

Physicochemical and biological dynamics in a coastal Antarctic lake as it transitions from frozen to open water

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Abstract: Pony Lake, at Cape Royds, Antarctica, is a shallow, eutrophic, coastal lake that freezes solid in the winter. Changes in Pony Lake's physicochemical parameters and microbial community were studied during the transition from ice to open water. Due to rising water temperatures, the progressive melt of the ice column and the gradual mixing of basal brines into the remaining water column, Pony Lake evolved physically and chemically over the course of the summer, thereby affecting the microbial community composition. Temperature, pH, conductivity, nutrients and major ion concentrations reached their maximum in January. Pony Lake was colonized by bacteria, viruses, phytoflagellates, ciliates, and a small number of rotifers. Primary and bacterial production were highest in mid-December (2.66 mg C l⁻¹ d⁻¹ and 30.5 µg C l⁻¹ d⁻¹, respectively). A 16S rRNA gene analysis of the bacterioplankton revealed 34 unique sequences dominated by members of the β - and γ -proteobacteria lineages. Cluster analyses on denaturing gradient gel electrophoresis (DGGE) banding patterns and community structure indicated a shift in the dominant members of the microbial community during the transition from winter ice, to early, and late summer lakewater. Our data demonstrate that temporal changes in physicochemical parameters during the summer months determine community dynamics and mediate changes in microbial species composition.

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Key words: Antarctica, microbes, seasonal transition

Introduction

Lakes and ponds are common features in the ice-free areas of the Antarctic continent. Often influenced by marine aerosols these limnetic systems possess a wide range of salinities and ionic compositions. Nutrients are typically supplied by local snow or glacial meltwaters, but highly enriched lakes and ponds can also be found in the vicinity of bird or seal colonies. The planktonic community in these lakes is dominated by microorganisms, including bacteria, phytoplankton and protozoa, with little or no metazoans typically present (Perriss & Laybourn-Parry 1997, Bell & Laybourn-Parry 1999, Laybourn-Parry *et al.* 2002). While most Antarctic lakes are perennially or seasonally ice-covered with liquid water present beneath the ice, shallow lakes and ponds freeze solid to the base during winter. The exclusion of salts and solutes from the expanding ice develops a saline, basal brine solution during freezing

(Healy *et al.* 2006, Hawes *et al.* 2011). During the summer, these shallow lakes and ponds can become partially or completely ice-free, thereby creating a unique and rapidly changing environment. The chemical stratification present within the ice can persist in the summer lakewater in the absence of strong winds to disrupt the salinity gradient. Thus, whether the pond remains chemically stratified or not after thawing will play an important role in structuring the microbial community. Organisms that persist throughout the course of the year in such an environment must be capable of surviving severe seasonal variations in chemical, redox and temperature conditions (Schmidt *et al.* 1991, Hawes *et al.* 2011, Webster-Brown *et al.* 2012). Most importantly, outlasting the hostile, frozen entrapment or the hypersaline basal liquids is essential for re-colonizing the lake when melt is initiated. This study describes the biological, physical, and chemical parameters of a shallow Antarctic lake during the annual transition from frozen solid to an open lake system.

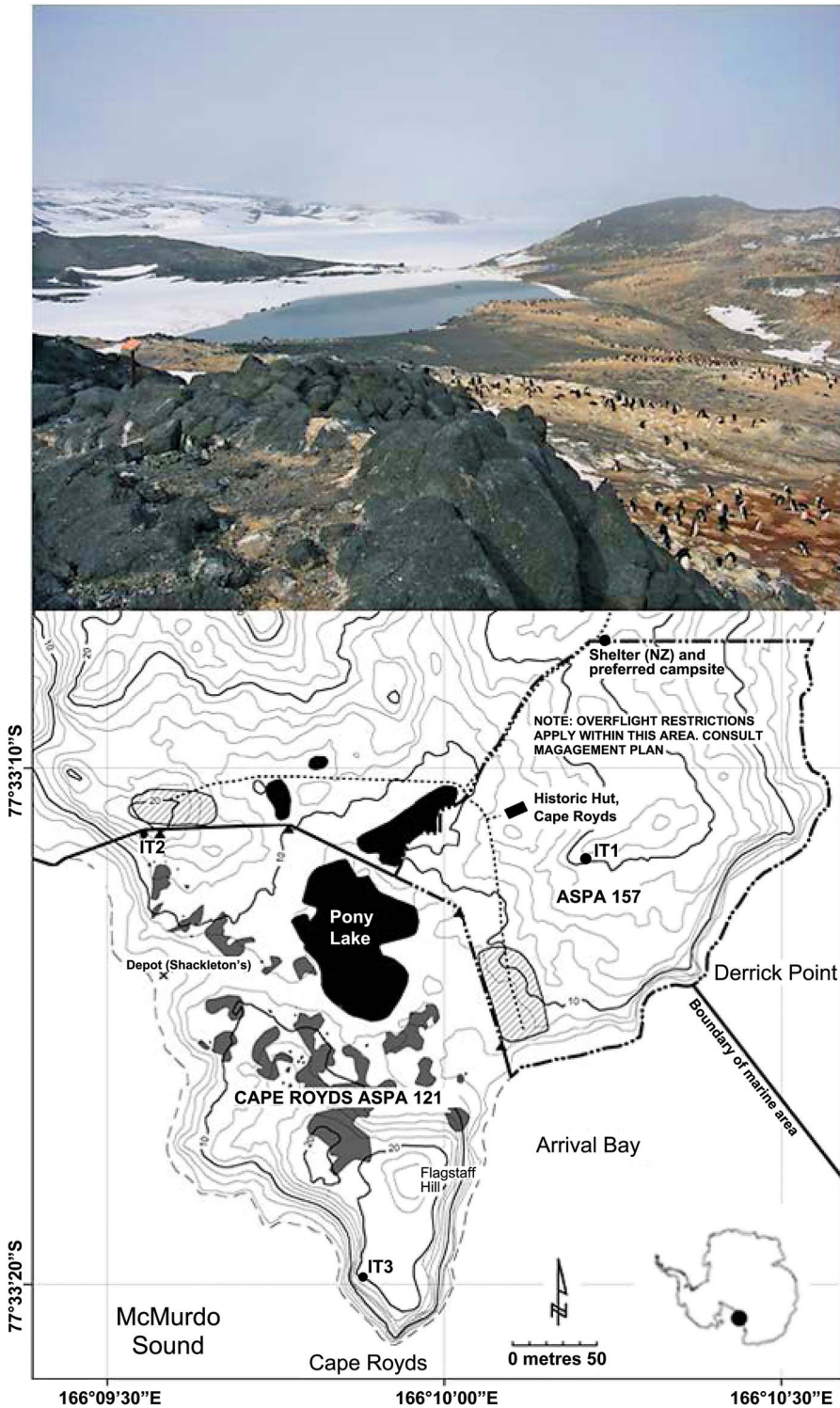


Fig. 1. Map of Cape Royds showing the location of Pony Lake (modified from www.scar.org, accessed October 2012). The inset image was taken in late December 2004 and shows Pony Lake partially ice-free.

Pony Lake is a coastal, eutrophic lake located on Cape Royds (77°33'S, 166°00'E), Ross Island, Antarctica (Fig. 1). The lake is *c.* 120 m long, 70 m wide, and 1–2 m deep. Except for midsummer, when warmer temperatures melt the ice-cover and strong winds cause thorough mixing of the water column, the lake is frozen solid. Ice melt typically

begins in mid-December, however, during years with heavy snowfall (e.g. 2005–06) the ice cover may persist throughout the summer season. The source of water to the lake is the accumulated snowpack, while water is lost by both sublimation of ice and evaporation. As a result of its proximity to the sea Pony Lake is brackish (5.5 ppt).

Previous studies have shown that Pony Lake may support very high dissolved organic carbon (DOC) concentrations (up to 100 mg C l⁻¹ in some seasons) (McKnight *et al.* 1994, Brown *et al.* 2004). There are no higher plants in the basin, but algal populations in the lake are abundant. *Chlamydomonas intermedia* Chodat has been reported as being the dominant chlorophyte algal species (McKnight *et al.* 1994), the same as found during Sir Ernest Shackleton's 1908 expedition (West & West 1911). Brown *et al.* (2004) reported that during the transition from ice-covered to ice-free conditions, the chlorophyte bloom in the lake was displaced by a cryptophyte bloom. Along the western shore of the lake lies an Adélie penguin rookery. However, runoff from the rookery into Pony Lake is believed to be minimal due to highly evaporative conditions and Brown *et al.* (2004) indicated that the organic matter in the lake is primarily derived from microbial sources. In a lacustrine system such as Pony Lake that exhibits low levels of proto- and metazooplankton, bacterioplankton will play a significant role in the flow of energy and nutrients.

Materials and methods

Sample collection

Water samples were collected during the summer season of 2004/05. Four time points were chosen to measure basic limnological parameters (biological, physical, and chemical) in Pony Lake during the transition from a frozen to ice-free lake. During our studies melting began along the edges of the lake in early December and continued into January. Strong winds allowed for frequent mixing in this shallow lake, which was reflected in a homogeneous conductivity profile of the water column. Thus, one representative depth was chosen after preliminary analyses, and instrument data and water samples were subsequently collected *c.* 30 cm beneath the water surface. Water samples (*c.* 5 l) were collected in acid washed, deionized water rinsed (x 6) Nalgene bottles and stored in coolers for transport back to Cray laboratory in McMurdo Station. Samples were processed within four hours of collection.

Physical parameters and chemical analyses

Conductivity, salinity, and temperature were measured with a portable multi-meter (Hydrolab minisonde). Dissolved oxygen (YSI) and pH (Beckman) were recorded for each sample. Samples were prepared for chemical analysis according to the protocols of the McMurdo Dry Valleys Long-term Ecological Research Group (Priscu & Wolf 2000). Samples for DOC and total nitrogen (TN) analyses were filtered under low vacuum (< 7 psi) in the dark through 25 mm pre-combusted GF/F filters, acidified with 6 N HCl to pH2 and analysed on a Shimadzu TOC-V and Shimadzu TNM-1 analyser, respectively. Filters from the

above analyses were wrapped in aluminum foil and kept frozen until extraction for chlorophyll *a* (chl *a*) analysis. Chlorophyll *a* was extracted in a 1:1 solution (90% acetone and dimethyl sulfoxide) for 12 hours under dark conditions at -20°C. Extracted chl *a* was analysed on a Turner 10-AU fluorometer. Samples for macronutrients were filtered through 25 mm pre-combusted GF/F filters and stored frozen until analysis (within a month of collection) on a Lachat autoanalyser. Samples for anion and cation determination were filtered through 0.4 µm 47 mm nucleopore filters. Deionized water was used as a filtration blank and samples were analysed on a Dionex DX-300 ion chromatography system. Dissolved inorganic carbon (DIC) was measured using infrared absorption following acidification and sparging of the sample with high purity N₂ gas. Peak areas were integrated and converted to mg l⁻¹ using a standard curve based upon a freshly prepared standard of NaHCO₃.

Productivity measurements

Bacterial productivity (BP) was measured via ³H-thymidine incorporation (20 nM final concentration) as described in Takacs & Priscu (1998). Five ³H-thymidine assays and triplicate formalin killed controls (5% final concentration, 30 min prior to ³H-thymidine addition) were incubated at 4°C for 20 hours. Samples were analysed using a liquid scintillation counter (Beckman LS 7200). Thymidine incorporation rates were converted to bacterial production rates using a conversion factor of 2.0 x 10¹⁸ cells mol⁻¹ TdR and a cell-to-carbon conversion factor of 11 fg C cell⁻¹ as outlined by Takacs & Priscu (1998).

Primary production (PPR) was measured via ¹⁴C-carbonate/bicarbonate incorporation (114.4 µCi ml⁻¹, pH ~ 9.5; ICN/MP Biomedicals) as described in Lizotte *et al.* (1996). Quadruple light assays and duplicate dark controls were incubated at 4°C in an illuminated chamber for 24 hours. Following incubation, samples were filtered through pre-combusted 25 mm GF/F filters in the dark. Filters were transferred into 20 ml scintillation vials, acidified with 500 µl 3 M HCL and dried before liquid scintillation counting (Beckman LS 7200).

Plankton analyses

Water samples for the determination of bacterial abundances were fixed with formalin (2% final concentration) and stained with a 25X solution of the fluorochrome SYBR[®] Gold (Invitrogen Inc) for 15 min following Lisle & Priscu (2004). Samples were filtered onto 25 mm 0.2 µm black polycarbonate filters with a 0.45 µm nitrocellulose backing filter under gentle vacuum. To reduce the possibility of contamination, filter towers were pre-combusted and all reagents used for the staining were passed through 0.2 µm sterile filters to remove extraneous particles and cells.

Bacterial cells were enumerated using a Zeiss Axioscop epifluorescence microscope with a final magnification of 1000x.

Samples for bacteriophage or virus like particle (VLP) enumeration were collected in sterile 125 ml screw cap flasks. All flasks were immediately flash frozen in liquid nitrogen and stored at -80°C . Prior to sample processing all flasks were removed from -80°C storage and allowed to thaw in the dark at room temperature overnight. Samples were pre-filtered through a $0.20\ \mu\text{m}$ pore size filter to remove bacteria. Filtrate from each sample was aseptically collected and filtered through a 25 mm diameter, $0.02\ \mu\text{m}$ pore size filter to retain the VLP. Virus like particles were stained with SYBR[®] Gold as described by Lisle & Priscu (2004) and counted using an Olympus BX51 epifluorescent microscope.

Samples (11) for phytoplankton (algae, diatoms, *Cyanobacteria*) and zooplankton analyses were fixed with Lugol's iodine (10 ml Lugol's) and concentrated by settling for one week in amber Nalgene bottles. After settling, the upper solution was gently siphoned off, leaving *c.* 60 ml of sample, which was then transferred to a clean 60 ml Nalgene bottle for transport to the UK. Subsamples were counted in a Sedgewick-Rafter counting chamber using phase microscopy at 320x magnification (Laybourn-Parry & Marshall 2003).

Environmental DNA extraction and DGGE

Water samples (70–100 ml) were filtered onto 47 mm Supor[®]-200 $0.2\ \mu\text{m}$ pore size, sterile membrane filters under low pressure ($< 7\ \text{psi}$). Filters were placed into 5 ml

cryovials filled with TES (100 mM Tris, 100 mM EDTA and 2% SDS) buffer, flash-frozen in liquid nitrogen, and stored at -80°C .

DNA was extracted from the Supor[®]-200 membrane filters using an Ultra Clean Soil DNA Kit (MoBio). Denaturing gradient gel electrophoresis (DGGE) was used as a molecular fingerprinting tool to characterize the microbial population structure and diversity. A portion of the 16S rRNA gene was amplified with primers 341F (5'-CCTACGGGAGGCAGCAG-3') and 534R (5'-AATA CCGCGGCTGCTGG-3') (Muyzer *et al.* 1996). A 40 base pair GC clamp was added to the 5' end of the 341F primer (CGCCCGCCGCGCGCGCGGGCGGGGCGGGGGCAC GGGGGG). The amplification protocol included a hot start (94°C for 4 min) and a touchdown programme that consisted of an initial annealing temperature of 65°C followed by a 1°C decrease for eight cycles down to 58°C , and 17 cycles with an annealing temperature of 58°C . A final elongation step occurred for 10 min at 72°C . Each $50\ \mu\text{l}$ reaction mixture contained $1.5\ \mu\text{l}$ of environmental DNA extract, MgCl_2 buffer (final concentration 1X), Taq Master (final concentration 1X), PCR nucleotide mix (final concentration $800\ \mu\text{M}$), and Taq DNA polymerase (final concentration $0.025\ \mu\text{l}^{-1}$) (all components from 5 Prime, Eppendorf), upstream and downstream primers (final concentration $0.5\ \mu\text{M}$), and nuclease free water (Promega). Polymerase Chain Reaction (PCR) amplifications were carried out in an automated thermal cycler (Mastercycler ep, Eppendorf). DGGE was performed with a BioRad D CodeTM system as

Table 1. General site characteristics, dissolved organic carbon (DOC), dissolved inorganic carbon (DIC), total inorganic nitrogen (TIN), nutrient and major ion concentrations for Pony Lake measured during the summer 2004/05.

	11 Dec 2004	21 Dec 2004	29 Dec 2004	14 Jan 2005
Temp. ($^{\circ}\text{C}$)	1.3	-	4.2	7.5
Salinity (ppt)	1.86	2.25	3.25	3.93
pH	8.07	8.66	9.71	9.77
Conductivity ($\mu\text{S cm}^{-1}$)	3429	4134	5895	7058
DO (mg l^{-1})	18.0	> 20	18.0	> 20
DOC (mg l^{-1})	11.75	12.35	28.27	28.62
DIC (mg l^{-1})	10.84	4.30	3.40	4.19
TIN (mg l^{-1})	19.02	14.3	17.23	25.91
Chl <i>a</i> ($\mu\text{g l}^{-1}$)	28.0	43.8	94.5	140.1
Nutrients (mg l^{-1})				
$\text{NH}_4^+\text{-N}$	16.64	11.58	13.76	23.53
$\text{NO}_3\text{-N}$	2.38	2.72	3.47	2.38
$\text{NO}_2\text{-N}$	$< \text{D.L.}$	$< \text{D.L.}$	$< \text{D.L.}$	$< \text{D.L.}$
PO_4^{3-}	2.99	2.70	2.93	4.36
Major ions (mg l^{-1})				
Cl ⁻	863	-	-	-
SO_4^{2-}	254	-	-	-
Na^+	547	708	1049	-
K^+	58.9	67.4	92.6	-
Mg^{2+}	62.7	72.9	95.7	-
Ca^{2+}	35	36.8	45.1	-

DO = dissolved oxygen, - = not analysed, D.L. = detection limit.

Table II. Planktonic community abundance and productivity in Pony Lake during the summer 2004/05.

	Bacteria ($\times 10^5$ cells ml^{-1})	BP ($\mu\text{g C l}^{-1} \text{d}^{-1}$)	VLP ($\times 10^5$ cells ml^{-1})	PPL ($\times 10^3$ cells ml^{-1})	PPR ($\text{mg C l}^{-1} \text{d}^{-1}$)	Ciliates (cells ml^{-1})
11 Dec 04	4.05 ± 0.57	27.7 ± 1.9	3.56 ± 0.99	0.16	0.67 ± 0.10	1.02^*
21 Dec 04	2.15 ± 0.82	30.5 ± 2.9	2.33 ± 0.60	0.75	2.66 ± 0.31	3.22
29 Dec 04	3.41 ± 1.08	19.7 ± 3.0	4.50 ± 1.15	1.39	1.49 ± 0.05	0.80
14 Jan 05	13.6 ± 2.66	20.3 ± 3.8	0.37 ± 0.10	3.24	1.23 ± 0.09	76.00

*including cysts.

BP = bacterial production, VLP = virus like particles, PPL = phytoplankton, PPR = primary production.

described by Murray *et al.* (1996). PCR products were loaded onto 8–12% polyacrylamide gels. The denaturing gradient contained 40–70% denaturant. The gels ran in 1X TAE at 60 V for 17 hours. Gels were stained with SYBR[®] Gold (Invitrogen) for 15 min and viewed with an Alpha Innotech FluorChem[™] 8800 system. Gel images were analysed using GelComparII software (Applied Math).

16S rRNA gene clone library

We constructed clone libraries for three different dates (early December = 11 December 2004 (ED), late December = 29 December 2004 (LD), and mid-January = 14 January 2005 (MJ)) during the transition period from ice-covered to ice-free conditions by amplifying the 16S rRNA gene with primers 9F (5'-GAGTTTGATCCTGGCTCAG-3') and 1492R (5'-GGTTACCTTGTACGACTT-3') (Stackebrandt & Liesack 1993). PCR products were cloned into pCR[®] 2.1-TOPO vectors (TOPO TA cloning kit, Invitrogen) following the manufacturers guidelines. From each sample 70 clones containing inserts were picked for further analyses. Bacterial clones were sent to Functional Bioscience Inc on LB agar plates for high throughput DNA preparation and DNA sequencing using primer M13F (20). Nucleotide sequences were edited using Sequencher 4.5 (Gene Code Corporation). For each sequence a National Centre for Biotechnology Information (NCBI) Basic Local Alignment Search Tool (BLAST) search was performed in order to determine the closest relatives (BLASTN 2.2.21, ncbi.nlm.nih.gov/BLAST/, accessed May 2009; Zhang *et al.* 2000).

Statistical analyses

Monothetic cluster analysis was applied as outlined by Foreman *et al.* (2011) and Greenwood (2012). This approach aimed to describe differences in the microbial community composition throughout the summer based on the binary presence/absence patterns of bacterial clones.

Results

Physical parameters and chemical analyses

Pony Lake was frozen solid to its base until late November 2004. By the end of November, peripheral moating

occurred at the west end of the lake and by 14 January 2005 c. 80% of the lake was ice-free. The remaining 20% of the ice layer was multi-year ice, but did not impede mixing of the main water column. Surface inflow or runoff from the surrounding hills was not observed. During the open water period the water column was fully mixed by strong winds.

Pony Lake was sampled on four different time points during summer 2004/05, corresponding to changes in the lake ice cover (Table I). During mid-December 2004 to mid-January 2005 Pony Lake was supersaturated with dissolved oxygen (DO). The water temperature increased gradually throughout the sampling period from 1.3°–7.5°C. Lake water was moderately to highly alkaline, ranging from a pH of 8.07–9.77. Pony Lake was Cl^- dominated, although Na^+ and SO_4^{2-} concentrations remained high. The suite of major ion concentrations increased between the initial melt period and mid-January. Dissolved inorganic carbon and DOC concentrations ranged between 3.40–10.84 mg l^{-1} and

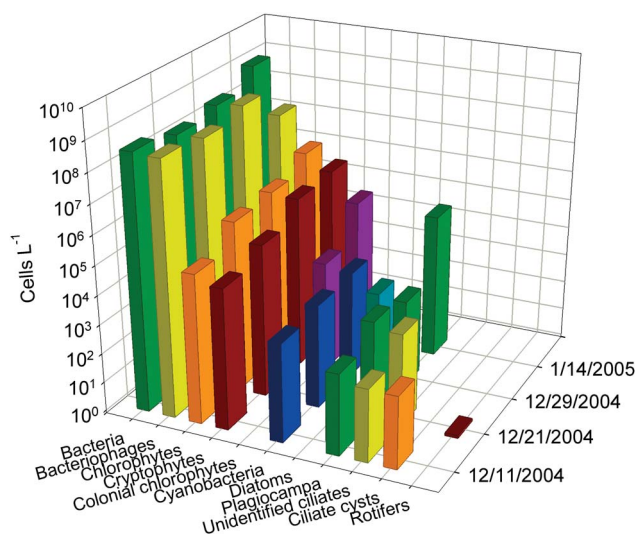


Fig. 2. Planktonic composition of Pony Lake during the 2004/05 summer. Changes in the planktonic composition are displayed by the gradual increase of algal species (chlorophytes and cryptophytes), the increasing abundance of ciliates (e.g. *Plagiocampa*) and the short-term appearance of *Cyanobacteria* and diatoms during the observation period.

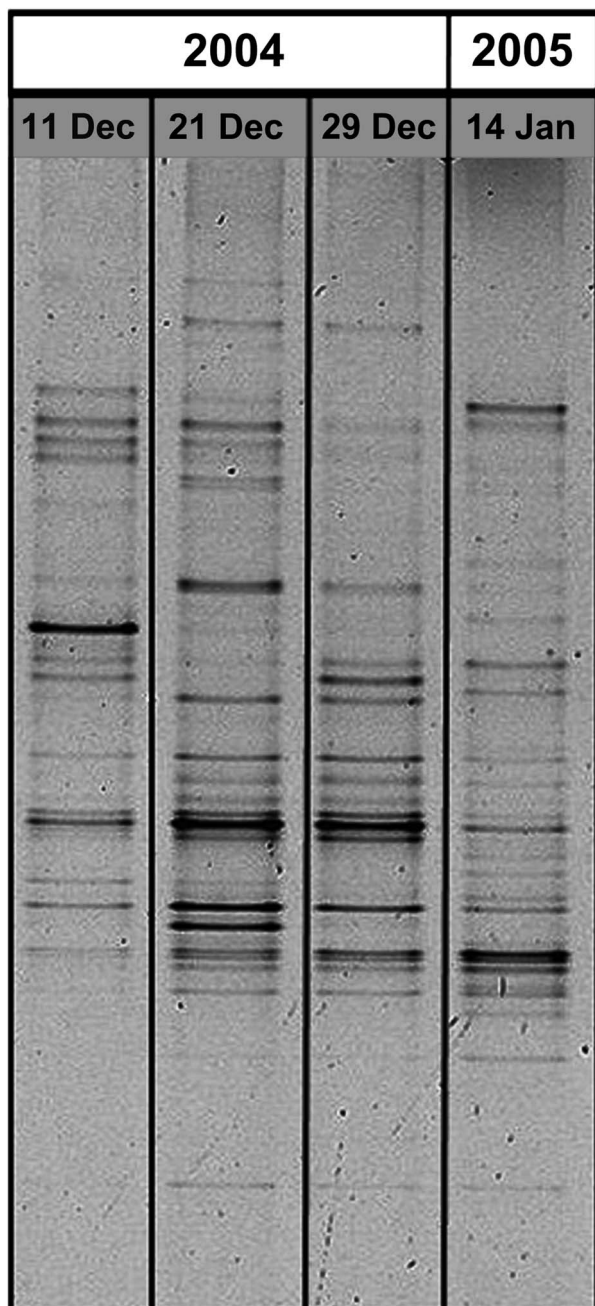


Fig. 3. Denaturing gradient gel electrophoresis (DGGE) profiles of microbial communities from different Pony Lake water samples collected between 11 December 2004 and 14 January 2005 show a shift in community structure. From left to right: 11 December 2004, 21 December 2004, 29 December 2004, 14 January 2005. Image colours were inverted on the camera, but not manipulated.

11.75–28.62 mg C l⁻¹, respectively. Characteristically for a eutrophic lake, inorganic nitrogen and phosphorous concentrations were elevated, with the exception of nitrite, which was below the detection limit (3 µg l⁻¹) throughout this study (Table I).

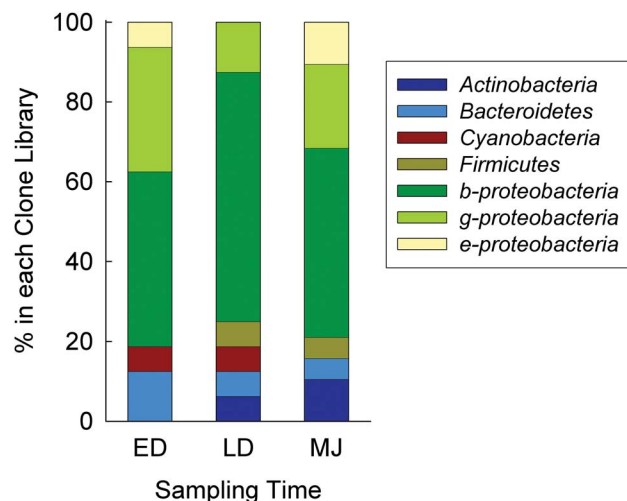


Fig. 4. Distribution of taxonomic classes within the Pony Lake water sample clone libraries from three sampling dates. ED (11 December 2004), LD (29 December 2004), and MJ (14 January 2005).

Productivity measurements and plankton analyses

Pony Lake was populated by bacteria, viruses, phytoflagellates and protozooplankton. Numbers of planktonic organisms are presented in Table II. Bacterial abundances and productivity ranged between 2.15×10^5 and 1.36×10^6 cells ml⁻¹ and from 19.7–30.5 µg C l⁻¹ d⁻¹. Virus like particles peaked by the end of December and abundances ranged from 3.72×10^4 – 4.5×10^5 cell ml⁻¹. Within Pony Lake the phototrophic nanoflagellate (PNAN) assemblage consisted of chlorophytes and cryptophytes (Fig. 2). These phyla were at their maximum of 2.48×10^3 and 7.60×10^2 cells ml⁻¹, respectively, in mid-January. In contrast, heterotrophic nanoflagellates (HNAN) were not detected. Maximum primary production did not coincide with highest phytoplankton numbers. Rather, rates were at their highest in mid-December and ranged from 0.67–2.66 mg C l⁻¹ d⁻¹. Ciliated protozoans were poorly represented in Pony Lake and comprised only one dominant morphotype, *Plagiocampa* sp. (Fig. 2). About 30% of the total number of ciliates remained unidentified (equivalent to ~ 1 cell ml⁻¹). Ciliated cysts were present during the initial melting stage. Numerically, *Cyanobacteria* and diatoms accounted for <0.5% of the total phytoplankton population in Pony Lake. Planktonic *Cyanobacteria* were rare (~ 3 cell ml⁻¹) throughout December 2004 and undetectable in mid-January 2005 (Fig. 2). The dominant *Cyanobacteria* found in Pony Lake belonged to the *Oscillatoria* and *Phormidium* genera. Diatoms were only present on one sampling date. Metazooplankton were sparse with rotifers (one organism l⁻¹; Fig. 2) present only in mid-December.

Table III. Affiliation of 16S rRNA gene phylotypes found in Pony Lake water during 2004/05. Clones labelled ED were from early December, LD from late December and MJ from mid-January in Pony Lake. All Pony Lake clones submitted to GenBank have the prefix ANTPL_.

Taxonomic phylum	Taxonomic class	16S rRNA gene identification(closest neighbour)	GenBank no.	Sampling date					
				11 Dec 2004 Clone	% id.	29 Dec 2004 Clone	% id.	14 Jan 2005 Clone	% id.
<i>Actinobacteria</i>	<i>Actinobacteridae</i>	<i>Candidatus aquiluna rubra</i> MWH-CanK2, AM999977	HM192970			LD03	98	MJ01	98
		<i>Salinibacterium xinjiangense</i> 0543, DQ515964	HM192971					MJ11	98
<i>Bacteroidetes</i>	<i>Flavobacteria</i>	<i>Flavobacterium weaverense</i> AT1042, AY581114	HM192972	ED02	96				
		<i>Cryomorphaceae</i> bacterium Haldis-1, FJ424814	HM192973			LD54	95		
	Uncultured	Uncult. <i>Flavobacteriaceae</i> bacterium F4C94, AY697925	HM192974	ED16	97				
		Uncult. <i>Bacteroidetes</i> bacterium 1D5, AJ627991	HM192975					MJ40	97
<i>Cyanobacteria</i>	<i>Oscillatoriales</i>	<i>Phormidium autumnale</i> Ant-Ph68, DQ493874	HM192976	ED03	99	LD39	99		
<i>Firmicutes</i>	<i>Clostridia</i>	<i>Clostridium acidurici</i> , M59084	HM192977			LD31	93		
	<i>Lactobacillales</i>	<i>Carnobacterium</i> sp. NJ-46, AM396913	HM192978					MJ13	98
<i>Proteobacteria</i>	β -proteobacteria	<i>Acidovorax</i> sp. G3DM-83, EU037287	HM192979	ED57	99	LD27	99	MJ53	99
		<i>Acidovorax</i> sp. BSB42, Y18617	HM192980			LD02	100		
		<i>Acidovorax delafieldii</i> DSM 50263, AJ420323	HM192981			LD05	100		
		<i>Acidovorax</i> sp. UFZ-B530, AF235013	HM192982					MJ30	99
		<i>Achromobacter</i> sp. EP24, AM403526	HM192983					MJ10	95
		<i>Bordetella trematum</i> DSM 11334 (T), AJ277798	HM192984	ED22	98	LD42	99	MJ28	98
		<i>Hermiimonas arsenicoxydans</i> , CU207211	HM192985					MJ59	99
		<i>Hydrogenophaga taeniospiralis</i> SE57, AY771764	HM192986	ED49	100	LD34	100	MJ09	99
		<i>Hydrogenophaga taeniospiralis</i> , AF078768	HM192987	ED06	98	LD41	98	MJ24	98
		<i>Hydrogenophaga atypical</i> BSB 41.8 T, AJ585992	HM192988	ED32	97	LD10	97	MJ25	97
	γ -proteobacteria	<i>Rhodoferrax antarcticus</i> , AF084947	HM192989	ED09	98	LD15	98		
			HM192990			LD36	96	MJ32	96
		<i>Rhodoferrax antarcticus</i> Fryx1, AY609198	HM192991	ED66	99				
		<i>Rhodoferrax ferrireducens</i> T118, CP000267	HM192992			LD07	100		
		<i>Glaciecola polariz</i> LMG 21857, AJ293820	HM192993	ED10	97				
			HM192994	ED68	95				
		<i>Glaciecola punicea</i> ACAM 611 T, U85853	HM192995	ED30	99	LD06	99	MJ48	99
		<i>Glaciecola pallidula</i> ACAM 615 T, U85854	HM192996					MJ02	93
		<i>Pseudomonas</i> sp. An18, AJ551156	HM192997					MJ27	99
		<i>Rheinheimera perlucida</i> BA131 T, AM183347	HM192998	ED48	97				
Uncultured	<i>Rhodanobacter lindaniclasticus</i> , L76222	HM192999					MJ14	95	
	<i>Shewanella baltica</i> NCTC10735, AJ000214	HM193000			LD26	99			
	Uncult. <i>Glaciecola</i> McMurdo.541, AF277554	HM193001	ED51	99					
	ϵ -proteobacteria	<i>Arcobacter cibarius</i> LMG 21997, AJ607392	HM193002	ED61	98			MJ35	98
	<i>Arcobacter cryaerophilus</i> , U25805	HM193003					MJ23	92	

Community structure analysis

DGGE banding patterns (Fig. 3), analysed with GelCompar II software, were used to compare the complex microbial community composition between sampling dates. We further compared the DGGE patterns from these Pony Lake water samples to the DGGE patterns from Pony Lake ice core samples, which were collected earlier in November 2004 when the lake was frozen solid to its base (for GelCompar II dendrogram see Dierer 2009). DGGE image analysis demonstrated that the dominant bacterial community structure in Pony Lake changed during the transition from ice-covered to ice-free conditions. Several different clusters were distinguishable. The water sample collected on 11 December 2004 was most closely related to the community found in the bottom half of the ice core, while samples collected later in the season clustered with

DGGE banding patterns obtained from the top ice core section. The samples collected on 21 December and 29 December grouped together, but were different from the samples collected from Pony Lake in January.

The relative distribution of major phylogenetic groups found within each clone library from the individual sampling dates is shown in Fig. 4. The phylogenetic relationship of Pony Lake clones to their closest neighbours, according to a BLAST search, is summarized in Table III. All clone sequences were submitted to GenBank and bear the prefix ANTPL_, for Antarctic Pony Lake clones. Accession numbers for the Pony Lake clones are from HM192934–HM193003. A total of 34 unique clones were identified from the three sampling dates, with β and γ -proteobacteria comprising the dominant fractions of the clone libraries. These two groups also showed the most compositional overlap between the three sampling dates.

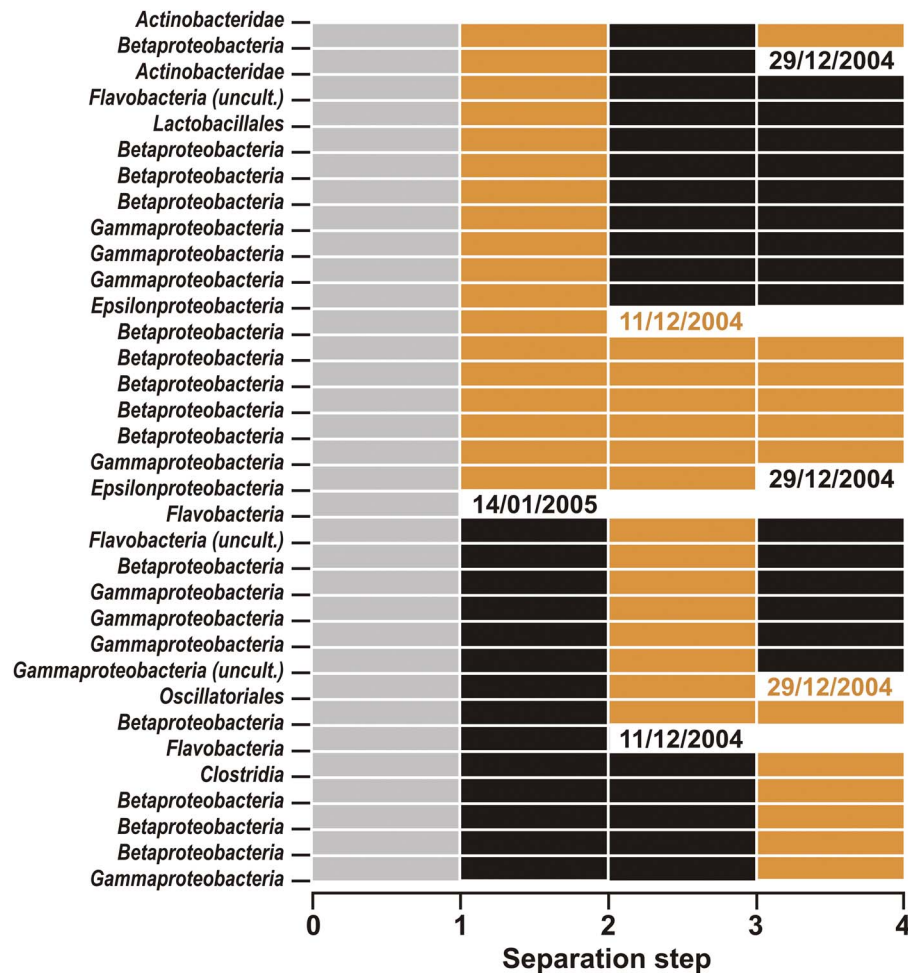


Fig. 5. Banner plot of the monothetic cluster analysis showing closely related taxonomic classes and phyla assigned to Pony Lake clones isolated from three different sampling dates. Orange indicates the presence and black the absence of a clone. Gray represents all of the clones before any clustering steps. The order of the sampling dates was chosen by the clustering algorithm. Thereby, each individual column best explains the responses in the column to its right. The plot is read from top to bottom. Two observations with the same response (absent or present) lead to a coloured bar between their lines. White cells indicate the transition between presence and absence, with the colour of the text indicating the presence or absence of the clone at the bottom of a cell. The plot highlights clusters of clones only found on a single date as well as the small degree of compositional overlap between dates.

For example, *Acidovorax* sp., *Bordetella* sp., *Hydrogenophaga* sp., and *Glaciecola* sp. were the closest reported relatives to clone sequences found throughout the summer season. *Actinobacteria*, *Bacteroidetes*, *Cyanobacteria*, *Firmicutes*, and *ε-proteobacteria* were described by only a small number of sequences. A statistical comparison based on the presence or absence of sequence types was used to discriminate differences between clone libraries for the individual sampling dates (Fig. 5). Distinct clusters, as well as overlap, can be seen for each individual clone library indicating that a shift in the bacterioplankton community did occur during the summer of 2004/05. Clone libraries from early December and mid-January consisted of large numbers of clones (~44% and 53% respectively) that were restricted to these individual sampling dates. Compositional overlap between clone libraries was small and more likely to occur

between consecutive sampling dates. Only six sequence types were present on all three sampling dates. Based upon the small number of clones sequenced we cannot account for all microbes, however our data suggest that a transition occurred among the dominant members of the community that we were able to identify.

Discussion

Physical parameters and chemical analyses

Evolution of the summer meltwaters of Antarctic ponds greatly depends on the extent of melt, wind derived mixing of the water column, and the characteristics of the density gradient after thaw. Typically basal brines, derived from ion exclusion during the freezing process, establish a

stratification of the meltwater pond with freshwater floating on top of layers of high salinity (Healy *et al.* 2006). Physical mixing of the water column can gradually introduce salts and nutrients into the top layers and oxygen into the reduced bottom waters. Consequently, the fate of the limnological structure of the ponds during summer will depend on whether or not this chemical stratification is maintained.

Pony Lake exhibits a steep chemical gradient within its ice column (Dieser 2009). During the summer of 2004/05 strong winds disrupted this chemical stratification after ice break-up and major ion concentrations along with dissolved organic matter and inorganic nutrients increased between the initial melt and late summer lakewater. Although previously attributed to evaporative concentration processes (Brown *et al.* 2004), wind induced mixing of the highly concentrated bottom waters as well as the area of the lake that becomes ice-free during summer more likely determined the chemical composition of the water column. Pony Lake was revisited during the 2005/06 season, however, due to extensive snow accumulation, there was only a small, peripheral ice-free area and mixing was insufficient to break down the salinity gradient. Thus, Pony Lake's water column remained chemically stable (Dieser 2009). The chemical composition of Pony Lake was similar to that found in other coastal Antarctic ponds (Torii *et al.* 1988, Schmidt *et al.* 1991). The brackish character of the lakewater is indicative of a legacy of evaporation and sublimation, the lack of inflows, and the accumulation of marine aerosols. It is important to note that the dissolution of soil salts could have potentially contributed to the increased ion concentrations.

Considerably lower (~three orders of magnitude) phytoplankton counts were measured in our study than those found in the mid 1990s by McKnight *et al.* (1994), which corresponds with a substantial decrease in DOC concentrations found in Pony Lake during this study (11.75–28.62 mg C l⁻¹) than previously reported (95–110 mg C l⁻¹, McKnight *et al.* 1994; 32.4–92.4 mg C l⁻¹, Brown *et al.* 2004). Several processes such as changes in phytoplankton blooms, progressive exploitation of the carbon pool by bacterioplankton (Bell & Laybourn-Parry 1999, Laybourn-Parry *et al.* 2002, Brown *et al.* 2004) or variable rates of dissolved organic matter (DOM) production by algal populations (Brown *et al.* 2004) may alter DOC concentrations. Our seasonal dataset suggests that a progressive exploitation of the DOC pool by bacteria seems unlikely in this instance as bacterial cell numbers and DOC concentrations peaked in January when production rates were markedly reduced compared to early and mid-December (Table II). Moreover, the decoupling of bacterial production and DOC maxima suggests changes in the DOM composition over the course of the summer. Generally, organic matter can undergo photochemically induced transformations that may render it more refractory and hence less bioavailable for

microbial uptake (Benner & Biddanda 1998). Thus, the observed increase in DOC concentration and the simultaneous decrease in bacterial production between early December and mid-January may be due to the accumulation of recalcitrant photolytic products. Mao *et al.* (2007) demonstrated that the fulvic acid fraction in Pony Lake had undergone significant humification from the original algal precursor material that generated highly cross-linked structures. While algal derived DOC is considered to be of young age, Mao *et al.* (2007) suggested that the organic matter in Pony Lake has been structurally altered ('aged') and would be of comparably poor substrate quality. Nonetheless, a steady exploitation of the rich DOC pool detected during previous studies over the past ten years seems evident (McKnight *et al.* 1994, Brown *et al.* 2004). Further, different phytoplankton blooms associated with changes in DOM production rates may have accounted for shifts in the DOC levels and the overall decrease in DOC concentrations (McKnight *et al.* 1994, Brown *et al.* 2004, this study). In past studies, algal blooms have correlated with DOC concentrations in Pony Lake. McKnight *et al.* (1994) reported high DOC concentrations during the chlorophyte bloom, and Brown *et al.* (2004) observed a decrease in DOC concentrations concurrent with a declining chlorophyte population and a shift in algal blooms, underscoring the argument that different algal species exhibit different DOC production rates.

Inorganic nitrogen and phosphorous concentrations in Pony Lake were elevated compared to those found in other eutrophic Antarctic lakes (Mataloni *et al.* 1998, Bell & Laybourn-Parry 1999, Butler 1999). Although runoff from the adjacent penguin rookeries was not observed, the high nitrogen and phosphorus levels in Pony Lake could potentially be derived from bird droppings (Vincent & Vincent 1982). Decreasing NH₄-N concentrations at the beginning of the growth season suggest that ammonium was the preferred nitrogen source for phytoplankton, as previously shown in several other Antarctic lakes (Hawes 1983). Maximum primary production rates were measured on 21 December 2004 when ammonium concentrations dropped markedly (Tables I & II). In contrast, while primary production rates gradually decreased between 21 December 2004 and mid-January 2005, the NH₄-N concentrations peaked during this same period. Furthermore, decomposition of organic matter by heterotrophic bacteria or photolytic reactions may have resulted in mineralization of the dissolved organic matter, thereby generating NH₄.

Productivity measurements and plankton analyses

In common with other continental Antarctic lakes, Pony Lake is populated by bacteria, viruses, phytoflagellates and protozooplankton. The high levels of inorganic nutrients and DOC concentrations (Table I) available throughout the study period imply that these factors should not limit

summer plankton growth and that other parameters such as temperature, ionic composition, and light intensity may be more important in triggering plankton blooms.

Bacterial cell abundances ranging from 2.15×10^5 – 1.36×10^6 cells ml⁻¹ were comparable to those reported from maritime or continental Antarctic lakes across different trophic levels (Takacs & Priscu 1998, Bell & Laybourn-Parry 1999, Butler 1999, Butler *et al.* 2000). Bacterial production measured in Pony Lake was considerably higher compared to oligotrophic Antarctic lakes (Takacs & Priscu 1998, Butler *et al.* 2000, Laybourn-Parry *et al.* 2002), but lower than reported in other saline or eutrophic Antarctic lakes (Laybourn-Parry *et al.* 2002). Importantly, DOC concentrations in these lakes (Laybourn-Parry *et al.* 2002) were similar to those found in Pony Lake. Bacterial growth is regulated by temperature, and the availability of inorganic nutrients (N and P) and organic substrates (DOC). It is unlikely that either temperature or nutrients would suppress summer bacterial production in Pony Lake. After an early season maximum, productivity decreased while water temperatures increased and nutrients were plentiful throughout the season. Although one might hypothesize that higher DOC concentrations would precede enhanced bacterial growth, this process is reversed in Pony Lake. Highly bioavailable DOC at the beginning of the season may not only be generated from a proliferating planktonic community, but may also be released from the melting ice. In shallow meltwater ponds, photosynthesis can persist beneath the ice during freeze-up when sufficient levels of incident light prevail (Hawes *et al.* 2011). Subsequently newly released DOC would become incorporated and preserved in the ice matrix, providing a rapidly available carbon pool when growth conditions turn more favourable. This mechanism has previously been suggested for Pony Lake due to the stimulated bacterial production in the mid-section of the winter ice column (Foreman *et al.* 2011). Conversely, increased DOC concentrations mixed in from basal brines later in the season did not enhance bacterial growth. This supports the view of a strong conservative behaviour of organic compounds and an accumulation of refractory organic material in lakewater (Howard-Williams & Hawes 2007, Hawes *et al.* 2011).

Virus like particle abundances reported from Pony Lake were in the same range as has been shown in other Antarctic lakes, such as Lake Druzhby, Crooked Lake, and Beaver Lake in Vestfold Hills or Lake Hoare and east lobe Lake Bonney in the McMurdo Dry Valleys (see Sävström *et al.* 2008 for a review). The virus to bacteria ratio (VBR) in polar inland lakes usually falls between 1 and 34, however exceptionally high ratios (> 120) have also been reported for the saline lakes in Vestfold Hills (Sävström *et al.* 2008). The VBR in Pony Lake ranged from 0.03–1.32. Low ratios can be explained in part by the autochthonous nature of carbon substrates in Antarctic

lakes (Sävström *et al.* 2008). Bacterial abundances are typically lower than VLP numbers due to the infectious interactions of VLP with the host cell causing viral induced cell lysis. However, low bacterial growth rates in polar waters do not sustain high infection rates. On average, only four viruses are released from each bacterial cell (Laybourn-Parry 2009). Viral processes and infectivity have also been related to a range of abiotic factors. Madan *et al.* (2005) reported a negative correlation between VLP and temperature, indicating lower decay rates under lower temperatures. Nutrient limitation can reduce bacterial proliferation, thereby indirectly affecting VLP numbers (Weinbauer 2004). Although the VBR was low in Pony Lake, the interaction between viruses and bacteria may play an important role in carbon and nutrient cycles. For instance, virus induced release of organic carbon from bacteria can provide a significant portion of the DOC pool and importantly short-circuit the carbon cycle before bacterial produced carbon is removed by protozoan grazing.

The dominance of chlorophytes and cryptophytes in Pony Lake has been previously reported (McKnight *et al.* 1994, Brown *et al.* 2004), and was also found during this study. It has been shown that these phyla dominate numerous Antarctic lakes (Butler *et al.* 2000, Roberts *et al.* 2004), while *Chlamydomonas* species are typically found in eutrophic Antarctic lakes (Mataloni *et al.* 1998, Butler 1999). While the dominant organisms have remained consistent across the years there is variability in the abundance of individual species and the timing of phytoplankton blooms (McKnight *et al.* 1994, Brown *et al.* 2004, this study). For example, Brown *et al.* (2004) showed that following the loss of the ice cover in mid-December 1997 the chlorophyte bloom was displaced by a bloom of cryptophytes, while the study by McKnight *et al.* (1994) reported the dominance of chlorophytes in Pony Lake at the end of January 1994. Dominance of a particular phylum or a distinct algal bloom was not observed in the current study, where both chlorophytes and cryptophytes increased gradually during the transition from an ice-covered to an ice-free lake. The dynamic nature of Antarctic phytoplankton populations has also been observed by Mataloni *et al.* (1998) in the highly eutrophic Otero Lake (unofficial name). Overall, phytoplankton was two to three orders of magnitude less abundant in Pony Lake during this study along with a 1.4 to 3 fold decrease in salinity. Laybourn-Parry *et al.* (2002) demonstrated a strong correlation between the productivity of the planktonic community and salinity in numerous lakes in Vestfold Hills, whereby productivity increased across the salinity spectrum (from brackish to hyper-saline).

Although phytoplankton numbers and chl *a* concentrations still increased throughout the season, primary production in Pony Lake peaked in mid-December. Chlorophyll *a* concentrations correlated with the phytoplankton maxima ($r = 0.97$), but were only weakly negatively correlated with

primary production rates ($r = -0.12$). This temporal decoupling of production and plankton maxima was also observed in several lakes in Vestfold Hills (Laybourn-Parry *et al.* 2002). In general, primary productivity in Pony Lake was higher than found in other more oligotrophic Antarctic lakes (Butler *et al.* 2000, Laybourn-Parry *et al.* 2002). However, similar or even larger primary productivity rates have been reported in several Antarctic lakes across trophic levels (Butler 1999, Laybourn-Parry *et al.* 2002). Variations in the photosynthetic communities or the underwater light climate between lakes (e.g. photosynthetically active radiation in an ice-covered vs ice-free system) may account for the differences in primary production.

Proto- and metazooplankton numbers in Pony Lake during this study were generally low with low species diversity. The dominant species, *Plagiocampa*, accounted for 70% of the total ciliate population. This differs from previous reports where Armitage & House (1962) primarily found *Euplotes* in Pony Lake. Ciliate cysts, also detected in Pony Lake ice cores (Foreman *et al.* 2011), were only present at the beginning of the transition period from ice-covered to ice-free. Since the growth season during the summer is typically short, cysts are produced as a strategy to survive unfavourable conditions during the winter (Mataloni *et al.* 1998, Bell & Laybourn-Parry 1999). Cysts will then lie dormant until conditions become suitable for vegetative growth. Similar to Armitage & House (1962) who identified *Philodina* as the dominant species, rotifers were rare and only detected in mid-December.

Although Pony Lake shows indications of a complex microbial food web, it seems unlikely that the small number of metazoan predators and the low abundance of raptorial ciliates (e.g. *Plagiocampa*) strongly control PNAN or bacteria by grazing. Mixotrophic cryptophytes could exert grazing pressure on bacteria, however, phagotrophy would seem to be more beneficial to support or supplement autotrophy during periods of light and nutrient limitation. As with other Antarctic lakes (Prisco *et al.* 1999, Roberts *et al.* 2004) there appears to be little top-down control over the microbial food web.

Community structure analysis

Overall, c. 68% of the Pony Lake clones closely matched sequences reported from geographically diverse Antarctic lake and marine environments (e.g. Van Trappen *et al.* 2002, Glatz *et al.* 2006) which indicates a high level of similarity for microorganisms inhabiting these cold ecosystems. An important goal of the phylogenetic analysis was to characterize any changes in the microbial community composition over the course of the summer in Pony Lake. Despite the inherent limitations of this technique, DGGE has been successfully employed to describe the minimum number of dominant phylotypes present (Pearce 2005, Villaescusa *et al.* 2010). Both gel

comparison of DGGE banding patterns and cluster analysis of representative clones from Pony Lake indicate a shift in the microbial community during the early and late summer open water period. Differences in the microbial composition were also apparent when comparing the open water to the lake ice community (Foreman *et al.* 2011). From the sequences found in the Pony Lake ice cores only eight of these sequence types could be identified in the Pony Lake water column as well. The highest degree of sequence overlap was found among members of the β -*proteobacteria* lineage. When assigned putative identifications these Pony Lake clones were shown to be most closely related to *Bordetella* sp. and *Hydrogenophaga* sp. Representatives from the anaerobic δ -*proteobacteria*, *Spirochaetes*, and *Verrucomicrobia* lineages were solely present in the ice core samples (Foreman *et al.* 2011). In contrast, the phylum *Actinobacteria* was only found in the Pony Lake water samples. Obvious differences were observed between taxonomic classes. Within the ice, the most dominant classes were *Bacteroidetes* and ϵ -*proteobacteria* (Foreman *et al.* 2011). Over the course of the summer sequence types diverged widely from those found in the initial ice community and members of the β - and γ -*proteobacteria* prevailed (Fig. 4). Physicochemical conditions in the Pony Lake ice were distinctly different (oxygen depletion and severe chemical stratification of nutrients, DOC and ions; Dieser 2009) from those in the summer lakewater. It seems plausible that the chemical stratification of the ice column and the oxygen depletion prior to complete freeze-up created an environment favouring microbes with high salt and low temperature tolerance, as well as anaerobes, as suggested by Foreman *et al.* (2011). Conversely, Pony Lake summer lakewater was well mixed and supersaturated with oxygen. Interestingly, despite these obvious differences in environmental conditions, the microbial community trapped within the ice column during winter is responsible for re-colonizing Pony Lake when melt is initiated. However, this process largely depends on the timing and extent of the ice cover loss. In mid-December 2004 the melt proceeded rapidly and physicochemical changes occurred in the water column (Table I), driven by higher air temperatures, strong winds, and proliferating algal communities. Pearce (2005) found that in three lakes on Signy Island, South Orkney Islands, physical factors affected the development of the microbial community during the seasonal transition from ice-covered to the summer ice-free, whereas biotic factors were believed to become more important later in the season. The establishment or disruption of a vertical stratification and, in particular, temperature critically affected community structure. Temperature differences of as little as 0.1°C have been shown to cause significant variations in the community composition (Pearce 2005). Although we believe that grazing on bacterioplankton by bacteriophagous protists, as well as mortality rates by virioplankton in

Pony Lake, were insufficient to offset bacterial abundance throughout the season, negative effects on individual group members of the community are still possible. Further, there is indication that the structuring of the microbial community is strongly correlated to the chemical progression of the water column. A distinct change in DGGE profiles was observed as thorough mixing gradually increased the salinity of Pony Lake. Conversely, when the water column of Pony Lake remained stable, the development of a different community throughout the open water period appeared to be minimal (Dieser 2009). In general, the introduction of new microorganisms from the deeper parts of the lake may also be considered. For instance, *Firmicutes* were confined to the bottom layer of the ice core (Foreman *et al.* 2011). Although absent at the beginning of the melt season, *Firmicutes* were present when Pony Lake became largely ice-free and fully mixed. Considering the extreme intra-seasonal alterations in environmental conditions in Pony Lake it does not seem surprising that we detected different, dominant members of the microbial community between winter ice, early, and late-summer lakewater. Depending on the lake area that becomes ice-free during summer it can be hypothesized that the microbial community within the ice initially seeds the development of the lakewater community. However, it is unclear whether the distribution of microbes within the ice is the effect of physical processes during freezing or other environmental factors. Hawes *et al.* (2011) showed that most of the lake/sediment area was frozen prior to the onset of more severe conditions, protecting planktonic and benthic organisms from stresses imposed by concentrated liquid brines. In this context it is noteworthy that while the presence of many community members is coupled with the physicochemical state of Pony Lake (e.g. redox conditions), more versatile species (e.g. clones that were related to *Bordetella* sp. and *Hydrogenophaga* sp.) find a perennial niche in this lake as well.

Conclusions

Pony Lake is a small, brackish, eutrophic lake at Cape Royds, Ross Island. Lake chemistry and nutrient levels are affected by the input of marine aerosols and the adjacent penguin rookery. The evolution of its microbial community appears to be a complex process, driven by seasonal physicochemical changes. Over the course of the summer progressive melt of the ice column, wind mixing, the lack of inflows, increases in water temperature, and changes in salt and nutrient concentrations stimulated a compositional shift in the dominant microbial community in Pony Lake. Aside from intra-seasonal changes, a clear decrease in the phytoplankton numbers was apparent in Pony Lake when compared to previous studies. In Pony Lake, where microorganisms form the basis of the food web and control biogeochemical cycles, such a prominent drop in phytoplankton numbers will alter the carbon pool, its fluxes

and transformation, and may compromise the entire lake ecosystem. Primary producers are the fundamental source of inorganic carbon fixation and autotrophic energy production, providing organic carbon and nutrients for subsequent trophic levels. Closely linked to primary production is the exudation of photosynthate as DOC, a major substrate for heterotrophic bacteria. It appears that over the past decade Pony Lake became less productive and its once rich DOC pool has been steadily exploited by the bacterioplankton community. Although bacteria are an essential link in the recycling of nutrients and transformation of organic carbon back into the food web, these organisms acted as a large carbon sink. The more labile DOC fractions would have been preferentially hydrolysed or re-mineralized by bacterioplankton while abiotic processes (e.g. solar radiation induced polymerization and condensation) can transfer the remaining DOC into more recalcitrant forms. Eventually what would emerge is a dissolved organic carbon pool, exhausted in rapidly, bioavailable substrates, but enriched in components that are less susceptible to enzymatic cleavage. Nonetheless, what triggered the decline in phytoplankton numbers in Pony Lake remains unclear. End member ecosystems, such as Pony Lake, can exhibit a rapid response to changes in their physicochemical environment. Highly enriched in nutrients, the microbial community in Pony Lake is probably controlled by factors such as air and water temperature, wind and salinity that determine the length of ice or open water period, the establishment of a stratified or mixed water column, and the ionic composition, respectively. Our seasonal dataset highlights the interaction between environmental parameters and changes in the microbial community composition. Thus, organisms which persist throughout the year in Pony Lake must be capable of surviving rapidly changing conditions.

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