 9 Bacteroidetes and Cvanobacteria OTUs found in the study Chitinophagaceae Chitinophagaceae Chitinophagaceae Chitinophagaceae Chitinophagaceae Chitinophagaceae Chitinophagaceae Chitinophagaceae Flexibacteraceae Family Sphingobacteriales Sphingobacteriales Sphingobacteriales Sphingobacteriales Sphingobacteriales Sphingobacteriales Sphingobacteriales Sphingobacteriales Sphingobacteriales Order **Oscillatoriophycideae** Sphingobacteriia Sphingobacteriia Sphingobacteriia Sphingobacteriia Sphingobacteriia Sphingobacteriia Sphingobacteriia Sphingobacteriia Sphingobacteriia Class Cyanobacteria Bacteroidetes **3***acteroidetes* **3***acteroidetes* **Bacteroidetes** Bacteroidetes Bacteroidetes Bacteroidetes **Bacteroidetes** Bacteroidetes Phylum 508 3388 3476 3635 461 2853 3081 3166 686 17

Significance of depth and vegetation on OTU abundance is indicated by *** p < 0.001; ** p < 0.01; * p < 0.05.. < 0.1., as determined by Kruskal–Wallis test. Significant Phormidiaceae effects were further tested by post hoc analyses which are provided in Online Resource 4 Oscillatoriales Oscillatoriophycideae Cyanobacteria 3453

**

Phormidium

Phormidiaceae

Oscillatoriales

Oscillatoriophycideae Oscillatoriophycideae Oscillatoriophycideae

Oscillatoriophycideae

Cyanobacteria Cyanobacteria Cyanobacteria Cyanobacteria

3313

779

4C0d-2

MLE1-12

Oscillatoriophycideae Oscillatoriophycideae

Cyanobacteria Cyanobacteria Cyanobacteria

[110[198[499[626[912

 Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
Article No. : 684		TYPESET
\$ MS Code : BIOC-D-13-00927	CP	🖌 DISK
		Biodivers Conse

Like the *Cyanobacteria*, *Bacteroidetes* were also significantly more abundant in the soil surface compared to the subsurface soil, however in this case the relationship held also for soils under tree and shrub canopies. Furthermore, whereas the cyanobacterial abundance was mostly achieved through a single OTU being very abundant in grass area BSCs, the *Bacteroidetes* phylum abundance is due to the collective abundance of many OTUs across all of the vegetation zones (Fig. 7). Overall the *Bacteroidetes* were slightly less abundant than the *Cyanobacteria*, comprising 5.1 % of sequences, but there were many more OTUs (297).

314 *Bacteroidetes* are ubiquitous in the soil and have a vast catabolic repertoire, particularly 315 in the breakdown of complex carbohydrates. Thomas et al. (2011) point out that many 316 carbohydrates are niche specific, and this seems a very likely scenario for BSCs because their constituent organisms are subject to unique survival challenges which will select for 317 318 production of specialist biomolecules. Bacteroidetes may therefore play a vital role in the 319 recycling of niche specific carbohydrates and associated molecules. It is clear that the soil 320 surface in grass and tree/shrub areas will be fundamentally different in terms of carbo-321 hydrate composition because only the grass interspace areas receive significant primary 322 production from Cyanobacteria and the stress of midday sun, therefore different Bacter-323 oidetes OTUs may be providing specialist degradation roles in these distinct niches. A key 324 role for BSC Bacteroidetes in carbon cycling is further supported by the work of Bailey 325 et al. (2013), which demonstrated a link between Chitinophagaceae family Bacteroidetes 326 abundance with β -glucosidase activity in soil. Eight of the top 9 Bacteroidetes in our 327 samples belonged to the Chitinophagaceae family.

328 In an attempt to ecologically classify soil bacteria, Fierer et al. (2007) have shown in a 329 meta-analysis that phylum abundances in soil are correlated with carbon availability. This 330 enabled them to broadly classify *Bacteroidetes* and β -*Proteobacteria* as copiotrophs, and 331 Acidobacteria as oligotrophs based on whether phylum abundance is positively or nega-332 tively correlated with carbon availability respectively. Previously Smit et al. (2001) had 333 made a similar suggestion: that the ratio of α - and γ -Proteobacteria to Acidobacteria could 334 provide an indication as to the nutritional status of the soil. Copiotrophic and oligotrophic 335 groupings are similar to the r- and K-strategists recognised in macroscopic ecology 336 (MacArthur and Wilson 1967). Copiotrophs therefore can be expected to maximise their 337 growth rates when resources are plentiful whereas oligotrophs are adapted for maximal 338 efficiency in the use of rare resources.

339 Based on correlation with total soil carbon (Fig. 4b, Online Resource 4) our data 340 identify *Bacteroidetes*, *Gemmatimonadetes* and all proteobacterial classes (α , β , δ , γ) as 341 potential copiotrophs, and Acidobacteria and candidate division WPS-2 as potential oli-342 gotrophs in Kalahari soils. Chloroflexi abundances were also negatively correlated with 343 carbon, however due to their carbon fixing abilities soil carbon data are not suitable to 344 attempt classification. In general, oligotrophs may be expected to function as primary 345 colonisers in the development of BSCs on nutrient-poor soils such as Kalahari Sand, and 346 copiotrophs may become more dominant as the crust develops and becomes more pro-347 ductive. The mean carbon content in our samples was 0.5 $\% \pm 0.1$ SE with a maximum of 348 2.2 %, which is low compared to typical mesic soils (Lal 2004) so an abundance of 349 oligotrophs might be expected, especially in the grass interspaces where there was least 350 carbon. However, the oligotroph/copiotroph classification is a continuum and may relate 351 not only to soil total carbon, but more specifically to available carbon and carbon turnover 352 rate. From a microbial growth perspective the available carbon fraction is determined by 353 the physiological capabilities of the community and this also defines the amount of soil 354 carbon which can be regarded as recalcitrant (Schmidt et al. 2011). High carbon turnover

306

307

308

309

310

311

312

	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684	□ LE	TYPESET
\$	MS Code : BIOC-D-13-00927	CP	🔽 DISK
Biodivers Conserv			

rate has previously been calculated for cryptogamic crusts (Elbert et al. 2012). This could
help copiotrophs to maintain high catabolic rates even when soil carbon is low, so long as
there is primary production or other nutrient input available.

358 Bacterial community diversity

359 Previous studies have found that dryland BSC and soil bacterial richness and diversity do 360 not vary with respect to depth or presence of plants, although the community composition 361 does vary (Garcia-Pichel et al. 2003; Saul-Tcherkas and Steinberger 2011; Steven et al. 362 2013). Furthermore Nagy et al. (2005) found no difference in community composition 363 between plant canopies and interspaces, interpreting this to indicate probable independence 364 of BSC communities for vascular plant resources. Conversely, our results show that 365 samples from different depths and near different plants have distinct microbial commu-366 nities (Fig. 4) which also differ in richness and diversity (Fig. 3). This is not necessarily a 367 contradiction, however, due to differences in methodology, edaphic factors and disturbance 368 regime in the previous and current studies. In our samples it seems most likely that the tree 369 and shrub bacterial communities are at least partly, if not significantly, dependent on 370 vascular plant and animal-derived resources because they lack phototrophs. Animal dis-371 turbance could be a key factor in differentiating soil microbial communities because severe 372 animal disturbance would lead to burial of phototrophs, selectively disadvantaging these 373 organisms but not significantly disadvantaging heterotrophs which can function at any 374 depth. In the case of our grass interspaces, animal disturbance is less concentrated and 375 plant and animal derived inputs are expected to be less, suggesting that the phototrophic 376 component of BSCs will be more resilient compared to those near trees and shrubs.

377 Niche partitioning

378 Our results clearly demonstrate niche partitioning of the microbial community between 379 BSCs and the subsurface soil, and between soils under different vegetation types. This is 380 evident from the phylum level breakdown (Fig. 4), the most abundant OTUs overall (Fig. 6), the OTUs within each phylum (Fig. 7, Online Resource 4), and the overall 381 382 community structure (Fig. 5). Although in principle different microbial communities can 383 be functionally identical, we expect that the differences observed between sites and depths 384 are at least partly driven by different environmental conditions requiring a different 385 functional response.

From our field observations and soil chemistry results (Fig. 2), we would expect the grass area BSCs to be functionally similar on the macroscopic scale because the conditions are similar. One difference is that the BSC patches in perennial grass interspaces are likely to be older than the patches in annual grass interspaces, because the interspaces have become stabilised near perennial grasses.

391 A remarkable result was the detection of a clear difference between the soil bacterial 392 community in the surface and below the surface under trees. At the time of sampling, it 393 appeared that the homogenisation of the soil resulting from animal disturbances, had 394 removed all possibility of retaining any depth dependent structure in the biological com-395 munity. In fact it can be seen clearly in Fig. 5 and other figures that the surface (0-1 cm 396 depth) and subsurface (1-2 cm depth) communities from under the trees were quite dif-397 ferent, and also that they cluster closely with the same respective communities under the 398 shrub which were not subject to disturbance by cattle. The separation of BSC and sub-399 surface soil communities for both tree/shrub and grass locations on axis 2 of the ordination

	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684	🗆 LE	TYPESET
\sim	MS Code : BIOC-D-13-00927	CP	🖌 DISK
			Biodivers Conse

in Fig. 5 suggests that there are community structure patterns characterising surface and
sub-surface microbial communities, which apply regardless of the presence of BSC or
nearby vegetation.
The clear identification of niche partitioning leads to questions about the functional

The clear identification of niche partitioning leads to questions about the functional significance of different communities, and the ecosystem services delivered by BSCs compared to the subsurface soil. The experimental design of this study means that we cannot thoroughly address these questions but it does highlight the need for future targeted studies to do so. We can use the taxonomic identification of sequences to infer possible functions based on existing knowledge about microbial function in the environment, but proof of function needs to be addressed separately.

410 Although we have taken microbial community niche partitioning as an indicator of functional differences between soil depths and vegetation zones, this does not imply that 411 412 BSC community structure is shaped only by deterministic factors. There is a growing 413 recognition of the applicability of neutral models in determining microbial community 414 structure. These models confer key roles to the stochastic processes of birth, death and 415 immigration (Sloan et al. 2006) whilst ignoring deterministic factors such as a species 416 adaptation to drought. Caruso et al. (2011) have suggested that both stochastic and 417 deterministic processes interact in the assembly of dryland microbial communities and that 418 the role of niche partitioning at fine scales in dryland ecosystems has been previously 419 underestimated. They suggest it is incorrect to assume that extreme conditions are the main 420 determinant of species distribution, because this would lead to the conclusion that com-421 munities in extreme environments should converge towards stable low diversity commu-422 nities. The most extreme micro-environments in our study were the BSCs in grassed areas, 423 but they should not necessarily be regarded extreme for adapted microbes. These areas did 424 have the lowest diversity in our study, although the community structure varied greatly 425 between individual samples. Whilst this variability in BSC community structure may be 426 explained in the context of classical ecology as resulting from unobserved environmental 427 heterogeneity and community succession, it is also quite plausible that stochastic processes 428 may be playing a major role in community assembly. As BSCs have been recognised as 429 ecosystem engineers (Bowker 2007), the prospect of stochastic community assembly is 430 very important because it may directly drive diversity of BSC function and soil properties, 431 thus diversifying the landscape itself.

432 Possible ecological significance of the most abundant OTUs

433 Three of the top 9 OTUs (Fig. 6; Table 2) were significantly associated with BSCs 434 (0-1 cm depth) based on frequency of sequence detection. These were a cyanobacterium of 435 the genus Phormidium (OTU 1912), a Methylobacterium species (OTU 1064) belonging to 436 the phylum Proteobacteria, and a Balneimonas species (OTU 1150) of the phylum Pro-437 teobacteria. In addition, sequences of proteobacterial OTU 2850 and actinobacterial OTU 438 3323 were found to be significantly more abundant in subsurface soil compared to BSC. It 439 can be seen in Fig. 6 that in several cases differential abundance between BSC and sub-440 surface soil appears to be affected by the vegetation classification, particularly grasses vs. 441 tree or shrub (e.g. OTU 1912). This is supported by the ADONIS results (Online Resource 442 3) which suggest that there is a significant interaction between depth and vegetation zone 443 driving microbial community structure.

Phormidium species have previously been described as typical *Cyanobacteria* of early
successional stage BSCs by Büdel et al. (2009), consistent with our present observation in
weakly developed type 1 and type 2 BSCs (Thomas and Dougill 2006; 2007). They are

404

405

406

407

408

· •	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684		TYPESET
\sim	MS Code : BIOC-D-13-00927	CP	🔽 DISK
Biodivers Conserv			

Author Proof

447

448

449

450

451

452

453

454

455

456

457

458

459

widespread filamentous *Cyanobacteria* which are found commonly in hot and cold arid soils but are also found in aquatic habitats including ultra-oligotrophic Antarctic seasonal lakes (Keskitalo et al. 2013), demonstrating a great plasticity for the contrasting environmental conditions of relevance to the Kalahari Sand soil surface.

Chen et al. (2012) found *P. tenue* to be potentially useful for the stabilisation of sand dunes by inoculation due to the desiccation tolerance afforded by its extracellular polysaccharide (EPS). The EPS of *P. tenue* has also been shown to promote the germination of seeds and the fitness of seedlings (Xu et al. 2013), which may have been related to numerous mechanisms including water retention, provision of nutrients, and protection from oxidative stress. Furthermore, Boopathi et al. (2013) showed that a *Phormidium* species associated with mangroves produces indole-3-acetic acid (IAA) which is an important plant hormone associated with diverse responses including enhanced germination and root growth.

460 BSCs have been associated with both enhancement and inhibition of plant growth 461 through a variety of mechanisms and with some controversy (e.g. Beyschlag et al. 2008; 462 Prasse and Bornkamm 2000). We tentatively suggest that microbial secretion of plant 463 hormones may be an as-yet unrecognised process of relevance to BSC-plant interactions, 464 and that *Phormidium* species could influence plant cover in the Kalahari if they are 465 secreting IAA similarly to the species studied by Boopathi et al. (2013). The recruitment of 466 plants is potentially at odds with the maintenance of a photosynthetic BSC due to com-467 petition, so one might expect the exclusion of plants rather than their promotion to be a 468 more successful ecological strategy for BSC communities. Another effect of IAA on plant 469 roots is to reduce cell wall integrity, causing the release of nutrients such as sugars which 470 can promote microbial populations near the root. Thus it is possible that in addition to 471 being a significant primary producer in BSCs, *Phormidium* species may also be able to 472 derive carbon from plants, reducing reliance on photosynthesis and potentially promoting 473 further plant-microbe interactions. Again, this supposition is dependent upon the pro-474 duction of plant hormones by the *Phormidium* species in the BSC which has not been 475 tested. The dominance of sequences assigned to this Phormidium OTU in our samples 476 strongly suggests a numerical dominance of the BSC bacterial community, and leads to the 477 expectation of a large amount of *Phormidium*-derived EPS in the BSCs and soils examined 478 in this study. The phyla Bacteroidetes and Cyanobacteria were both more abundant in BSC 479 compared to subsurface soil, however unlike the Bacteroidetes which were represented by 480 many OTUs, the Cyanobacteria were dominated by the single Phormidium OTU 1912. 481 This may be a reflection of the different life strategies of the phyla. The Bacteroidetes 482 likely being specialist degraders in BSCs as discussed earlier, are strongly dependent on 483 the soil makeup as determined by the life history and current activity in the soil they 484 inhabit, so a large genetic (functional) diversity is called for. On the other hand Cyano-485 *bacteria* being principally photosynthetic are probably less sensitive to soil life history, but 486 their distribution is likely to be controlled more by abiotic factors such as weather and 487 hydrology. Since the Kalahari Sand substratum is so uniform over very large areas this 488 suggests a possibility for one or a few *Phormidium* species to be strongly influencing the 489 Kalahari soil surface on a very large scale far exceeding our study site.

490 The *Balneimonas* (OTU 1150) abundance was positively correlated to soil carbon, 491 suggesting it could be functioning as a copiotroph. Six of the other abundant OTUs which 492 were not significantly associated with BSCs were negatively correlated with soil carbon, so 493 although potential oligotrophs are not significantly increased in crusts, they do appear to 494 form a large fraction of the BSC and subsurface soil communities.

~	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684		TYPESET
\$	MS Code : BIOC-D-13-00927	CP	V DISK
			Biodivers Cons

Balneomonas was recognised as a new genus in 2004 (Takeda et al. 2004), however it has since been proposed for reclassification as *Microvirga* by Weon et al. (2010). In any case the original description of the type species is of a thermophilic (40–45 °C optimum) cellulose producing species isolated from a bath fed by a hot spring in Japan and related to *Methylobacterium* which is the genus of our other OTU of interest here. Cellulose production at high temperature was identified as a remarkable property of the type species *B. flocculans*, and noted for causing flocculation of the cells via adhesive cellulose fibrils similar to those thought to be involved in nonspecific binding of other rhizobial bacteria to plant host cells. The production of EPS is recognised as an important factor in the development and survival of BSCs, and is normally attributed to cyanobacteria (Mager and Thomas 2011). This result suggests that BSC stabilisation by EPS may also be facilitated by heterotrophic bacteria, and a role for *Balneomonas* species as BSC pioneers in advance of cyanobacterial establishment seems quite plausible.

508 Our Methylobacterium OTU was significantly associated with BSCs but not with any 509 particular vegetation zone, suggesting a role in the soil surface which is not related to the 510 presence of a particular vegetation type. The type species *M. organophilum* (Patt et al. 511 1976) and others are claimed to be facultative methylotrophs-capable of growth on 512 methane or other C1 carbon sources, however although facultative methylotrophy has now 513 been proved for some organisms there has been some controversy over this (Theisen and 514 Murrell 2005). It is thought that 50-90 % of methane released in soils is oxidised by 515 methylotrophs before reaching the atmosphere (Nazaries et al. 2013), therefore it is an 516 important ecosystem function with relevance to climate change. None of the known 21 517 obligate methane oxidising genera (Nazaries et al. 2013) were detected in our samples.

518 The most widely recognised role for Methylobacterium species is as epiphytes which 519 consume methanol emitted from stomata and secrete plant growth promoting hormones 520 (Lidstrom and Chistoserdova 2002). Several Methylobacterium isolates have been con-521 firmed as aerobic anoxygenic phototrophs in BSCs (Csotonyi et al. 2010), meaning that 522 their photosynthetic pathway does not produce oxygen, and that they are obligate aerobes 523 (in contrast to most other anoxygenic phototrophs). This is an important finding because it 524 increases the potential light harvesting efficiency of BSCs. Csotonyi et al. (2010) found 525 that anoxygenic phototrophs represented up to 5.9 % of the cultivable BSC bacterial 526 community, and we found Methylobacterium OTU 1064 alone represented 3 % of 527 sequences in annual grass zone BSCs (Online Resource 4).

528 Methodological limitations

529 The determination of bacterial community structure through analysis of ribosomal RNA 530 genes as carried out in this study has some limitations which should be kept in mind when 531 interpreting results and planning future studies. Methodological aspects including DNA 532 extraction efficiency differences, PCR bias, and primer specificity can affect observed 533 sequence frequencies. Furthermore, natural rRNA gene copy number variation will affect 534 results. For instance, it is known that oligotrophs in general carry fewer rRNA gene copies 535 than copiotrophs (Klappenbach et al. 2000), which could lead to under-representation of 536 oligotrophs in our study.

537 Implications for land management

The management of BSCs through protection, restoration, or engineering has the potentialto deliver environmental benefits from local to global scales, and is relevant to numerous

495

496

497

498

499

500

501

502

503

504

505

506

	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684	□ LE	TYPESET
\$	MS Code : BIOC-D-13-00927	CP	🔽 DISK
Biodivers Conserv			

540

541

542

543

544

545

546

547

548

549

550

551

552

land management challenges. Landscape changes associated with moderate grazing in the Kalahari include a shift from perennial grass to annual grass species and the promotion of thorny shrubs (Skarpe 1990; Ward 2009). We characterised the microbial communities from soils near these plants and found that in all cases a surface specific community can be detected, even beneath the *Acacia* canopy where there was no consolidated BSC. This indicates a strong tendency of the soil community to become vertically stratified and suggests that once grazing pressure is removed a rapid recovery of the BSC is likely. Soil communities beneath tree/shrub canopies were distinct from those in grass interspaces as has been shown previously (Saul-Tcherkas and Steinberger 2011), showing that typical grazing induced vegetation change can be associated with a change in soil microbiota, notably a loss of *Cyanobacteria*. The extreme reduction in cyanobacterial inoculum may retard BSC establishment in heavily shrub-encroached areas, for instance after fire or attempts to reclaim the land by removal of shrubs.

553 The deliberate rehabilitation of BSCs to restore ecosystem function has been discussed 554 by Bowker (2007) and recently confirmed by cyanobacterial inoculation of shifting sand 555 dunes to establish BSCs which facilitated vascular plant succession from early nitrogen 556 fixing legumes to latter successional grasses (Lan et al. 2014). A problem with the inoc-557 ulation approach is that it usually relies upon a sacrifice zone, and may not work if there are 558 nutrient or stability limitations. These problems can potentially be overcome by the 559 industrial preparation of a designed mixed inoculum. Candidates for a BSC inoculum in the 560 Kalahari rangelands could be identified for testing based upon our data, which is ideal for 561 this purpose because early stage BSCs are likely to still contain in large number the pioneer 562 species which helped them to become established. The Phormidium genus of Cyanobac-563 teria (P. tenue specifically) has been shown by Chen et al. (2012) to possess excellent 564 qualities for the stabilisation of sand, and its abundance in our study suggests that delib-565 erate establishment on degraded land in the Kalahari could be feasible. A suitable inoc-566 ulum to help the establishment of *Cyanobacteria* in general might include oligotrophic 567 organisms such as Balneomonas species which are able to utilise recalcitrant carbon and 568 quickly stabilise the soil by release of EPS, including beneath the photic zone. The idea 569 that the early stages of BSC recovery or development could be helped by oligotrophic non-570 photosynthetic bacteria has been confirmed by Wu et al. (2010) in laboratory and field 571 experiments, however the identity of organisms involved was not known.

572 Conclusions

573 We found that the bacterial OTU (approximate species level) diversity was greater in 574 subsurface soil (1-2 cm depth) compared to the BSC at the surface (0-1 cm depth), and 575 community composition exhibited clear spatial patterns. Crusted grass interspaces were 576 dominated by a single cyanobacterial OTU from the genus Phormidium, but Cyanobac-577 teria were very rare in tree and shrub areas. The BSC community structure was not defined 578 by the presence of *Cyanobacteria* alone nor by other phototrophs as is often presumed. In 579 all areas a characteristic but variable BSC bacterial community was present, and this was 580 defined in part by non-cyanobacterial OTUs which were associated with the soil surface, 581 and especially the Bacteroidetes phylum. The functions of these BSC specific bacteria are 582 unknown, but we speculated possible roles for the most abundant ones in soil stabilisation, 583 carbohydrate catabolism, photosynthesis, and plant interactions based on similar species 584 reported in the literature. Our data suggest that shrub encroachment in Kalahari rangelands 585 can almost eliminate Cyanobacteria from soil surfaces in some circumstances, depriving

	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No.: 684	🗆 LE	□ TYPESET
\sim	MS Code : BIOC-D-13-00927	CP	V DISK
			Biodivers (

586 soils of the ecosystem services delivered by well-developed phototrophic crusts and lim-587 iting the potential for natural BSC regeneration.

Acknowledgments Financial support was provided by the Leverhulme Trust (F/00 426/H) and Manchester Metropolitan University. Research in Botswana was conducted with the Republic of Botswana Research Permit No. EWT8/36/4 VIII(4). We would like to thank Jill Thomas of Berrybush farm for her considerable support and access to her land. We also thank Peter Harding for assistance with computer resources and configuration. The constructive feedback of an anonymous reviewer and Matthew Bowker is greatly appreciated.

596 References

- Aranibar JN, Anderson IC, Ringrose S, Macko SA (2003) Importance of nitrogen fixation in soil crusts of southern African arid ecosystems: acetylene reduction and stable isotope studies. J Arid Environ 54:345–358
 - Bailey VL, Fansler SJ, Stegen JC, McCue LA (2013) Linking microbial community structure to β-glucosidic function in soil aggregates. ISME J 7(10):2044–2053. doi:10.1038/ismej.2013.87
 - Belnap J (2006) The potential roles of biological soil crusts in dryland hydrologic cycles. Hydrol Process 20(15):3159–3178
 - Benjamini Y, Hochberg Y (1995) Controlling the false discovery rate: a practical and powerful approach to multiple testing. J R Stat Soc B 57:289–300
 - Beyschlag W, Wittland M, Jentsch A, Steinlein T (2008) Soil crusts and disturbance benefit plant germination, establishment and growth on nutrient deficient sand. Basic Appl Ecol 9(3):243–252
 - Boopathi T, Balamurugan V, Gopinath S, Sundararaman M (2013) Characterization of IAA production by the mangrove *cyanobacterium phormidium sp.* MI405019 and its influence on tobacco seed germination and organogenesis. J Plant Growth Regul 32(4):758–766
 - Bowker MA (2007) Biological soil crust rehabilitation in theory and practice: an underexploited opportunity. Restor Ecol 15(1):13–23
 - Bowker MA, Maestre FT, Escolar C (2010) Biological crusts as a model system for examining the biodiversity–ecosystem function relationship in soils. Soil Biol Biochem 42(3):405–417
 - Brussard L (2012) Ecosystem services provided by soil biota. In: Diana H Wall (ed) Soil ecology and
 ecosystem services. Oxford University Press, Oxford, pp 45–58
 - Büdel B, Darienko T, Deutschewitz K, Dojani S, Friedl T, Mohr KI, Salisch M, Reisser W, Weber B (2009) Southern African biological soil crusts are ubiquitous and highly diverse in drylands, being restricted by rainfall frequency. Microb Ecol 57(2):229–247. doi:10.1007/s00248-008-9449-9
 - Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, Fierer N, Pena AG, Goodrich JK, Gordon JI, Huttley GA, Kelley ST, Knights D, Koenig JE, Ley RE, Lozupone CA, McDonald D, Muegge BD, Pirrung M, Reeder J, Sevinsky JR, Turnbaugh PJ, Walters WA, Widmann J, Yatsunenko T, Zaneveld J, Knight R (2010) QIIME allows analysis of high-throughput community sequencing data. Nat Methods 7:335–336
- 625 Caruso T, Chan Y, Lacap DC, Lau MCY, McKay CP, Pointing SB (2011) Stochastic and deterministic
 626 processes interact in the assembly of desert microbial communities on a global scale. ISME J
 627 5:1406–1413
- Castillo-Monroy AP, Bowker MA, Maestre FT, Rodríguez-Echeverría S, Martínez I, Barraza-Zepeda CE, Escolar C (2011) Relationships between biological soil crusts, bacterial diversity and abundance, and ecosystem functioning: insights from a semi-arid Mediterranean environment. J Veg Sci 22:165–174
- Chen L, Yang Y, Deng S, Xu Y, Wang G, Liu Y (2012) The response of carbohydrate metabolism to the fluctuation of relative humidity (RH) in the desert soil *cyanobacterium Phormidium tenue*. Euro J Soil Biol 48:11–16
- 634 Csotonyi JT, Swiderski J, Stackebrandt E, Yurkov V (2010) A new environment for aerobic anoxygenic
 635 phototrophic bacteria: biological soil crusts. Environ Microbiol Rep 2(5):651–656
- beSantis TZ, Hugenholtz P, Larsen N, Rojas M, Brodie EL, Keller K, Huber T, Dalevi D, Hu P, Andersen GL (2006) Greengenes, a chimera-checked 16S rRNA gene database and workbench compatible with ARB. Appl Environ Microbiol 72:5069–5072

588

589

590

591

592

593

	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684	□ LE	TYPESET
\sim	MS Code : BIOC-D-13-00927	CP	🖌 DISK

- Dojani S, Kauff F, Weber B, Büdel B (2013) Genotypic and phenotypic diversity of cyanobacteria in biological soil crusts of the succulent karoo and nama karoo of southern Africa. Microb Ecol. doi:10. 1007/s00248-013-0301-5
- Dowd S, Callaway T, Wolcott R, Sun Y, McKeehan T, Hagevoort R, Edrington T (2008) Evaluation of the bacterial diversity in the feces of cattle using 16S rDNA bacterial tag-encoded FLX amplicon pyrosequencing (bTEFAP). BMC Microbiol 8(1):125
- Edgar RC (2010) Search and clustering orders of magnitude faster than BLAST. Bioinformatics 26(19):2460-2461. doi:10.1093/bioinformatics/btq461
- Elbert W, Weber B, Burrows S, Steinkamp J, Büdel B, Andreae MO, Pöschl U (2012) Contribution of cryptogamic covers to the global cycles of carbon and nitrogen. Nat Geosci 5(7):459-462
- Eldridge DJ, Greene RSB (1994) Microbiotic soil crusts: a review of their roles in soil and ecological processes in the rangelands of Australia. Aust J Soil Res 32:389-415
- FAO (1990) Guidelines for soil description. 3rd edn. (revised). Soil resources, management and conservation service, land and water development division, FAO, Rome
- Fierer N, Bradford MA, Jackson RB (2007) Toward an ecological classification of soil bacteria. Ecology 88(6):1354-1364
- Fierer N, Lauber CL, Ramirez KS, Zaneveld J, Bradford MA, Knight R (2012a) Comparative metagenomic, phylogenetic and physiological analyses of soil microbial communities across nitrogen gradients. ISME J 6:1007-1017
- Fierer N, Leff JW, Adams BJ, Nielsen UN, Bates ST, Lauber CL, Owens S, Gilbert JA, Wall DA, Caporaso JG (2012b) Cross-biome metagenomic analyses of soil microbial communities and their functional attributes. Proc Natl Acad Sci USA 109:21390-21395
- Garcia-Pichel F, Johnson SL, Youngkin D, Belnap J (2003) Small-scale vertical distribution of bacterial biomass and diversity in biological soil crusts from arid lands in the Colorado plateau. Microb Ecol 46(3):312-321
- Gundlapally SR, Garcia-Pichel F (2006) The community and phylogenetic diversity of biological soil crusts in the Colorado plateau studied by molecular fingerprinting and intensive cultivation. Microb Ecol 52(2):345-357
- Haas BJ, Gevers D, Earl AM, Feldgarden M, Ward DV, Giannoukos G, Ciulla D, Tabbaa D, Highlander SK, Sodergren E, Methé B, DeSantis TZ; Human Microbiome Consortium, Petrosino JF, Knight R, Birren BW (2011) Chimeric 16S rRNA sequence formation and detection in Sanger and 454-pyrosequenced PCR amplicons. Genome Res 21(3):494-504. doi:10.1101/gr.112730.110
- Keskitalo J, Leppäranta M, Arvola L (2013) First records of primary producers of epiglacial and supraglacial lakes in western dronning maud land, Antarctica. Polar Biol 36(10):1441-1450
- Klappenbach JA, Dunbar JM, Schmidt TM (2000) rRNA operon copy number reflects ecological strategies of bacteria. Appl Environ Microbiol 66(4):1328-1333
- Lal R (2004) Carbon sequestration in dryland ecosystems. Environ Manage 33(4):528-544
- Lan S, Zhang Q, Wu L, Liu Y, Zhang D, Hu C (2014) Artificially accelerating the reversal of desertification: 677 cyanobacterial inoculation facilitates the succession of vegetation communities. Environ Sci Technol. 678 doi:10.1021/es403785j
- 679 Lidstrom ME, Chistoseerdova L (2002) Plants in the pink: cytokinin production by Methylobacterium. 680 J Bacteriol 184:1818
- 681 MacArthur RH, Wilson E (1967) The theory of island biogeography. Princeton University Press, Princeton
- 682 Mager DM, Thomas AD (2011) Extracellular polysaccharides from cyanobacterial soil crusts: a review of 683 their role in dryland soil processes. J Arid Environ 75(2):91-97
- 684 McMurdie PJ, Holmes S (2013) phyloseq: an r package for reproducible interactive analysis and graphics of 685 microbiome census data. PLoS ONE 8(4):e61217
- 686 Menon M, Yuan Q, Jia X, Dougill AJ, Hoon SR, Thomas AD, Williams RA (2011) Assessment of physical 687 and hydrological properties of biological soil crusts using X-ray microtomography and modeling. 688 J Hydrol 397(1):47-54
- 689 Meyer F, Paarmann D, D'Souza M, Olson R, Glass EM, Kubal M et al (2008) The metagenomics RAST 690 server-a public resource for the automatic phylogenetic and functional analysis of metagenomes. BMC 691 Bioinform 9(1):386
- 692 Muyzer G, De Waal EC, Uitterlinden AG (1993) Profiling of complex microbial populations by denaturing 693 gradient gel electrophoresis analysis of polymerase chain reaction-amplified genes coding for 16S 694 rRNA. Appl Environ Microbiol 59:695-700
- 695 Muyzer G, Brinkhoff T, Nübel U, Santegoeds C, Schäfer H, Waver C. (1998). Denaturing gradient gel 696 electrophoresis (DGGE) in microbial ecology. In: Akkermans ADL, van Elsas JD, de Bruijn FJ (eds). 697 Molecular microbial ecology manual. Kluwer Academic Publishers, Dordrecht

639

640

641

642

643

644

645

646

647

649

651

652

653

Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
Article No. : 684	🗆 LE	□ TYPESET
MS Code : BIOC-D-13-00927	CP	🖌 DISK
		Biodivers Co

- 698 Nagy ML, Pérez A, Garcia-Pichel F (2005) The prokaryotic diversity of biological soil crusts in the sonoran 699 desert (organ pipe cactus national monument, AZ). FEMS Microbiol Ecol 54(2):233-245 700
 - Nazaries L, Murrell JC, Millard P, Baggs L, Singh BK (2013) Methane, microbes and models: fundamental understanding of the soil methane cycle for future predictions. Environ Microbiol 15(9):2395-2417. doi:10.1111/1462-2920.12149
- 70 Age Oksanen J, Blanchet FG, Kindt R, Legendre P, Minchin PR, O'Hara RB, Simpson GL, Solymos P, Stevens 704HMH, Wagner H (2013) Vegan: community ecology package. R package version 2.0–7. http://CRAN. 705 R-project.org/package=vegan 706
 - Patt TE, Cole GC, Hanson RS (1976) Methylobacterium, a new genus of facultatively Methylotrophic bacteria. Int J Syst Bacteriol 26:226-229
 - Pereira MC, Dias ACF, vanElsas JD, Salles JF (2012) Spatial and temporal variation of archaeal, bacterial and fungal communities in agricultural soils. PLoS ONE 7:e51554. doi:10.1371/journal.pone.0051554
 - Phosri C, Polme S, Taylor AFS, Koljalg U, Suwannasai N, Tedersoo L (2012) Diversity and community composition of ectomycorrhizal fungi in a dry deciduous dipterocarp forest in Thailand. Biodivers Conserv 21:2287-2298
 - Prasse R, Bornkamm R (2000) Effect of microbiotic soil surface crusts on emergence of vascular plants. Plant Ecol 150(1-2):65-75
- 71 A03 R Core Team (2012) R: a language and environment for statistical computing. In: R foundation for statistical computing, Vienna, ISBN 3-900051-07-0, URL http://www.R-project.org/
 - Reynolds JF, Smith DMS, Lambin EF, Turner BL, Mortimore M, Batterbury SP et al (2007) Global desertification: building a science for dryland development. Science 316(5826):847-851
 - Rousk J, Bååth E, Brookes PC, Lauber CL, Lozupone C, Caporaso JG, Knight R, Fierer N (2010) Soil bacterial and fungal communities across a pH gradient in an arable soil. ISME J 4:1340-1351
 - Saul-Tcherkas V, Steinberger Y (2011) Soil microbial diversity in the vicinity of a negev desert shrub-Reaumuria negevensis. Microb Ecol 61(1):64-81. doi:10.1007/s00248-010-9763-x
 - Schmidt MW, Torn MS, Abiven S, Dittmar T, Guggenberger G, Janssens IA, Kleber M, Kögel-Knabner I, Lehmann J, Manning DA, Nannipieri P, Rasse DP, Weiner S, Trumbore SE (2011) Persistence of soil organic matter as an ecosystem property. Nature. doi:10.1038/nature10386
 - Skarpe C (1990) Structure of the woody vegetation in disturbed and undisturbed arid savanna Botswana. Vegetatio 87(1):11-18
 - Sloan WT, Lunn M, Woodcock S, Head IM, Nee S, Curtis TP (2006) Quantifying the roles of immigration and chance in shaping prokaryote community structure. Environ Microbiol 8(4):732–740
 - Smit E, Leeflang P, Gommans S, van den Broek J, van Mil S, Wernars K (2001) Diversity and seasonal fluctuations of the dominant members of the bacterial soil community in a wheat field as determined by cultivation and molecular methods. Appl Environ Microbiol 67(5):2284–2291
 - Stackebrandt E, Goebel BM (1994) A place for DNA-DNA reassociation and 16S ribosomal-RNA sequence-analysis in the present species definition in bacteriology. Int J Syst Evol Microbiol $44 \cdot 846 - 849$
 - Steven B, Gallegos-Graves LV, Belnap J, Kuske CR (2013) Dryland soil microbial communities display spatial biogeographic patterns associated with soil depth and soil parent material. FEMS Microbiol Ecol 86(1):101-113. doi:10.1111/1574-6941.12143
 - Stringer L (2008) Can the UN Convention to combat desertification guide sustainable use of the world's soils? Front Ecol Environ 6(3):138-144
 - Stringer LC, Dougill AJ, Thomas AD, Spracklen DV, Chesterman S, Speranza CI, Rueff H et al (2012) Challenges and opportunities in linking carbon sequestration, livelihoods and ecosystem service provision in drylands. Environ Sci Policy 19:121–135
 - Takeda M, Suzuki I, Koizumi JI (2004) Balneomonas flocculans gen. nov., sp. nov., a new celluloseproducing member of the α-2 subclass of Proteobacteria. Syst Appl Microbiol 27(2):139-145
 - Theisen AR, Murrell JC (2005) Facultative methanotrophs revisited. J Bacteriol 187(13):4303–4305
 - Thomas AD (2012) Impact of grazing intensity on seasonal variations of soil organic carbon and soil CO2 efflux in two semi-arid grasslands in southern Botswana. Philos Trans Royal Soc B 367:3076-3086
 - Thomas AD, Dougill AJ (2006) Distribution and characteristics of cyanobacterial soil crusts in the Molopo Basin, South Africa. J Arid Environ 64:270-283
 - Thomas AD, Dougill AJ (2007) Spatial and temporal distribution of cyanobacterial soil crusts in the Kalahari: implications for soil surface properties. Geomorphology 85:17-29
 - Thomas F, Hehemann J, Rebuffet E, Czjzek M, Michel G (2011) Environmental and gut Bacteroidetes: the food connection. Frontiers Microbiol 2. doi:10.3049/fmicb.2011.00093
 - Thomas AD, Hoon SR, Dougill AJ, Mairs H (2012) Soil organic carbon in deserts: examples from the kalahari. In: Mol L, Sternberg T (eds) changing deserts: integrating environments, people and challenges. White Horse Press, Cambridge

701

702

707

708

709

710

711

712

713

714

716

717

718

719

720

721

 $\dot{7}\bar{2}\dot{2}$

723

724 725

726

727 728

729

🕗 Springer

(I)	Journal : Small 10531	Dispatch : 1-4-2014	Pages : 25
	Article No. : 684	🗆 LE	TYPESET
	MS Code : BIOC-D-13-00927	CP	🖌 DISK

- Wang Q, Garrity GM, Tiedje JM, Cole JR (2007) Naive Bayesian classifier for rapid assignment of rRNA sequences into the new bacterial taxonomy. Appl Environ Microbiol 73:5261-5267
- Ward D (2009) The Biology of deserts. Oxford University Press, Oxford
- Weon HY, Kwon SW, Son JA, Jo EH, Kim SJ, Kim YS, Kim BY, Ka JO (2010) Description of Microvirga aerophila sp. nov. and Microvirga aerilata sp. nov., isolated from air, reclassification of Balneimonas flocculans takeda, et al. 2004 as Microvirga flocculans comb nov. and emended description of the genus Microvirga. Int J Syst Evol Microbiol 60(11):2596-2600
- Wu N, Zhang YM, Pan HX, Zhang J (2010) The role of nonphotosynthetic microbes in the recovery of biological soil crusts in the gurbantunggut desert, northwestern China. Arid Land Res Manage 24(1):42-56
- Xu Y, Rossi F, Colica G, Deng S, De Philippis R, Chen L (2013) Use of cyanobacterial polysaccharides to promote shrub performances in desert soils: a potential approach for the restoration of desertified areas. Biol Fertil Soils 49(2):143-152

758 759 Author Proof

760

761

762

763

764

765

766

767

768

769