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Data Article

# Seasonal Ely Copper Mine Superfund site shotgun metagenomic and metatranscriptomic data analysis



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# ABSTRACT

High throughput sequencing data collected from acid rock drainage (ARD) communities can reveal the active taxonomic and functional diversity of these extreme environments, which can be exploited for bioremediation, pharmaceutical, and industrial applications. Here, we report a seasonal comparison of a microbiome and transcriptome in Ely Brook (EB-90M), a confluence of clean water and upstream tributaries that drains the Ely Copper Mine Superfund site in Vershire, VT, USA. Nucleic acids were extracted from EB-90M water and sediment followed by shotgun sequencing using the Illumina NextSeq platform. Approximately 575,933 contigs with a total length of 1.54 Gbp were generated. Contigs of at least a size of 3264 (N50) or greater represented 50% of the sequences and the longest contig was 488,568 bp in length. Using Centrifuge against the NCBI "nt" database 141 phyla, including candidate phyla, were detected. Roughly 380,000 contigs were assembled and ~1,000,000 DNA and ~550,000 cDNA sequences were identified and function-

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ally annotated using the Prokka pipeline. Most expressed KEGG-annotated microbial genes were involved in amino acid metabolism and several KEGG pathways were differentially expressed between seasons. Biosynthetic gene clusters involved in secondary metabolism as well as metal- and antibiotic-resistance genes were annotated, some of which were differentially expressed, colocalized, and coexpressed. These data can be used to show how ecological stimuli, such as seasonal variations and metal concentrations, affect the ARD microbiome and select taxa to produce novel natural products. The data reported herein is supporting information for the research article "Characterization of an acid rock drainage microbiome and transcriptome at the Ely Copper Mine Superfund site" by Giddings et al. [1].

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# **Specifications Table**

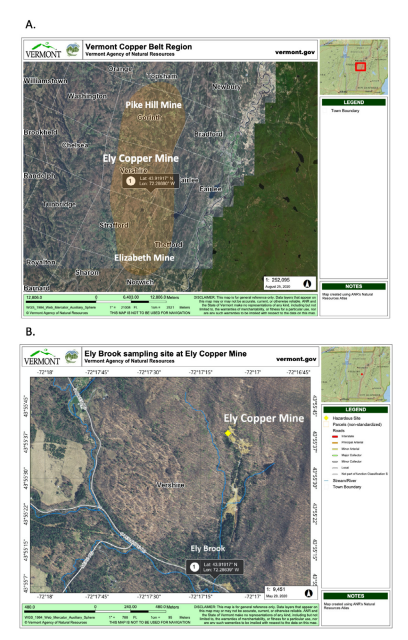
Subject	Microbial Ecology, Genomics and Molecular Biology
Specific subject area	Metagenomics
Type of data	Tables, figures, raw data
How data were acquired	Shotgun metagenomic and metatranscriptomic sequence data were acquired using an Illumina NextSeq500 instrument. Centrifuge was used to perform a read-based taxonomic analysis of metagenomic data. Prokka was used to detect and functionally annotated open reading frames. The predicted amino acid sequence was searched against Swiss-Prot database using DIAMOND. KEGG orthology annotations were predicted for open reading frames. All differential and statistical analyses on taxonomic summaries were performed in edgeR [2]. BacMet [3], antiSMASH 5.0 [4], ARTS version 2.0 [5] databases were used to annotate genes.
Data format	Annotated data, Bray-Curtis dissimilarity matrices, Non-metric multidimensional scaling (NMDS) plots, principal component analysis (PCA) plots, heat map and hierarchal clustering, raw count data, and gradient plots.
Parameters for data collection	Seasonal environmental water and sediment samples were collected and sequenced. Five water and three sediment samples from summer as well as three sediment samples from winter.
Description of data collection	Shotgun metagenomic and metatranscriptomic sequencing was performed using an Illumina NextSeq500 instrument.
Data source location	Sediment (July 28th, 2017 and January 14th, 2018) and water (July 14th, 2017 and July 28th, 2017) samples were collected 90 m upstream from the mouth of Ely Brook (EB-90M) at Ely Copper Mine, Vershire, VT, USA (43°55'9″ N, 72°17'11″ W).
Data accessibility	Data are shown in this article. Raw metagenomic and metatranscriptomic data have been deposited in the Sequence Read Archive of the National Center for Biotechnology Information (BioProject identifier, PRJNA540505). Taxonomic and functional annotations as well as normalized count data used for all analyses are available in a public repository: Repository name: FigShare Data identification number: 10.6084/m9.figshare.c.4864863 Direct URL to data: https://doi.org/10.6084/m9.figshare.c.4864863
Related research article	L-A. Giddings, G. Chlipala, K. Kunstman, S. Greene, K. Morillo, K. Bhave, H. Peterson, H. Driscoll, M. Maienschein-Cline, Characterization of an acid rock drainage microbiome and transcriptome at Ely Copper Mine Superfund site, PLoS One, 15(8) (2020) e0237599. https://doi.org/10.1371/journal.pone.0237599

# Value of the Data

- This is the first characterization of an acid rock drainage (ARD) metagenome and transcriptome within the Vermont copper belt region, USA, which is comprised of Ely Copper Mine, Elizabeth Mine, and Pike Hill Copper Mine.
- The metagenomic data provide seasonal taxonomic profiles of the microbial diversity in the sediment and water of EB-90M.
- Active taxa in ARD environments are understudied and the metagenomic and metatranscriptomic data provide insight into their seasonal functional roles within these acidic, metal-rich environments.
- These data can be used to perform comparative taxonomic and functional analyses with other ARD metagenomes.
- These data can be used to bioprospect enzymes that can be exploited for the bioremediation of metal polluted environments.
- These data can be used to identify novel genes encoding proteins involved in the production of bioactive secondary metabolites, which can be used for pharmaceutical and industrial applications.

# 2. Data Description

Ten water and six sediment samples at Ely Brook (EB-90M) (Fig. 1), Ely Copper Mine Superfund site were collected in July 2017 and January 2018. Shotgun metagenomic sequencing of nucleic acids extracted from water and sediment samples generated  $\sim$ 31,545,991 reads with an average length of 147 bp and a total length of 1.54 Gb for 11 samples. Samples of the same sample type (i.e., water or sediment) or season (i.e., summer or winter) were treated as biological replicates. Summer water samples were denoted as July\_Water1, July\_Water2, July\_Water3, July\_Water4, July\_Water5. Summer sediment samples were denoted as July\_Sed1, July\_Sed2, and July\_Sed3. Winter sediment samples were denoted as Jan\_Sed1, Jan\_Sed2, and Jan\_Sed3. All winter water samples (five samples) did not yield viable sequencing data. Of the remaining 11 samples,  $\sim$ 12 Gb of data (50 M clusters) were produced per sample with an average of 25,181,359 reads per sample over a range of 8,657,966 and 44,323,783 reads for both metagenomic and metatranscriptomic data. Contigs of  $\geq$  3264 bp (N50) represented 50% of data and the longest contig was 488,568 bp in length. Using Centrifuge [6] to perform read-based taxonomic annotation, 141 distinct phyla were annotated, including candidate phyla (Table 1). Taxonomic differences across season and sample type were observed by NMDS and PCA analyses of normalized count data (i.e., counts per million) between the bacteria, archaea, and fungi in samples as well as molecule types (Figs. 2–8). Differences between molecule type (i.e., DNA or RNA) across sample type and season were assessed by multivariate principal component analyses (PCA) (Fig. 9). Using Prokka-annotated open reading frames [7], Kyoto Encyclopedia of Genes and Genomes (KEGG) reference pathways [8] were annotated and quantified (Table 2). Significantly differentially expressed KEGG pathways and genes in winter versus summer were defined as having winter/summer RNA p-values  $\leq 0.05$  for the interaction of season and molecule type followed by false discovery rate (FDR) corrections [9] (q-values)  $\leq 0.05$ (Figs. 10–12). Secondary metabolite gene clusters (Table 3), metal resistance genes (Table 4), and antibiotic resistance genes were identified (Table 5). Approximately 288 metal resistance genes were differentially expressed between winter and summer seasons (Fig. 13). Furthermore, some of these genes were colocalized and coexpressed with genes involved in secondary metabolism (Table 6; Figs. 14–18).



**Fig. 1.** *Vermont copper belt.* A) Map of Vermont copper belt (highlighted in yellow), which includes Ely Copper Mine (sampling site), Pike Hill Mine, and Elizabeth Mine. B) Map of Ely Brook sample site, which drains Ely Copper Mine. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

#### Table 1

Taxonomic annotation. List of 141 unique phyla across water and sediment metagenomic samples at EB-90 M. sk, superkingdom; k, kingdom; p, phylum. Incertae sedis represents kingdoms that have not been assigned.

Unique phyla across water and sediment metagenomic sam	nples
sk_Archaea;kArchaea incertae sedis;pArchaea	sk_Bacteria;kBacteria incertae
incertae sedis	sedis;p_Planctomycetes
sk_Archaea;kArchaea incertae sedis;pCandidatus	sk_Bacteria;kBacteria incertae
Korarchaeota	sedis;p_Proteobacteria
sk_Archaea;kArchaea incertae sedis;pCandidatus	sk_Bacteria;kBacteria incertae
Micrarchaeota	sedis;pSpirochaetes
sk_Archaea;kArchaea incertae sedis;pCandidatus	sk_Bacteria;k_Bacteria incertae
Nanohaloarchaeota	sedis;p_Synergistetes
sk_Archaea;k_Archaea incertae sedis;pCandidatus	sk_Bacteria;k_Bacteria incertae sedis;p_Tenericutes
Parvarchaeota	sk_bacteria, A_bacteria incertae seais, p_renericates
sk_Archaea;kArchaea incertae	sk_Bacteria;kBacteria incertae
sedis;p_Crenarchaeota	sedis;pThermodesulfobacteria
sk_Archaea;kArchaea incertae	sk_Bacteria;k_Bacteria incertae
sedis;p_Euryarchaeota	sedis;pThermotogae
sk_Archaea;k_Archaea incertae	sk_Bacteria;k_Bacteria incertae
sedis;p_Nanoarchaeota	sedis;p_Verrucomicrobia
sk_Archaea;kArchaea incertae	sk_Bacteria;k_Bacteria incertae sedis;p_candidate
sedis;p_Thaumarchaeota	division CPR2
sk_Bacteria;kBacteria incertae	sk_Bacteria;k_Bacteria incertae sedis;p_candidate
sedis;p_Acidobacteria	division CPR3
sk_Bacteria;kBacteria incertae	sk_Bacteria;k_Bacteria incertae sedis;p_candidate
sedis;p_Actinobacteria	division NC10
sk_Bacteria;k_Bacteria incertae sedis;p_Aquificae	sk_Bacteria;k_Bacteria incertae sedis;p_candidate division WPS-2
sk_Bacteria;k_Bacteria incertae	sk_Bacteria;kBacteria incertae sedis;pcandidate
sedis;p_Armatimonadetes	division WWE3
sk_Bacteria;k_Bacteria incertae sedis;p_Bacteria	sk_Eukaryota;kEukaryota incertae
incertae sedis	sedis;pApicomplexa
sk_Bacteria;kBacteria incertae	sk_Eukaryota;kEukaryota incertae
sedis;p_Bacteroidetes	sedis;pBacillariophyta
sk_Bacteria;k_Bacteria incertae	sk_Eukaryota;kEukaryota incertae
sedis;p_Balneolaeota	sedis;p_Bolidophyceae
sk_Bacteria;k_Bacteria incertae sedis;p_Caldiserica	sk_Eukaryota;kEukaryota incertae
;;F;F	sedis;p_Chromerida
sk_Bacteria;k_Bacteria incertae	sk_Eukaryota;k_Eukaryota incertae
sedis;p_Calditrichaeota	sedis;p_Colponemidia
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus	sk_Eukaryota;kEukaryota incertae
Acetothermia	sedis;p_Euglenida
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus	sk_Eukaryota;kEukaryota incertae
Adlerbacteria	sedis;pEukaryota incertae sedis
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus	sk_Eukaryota;k_Eukaryota incertae
Amesbacteria	
	sedis;p_Eustigmatophyceae
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus	sk_Eukaryota;kEukaryota incertae
Atribacteria	sedis;p_Haplosporidia
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus	sk_Eukaryota;kEukaryota incertae
Azambacteria	sedis;p_Phaeophyceae
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus	sk_Eukaryota;kEukaryota incertae sedis;pPicozoa
Beckwithbacteria	
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus	sk_Eukaryota;kEukaryota incertae
Berkelbacteria	sedis;pPinguiophyceae
sk_Bacteria;kBacteria incertae sedis;pCandidatus	sk_Eukaryota;kEukaryota incertae
Campbellbacteria	sedis;p_Xanthophyceae
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus	sk_Eukaryota;kFungi;pAscomycota
Cloacimonetes	
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus	sk_Eukaryota;kFungi;pBasidiomycota
Collierbacteria	
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus	sk_Eukaryota;kFungi;pBlastocladiomycota
Curtissbacteria	-

# Table 1 (continued)

Unique phyla across water and sediment metagenomic sa	amples
sk_Bacteria;kBacteria incertae sedis;pCandidatus Daviesbacteria	sk_Eukaryota;kFungi;pChytridiomycota
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Falkowbacteria	sk_Eukaryota;kFungi;pCryptomycota
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Giovannonibacteria	sk_Eukaryota;kFungi;pEntorrhizomycota
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Gottesmanbacteria	sk_Eukaryota;kFungi;pFungi incertae sedis
S_Bacteria;k_Bacteria incertae sedis;p_Candidatus Gracilibacteria	sk_Eukaryota;kFungi;pMicrosporidia
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Jorgensenbacteria	sk_Eukaryota;kFungi;pMucoromycota
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Kaiserbacteria	sk_Eukaryota;kFungi;pZoopagomycota
K_Bacteria;k_Bacteria incertae sedis;p_Candidatus Kuenenbacteria	sk_Eukaryota;kMetazoa;pAcanthocephala
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Levybacteria	sk_Eukaryota;kMetazoa;pAnnelida
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Magasanikbacteria	sk_Eukaryota;kMetazoa;pArthropoda
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Melainabacteria	sk_Eukaryota;kMetazoa;pBrachiopoda
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Moranbacteria	sk_Eukaryota;kMetazoa;pBryozoa
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Nomurabacteria	sk_Eukaryota;kMetazoa;pChaetognatha
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Omnitrophica	sk_Eukaryota;kMetazoa;pChordata
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Pacebacteria	sk_Eukaryota;kMetazoa;pCnidaria
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Parcubacteria	sk_Eukaryota;kMetazoa;pCtenophora
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Peregrinibacteria	sk_Eukaryota;kMetazoa;pCycliophora
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Roizmanbacteria	sk_Eukaryota;kMetazoa;pEchinodermata
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Saccharibacteria	sk_Eukaryota;kMetazoa;pEntoprocta
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Shapirobacteria	sk_Eukaryota;kMetazoa;pGastrotricha
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Tectomicrobia	sk_Eukaryota;kMetazoa;pGnathostomulida
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Uhrbacteria	sk_Eukaryota;kMetazoa;pHemichordata
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Woesebacteria	sk_Eukaryota;kMetazoa;pKinorhyncha
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Wolfebacteria	sk_Eukaryota;kMetazoa;pMetazoa incertae sedis
sk_Bacteria;k_Bacteria incertae sedis;p_Candidatus Yanofskybacteria	sk_Eukaryota;kMetazoa;pMollusca
sk_Bacteria;k_Bacteria incertae sedis;p_Chlamydiae sk_Bacteria;k_Bacteria incertae sedis;p_Chlorobi	sk_Eukaryota;kMetazoa;pNematoda sk_Eukaryota;kMetazoa;pNematomorpha
sk_Bacteria;k_Bacteria incertae sedis;p_Chloroflexi sk_Bacteria;k_Bacteria incertae	sk_Eukaryota;k_Metazoa;p_Nemertea sk_Eukaryota;k_Metazoa;p_Onychophora
sedis;pChrysiogenetes sk_Bacteria;kBacteria incertae	sk_Eukaryota;kMetazoa;pPlacozoa
sedis;p_Coprothermobacterota sk_Bacteria;k_Bacteria incertae	sk_Eukaryota;kMetazoa;pPlatyhelminthes
sedis;p_Cyanobacteria	

(continued on next page)

#### Table 1 (continued)

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Unique phyla across water and sediment metagenomic samples								
sk_Bacteria;k_Bacteria incertae sedis;p_Deferribacteres	sk_Eukaryota;kMetazoa;pPorifera							
sk_Bacteria;k_Bacteria incertae sedis;pDeinococcus-Thermus	sk_Eukaryota;kMetazoa;pPriapulida							
sk_Bacteria;k_Bacteria incertae sedis;p_Dictyoglomi	sk_Eukaryota;kMetazoa;pRhombozoa							
sk_Bacteria;k_Bacteria incertae sedis;p_Elusimicrobia	sk_Eukaryota;kMetazoa;pRotifera							
sk_Bacteria;k_Bacteria incertae sedis;pFibrobacteres	sk_Eukaryota;kMetazoa;pTardigrada							
sk_Bacteria;k_Bacteria incertae sedis;p_Firmicutes	sk_Eukaryota;kMetazoa;pXenacoelomorpha							
sk_Bacteria;k_Bacteria incertae sedis;p_Fusobacteria	sk_Eukaryota;kViridiplantae;pChlorophyta							
sk_Bacteria;k_Bacteria incertae sedis;pGemmatimonadetes	sk_Eukaryota;kViridiplantae;pStreptophyta							
sk_Bacteria;k_Bacteria incertae sedis;p_Ignavibacteriae	<pre>sk_Viroids;k_Viroids incertae sedis;p_Viroids incertae sedis</pre>							
sk_Bacteria;k_Bacteria incertae sedis;p_Kiritimatiellaeota	sk_Viruses;kViruses incertae sedis;pViruses incertae sedis							
sk_Bacteria;kBacteria incertae sedis;pNitrospirae								

### Table 2

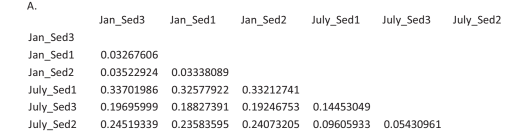
BRITE level 1 annotation statistics. Average percentages of normalized counts that were annotated at BRITE level 1 using the KEGG database.

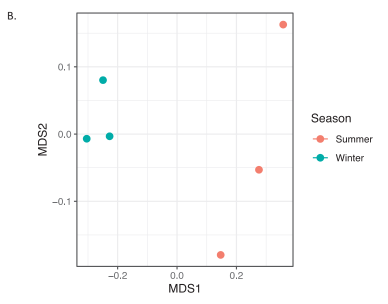
Average BRITE level 1 Observations Across All Sediment Samples	Percentage, %
09100 Metabolism	0.1726149
09120 Genetic Information Processing	0.036885
09130 Environmental Information Processing	0.0244681
09140 Cellular Processes	0.0197084
09150 Organismal Systems	0.0107033
09160 Human Diseases	0.020198
09180 BRITE Hierarchies	0.1912439
09190 Not Included in Pathway or BRITE	0.0202365
Unassigned	0.5039421

#### Table 3

antiSMASH annotation. Summary of the number of genes and gene clusters annotated by antiSMASH 5.0 as well as those that match the Prokka-annotated data.

Total count of contigs	575,933
Total number of contigs annotated by antiSMASH	1589
Total number of contigs not annotated by antiSMASH	574,344
antiSMASH annotated genes	10,579
antiSMASH annotated genes that aligned with PROKKA analyzed data	4977
antiSMASH annotated genes that did not align with PROKKA analyzed data	5602
antiSMASH annotated gene clusters that align with PROKKA analyzed data	1349
antiSMASH annotated gene clusters that did not align with PROKKA analyzed	240
data	
antiSMASH annotated gene clusters that aligned with PROKKA analyzed data	449
and met the criteria of a sum of at least 100 counts across all samples and	
10 counts in three samples	
Annotated gene clusters that remain after filtering by $p$ -interaction value	176
Annotated gene clusters that remain after subsequent filtering by	65
q-winter/summer RNA value	





**Fig. 2.** *Bray-Curtis dissimilarity indices for archaea in sediment.* A) Matrix of dissimilarity indices calculated for genera of archaea in sediment samples using the Bray-Curtis method. 'Sed' = sediment. B) NMDS plot to visualize the dissimilarity between genera of archaea in summer (July\_Sed1, July\_Sed2, and July\_Sed3 in orange) and winter (Jan\_Sed1, Jan\_Sed2, and Jan\_Sed3 in blue) sediment collected at EB-90M. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

#### Table 4

*Metal resistance gene annotation.* Statistics on metal resistance genes identified using the BacMet database. A gene identifier (i.e., gene ID) is defined as a gene symbol plus a number, for example, copR\_X, where X is a number. The eight missing gene IDs that were not expressed, include copR\_13, corC\_121, cusR\_32, czcA\_647, nikE\_38, pstC\_144, ruvB\_54, Int\_122. Differentially expressed features were defined based on 1) the interaction term *p*-value (Type:Season) of 0.05 or less in combination with 2) the pairwise seasonal comparison of RNA expression ('Winter.rna/Summer.rna') FDR-adjusted *p*-value (*q*-value) of 0.05 or less.

DNA and RNA	296,476
DNA and RNA with gene IDs	161,984
Number of gene symbols found in DNA and RNA	5579
Number of gene symbols found in DNA and RNA in	133
BacMet database	
Number of gene IDs from DNA and RNA found in	7021
BacMet	
Number of gene IDs from DNA in BacMet database that	8 (copR_13, corC_121, cusR_32, czcA_647, nike_38,
are not found in RNA	pstC_144, ruvB_54, Int_122)
Number of gene IDs that are differentially expressed	947

#### Table 5

ARTS annotated contigs. ARTs (https://arts3.ziemertlab.com) annotated contigs using Actinobacteria and Alphaproteobacteria reference sets. Phylogeny is not applicable (N/A) to this metagenomic dataset. These data are also located on Figshare; DOI: 10.6084/m9.figshare.c.11879226. URL – https://doi.org/10.6084/m9.figshare.c.11879226).

																		Totals	
Contigs	1-3712	3713-	4242-	10,000-	15,485-	25,001-	35,574-	45,001-	66,478-	85,001-	110,410-	135,001-	169,689-	240,000-	330,000-	440,000-	501,400-		
		4241	9999	15,484	25,000	35,573	45,000	66,477	85,000	110,409	135,000	169,688	239,999	329,999	439,999	501,399	579,964		
Total Genes	162,298	14,102	98,404	65,200	88,082	78,662	59,473	115,925	84,874	101,037	86,501	108,789	188,391	205,523	218,732	111,940	130,993	Total Genes	1,918,926
Core Essential Genes	395	303	387	387	387	383	381	392	377	384	379	382	399	391	370	333	328	Core Essential Genes	6358
Total BGC Hits	136	9	98	81	90	74	70	119	74	95	75	80	134	142	139	92	81	Total BCGC Hits	1589
Known Resistance Models	944	71	595	411	526	420	331	580	406	473	332	474	683	744	742	346	423	Known Resistance Models	8501
Gene Duplication	364	198	354	342	351	344	336	350	334	344	334	345	355	347	324	289	284	Gene Duplication	5595
BGC Proximity	198	0	44	28	20	10	11	12	2	4	1	2	1	0	0	0	1	BGC Proximity	334
Phylogeny/ HGT	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Phylogeny/ HGT	0
2 or more	128	0	36	25	16	8	9	11	2	4	1	2	1	0	0	0	1	2 or more	244
3 or more	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	3 or more	0
Reference Set: Alpha Proteobacteria																			
Nodes	1-3712	3713-	4242-	10,000-	15,485-	25,001-	35,574-	45,001-	66,478-	85,001-	110,410-	135,001-	169,689-	240,000-	330,000-	440,000-	501,400-		
		4241	9999	15,484	25,000	35,573	45,000	66,477	85,000	110,409	135,000	169,688	239,999	329,999	439,999	501,399	579,964		
Total Genes	162,298	14,102	98,404	65,200	88,082	78,662	59,473	115,925	84,874	101,037	86,501	108,789	188,391	205,523	218,732	111,940	130,993	Total Genes	1,918,926
Core Essential Genes	516	359	506	495	506	505	504	517	492	509	488	502	510	504	488	444	444	Core Essential Genes	8289
Total BGC Hits	136	9	98	81	90	74	70	119	74	95	75	80	134	142	139	92	81	Total BCGC Hits	1589
Known Resistance	944	71	595	411	526	420	331	580	406	473	332	474	683	744	742	346	423	Known Resistance	8501
Models																		Models	
																			0
Gene Duplication	486	216	470	444	461	459	439	473	441	470	437	449	478	472	449	380	371	Gene Duplication	7395
BGC Proximity	220	1	71	40	22	21	13	23	5	8	5	5	2	3	2	0	1	BGC Proximity	442
Phylogeny/ HGT	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Phylogeny/ HGT	0
2 or more	138	1	56	34	17	12	10	14	3	6	3	4	1	1	2	0	1	2 or more	303
3 or more	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	3 or more	0

#### Table 6

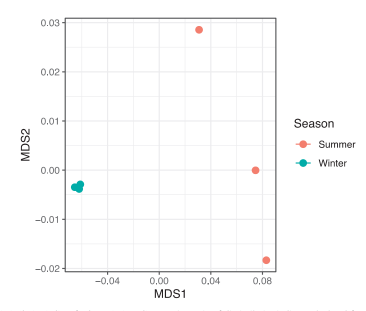
Colocalized and/or coexpressed genes. Colocalized and/or coexpressed BacMet genes with BGCs. Differentially expressed features were defined based on 1) the interaction term p-value (Type:Season) (p-interaction) of 0.05 or less in combination with 2) the pairwise seasonal comparison of RNA expression ('Winter.rna/Summer.rna') FDR-adjusted p-value (q-value) of 0.05 or less.

	Contig #	Genus	Percent match to genus	Gene ID	Gene	Funtion	p-interaction value	p-winter RNA/summer RNA value	q-winter RNA/summer RNA value	Winter RNA/summer RNA Log2-fold change
Cluster 1										
Metal resistance	4689			FHBHJPKI_167716	pitA_11	L-methionine sulfoximine/L-methionine sulfone acetyltransferase	4.37E-05	0.000733	0.00468	-2.94
Secondary metabolite	4689			FHBHJPKI_167725		Involved in synthesis of homoserine lactone-nonribosomal peptide	1.93078E-11	2.85272E-09	5.90633E-08	-3.500187665
Cluster 2										
Metal resistance	80	Candidatus Solibacter usitatus Ellin6076	100	FHBHJPKI_12377	mgtA_4	Magnesium-transporting ATPase-2C P-type 1	0.00701	0.0182	0.0167	-2.01
Secondary metabolite	80	Candidatus Solibacter usitatus Ellin6076	100	FHBHJPKI_12365	lgrD_9	Linear gramicidin synthase subunit D	0.841648661	0.004247211	0.021296771	-2.463862034
Cluster 3				-						
Metal resistance	12,335	Acidobacterium capsulatum ATCC 51,196	33	FHBHJPKI_283288	mdtA_189	Multidrug resistance protein MdtA	3.68E-10	1.23E-16	8.92E-15	-4.71
Secondary metabolite	12,335	Acidobacterium capsulatum ATCC 51,196	33	FHBHJPKI_283295	crtB_77	All-trans-phytoene synthase	0.15602993	0.006446359	0.030262887	-2.056268682
Cluster 4										
Metal resistance	214	Ralstonia solanacearum CMR15	22	FHBHJPKI_24632	smtB_5	Succinyl-CoA-L-malate CoA-transferase beta subunit	0.042	0.000392	0.0027	-4.18
Secondary metabolite	214	Ralstonia solanacearum CMR15	22	FHBHJPKI_24627	shc_2	Squalene-hopene cyclase	0.067593728	0.000537894	0.003591324	-2.489137791
Cluster 5				,						
Metal resistance	185	Candidatus Koribacter versatilis Ellin345	100	FHBHJPKI_22308	czcA_9	Cobalt-zinc-cadmium resistance protein CzcA	0.0321	0.0017	0.00968	-2.31
Secondary metabolite	185	Candidatus Koribacter versatilis Ellin345	100	FHBHJPKI_22329		Putative ligase/MSMEI_5285	0.270379299	0.031944273	0.111550994	-1.879407266
Cluster 6				,						
Metal resistance	4698			FHBHJPKI_167937	mdtA_99	Multidrug resistance protein MdtA	0.0292	0.0299	0.106	-3.16
Secondary metabolite	4698			FHBHJPKI_167934	ppsE_5	Involved in synthesis of Phthiocerol/phenolphthiocerol polyketide	0.01973596	0.001581153	0.009156685	-1.99280005



	Jan_Sed3	Jan_Sed1	Jan_Sed2	July_Sed1	July_Sed3	July_Sed2
Jan_Sed3						
Jan_Sed1	0.01126638					
Jan_Sed2	0.01126957	0.01335446				
July_Sed1	0.05546984	0.05502813	0.05731778			
July_Sed3	0.08193703	0.08160468	0.08391454	0.03925216		
July_Sed2	0.07715887	0.0767676	0.0790924	0.03172766	0.01837441	

Β.



**Fig. 3.** Bray-Curtis dissimilarity indices for bacteria in sediment. A) Matrix of dissimilarity indices calculated for genera of bacteria in sediment samples using the Bray-Curtis method. 'Sed' = sediment. B) NMDS plot to visualize the dissimilarity between genera of bacteria in summer (July\_Sed1, July\_Sed2, and July\_Sed3 in orange) and winter (Jan\_Sed1, Jan\_Sed2, and Jan\_Sed3 in blue) sediment collected at EB-90M. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

# 3. Experimental Design, Materials and Methods

# 3.1. Sample collection

On July 28th, 2017 and January 14th, 2018, Ely Brook (43°55'9″ N, 72°17'11″ W), 90 m upstream from the mouth of the brook (EB-90M), was sampled along with unsaturated sediment (10 cm deep). The physicochemical properties, nucleic acid extraction, library preparation, and metatranscriptomic and metatranscriptomic sequencing, taxonomic annotation of raw reads, metagenomic assembly, and functional annotations of these samples were reported by Giddings et al. [1]. Α.

Jan_Sed3	Jan_Sed1	Jan_Sed2	July_Sed1	July_Sed3	July_Sed2
0.02604939					
0.02281105	0.02776086				
0.08264324	0.0775801	0.08953024			
0.0761283	0.06914731	0.07990833	0.0259954		
0.07445164	0.06903036	0.07944745	0.03169012	0.0253259	
	- 0.02604939 0.02281105 0.08264324 0.0761283	0.02604939           0.02281105         0.02776086           0.08264324         0.0775801           0.0761283         0.06914731	0.02604939           0.02281105         0.02776086           0.08264324         0.0775801         0.08953024           0.0761283         0.06914731         0.07990833	0.02604939 0.02281105 0.02776086 0.08264324 0.0775801 0.08953024 0.0761283 0.06914731 0.07990833 0.0259954	0.02604939 0.02281105 0.02776086 0.08264324 0.0775801 0.08953024 0.0761283 0.06914731 0.07990833 0.0259954



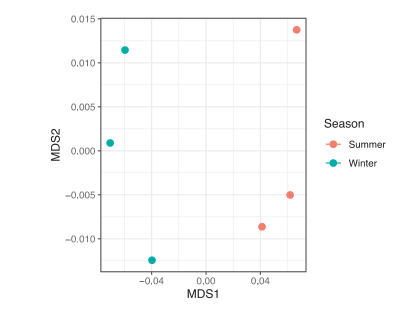
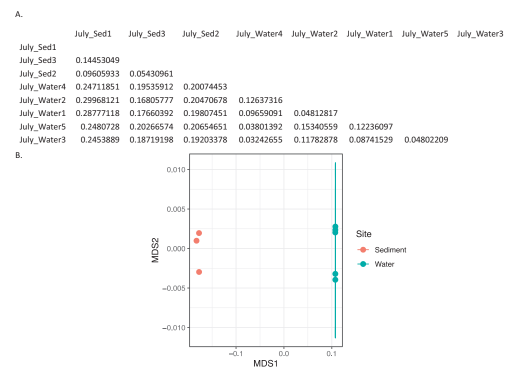


Fig. 4. Bray-Curtis dissimilarity indices for eukaryota in sediment. A) Matrix of dissimilarity indices calculated for genera of eukaryota in sediment samples using the Bray-Curtis method. 'Sed' = sediment. B) NMDS plot to visualize the dissimilarity between genera of eukaryota in summer (July\_Sed1, July\_Sed2, and July\_Sed3 in orange) and winter sediment (Jan\_Sed1, Jan\_Sed2, and Jan\_Sed3 in blue) collected at EB-90M. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

# 3.2. Statistical comparison of microbial community, DNA, and RNA

EB-90M samples of the same sample type or season were treated as biological replicates. Subsets (i.e., season or sample type) of data were compared to each other in statistical analyses. Beta diversity was evaluated via Bray-Curtis measure of dissimilarity [10] using default parameters in R in the vegan library [11]. Prior to analysis, data were  $\log_{10}(x+1)$  transformed and the resulting dissimilarity indices were used to generate NMDS in R using the metaMDS functions in vegan and ggplot2 library [11, 12]. Multivariate PCAs were performed in Partek Flow software v8.0 to assess sample group variation based on genera using normalized read counts from read-based taxonomic annotations and quantification. Feature counts (e.g., taxon) were standardized prior to the PCA so that the contribution of each feature did not depend on its variance. PCA

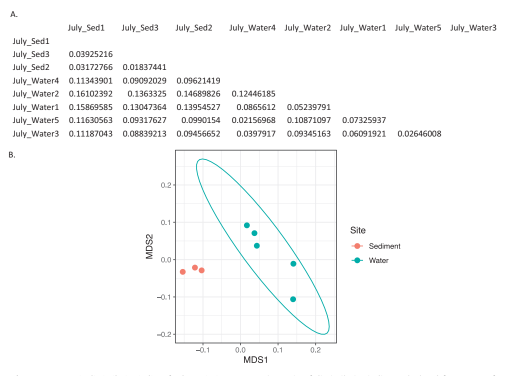


**Fig. 5.** *Bray-Curtis dissimilarity indices for archaea in summer.* A) Matrix of dissimilarity indices calculated for genera of archaea in summer samples using the Bray-Curtis method. 'Sed' = sediment. B) NMDS plot to visualize the dissimilarity between genera of archaea in summer sediment (July\_Sed1, July\_Sed2, and July\_Sed3 in orange) and water (July\_Water1, July\_Water2, July\_Water3, July\_Water4, and July\_Water5 in blue) collected at EB-90M. The ellipse indicates a clustering of more than 3 samples. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

plots were generated for DNA and RNA using 1) normalized read counts (i.e., fractions for relative abundance) from the metagenomic assembly and 2) normalized read counts from the metatranscriptome, respectively. Heat maps and hierarchal clusters were generated in Partek Flow v8.0 using the following, respectively: 1) normalized counts of taxa from the metagenome and predicted open reading frames (ORFs) across samples and 2) the Euclidean dissimilarity index and average linkage method to cluster similar expression patterns and taxon abundances. The normalized data were standardized to a mean of zero and a standard deviation of 1 prior to hierarchal clustering.

#### 3.3. Differential expression and visualization of KEGG pathways

Differentially expressed KEGG pathways were represented by color gradation maps (Figs. S14–S15). Log<sub>2</sub>fold-changes from gene expression analysis results were converted to a color gradation using KEGG Mapper – Color Pathway tool (https://www.genome.jp/kegg/tool/map\_pathway3.html), where blue denotes decreased expression in the winter (RGB color code #6363F7) and red denotes increased expression in the winter (RGB color code #FF000). Genes with no change in expression are shaded in light gray (RGB color code #D3D3D3). Genes shaded

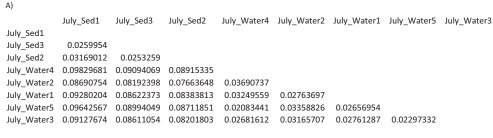


**Fig. 6.** *Bray-Curtis dissimilarity indices for bacteria in summer.* A) Matrix of dissimilarity indices calculated for genera of bacteria in summer samples using the Bray-Curtis method. 'Sed' = sediment. B) NMDS plot to visualize the dissimilarity between genera of bacteria in summer sediment (July\_Sed1, July\_Sed2, and July\_Sed3 in orange) and water (July\_Water1, July\_Water2, July\_Water3, July\_Water4, and July\_Water5 in blue) collected at EB-90M. The ellipse indicates a clustering of more than 3 samples. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

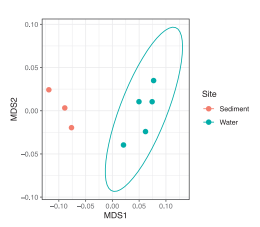
in white indicates that the gene was undetected in the dataset. The numbers in boxes refer to enzyme nomenclature from the KEGG database. Expression data (i.e., normalized counts) for sediment were fit to a linear model, assuming a negative binomial distribution, that included season (i.e., winter versus summer), molecule type (i.e., RNA versus DNA), as well as the interaction of season and molecule type (*p*-interaction). Pairwise comparison tests of season were performed within and between each data type and *p*-values were FDR-corrected [9]. Significant differentially expressed genes met the following criteria: a molecule type-season interaction term *p*-value of 0.05 or less in combination with an FDR-adjusted *p*-value (*q*-value) of 0.05 or less for the pairwise comparison of winter RNA versus summer RNA. Significant data were indicated by an orange star; however, the overall expression of a node may include other genes.

# 3.4. Analysis of genes involved in natural product biosynthesis, metal resistance, and antibiotic resistance

Contigs were mined for secondary metabolite biosynthetic gene clusters (BGCs) in the bacterial and fungal antiSMASH 5.0 [4] database using default parameters. The BacMet database was used to mine DNA and RNA for experimentally validated metal resistance genes [3]. After filtering annotated-BGCs and BacMet genes that had  $\geq$  100 raw counts in each sample and at least

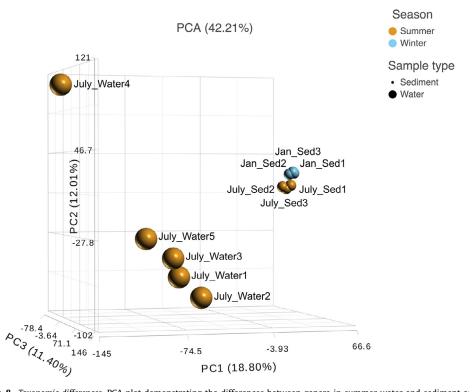


B)



**Fig. 7.** *Bray-Curtis dissimilarity indices for eukaryota in summer.* A) Matrix of dissimilarity indices calculated for genera of eukaryota in summer samples using the Bray-Curtis method. 'Sed' = sediment. B) NMDS plot to visualize the dissimilarity between genera of eukaryota in summer sediment (July\_Sed1, July\_Sed2, and July\_Sed3 in orange) and water (July\_Water1, July\_Water2, July\_Water3, July\_Water4, and July\_Water5 in blue) collected at EB-90M. The ellipse indicates a clustering of more than 3 samples. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

10 counts in three or more samples, relative BGC and BacMet gene expression was assessed by comparing the counts of Prokka-annotated transcripts to those of DNA using the criteria described by Giddings et al. [1]. Gradient plots were generated in Partek Flow v8.0 for differentially expressed BGCs and those co-expressed with metal resistance genes. Contigs were also mined for antibiotic resistance genes that were within close proximity or colocalized with BGCs using the Antibiotic Resistant Target Seeker (ARTS) version 2 [5] using default parameters. Duplication and BGC proximity, resistance model screens, and genomes that mapped to the following reference phyla were selected: Actinobacteria and Alphaproteobacteria.



**Fig. 8.** *Taxonomic differences.* PCA plot demonstrating the differences between genera in summer water and sediment as well as summer (orange) and winter (blue) sediment. Plot is based on normalized read counts at the genus level from the taxonomic annotation and quantification of paired-end reads. The sample name notation is based on the month the sample was collected, the sample type (i.e., sediment or water), and individual sample number. 'Sed' = sediment. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

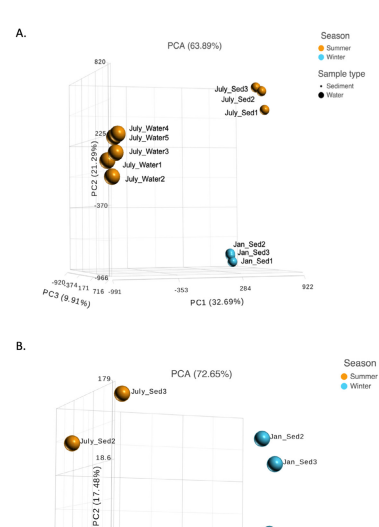


Fig. 9. Differences in DNA and RNA. PCA plots of A) DNA in water and sediment and B) RNA present in summer and winter sediment based on normalized counts of all functionally annotated genes from the metagenomic assembly, demonstrating differences between sample type. Each gene's normalized read count contributes equally to the PCA. The sample name notation is based on the month the sample was collected, the sample type (i.e., sediment or water), and individual sample number. 'Sed' = sediment.

-99.1

-142

-303

-263<sub>-89.8</sub> 83.7 257 -285

PC3 (15.28%)

July\_Sed1

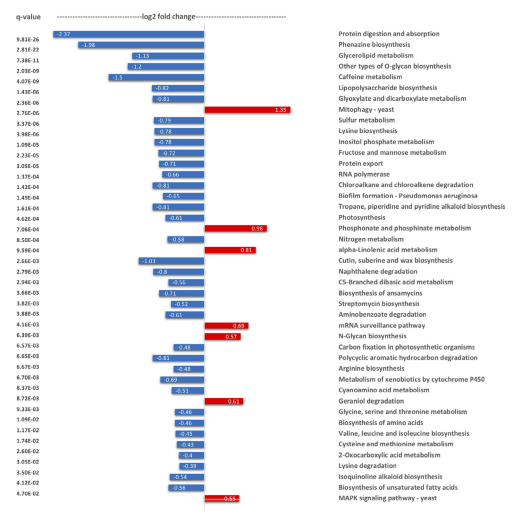
Jan\_Sed1

273

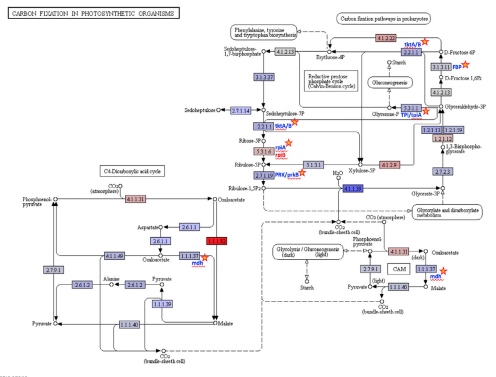
87.1

PC1 (39.89%)



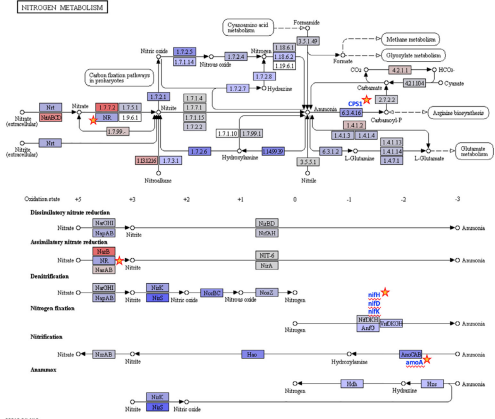


**Fig. 10.** Significantly differentially xpressed KEGG pathways. Bar graph of select significantly differentially expressed KEGG pathways in winter versus summer. Differentially expressed pathways were defined based on an unadjusted p-value  $\leq$  0.05 for the interaction term (molecule type-season) in combination with a q-winter/summer RNA value  $\leq$  0.05, respectively. Red and blue represent increased and decreased expression in winter, respectively. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



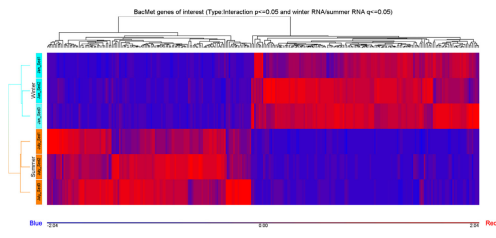
00710 5/29/19 (c) Kanehisa Laboratories

**Fig. 11.** *Carbon fixation in photosynthetic organisms.* Carbon metabolism KEGG reference pathway map (https://www. kegg.jp/pathway/map00710) with color gradation highlighting KEGG genes that change significantly between seasons. Log<sub>2</sub>fold-changes from gene expression analyses were converted to a color gradation using the KEGG Mapper – Color Pathway tool, where blue denotes decreased expression in the winter (RGB color code #6363F7) and red denotes increased expression in the winter (RGB color code #FF000). The Log<sub>2</sub>fold-changes range from –2.33 (blue) to +1.88 (red). Genes with no change in expression are shaded in light gray (RGB color code #D3D3D3) and genes shaded white were undetected in the dataset. Significantly differentially expressed genes are indicated by a star and met the following criteria: *p*-interaction value  $\leq$  0.05 in combination with a *q*-winter/summer RNA value  $\leq$  0.05, respectively. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

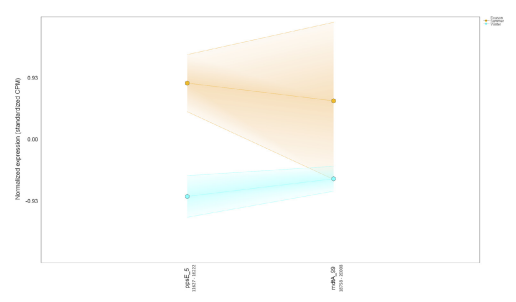


<sup>00910 8/16/18</sup> (c) Kanehisa Laboratories

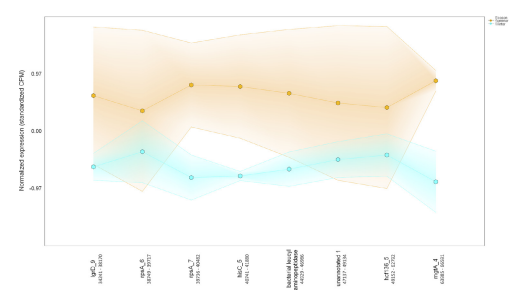
**Fig. 12.** Nitrogen metabolism gene expression. Nitrogen metabolism KEGG reference pathway map diagram (https://www. kegg.jp/pathway/map00910) with color gradation highlighting KEGG genes that change significantly between seasons. Log<sub>2</sub>fold-changes from gene expression analyses were converted to a color gradation using the KEGG Mapper – Color Pathway tool, where blue denotes decreased expression in the winter (RGB color code #6363F7) and red denotes increased expression in the winter (RCB color code #FF000). The Log<sub>2</sub>fold-changes range from -3.92 (blue) to +1.91 (red). Genes with no change in expression are shaded in light gray (RGB color code #D3D3D3) and genes shaded white were undetected in the dataset. Significantly differentially expressed genes are indicated by a star and met the following criteria: *p*-interaction value  $\leq 0.05$  in combination with a *q*-winter/summer RNA value  $\leq 0.05$ , respectively. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



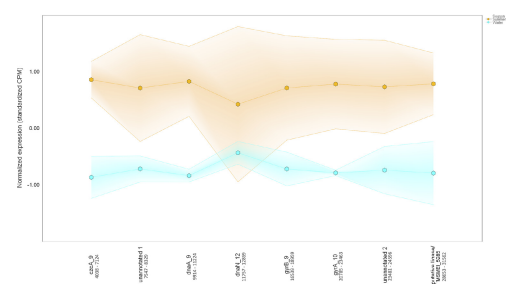
**Fig. 13.** *Metal resistance gene expression.* Hierarchical clustering and heat map of differentially expressed select (288) genes (e.g., *dnaK*, *copA*, *copB*, *copD*, *pst5*, *cusA*, *cusB*, *mdtA*, *mdtB*, *mdtC*, *actP*, *mco*, *ycnJ*, *corA*, *csoR*, and *copZ*) from the BacMet database across sediment samples. Increases or decreases in gene expression range from -2.04 (blue) to +2.04 (red). All data met the following criteria: *p*-interaction value  $\leq 0.05$  in combination with a *q*-winter/summer RNA value  $\leq 0.05$ , respectively. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



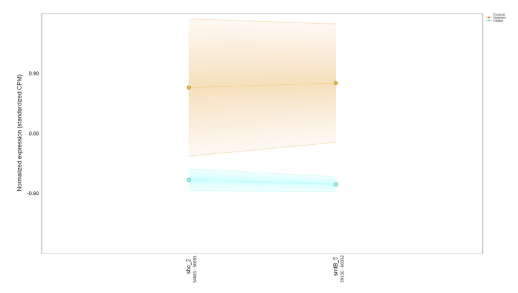
**Fig. 14.** Colocalization and coexpression of metal resistance and secondary metabolite genes. Gradient plot demonstrating the differential coexpression of *mdtA*, a metal resistance gene encoding a multidrug resistance protein, with a gene (*ppsE*) annotated to be involved in phthiocerol/phenolphthiocerol polyketide biosynthesis in contig 4698 (20,390 nucleotides long) in summer (orange) and winter (blue). The lines on the y-axis represent the maximum, minimum, and mean of the standardized expression values (i.e., counts per million). All data met the following criteria: *p*-interaction  $\leq$  0.05, respectively. Nucleotide positions in contig are shown below gene IDs. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



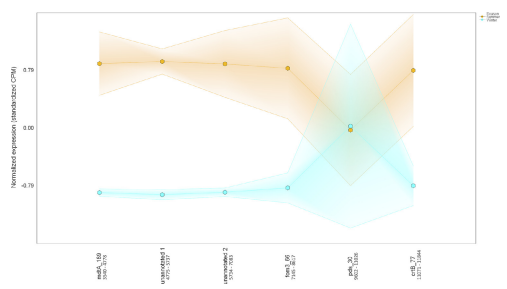
**Fig. 15.** Colocalization and coexpression of metal resistance and secondary metabolite genes. Gradient plot demonstrating the differential coexpression of *mgtA*, a metal resistance gene encoding a cation transport ATPase that mediates magnesium influx into the cytosol, with genes (*lgrD*) annotated to be involved in gramicidin biosynthesis in contig 80 (113,676 nucleotides long) in summer (orange) and winter (blue). The lines on the y-axis represent the maximum, minimum, and mean of the standardized expression values (i.e., counts per million). Only *mgtA* met the following criteria: *p*-interaction  $\leq 0.05$  in combination with a *q*-winter/summer RNA  $\leq 0.05$ , respectively. Nucleotide positions in contig are shown below gene IDs. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



**Fig. 16.** Colocalization and coexpression of metal resistance and secondary metabolite genes. Gradient plot demonstrating the differential coexpression of *czcA*, a metal resistance gene encoding a cobalt-zinc-cadmium resistance protein, with a ligase/MSMEL5285 gene annotated to be involved in the biosynthesis of a polyketide in contig 185 (85,942 nucleotides long) in summer (orange) and winter (blue). The lines on the y-axis represent the maximum, minimum, and mean of the standardized expression values (i.e., counts per million). Only *czcA* met the following criteria: *p*-interaction  $\leq$  0.05 in combination with a *q*-winter/summer RNA  $\leq$  0.05, respectively. Nucleotide positions in contig are shown below gene IDs. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



**Fig. 17.** Colocalization and coexpression of metal resistance and secondary metabolite genes. Gradient plot demonstrating the differential coexpression of *smtB*, a zinc-resistance gene encoding a repressor protein of the metallothionein gene *smtA*, with a gene annotated to be involved in the biosynthesis of a terpene in contig 214 (80,995 nucleotides long) in summer (orange) and winter (blue). The lines on the y-axis represent the maximum, minimum, and mean of the standardized expression values (i.e., counts per million). Only *SmtB* met the following criteria: *p*-interaction  $\leq$  0.05 in combination with a *q*-winter/summer RNA  $\leq$  0.05, respectively. Nucleotide positions in contig are shown below gene IDs. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



**Fig. 18.** Colocalization and coexpression of metal resistance and secondary metabolite genes. Gradient plot demonstrating the differential coexpression of *mdtA*, a metal resistance gene encoding multidrug resistance protein, with genes annotated to be involved in the biosynthesis of a terpene in contig 12,335 (11,958 nucleotides long) in summer (orange) and winter (blue). The lines on the y-axis represent the maximum, minimum, and mean of the standardized expression values (i.e., counts per million). Only *mdtA* met the following criteria: *p*-interaction  $\leq$  0.05, in combination with a *q*-winter/summer RNA  $\leq$  0.05, respectively. Nucleotide positions in contig are shown below gene IDs. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

# **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships which have, or could be perceived to have, influenced the work reported in this article.

#### Acknowledgments

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# Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi: 10.1016/j.dib.2020.106282.

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