

Proteomic data of dilution series of bacteria and diatoms using MS (DDA).

Website: <https://www.bco-dmo.org/dataset/719122>

Version:

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Project

» [Collaborative Research: Proteins as functional biomarkers: integrating organic characterization with proteomics to track routes for carbon and nitrogen recycling and preservation](#) (Proteins as Biomarkers)

Contributors	Affiliation	Role
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Dataset Description

To mimic a complex marine sample, a dilution series of *R. pomeroyi* and *T. pseudonana* was created at different cellular ratios. These mixtures were filtered and proteins were extracted from the filter for tryptic digestion and LC-MS/MS analysis.

Biological fractions were lysed, digested and analyzed using proteomic mass spectrometry.

Data are available for download at the EBI PRIDE Archive.

Homepage: <https://www.ebi.ac.uk/pride/archive>

Project URL: <https://www.ebi.ac.uk/pride/archive/projects/PXD004799>

Data URL: <https://www.ebi.ac.uk/pride/archive/projects/PXD004799/files>

Methods & Sampling

Data Dependent Acquisition on QExactive

Samples were analyzed in technical duplicates (triplicates for some) for 3 biological replicates for each cellular dilution in data dependent analysis mode on the top 20 precursor ions. The analytical column was 20 cm long (packed with C18) and the trap was 3 cm (packed with C12). Peptides were eluted using a 5-35% CAN gradient over 60 minutes with a 300 nl/min flow rate. MS1 ions were collected in the scan range of 400-1400 m/z.

DDA data.

Instrument(s): QExactive Thermo Finnegan: DDA data

Data Processing Description

Completed proteomes for both *R. pomeroyi* and *T. pseudonana* were downloaded from Uniprot for searches using Comet and the Trans Proteomic Pipeline. These data were used to create selected reaction monitoring assays (see accompanying datasets; in review).

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Data Files

File
Bacterial_detectionDDA_Timmins.csv (Comma Separated Values (.csv), 193 bytes) MD5:be1ebbd4243c527e5be9079f56e93905
Primary data file for dataset ID 719122

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Parameters

Parameter	Description	Units
Repository	Name of database where data are currently served	unitless
Project	Unique project identifier for the database where data are currently served	unitless
Project_URL	Link to project page at the database where data are currently served	unitless

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Instruments

Dataset-specific Instrument Name	QExactive Thermo Finnegan
Generic Instrument Name	Mass Spectrometer
Generic Instrument Description	General term for instruments used to measure the mass-to-charge ratio of ions; generally used to find the composition of a sample by generating a mass spectrum representing the masses of sample components.

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Project Information

Collaborative Research: Proteins as functional biomarkers: integrating organic characterization with proteomics to track routes for carbon and nitrogen recycling and preservation (Proteins as Biomarkers)

Coverage: Multiple locations: Canadian and US Beaufort Sea

NSF Award Abstract:

Proteins are a major contributor to organic carbon and nitrogen in the ocean and their amino acid building blocks comprise a major fraction of the total nitrogen identified in coastal and oceanic waters and sediments. Over the last decade, tandem mass spectrometry methods developed for the analysis of peptides and protein reconstruction together with informatics developments have re-assembled the amino acid signatures and unlocked the information inherent in the proteins they represent. This interdisciplinary research team has the combined expertise in marine organic biogeochemistry, proteomics, and bioinformatics to help determine the fate of individual proteins during degradation and likely mechanisms for their preservation. This research aims to identify and quantify the proteins responsible for chemical transformations of organic matter in the ocean, thereby exposing how microbial communities control and contribute to the carbon and nitrogen cycle and to organic matter preservation. This project has the potential to trigger a fundamental change in how we view, analyze, and model microbial degradation processes.

Recent findings from microbial ecologists show that bacteria with broad responses to various nutrients (i.e. the generalists) can fine-tune the gene expression of various proteins as a reflection of substrate availability; the proponents of this project have observed a suite of bacterial proteins present during organic matter recycling and that this protein expression changes as degradation proceeds. These observations hint that bacterial proteins have potential as organic biomarkers to reflect the taxonomic distribution of the functioning catalysts in degradation processes. The investigators propose to build on these results to investigate the potential for organic matter preservation to be linked to proteomic expression of the microbial community that drive carbon and nitrogen cycling. The project objectives are: 1) to address the relationship between marine organic matter composition and protein expression by the bacterial community (metaproteome) which act as catalysts for degradation; 2) to detail protein expression in the context of detailed organic matter characterization and 3) to distinguish how proteins present in oceanic sediments derived from different kingdoms of life (e.g. eukaryotic or bacterial) reflect the complex process of organic matter preservation. The overall goal is to capture proteins as critical markers of both presence and process in marine systems. The investigators will link the rapid advances in protein identification to track bacterial proteins to act as "functional biomarkers" and indicators of carbon and nitrogen utilization.

The project would provide multiple opportunities for interdisciplinary student training in marine geochemistry and proteomics, and address the goal of disseminating results and tools to a broad audience. In the more traditional role, the project will expand the career for a female research faculty member in marine proteomics and support both graduate and undergraduate student research at each participating institution. On the broader community level, both Harvey and Nunn are also heavily involved in high school outreach programs.

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Funding

Funding Source	Award
NSF Division of Ocean Sciences (NSF OCE)	OCE-1633939

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