

# Pfam differential expression data for diatoms from an iron incubation experiment during the PUPCYCLE II R/V Sally Ride cruise within the California Current System in May and June of 2023

**Website:** <https://www.bco-dmo.org/dataset/1004674>

**Version:** 1

**Version Date:** 2026-08-13

## Project

» [CAREER: An integrated molecular and physiological approach to examining the dynamics of upwelled phytoplankton in current and changing oceans](#) (Upwelled Phytoplankton Dynamics)

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|--------------------------------------|---------------------------------------------------------------|------------------------|
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## Coverage

**Location:** California Current System, off the southern coast of Oregon at 43°02'42.7"N, 124°33'07.2"W, depth 55m

**Spatial Extent:** N:44.62701922 E:-123.867405 S:37.71706162 W:-125.492809

**Temporal Extent:** 2023-05-29 - 2023-06-10

## Dataset Description

This dataset is one of many generated from an iron incubation experiment conducted as part of the PUPCYCLE II cruise in May and June 2023. The different analyses are listed here, with links to other datasets from this study in the Related Datasets section.

Analyses include:

- Physiology (dataset 994607)
  - Dissolved inorganic nutrients
  - Size-fractionated chlorophyll a
  - Size-fractionated isotope uptake rates
  - Flow cytometry
  - FlowCAM analysis
  - FLP experiments for mixotroph grazing
- RNA taxonomy data (dataset 1003525)
- Differential expression: Pfam
  - Mixotroph Pfam (dataset 1004659)
  - **Diatom Pfam (this dataset 1004674)**
- Differential expression: KEGG gene (dataset 1004689)
- Differential expression: KEGG pathways (dataset 1004704)
- Dissolved iron concentrations (dataset 1004007)

See Related Datasets section below for links to above mentioned datasets.

## Methods & Sampling

Data collection took place on the R/V Sally Ride from May 29th to June 10th, 2023. To simulate upwelling conditions under different iron treatments, an onboard incubation experiment was conducted. Seawater for the incubation experiment was collected within the northern CCS, off the southern coast of Oregon at 43°02'42.7"N, 124°33'07.2"W. The collected seawater was deemed as freshly upwelled water and taken from a depth of 55 m – which corresponds to a depth slightly below the euphotic zone receiving less than 1% irradiance – using trace-metal clean techniques on May 30th, 2023, 13:30 GMT. The seawater was pumped and homogenized using trace metal clean techniques, then transferred into a total of thirty 20 L low-density polyethylene cubitainers. Three cubitainers were immediately harvested for the initial timepoint (T0). The remaining twenty-seven cubitainers were assigned treatments: nine were unamended (Ctrl), nine were amended with 5 nM FeCl<sub>2</sub> (+Fe), and nine were amended with 200 nM desferrioxamine B, a strong iron chelator that inhibits dissolved iron uptake (DFB). The cubitainers were placed in an on-deck incubator covered with two layers of neutral density screening to simulate 26% of incidence irradiance supplied with flow-through surface seawater to maintain ambient surface temperature. Three cubitainers from each treatment were harvested for each of the three subsequent timepoints: 48 hours (T1), 168 hours (T2), and 264 hours (T3) after incubation.

**RNA Collection and Metatranscriptomics.** Cubitainers were harvested for RNA by collecting and filtering approximately 2.5 to 4 L of each sample onto 0.8 µm Pall Supor filters (142 mm) using a peristaltic pump; samples were then immediately flash frozen in liquid nitrogen and stored at -80°C. RNA was extracted using the RNAqueous-4PCR kit, per instructions from the manufacturer with few modifications – filters were cut up due to their large size, 200 µL of glass beads were added, and 3 mL of lysis buffer was added. RNA samples were sent to GENEWIZ for library preparation and sequencing with PolyA tail selection. Sequencing was performed on an Illumina HiSeq 4000 with a 2x150 bp configuration. GENEWIZ provided raw paired-end read sequences for each sample.

Raw reads were trimmed using Trim Galore v0.6.10. and quality control was assessed through FastQC (1). A *de novo* metatranscriptome assembly was conducted using rnaSPAdes v3.15.5 for individual assemblies (2) and CD-HIT v4.8.1 for a grand assembly (3). TransDecoder v5.7.1 was used to determine the most likely open reading frame (ORF) for each contig (4). ORFs were annotated using the Marine Functional Eukaryotic Reference Taxa (MarFERRet) database v1.1.1 (e-value < 1e<sup>-06</sup>) (5) for NCBI taxonomies (6), PR<sup>2</sup> taxonomies (7) and Pfam 34.0 functions (8). To remove any potential bacterial contamination, ORFs were also annotated against the PhyloDB v1076 database using EUKulele (e-value < 1e<sup>-06</sup>) (9). Any ORF with a higher bitscore for a bacteria or virus compared to its MarFERRet annotation was removed (resulting in < 3% of ORFs removed). To maximize functional annotation coverage, ORFs were annotated using eggNOG-mapper v2.1.12 (10) to obtain annotations from the Kyoto Encyclopedia of Genes and Genomes (KEGG) (11). Trimmed samples were then aligned to the grand assembly using Salmon v1.10.3 (12) and exported into a comprehensive counts table using tximport (13). Reads not mapping to a protist group as defined by MarFERRet annotations were removed before downstream analysis.

For taxonomic composition, all protist reads were normalized using DESeq2's median of ratios methods (14). The mean and standard deviation for the number of reads mapping to each protist group and the proportion of reads mapping to each protist group were calculated for each timepoint/treatment. Mean/standard deviation

for number and proportion of protist reads specifically mapping to mixotroph genera and diatom genera were also calculated. Mixotrophs were identified based on constitutive mixotroph genera within the Mixoplankton Database (MDB) (15) and diatoms genera were identified based on mapping to the lineage Bacillariophyta.

## Data Processing Description

The metatranscriptomic pipeline code used for this project is publicly available at <https://github.com/emilyspeciale/Speciale-Metatranscriptomics>.

Trim Galore v0.6.10  
CD-HIT v4.8.1  
TransDecoder v5.7.1  
Marine Functional Eukaryotic Reference Taxa (MarFERReT) database v1.1.1  
PhyloDB v1076 database using EUKulele  
eggNOG-mapper v2.1.12  
Salmon v1.10.3  
tximport  
DESeq2

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## Related Datasets

### IsRelatedTo

Speciale, E., Marchetti, A., Lim, P., Jeong, Y., Cook, C., Cohen, N., McClure, W., Schnetzer, A. (2026) **KEGG gene differential expression data for mixotrophs and diatoms**. Biological and Chemical Oceanography Data Management Office (BCO-DMO). (Version 1) Version Date 2026-08-13 <http://lod.bco-dmo.org/id/dataset/1004689> [[view at BCO-DMO](#)]

Speciale, E., Marchetti, A., Lim, P., Jeong, Y., Cook, C., Cohen, N., McClure, W., Schnetzer, A. (2026) **KEGG pathway differential expression data for mixotrophs and diatoms**. Biological and Chemical Oceanography Data Management Office (BCO-DMO). (Version 1) Version Date 2026-08-14 <http://lod.bco-dmo.org/id/dataset/1004704> [[view at BCO-DMO](#)]

Speciale, E., Marchetti, A., Lim, P., Jeong, Y., Cook, C., Cohen, N., McClure, W., Schnetzer, A. (2026) **Pfam differential expression data for mixotrophs from an iron incubation experiment during the PUPCYCLE II R/V Sally Ride cruise within the California Current System in May and June of 2023**. Biological and Chemical Oceanography Data Management Office (BCO-DMO). (Version 1) Version Date 2026-08-12 <http://lod.bco-dmo.org/id/dataset/1004659> [[view at BCO-DMO](#)]

Speciale, E., Marchetti, A., Lim, P., Jeong, Y., Cook, C., Cohen, N., McClure, W., Schnetzer, A. (2026) **Physiological measurements from iron incubation experiment using upwelled waters in the California Current System during the PUPCYCLE II cruise in May and June 2023**. Biological and Chemical Oceanography Data Management Office (BCO-DMO). (Version 1) Version Date 2026-08-11 <http://lod.bco-dmo.org/id/dataset/994607> [[view at BCO-DMO](#)]

Speciale, E., Marchetti, A., Lim, P., Jeong, Y., Cook, C., Cohen, N., McClure, W., Schnetzer, A. (2026) **RNA taxonomy data from an iron incubation experiment during the PUPCYCLE II cruise (R/V Sally Ride) within the California Current System in May and June of 2023**. Biological and Chemical Oceanography Data Management Office (BCO-DMO). (Version 1) Version Date 2026-08-12 <http://lod.bco-dmo.org/id/dataset/1003525> [[view at BCO-DMO](#)]

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## Parameters

*Parameters for this dataset have not yet been identified*

## Instruments

|                                         |                                                                                                                    |
|-----------------------------------------|--------------------------------------------------------------------------------------------------------------------|
| <b>Dataset-specific Instrument Name</b> | Illumina HiSeq 4000                                                                                                |
| <b>Generic Instrument Name</b>          | Automated DNA Sequencer                                                                                            |
| <b>Dataset-specific Description</b>     | Sequencing was performed on an Illumina HiSeq 4000 with a 2x150 bp configuration.                                  |
| <b>Generic Instrument Description</b>   | A DNA sequencer is an instrument that determines the order of deoxynucleotides in deoxyribonucleic acid sequences. |

|                                         |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |
|-----------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <b>Dataset-specific Instrument Name</b> | Sea Bird SBE 9 plus CTD                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
| <b>Generic Instrument Name</b>          | Sea-Bird SBE 9plus CTD                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |
| <b>Generic Instrument Description</b>   | High precision and accuracy CTD comprising an SBE 9plus underwater unit (SBE 3plus temperature, SBE 4C conductivity, and Paroscientific Digiquartz pressure sensors, and an SBE 5T submersible pump). Can be used for either real-time data acquisition or for autonomous operations at a sampling speed of up to 24 Hz. The instrument package also includes a TC duct, to reduce salinity spiking caused by ship heave for improved resolution of water column features, and to ensure that temperature and conductivity measurements are made on the same parcel of water. Supplied with both an aluminium and titanium main housing, allowing for use up to 6800 and 10,500 metre depths respectively. Also capable of measuring from eight auxiliary sensors. |

## Deployments

### SR2311

|                    |                                                                                                       |
|--------------------|-------------------------------------------------------------------------------------------------------|
| <b>Website</b>     | <a href="https://www.bco-dmo.org/deployment/994610">https://www.bco-dmo.org/deployment/994610</a>     |
| <b>Platform</b>    | R/V Sally Ride                                                                                        |
| <b>Start Date</b>  | 2023-05-29                                                                                            |
| <b>End Date</b>    | 2023-06-10                                                                                            |
| <b>Description</b> | California Current System, off the southern coast of Oregon at 43°02'42.7"N, 124°33'07.2"W, depth 55m |

## Project Information

**CAREER: An integrated molecular and physiological approach to examining the dynamics of upwelled phytoplankton in current and changing oceans (Upwelled Phytoplankton Dynamics)**

**Coverage:** California Upwelling Zone

#### *NSF Award Abstract:*

Upwelling zones are hotspots of photosynthesis that are very dynamic in space and time. Microscopic algae, known as phytoplankton, bloom when deep, nutrient-rich waters are upwelled into sunlit surface layers of the ocean, providing nourishment that supports productive food webs and draws down carbon dioxide (CO<sub>2</sub>) from the atmosphere to the deep ocean. Photosynthetic microbes in these regions must constantly adapt to changes in their chemical and physical environments. For example, subsurface populations respond to changes in light as they approach the surface. When upwelled waters move offshore, cells sink out of the illuminated zone, establishing seed populations that remain inactive until the next upwelling event. This process is called the upwelling conveyor belt cycle (UCBC). How phytoplankton respond to these changes in environmental conditions and how they may influence their nutrient requirements remains unknown. With future ocean changes predicted to alter seawater chemistry, including ocean acidification and decreased iron availability, some phytoplankton groups may be more vulnerable than others. Accompanying educational activities provide learning experiences to enhance understanding and awareness of marine microbes. The development of a research hub at UNC aims to provide infrastructure and support for scientists and students conducting research on environmental genomics. A laboratory component for an upper-level undergraduate course focused on marine phytoplankton is being developed. Educational outreach activities to broader communities include creation of a lesson plan on phytoplankton in upwelling zones and a virtual research cruise experience for middle-school students, as well as a hands-on lab activity for a local museum focused on marine phytoplankton and the important roles they play in shaping our planet.

The project examines how phytoplankton respond at the molecular and physiological level to the different UCBC stages, which seed populations (i.e., surface versus subsurface) contribute most to phytoplankton blooms during upwelling events of varying intensity, how phytoplankton elemental compositions are altered throughout UCBC stages, and how future predicted ocean conditions will affect the phytoplankton responses to UCBC conditions. This project contains both laboratory and fieldwork. In the laboratory, phytoplankton isolates recently obtained from upwelling regions are exposed to simulated UCBC conditions to examine changes in gene expression, growth and photosynthetic characteristics and elemental composition. Cultures are subjected to both current and future ocean conditions, including reduced iron availability and higher CO<sub>2</sub>. In the field, research cruises within upwelling regions study the dynamics of natural phytoplankton communities (both surface and subsurface) experiencing upwelling and relaxation and within simulated upwelling incubation experiments. Knowledge of how phytoplankton are affected by UCBC conditions at an integrated molecular, physiological and elemental level under both current and future scenarios is imperative for the proper conservation and management of these critically important ecosystems.

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## **Funding**

| <b>Funding Source</b>                                    | <b>Award</b>                |
|----------------------------------------------------------|-----------------------------|
| <a href="#">NSF Division of Ocean Sciences (NSF OCE)</a> | <a href="#">OCE-1751805</a> |

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