

Shifts in DNA yield and biological community composition in stored sediment: implications for paleogenomic studies

Katie A. Brasell^{1,2}, Xavier Pochon^{1,2}, Jamie Howarth³, John K. Pearman¹, Anastasija Zaiko^{1,2}, Lucy Thompson¹, Marcus J. Vandergoes⁴, Kevin S. Simon², Susanna A. Wood¹

¹ Cawthron Institute, 98 Halifax Street East, Nelson, New Zealand

² University of Auckland, 23 Symonds Street, Auckland, New Zealand

³ Victoria University of Wellington, Kelburn Parade, Wellington, New Zealand

⁴ GNS Science, 1 Fairway Drive, Lower Hutt, New Zealand

Corresponding author: Katie A Brasell (kbrasell@gmail.com)

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Abstract

Lake sediments hold a wealth of information from past environments that is highly valuable for paleolimnological reconstructions. These studies increasingly apply modern molecular tools targeting sedimentary DNA (sedDNA). However, sediment core sampling can be logistically difficult, making immediate subsampling for sedDNA challenging. Sediment cores are often refrigerated (4 °C) for weeks or months before subsampling. We investigated the impact of storage time on changes in DNA (purified or as cell lysate) concentrations and shifts in biological communities following storage of lake surface sediment at 4 °C for up to 24 weeks. Sediment samples (~0.22 g, in triplicate per time point) were spiked with purified DNA (100 or 200 ng) or lysate from a brackish water cyanobacterium that produces the cyanotoxin nodularin or non-spiked. Samples were analysed every 1–4 weeks over a 24-week period. Droplet digital PCR showed no significant decrease in the target gene (*nodularin synthetase* – subunit F; *ndaF*) over the 24-week period for samples spiked with purified DNA, while copy number decreased by more than half in cell lysate-spiked samples. There was significant change over time in bacteria and eukaryotic community composition assessed using metabarcoding. Amongst bacteria, the cyanobacterial signal became negligible after 5 weeks while Proteobacteria increased. In the eukaryotic community, Cercozoa became dominant after 6 weeks. These data demonstrate that DNA yields and community composition data shift significantly when sediments are stored chilled for more than 5 weeks. This highlights the need for rapid subsampling and appropriate storage of sediment core samples for paleogenomic studies.

Key Words

chilled storage, droplet digital PCR, extracellular DNA, metabarcoding, sedimentary DNA (sedDNA)

Introduction

Molecular techniques, in particular metabarcoding, are increasingly being used to monitor biological communities and assess ecological conditions in aquatic environments (Pawlowski et al. 2014; Rawlence et al. 2014; Domaizon et al. 2017; Pochon et al. 2017; Baho et al. 2019; Ellegaard et al. 2020; Tsukamoto et al. 2021). Paleolimnological studies are also benefiting from the advances in affordable and high-throughput molecular techniques.

Molecular analysis of sedimentary DNA (sedDNA; intra- and extra-cellular DNA from bulk sediment samples) is now being used in paleogenomics to supplement traditional proxies through the provision of high-resolution community composition data (Capo et al. 2017a; Ficetola et al. 2018). For example, Capo et al. (2019) detected changes in lake microbial communities that corresponded with seasonal shifts in landscape and climate at an inter-annual scale over a 40-year period using a combination of molecular techniques targeting sedDNA, sediment

geochemistry, biomarkers and meteorological data. Molecular techniques have also been used to reconstruct historic phytoplankton communities, allowing the extension of lake water quality records beyond limited historical monitoring periods (Ibrahim et al. 2021).

Robust molecular analysis in paleogenomics relies on high quality sedDNA. Stability of sedDNA is dependent on multiple factors, including temperature, light and chemical bonding to sediment particles (Strickler et al. 2014; Seymour et al. 2017). Lake sediments contain DNA from both prokaryotes (bacteria and archaea) and eukaryotes (micro-eukaryotes and larger organisms), which can be intra- or extra-cellular in origin. In sediment samples, intra-cellular DNA is protected from light and enzyme activity when cell membranes are intact, such as in resting cysts of algae or rigid cell walls of diatoms, but these can be exposed to harsh conditions once cells are lysed. In the environment, nucleases within lysed cells and those produced by heterotrophic bacteria in sediments can quickly degrade DNA (Dell'Anno and Corinaldesi 2004; Corinaldesi et al. 2008). However, the majority of extra-cellular DNA is bound to sediment particles that help to protect it from nucleases and reduce the bioavailability of these molecules (Nielsen et al. 2006; Pietramellara et al. 2009; Ellegaard et al. 2020). There is a clear need to better understand DNA preservation and the influence of environmental factors on the amount and composition of sedDNA archived in sediment cores (Brunson and Reich 2019; Lindqvist and Rajora 2019).

Ideally, subsampling of sediment cores and freezing of samples should be undertaken immediately following fieldwork. However, sediment core sampling can be logistically difficult, especially in remote locations, making immediate subsampling for sedDNA challenging. There are an increasing number of large-scale projects collecting sediment cores from hundreds of lakes, such as Lake Pulse (Canada; <https://lakepulse.ca/>) and Lakes380 (New Zealand; www.lakes380.com), which operate in remote locations and make immediate subsampling impossible. Given the high sensitivity of molecular methods, extreme care is also needed during the subsampling stage to ensure there is no contamination. It is strongly recommended that subsampling is undertaken in a dedicated laboratory and not in the field (Domaizon et al. 2017; Taberlet et al. 2018). Sediment cores could be frozen upon arrival at the laboratory; however, freezing cores makes subsampling mechanically challenging and stratigraphic integrity can be compromised during freezing of cores with high water content. Freezer storage is also energy intensive and the capacity for storing large bulk samples, such as 2–6 m long sediment cores, is often limited. Therefore, the most common approach is to refrigerate sediment cores at 4 °C in the dark until subsampling and processing are logistically feasible (see Brasell et al. 2021 and Pearman et al. 2021a). While it is assumed that the dark/cold storage conditions mimic the environmental conditions found in the sedimentary layers of a lake, knowledge on whether sedDNA degrades or if community composition changes in stored sediment is very limited.

Numerous studies describe the effects of various *in situ* environmental factors on environmental DNA isolation (Corinaldesi et al. 2008; Turner et al. 2015; Eichmiller et al. 2016; Capo et al. 2017b; Bowers et al. 2021) and the chemical preservation of small subsamples (Rissanen et al. 2010). However, very few studies have explored how temporal storage of bulk sediment samples might impact the overall DNA yield and changes in biological community composition. Studies of blood (Li et al. 2018) and urine samples (Ng et al. 2018) found little difference in DNA yield between frozen and chilled storage treatments over 20 and 100 days, respectively. Studies of DNA yield from stored sediments and soils have mostly focused on frozen or dried preservation (Wallenius et al. 2010). In a study of decadal-centennial fish abundance in marine sediments, Kuwae et al. (2020) found that sediment cores stored chilled for 6 days or 4 weeks had only minor degradation (DNA concentration) compared to a core frozen within 48 hrs. Less is known about changes to DNA of environmental samples in long-term storage, where most extracellular DNA is sediment-bound.

Bacteria can degrade and recycle extracellular DNA in marine sediments via extracellular nucleases (Wasmund et al. 2021), indicating there is potential for decreased DNA yields if sediment bacteria survive during long-term storage. Furthermore, some microbes can survive in low energy conditions, such as subsurface sediments (Ciobanu et al. 2014) and they may influence the community composition by consuming or outcompeting other microbes. Some common protist taxa consume bacterial and fungal communities in terrestrial soils and marine sediments (Beaudoin et al. 2016; Dumack et al. 2019, 2020; Khanipour Roshan et al. 2021), but there is little information about their impact on sedDNA integrity and composition during chilled sediment storage.

Our aim was to determine the effects of chilled storage on DNA yield and compositional shifts of bacterial and eukaryotic communities in lake sediment samples. Due to the need for replication and the multiple treatments tested, it was not possible to test samples from multiple layers within a sediment core. As such, surface lake sediments (top 1 cm) were selected for this study. Surface samples contain a mixture of live organisms, as well as extra- and intra-cellular DNA (Pearman et al. 2021b); therefore, they provide a worst-case scenario (compared to deeper sediment layers) as these are environments where DNA degradation and community compositional shifts are likely to occur. As such, surface sediment assessments are important for broader paleolimnological and ecological studies; they represent the current condition of lakes and it is paramount that data, generated from these samples, are accurate.

In our study, sediment samples were collected from a eutrophic lake and subsamples were artificially spiked with either purified DNA (100 or 200 ng) or lysate from a brackish water cyanobacteria (*Nodularia spumigina*) or not spiked. We chose to spike with *Nodularia spumigina* DNA as it would not be found in these samples and this species produces the cyanotoxin nodularin. The enzyme

cluster responsible for nodularin synthesis has been characterised (Moffitt and Neilan 2004) and PCR-based assay developed to quantify an abundance-specific gene (Koskenniemi et al. 2007), one of which we targeted in this study. Samples were stored chilled (4 °C) in the dark for up to 24 weeks. The spiked samples were analysed using a targeted droplet digital PCR (ddPCR) assay and the non-spiked samples used for metabarcoding of bacterial (16S ribosomal RNA) and eukaryotic (18S ribosomal RNA) genes to assess sediment community composition. We hypothesised that, across a 24-week period, copy numbers of the target gene (*nodularin synthetase*) would: *i*) not change in samples spiked with purified DNA, but *ii*) a decrease would be detectable in samples spiked with cell lysate due to the additional cellular enzymes introduced and *iii*) the bacterial and eukaryotic composition would not change significantly because the cold, dark conditions are similar to the lake-bed environment.

Methods

Site and sample collection

Lake Pounui is a small (46 ha), shallow (max. depth 9.6 m), lowland coastal lake situated 14 m above sea level, 30 km northeast of Wellington, New Zealand

(41°20'34"S, 175°6'48"E). Lake Pounui's catchment (627 ha) reaches an elevation of 470 m in the foot hills of the Rimutaka Ranges and is covered by unmodified indigenous beech-podocarp forest (96%), with the remainder in pastoral land cover (Perrie and Milne 2012). Although Lake Pounui has a forested catchment, it now regularly experiences cyanobacterial blooms dominated by *Microcystis* spp. and *Dolichospermum* spp.

Triplicate ponar grab samples were taken at a central point in the Lake. Using a sterile spatula, the top 1 cm of the undisturbed surface sediment layer of each grab was placed in separate sterile 400 ml containers and transported chilled to the laboratory. Samples were stored chilled for two days and then processed. The samples were homogenised by gentle hand shaking and 53 aliquots (ca. 0.22 g) from each container were transferred into pre-weighed microtubes (1.7 ml; Qiagen) with a 1000 µl mechanical pipette set to 200 µl and using a sterile wide bore pipette tip (step 1 in Fig. 1). The microtubes were then weighed and sediment weights calculated by subtracting the weight of empty microtubes from the filled tube weight (target weight 0.22 g +/- 0.03 g).

Experimental set-up

DNA and cell lysate from the brackish-water cyanobacterium *Nodularia spumigina* were used for the experiment

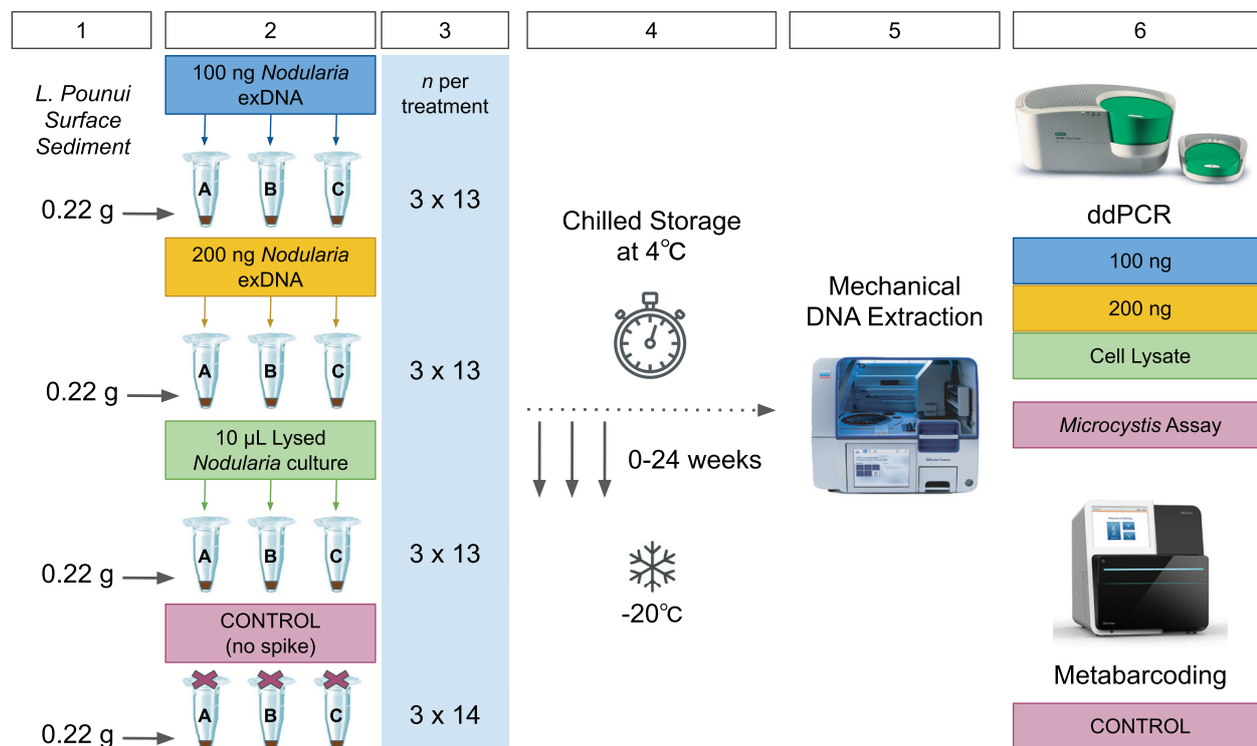


Figure 1. Schematic diagram of the experimental set up and analytical process used in this study, broken down into six steps. 1) Aliquoting of three independent surface sediment grabs, 2) Addition of DNA treatments, 3) Chilling samples in sets of 3 reps × 13 time points (or 14 time points for control samples, which include a time 0 control), 4) The storage phase where triplicates of each treatment are removed from fridge at specific intervals and frozen for later analysis, 5) DNA extraction of all treatments including controls, using a QiaCube (QIAGEN, Germany), 6) DNA amplification of *N. spumigina*-spiked treatments and a specific investigation of *Microcystis* abundance via droplet digital PCR (ddPCR) and metabarcoding of the non-spiked control samples for assessing community structure.

as this species is not present in Lake Pounui. Three spiking treatments were used: 100 ng and 200 ng of DNA extract and 10 μ l cell lysate (step 2 in Fig. 1).

To obtain extracellular DNA (exDNA) for spiking, DNA was extracted from a non-axenic culture of *N. spumigina* (CAWBG21; <http://cultures.cawthron.org.nz/>). Cultured material (1 ml) was siphoned off into three microtubes and centrifuged (10 mins, 3,000 \times g). The remaining liquid media was decanted and the cells transferred into six bead tubes of a DNEasy PowerSoil DNA Isolation Kit (QIAGEN, USA). DNA extraction was automated using a QIAcube (QIAGEN, USA), with DNA eluted in 100 μ l volumes. DNA extracts were pooled and quantified with a NanoPhotometer NP80 (Implen GmbH, Munich, Germany) to calculate the required volume for spiking to achieve concentrations of 100 ng/ μ l (low concentration spike) and 200 ng/ μ l (high concentration spike). To obtain cell lysate for spiking, 10 ml of *N. spumigina* (CAWBG21) culture with liquid media was sonicated (30 secs) and filtered (3 μ m; filter type/brand/brand) to remove excess non-DNA cellular contents.

For each treatment, 13 sediment aliquots were spiked in triplicate (step 3 in Fig. 1). The DNA or cell lysate was mixed with the sediment by pipetting the mixtures up and down three times. The samples were stored in the dark in a polystyrene container at 4 $^{\circ}$ C ($n = 39$ samples per treatment). Triplicate non-spiked control samples for each time point were chilled alongside the treatments, including a set of triplicate samples for time zero that were immediately frozen at -20 $^{\circ}$ C ($n = 49$ control samples). Triplicate samples were removed from chilled storage and frozen (-20 $^{\circ}$ C) at 1, 2, 3, 4, 5, 6, 8, 10, 12, 14, 16, 20 and 24 weeks (step 4 in Fig. 1).

Post-storage DNA extraction and nodularin-specific droplet digital PCR assay

Each step of the molecular analysis, including DNA extraction, PCR set-up, template addition, PCR amplification, amplicon purification and quantification, was conducted at the Cawthron Institute (Nelson, New Zealand) in separate sterile rooms dedicated to that step, with sequential workflow to prevent cross-contamination. Rooms dedicated to DNA extraction, PCR set-up and template addition were sterilised with ultra-violet light for a minimum of 15 mins before and after each use. The PCR set-up and template addition were undertaken in laminar flow cabinets with HEPA filtration. Aerosol barrier tips (Eppendorf, Germany) were used throughout.

The sediment from each time point ($n = 53$) was transferred from microtubes into bead tubes of a DNEasy PowerSoil DNA Isolation Kit (QIAGEN, USA) using a pipette and the bead tube liquid to assist collection of sediment from the microtubes. DNA was then extracted using a QIAcube (QIAGEN, USA), with DNA eluted in 100 μ l volumes (step 5 in Fig. 1). Extractions were performed in batches of ten to twelve samples, including a negative control (i.e. extraction blank containing all the

reagents, but no sediment) every two batches. DNA concentration and quality were measured using a NanoPhotometer NP80 (Implen GmbH, Munich, Germany).

The genus *Nodularia* is the only known producer of the hepatotoxin nodularin, which is produced via the *nodularin synthetase* enzyme complex (Koskenniemi et al. 2007). To quantify spiked extracellular DNA and cell lysate, absolute concentrations of the *nodularin synthetase* gene subunit F (*ndaF*) were measured in all samples using a BioRad QX200 Droplet Digital PCR (ddPCR) system (step 6 in Fig. 1). Primers used were the *ndaF*8452 forward primer 5'-GTG ATT GAA TTT CTT GGT CG-3' and the *ndaF*8640 reverse primer 5'-GGA AAT TTC TAT GTC TGA CTC AG-3' (Koskenniemi et al. 2007). Each ddPCR reaction included 450 nM of forward and reverse primer, 10 μ l of 2 \times BioRad ddPCR EvaGreen Supermix, 3 μ l DNA and sterile water for a total reaction volume of 22 μ l. The BioRad QX200 droplet generator partitioned each reaction mixture into nanodroplets by combining 20 μ l of the reaction mixture with 70 μ l of BioRad droplet oil. After processing, this resulted in a total nanodroplet volume of 40 μ l, which was transferred to a PCR plate for amplification using the following cycling protocol; hold at 95 $^{\circ}$ C for 5 min, 40 cycles of 95 $^{\circ}$ C for 30 s and 60 $^{\circ}$ C 1 min, a cooling step of 4 $^{\circ}$ C for 5 min and a final enzyme deactivation step at 98 $^{\circ}$ C for 5 min. The plate was then analysed on the QX200 Droplet Reader. For each ddPCR plate run, one negative control (RNA/DNA-free water; Life Technologies) per column of samples and one positive control (genomic DNA extracted from a sample known to contain *N. spumigina*) per plate were included. When samples were too concentrated (lacking clear negative droplets), these were diluted (1:10) with RNA/DNA-free water (UltraPureTM, Life Technologies, CA, USA) and re-analysed. The results were then converted to copies per gram using the following formula: number of copies per μ l \times 22 μ l (the initial volume of the PCR reaction) \times 33.333 (proportion of total eluted DNA from the extraction step; 100 μ l / 3 μ l) \times 10 (if 1:10 dilution was used in PCR reaction) \times 0.22 (sediment wet weight in grams).

Metabarcoding analysis

To characterise the bacterial prokaryote and eukaryote composition over the 24 weeks of storage, the V3-V4 regions of the prokaryote 16S rRNA gene and the V4 region of the eukaryote 18S rRNA gene were amplified by PCR, using the prokaryote-specific primers 341F: 5'-CCT ACG GGN GGC WGC AG-3' and 805R: 5'-GAC TAC HVG GGT ATC TAA TCC-3 (Herlemann et al. 2011; Klindworth et al. 2013) and eukaryote primers Uni18SF: 5'-AGG GCA AKY CTG GTG CCA GC-3' and Uni18SR: 5'-GRC GGT ATC TRA TCG YCT T-3' (Zhan et al. 2013), each including adapter tails for dual-index sequencing (Kozich, Westcott, Baxter, Highlander and Schloss 2013). PCR reactions were performed in 25 μ l volumes for each sample with the reaction mixture

containing 12 μl of MyFi Mix, 5 μl of RNA/DNA free water (UltraPure, ThermoFisher Scientific), 1 μl each of forward and reverse primer (10 μM) and 6 μl of template DNA. The reaction mixture was held at 95 °C for 1 min followed by 35 cycles of 95 °C for 15 s, 50 °C for 15 s, 72 °C for 15 s, with a final extension step at 72 °C for 7 min. Negative PCR controls and DNA extraction controls were run alongside the samples for each PCR analysis. PCR products from the triplicates were pooled and 25 μl of the pooled product was cleaned and normalised using SequalPrep Normalization plates (ThermoFisher Scientific, USA). This resulted in a cleaned DNA concentration of $\sim 1 \text{ ng } \mu\text{l}^{-1}$. Samples were sent to Auckland Genomics Facility for sequencing on an Illumina Miseq™ platform (step 6 in Fig. 1). The samples were prepared for sequencing following the Illumina 16S metagenomics library prep manual with the exception that, after the indexing PCR, 5 ml of each sample (including water samples acting as sequencing blank) was pooled and a single clean-up was undertaken on the pool instead of samples being individually cleaned. The library pool quality control was undertaken on a bioanalyser before the library was diluted to 4 nM, denatured and diluted to a final loading concentration of 7 pM with a 15% PhiX spike. Raw sequence reads are deposited in the NCBI short read archive under the accession number: PRJNA780583.

Bioinformatics

Bioinformatic pipelines for both the 16S and 18S rRNA genes were identical unless otherwise stated. Primers were removed from the raw reads using *cutadapt* with one mismatch allowed (Martin 2011) and subsequently processed using the *DADA2* package (Callahan et al. 2016) within R (R Core Team 2018). Reads were truncated (230 and 228 base pairs (bp) for forward and reverse reads, respectively) and filtered with a maximum number of “expected errors” (maxEE) threshold of two forward reads and four reverse reads; reads not matching this criterion were discarded from further analysis. A parametric error matrix was constructed, based on the first 10^8 bp of the sequences. Sequence variants for the forward and reverse reads were inferred, based on this error matrix following sequence dereplication. Singletons were discarded and remaining paired-end reads were merged with a maximum mismatch of 1 bp and a required minimum overlap of 10 bp. The chimeric sequences were removed using the *removeBimeraDenovo* command within the *DADA2* package.

The resulting Amplicon Sequence Variants (ASVs) were used for taxonomic classification against the SILVA 138 (Pruesse et al. 2007) and PR2 (Guillou et al. 2013) databases for the 16S and 18S rRNA gene datasets, respectively. We chose to use the PR2 database for the classification of eukaryotes because it provides better classification of protists, which are likely to be the most important and abundant organisms in these samples, compared to other available databases, for example, SILVA.

The sequences were classified, based on the *rdp* classifier (Wang et al. 2007) with a bootstrap of 50 to allow classifications at higher taxonomic levels. The results were combined into a *phyloseq* object (McMurdie and Holmes 2013) and sequences assigned as eukaryotes, chloroplasts and mitochondria were removed from the 16S rRNA dataset, while bacterial sequences were removed from the 18S rRNA dataset. Read numbers for ASVs found in negative control samples were removed from the dataset via subtraction. For comparisons between samples, subsampling to an even depth was undertaken: 13,000 and 8,000 reads for bacterial and eukaryotic datasets, respectively.

Microcystis-specific droplet digital PCR assay

To investigate abundances of the cyanobacterial genus *Microcystis* in the non-spiked samples, a ddPCR assay, targeting the 16S rRNA gene, was used. Absolute concentrations of the 16S rRNA gene were measured in all samples using a BioRad QX200 ddPCR system, Cyanobacteria-specific primers (MICR-184 forward primer, 5'-GCC GCR AGG TGA AAM CTA A-3'; MICR-431 reverse primer, 5'-AAT CCA AAR ACC TTC CTC CC-3'; Neilan et al. 1997) and a *Microcystis*-specific probe (MICR-228 probe FAM-AAG AGC TTG CGT CTG ATT AGC TAG T-BHQ-1; Rinta-Kanto et al. 2005). Each ddPCR reaction included 450 nM (1 μl) of each primer and probe, 10 μl of $2 \times$ BioRad ddPCR Probe Supermix, 2 μl DNA and sterile water for a total reaction volume of 22 μl . PCR set-up and droplet reading settings and equations used to determine concentrations were as described above for the *nodularin* assay. The PCR conditions were as follows: hold at 95 °C for 10 min, 40 cycles of 94 °C for 30 s and 60 °C 1 min and a final enzyme deactivation step at 98 °C for 10 min. All samples were diluted (1:100) with RNA/DNA-free water (UltraPure™, Life Technologies, CA, USA). The plate was then analysed on the QX200 Droplet Reader. For each ddPCR plate run, one negative control (RNA/DNA-free water; Life Technologies) per column of samples and one positive control (genomic DNA extracted from a sample known to contain *Microcystis*) per plate were included. The results were then converted to copies per gram using the following formula: number of copies per μl $\times 22 \mu\text{l}$ (the initial volume of the PCR reaction) $\times 50$ (proportion of total eluted DNA from the extraction step; 100 μl / 2 μl) $\times 100$ (dilution used in PCR reaction) $\times 0.22$ (sediment wet weight in grams).

Statistical analysis

Statistical analysis of the data was undertaken within R (R Core Team 2018). Linear regression was used to assess the relationship between concentrations of the *ndaF* gene (copies per gram as determined with ddPCR) and storage time (weeks), for each treatment (spiking with 100 ng, 200 ng and cell lysate). A separate linear regression was used to assess the relationship between

concentrations of *Microcystis*-specific 16S rRNA gene (copies per gram as determined with ddPCR) and storage time (weeks).

Community diversity and composition analyses were conducted for both bacteria (16S rRNA) and eukaryote (18S rRNA) metabarcoding datasets from the non-spiked samples. Observed ASV richness, Shannon and Simpson Indices were considered and linear regressions were used to assess the relationship between each diversity measure and storage time (weeks). Bray-Curtis dissimilarities of individual samples were compared using Principal Components Analysis (PCoA), plotted using the `plot_ordination` function within the `phyloseq` package in R (McMurdie and Holmes 2013). Compositions of bacterial phyla, eukaryotic phyla, cercozoan classes and cyanobacterial genera for each week of storage were visualised using the `plot_bar` function within `phyloseq`. Relative abundances were calculated for each week of storage by summing the three replicates. Statistical differences in taxonomic composition grouped by storage time (weeks) were tested via permutational multivariate analysis of the Bray-Curtis dissimilarities, using the `adonis` function within the `vegan` package (Oksanen et al. 2019).

Results

Changes in copy numbers in spiked DNA and cell lysate samples

Copy numbers of the *ndaF* gene did not increase or decline over the 24 weeks of storage for the 100 ng and 200 ng spiked treatments ($R^2 = 0.067$, $p = 0.11$ and $R^2 = 0.058$, $p = 0.13$, Fig. 2a, b). However, average copy numbers declined in the cell lysate spiked treatment from 315,777 (SD = 90,737, SE = 52,387) copies per gram to 151,360 (SD = 42,334, SE = 24,442) at week 12 and 40,325 (SD = 37,542, SE = 21,675) by week 24, equating to 52% and 87% reductions from initial yields, respectively. ($R^2 = 0.42$, $p < 0.001$, Fig. 2c).

Shifts in bacterial community composition

A total of 17,548 ASVs were reduced to 13,096 ASVs (ranging from 1,987–2,569 ASVs across 37 samples) after quality control, taxonomic filtering and rarefaction. Unassigned sequences at phylum level were retained and comprised 4.1% of remaining ASVs. Over the 24-week period, mean observed bacterial richness decreased from 2,364 (SD = 73.1, SE = 42.2, $n = 3$) to 2,081 (SD = 62, SE = 35.8, $n = 3$). There was a weak negative relationship between observed richness and sampling time ($R^2 = 0.23$, $p = 0.0025$, Suppl. material 1: Fig. S1). In contrast, Shannon and Simpson Indices did not change significantly over the study period ($R^2 = 0.041$, $p = 0.23$, $R^2 = 0.007$, $p = 0.61$, respectively, Suppl. material 1: Fig. S1).

Bacterial phyla comprising > 10% of retained sequences (summed across replicates) were Proteobacteria,

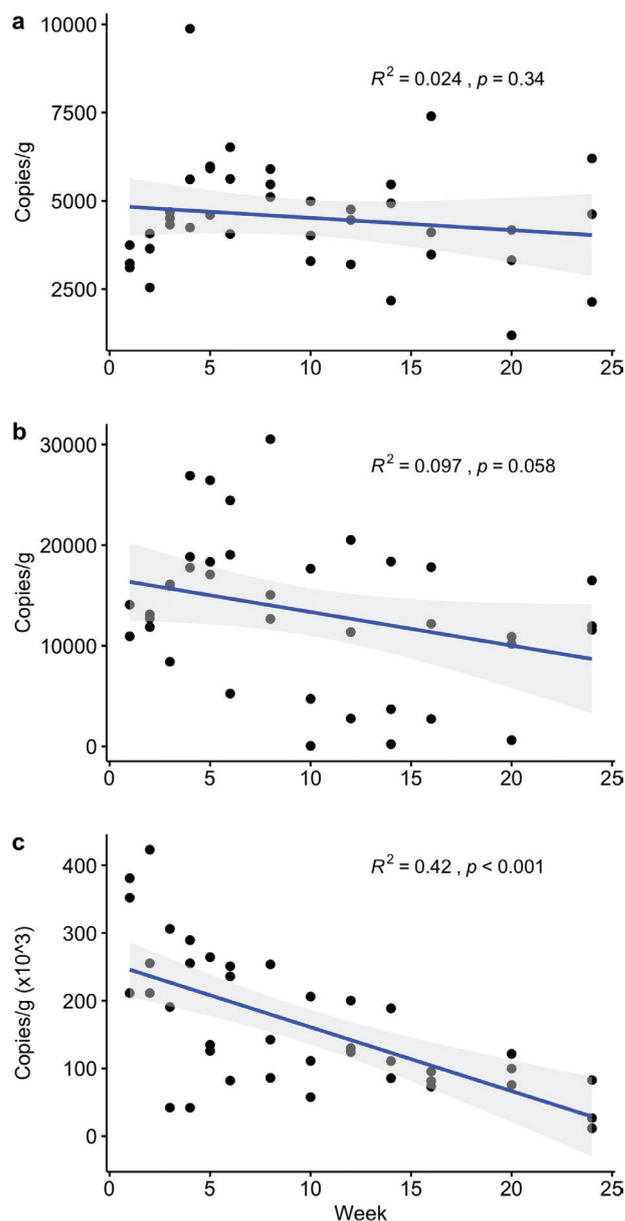


Figure 2. Linear regression between concentration (copies per g of wet weight) and storage time for the *ndaF* gene. Surface sediment samples were spiked with 100 ng (a) or 200 ng (b) of DNA or cell lysate (c). Samples were stored chilled (4 °C) in the dark and sampled at weekly to bi-weekly intervals and assessed using droplet digital PCR over a 24-week period.

Verrucomicrobiota, Bacteroidota (Fig. 3). The PCoA analysis showed a clear shift in the bacterial community during the experiment and replicates clustered closely together (Fig. 4a). Multivariate analysis identified significant differences in the bacterial composition across 24 weeks ($F = 1.366$, DF = 13, 23, $R^2 = 0.435$, $p = 0.001$; Suppl. material 2: Table S1). Changes in phyla relative abundances were visible with a noticeable decrease in cyanobacteria, which dropped from 9.6% at the beginning of the experiment to 0.4% at week 8 and $\leq 0.3\%$ from week 10 to 24 (Fig. 3). Cyanobacteria taxa predominantly comprised the genus *Microcystis* (57–99% of cyanobacterial sequence reads between weeks 0 and 14 and

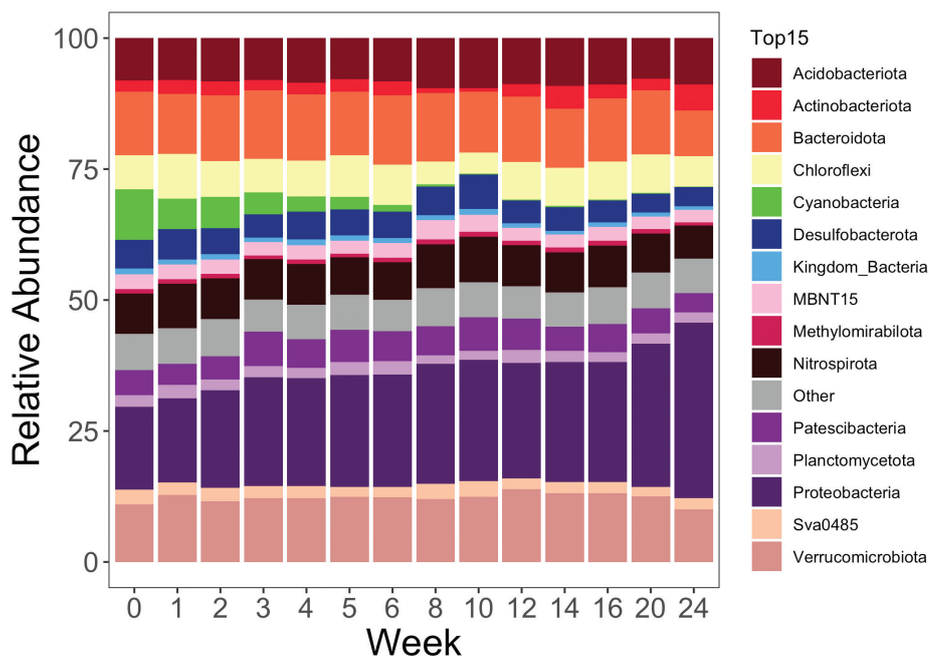


Figure 3. Bacterial phyla (relative abundance of 16S rRNA gene sequences, summed across replicates, $n = 3$) of non-spiked samples over 24 weeks of chilled (4 °C, dark) storage determined using metabarcoding. Sequences were rarefied to 13,000 reads. The 15 most abundant phyla are shown with the remainder grouped as ‘Other’.

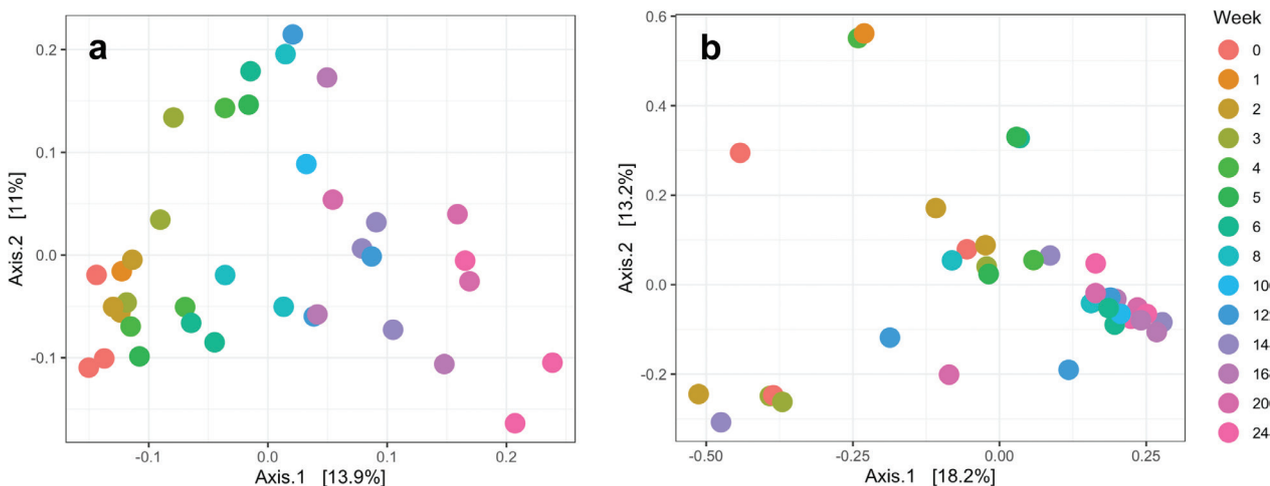


Figure 4. Principal Coordinates Analysis (PCoA) of bacterial 16S rRNA gene (a) and eukaryotic 18S rRNA gene (b) metabarcoding data using Bray-Curtis dissimilarity coloured by storage time (weeks).

24–57% between weeks 16 and 24) and read numbers decreased from 3,738 to 115 between weeks 0 and 8 (Suppl. material 1: Fig. S2). Proteobacteria noticeably increased from 15% at week 0 to 33% at week 24 (Fig. 3). Within this phylum, Gammaproteobacteria were the dominant class ($\geq 84\%$) throughout the 24 weeks and, together with Alphaproteobacteria, these comprised $\geq 99\%$ of sequence reads. Of the Gammaproteobacteria, Burkholderiales (47–73%) was consistently dominant throughout the experiment and Xanthomonadales increased from week 12 to dominate the Proteobacteria sequence counts in week 24 (48%; Suppl. material 1: Fig. S3).

Microcystis 16S rRNA gene copies declined in a non-linear fashion over time ($R^2 = 0.78$, $p < 0.001$; Fig. 5). From week 0, average gene copies (2.2 mil-

lion, SD = 725,736, SE = 419,004, $n = 3$) declined by 60% to 911,150 after 3 weeks and 99% to 12,736 after 24 weeks.

Shift in eukaryote community structure

Linear regression of eukaryote taxa in non-spiked samples did not show any relationship between observed richness and time, but there was a very weak positive relationship when using Shannon and Simpson Indices ($R^2 = 0.2$, $p = 0.0053$ and $R^2 = 0.22$, $p = 0.0033$, respectively, Suppl. material 1: Fig. S1). Mean Shannon Index increased by 20% from 2.75 at week 0 to 3.29 at week 24 and mean Simpson Index increased 13% from 0.77 at week 0 to 0.87 at week 24.

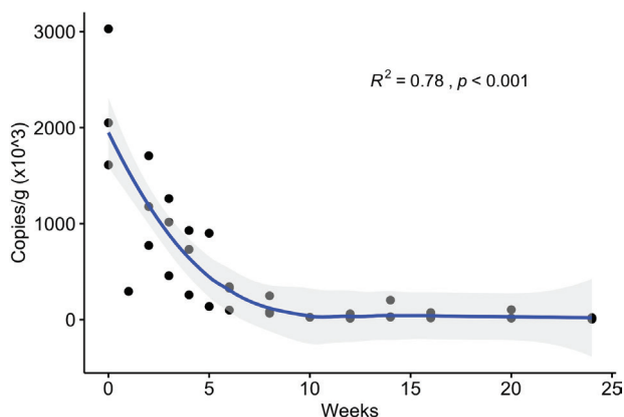


Figure 5. Concentrations (copies per g of wet weight) of the *Microcystis* 16S rRNA gene in non-spiked samples over 24 weeks of chilled (4 °C, dark) storage (Loess smoothed trend-line with 95% confidence intervals).

A total of 3,462 ASVs decreased to 2,736 ASVs (ranging from 108–369 ASVs across 37 samples) following quality control, taxonomic filtering and rarefaction. Unassigned sequences at phylum level were retained and comprised 13.4% of remaining ASVs. Eukaryote phyla comprising > 20% of retained sequences (summed across replicates) were Annelida, Arthropoda, Cercozoa and Dinoflagellata (Fig. 6). The PCoA analysis showed convergence of the eukaryotic community in samples stored for longer periods (Fig. 4b). Permutational multivariate analysis showed significant differences ($F = 1.314$, $DF = 13, 23$, $R^2 = 0.426$, $p = 0.012$; Suppl. material 2: Table S1) in eukaryotic composition across 24 weeks of storage. Changes in the relative abundances of eukaryotic phyla were visible with a noticeable increase in cercozoan sequences, which increased from 2.95% at the on-

set of the experiment to 53.4% at week 10 and 30.5% by week 24 (Fig. 6). Relative abundance of phylum Annelida and Arthropoda fluctuated across the 24 weeks of storage, ranging from 0.03–59% for the first 5 weeks and reducing to 0–43% for the remainder of the experiment. Within Cercozoa, Sarcomonadea and Thecofilosea were the dominant classes throughout the experiment (Fig. 7). Sarcomonadea comprised > 50% of reads for all weeks, except weeks 5, 6, 8 and 14. Thecofilosea increased in relative abundance from 9% at week 0 to 54% at week 6 and remained above 34% for the rest of the experiment (excluding week 20). Haptotaxida were the dominant order of Annelida (> 99% across all time points) and Calanoida was the dominant order of Arthropoda (> 99%) for nine of the thirteen time points (Suppl. material 1: Fig. S4).

Discussion

Freezing samples for preservation until analysis is ideal for maintaining sedDNA integrity. However, freezing whole sediment cores can be logistically challenging, makes subsampling difficult and can compromise core stratigraphy. Chilled storage of sediments is a simple and cost-effective way to maintain samples at conditions similar to their source environment. However, at temperatures above freezing, biological activity can continue. Understanding how long-term chilled storage affects the concentration and composition of sedDNA is important to aid in the interpretation of paleogenomic data. As evidenced by our study, chilled sediment stored for more than 5 weeks has implications for DNA yields and community composition.

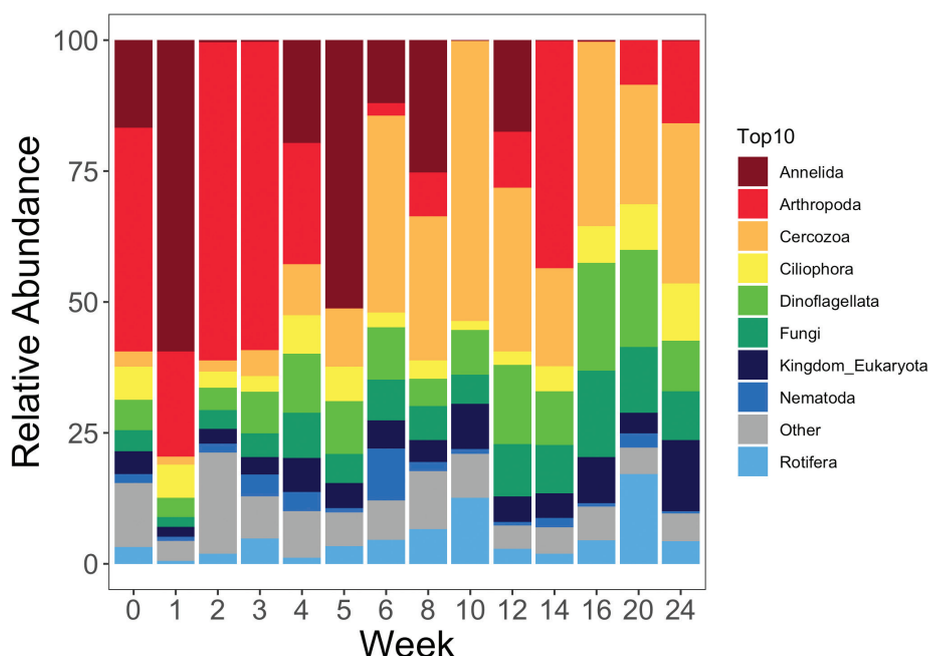


Figure 6. Eukaryote phyla (relative abundance of 18S rRNA gene sequences, summed across replicates, $n = 3$) of non-spiked samples over 24 weeks of chilled (4 °C, dark) storage determined using metabarcoding. Sequences were rarefied to 8,000 reads. The 15 most abundant phyla are shown with the remainder grouped as ‘Other’.

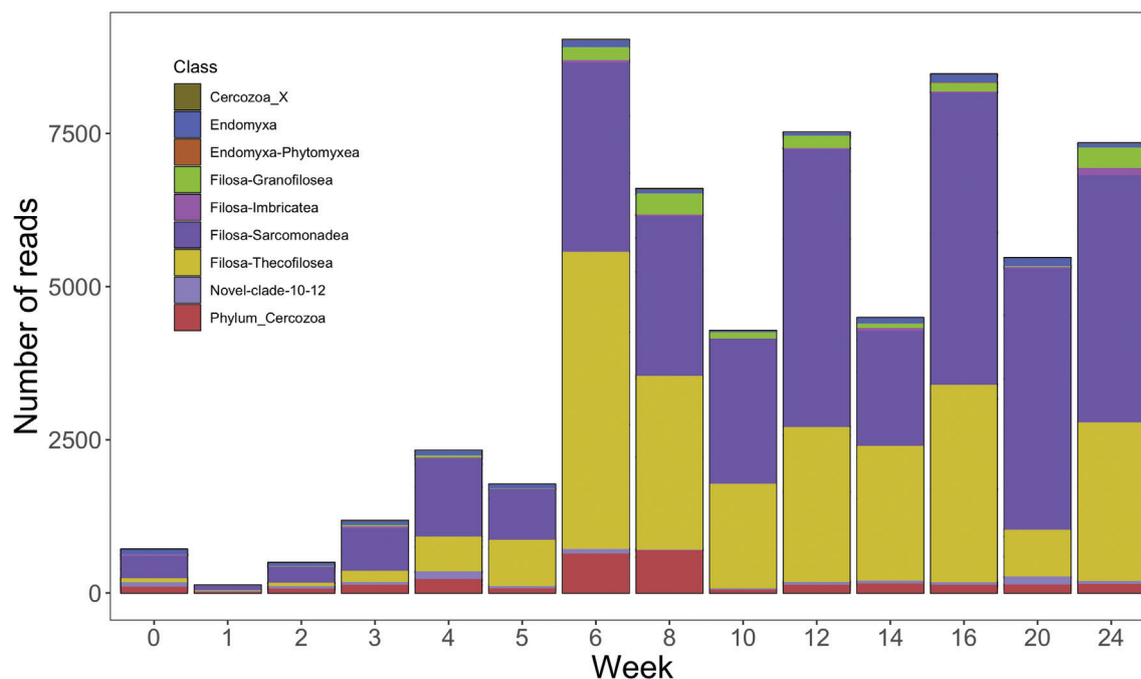


Figure 7. Cercozoa composition and number of reads as determined by metabarcoding of the eukaryotic 18S rRNA of non-spiked samples over 24 weeks of chilled (4 °C, dark) storage.

Storage effects on sedimentary DNA

We found differential effects of chilled storage on DNA yields from sediments artificially spiked with purified DNA and DNA from freshly lysed cells. Consistent with our first hypothesis, we observed no change to DNA yields for purified DNA when spiked at either 100 or 200 ng per sample during the 24 weeks of storage. Similar results have been found in previous studies with little change observed in recovered environmental DNA (eDNA) from soil samples stored for 14 days (Lauber et al. 2010). Our results suggest the purified extracellular DNA quickly became sediment-bound, likely reducing DNA degradation. This is in agreement with experimental studies that found: *i*) when DNA was added to non-sterile soils, 80% of it bound to soil within 1 hr (Blum et al. 1997) and *ii*) maximum DNA absorption on to quartz sand was reached after 2 h (Lorenz and Wackernagel 1987). Once DNA molecules are bound to sediment, they can be difficult to cleave off (Romanowski et al. 1991).

Consistent with our second hypothesis, yields of DNA from freshly lysed cells declined steadily across the 24 weeks, with concentrations reducing 87% over the experiment. Compared to the purified DNA-spiked samples, higher starting volumes of DNA in the lysate-spiked samples may have led to an overabundance of DNA versus sediment binding sites. Additionally, cell lysate also contains cellular nucleases that may contribute to DNA degradation before some molecules become sediment bound. Bacteria surviving in chilled sediments can also produce or take advantage of nucleases in the sediment matrix, allowing them to feed on depurinated DNA and subsequent degradation products (Wasmund et al. 2021). Proteobacteria, which increased in abundance during the experi-

ment (see discussion below) are known to have genes that enable consumption of DNA (Finkle and Kolter 2001) and this could be another possible mechanism for reduction in DNA of lysate-spiked samples. These cell lysate samples represented a realistic scenario of a senescing algal bloom, where cells gradually die and settle on the sediment surface. Our results highlight the need for rapid sampling, particularly for eutrophic lakes where blooms are a regular occurrence.

Storage effects on biological communities

Biological communities, detected in sediment samples, can be a combination of living cells, dormant cells and extracellular DNA (Taberlet et al. 2012). Some bacterial and eukaryotic microorganisms can survive in low energy, anoxic conditions and proliferate by feeding on each other or nutrient compounds in the sediment matrix (Koizumi et al. 2003; Compte-Port et al. 2020; Pearman et al. 2021a). Based on short-term storage studies, we hypothesised that maintaining samples in chilled and dark storage conditions would result in limited shifts in biological communities because the environment is similar to that from which they were collected. The number of bacterial taxa (observed ASV richness) showed a weakly associated decrease with time, but there was no significant change in either diversity index (Shannon and Simpson). This aligns with Lauber et al. (2010), who found similar phylogenetic structure and diversity of soil bacterial communities across storage temperatures.

However, there were several exceptions to this pattern with the most notable being the cyanobacterial genus *Microcystis*; a bloom-forming microorganism able

to produce potent hepatotoxins (Harke et al. 2016). The relative abundance of this genus halved by week 3. To confirm that the reduction of 16S rRNA sequence abundances in the metabarcoding data was not due to sequencing bias, which can occur when community composition changes over time (Lee et al. 2012; Pochon et al. 2013), a *Microcystis*-specific ddPCR assay was applied and an equivalent decrease in copy numbers was detected. Our results also showed a shift in eukaryotic community composition around 5–6 weeks of storage. This was mainly driven by an increase in Cercozoa; an ubiquitous phylum of single-celled eukaryotes that are common in sediment and functionally and morphologically diverse, including both autotrophs and free-living predators (Bass and Cavalier-Smith 2004; Dumack et al. 2020). The observed change in composition indicates the persistence of live organisms in sediment samples at 4 °C and suggests that they may feed on other taxa and proliferate over time. Although, to the best of our knowledge, Cercozoa have not specifically been shown to feed on *Microcystis*, some constituents can be bacterivorous or omnivorous (Flues et al. 2017; Dumack et al. 2020). Other unicellular eukaryotes have been shown to feed on *Microcystis* blooms in both field and laboratory experiments (Zhang et al. 2008; Van Wichelen et al. 2010). The detected increase in Cercozoa suggests a biotic mechanism for the decrease in *Microcystis* sequences by preferential grazing. Unlike some Cyanobacteria, *Microcystis* does not produce akinetes, which can be larger than vegetative cells and have thickened cell walls and this may make *Microcystis* cells more vulnerable to grazing. For studies investigating the history of Cyanobacteria proliferations (for example, Pal et al. 2015 and Monchamp et al. 2016), their abundance could be under-represented if samples are chill-stored for more than two weeks.

Gammaproteobacteria were the dominant class of proteobacterial phyla throughout the experiment, with the order Xanthomonadales notably increasing in abundance after week 12. Similar patterns of increased Gammaproteobacteria abundance were found in experimental liquid microcosms (Flues et al. 2017), where Gammaproteobacteria were positively affected by Cercomonads grazing on the bacterial community and the authors attributed this to the distinct changes in bacterial community composition. We suggest a similar positive effect on Xanthomonadales occurred in our samples, as members of this order are reported to possess anti-predation cell structures that confer a competitive advantage over other bacteria (Bayer-Santos et al. 2019).

Amongst the eukaryote data, there were several other notable patterns including variable numbers of Annelida sequences. As larger organisms, they are likely to have a patchier distribution and it is probable they were living in the sediment samples at the time of collection. Given that there were no discernible trends, we speculate that uneven numbers of them were contained in the sediment during the subsampling process. Calanoida, commonly

found as zooplankton, were the predominant order of arthropod detected in most samples and their abundance was also uneven. It is unlikely these arthropods were alive in the sediment samples; therefore, most of the DNA we detected was likely extracellular. The patchy nature of these organisms highlights the need for larger sediment samples when working with larger organisms for precise biodiversity characterisation.

Implications for sediment core archives

Considering the clear changes in bacterial and eukaryotic community compositions observed in this study, we recommend subsampling and freezing sediment core samples, particularly those near the surface where microbes may be more active, within 2–4 weeks of collection and chilled storage. Immediate freezing of subsamples within 1–2 weeks is recommended for studies where low abundance or rare taxa are important and for assessments of microorganisms (bacteria, cyanobacteria and phytoplankton). Time-frames for broad-scale community structure assessments may still be possible up to 5 weeks after sampling with chilled storage.

Our analysis of surface sediment is relevant for deeper core deposits as these surface processes (such as DNA binding, degradation or consumption) are part of the wider burial process. However, the small (0.22 g) samples were stored in isolation rather than as whole core samples and may not be affected by processes occurring in a larger sediment matrix, such as microbial recruitment from the adjacent sediment. Our interpretation is limited to lakes of similar trophic level and sediment type. The interplay of sediment binding sites and lower algal productivity in oligotrophic lakes may produce differing scenarios and further studies are required.

Conclusions

In conclusion, we showed that sedDNA inventories are affected by long-term chilled storage (up to 6 months), with the potential loss of DNA yield and changes in microbial community composition. Considering the reduction of cyanobacteria and concurrent increase in cercozoan taxa, we suggest that protist grazing activity can continue in chilled storage conditions and strongly recommend prompt subsampling and freezing of sediment cores for paleogenomic studies.

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Competing interest statement

The authors declare there are no competing interests.

Data resources

Raw sequence reads are deposited in the NCBI short read archive under accession number PRJNA780583.

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Supplementary material 1 Figures S1–S4

Author: Katie A. Brasell, Xavier Pochon, Jamie Howarth, John K. Pearman, Anastasija Zaiko, Lucy Thompson, Marcus J. Vandergoes, Kevin S. Simon, Susanna A. Wood

Data type: Images (pdf file)

Explanation note: Figure S1. Linear regressions with fitted Loess smoothed trend lines (blue) and 95% confidence intervals (grey), assessing the relationship between three different diversity measures at ASV level; Observed richness (a, d), Shannon Index (b, e) and Simpson Index (c, f) and time (weeks) for samples stored at 4 °C in the dark. Bacteria (a-c) and eukaryotes (d-f). Data were derived from the bacterial 16S rRNA and eukaryotic 18S rRNA metabarcoding analysis. Figure S2. Cyanobacteria composition and number of reads as determined by metabarcoding of the bacterial 16S rRNA of non-spiked samples over 24 weeks of chilled (4 °C, dark) storage. Note each week has 3 replicates, except week 1 and 10 that have single replicates and week 5 has two replicates. Figure S3. Raw proteobacteria sequence abundance coloured by order. Bars show sum of replicates per week. Figure S4. Raw sequence abundance for phylum Annelida (left) and Arthropoda (right) coloured by order. Bars show sum of replicates per week.

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Supplementary material 2**Table S1**

Author: Katie A. Brasell, Xavier Pochon, Jamie Howarth, John K. Pearman, Anastasija Zaiko, Lucy Thompson, Marcus J. Vandergoes, Kevin S. Simon, Susanna A. Wood

Data type: Table (pdf file)

Explanation note: Permutational multivariate analysis of Bray-Curtis dissimilarity for bacterial and eukaryote community composition (determined via 16S rRNA & 18S rRNA gene sequencing, respectively) across 24 weeks of chilled storage.

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