EXPORTS Measurements and Protocols for the NE Pacific Campaign

Editors:
Ivona Cetinić and Inia Soto Ramos
GESTAR/Universities Space Research Association, Columbia, Maryland

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Chapter 1 Introduction to the EXPORTS Parameter Working Groups

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Background

EXport Processes in the Ocean from Remote Sensing (EXPORTS) is a large-scale NASA-led and NSF co-funded field campaign that will provide critical information for quantifying the export and fate of upper ocean net primary production (NPP) using satellite information and state of the art technology.

The goal of EXPORTS is to develop a predictive understanding of the export and fate of global ocean net primary production (NPP) and its implications for the Earth’s carbon cycle in present and future climates (oceanexports.org). To develop this quantitative understanding, EXPORTS is measuring and modeling the export pathways that remove fixed organic carbon from the upper ocean and drive the attenuation of these vertical fluxes within the ocean interior.

EXPORTS datasets will be used to develop and test numerical predictive and satellite-data diagnostic models of NPP fates and their carbon cycle impacts. EXPORTS builds on decades of NASA-funded research on developing and validating satellite data-driven models of regional to global NPP and hence, EXPORTS will contribute to NASA’s upcoming Plankton, Aerosol, Cloud and ocean Ecosystem (PACE) mission (Werdell et al., 2019). NSF segment in the program stems from the long-term investment in the science of biological pump, culminating with the NSF Biology of the Biological Pump Workshop (BoBP) in 2016 (Burd et al., 2016).

The initial stages of EXPORTS date back to COOPEX, a NASA ROSES (2012, A.3) funded project which final objective was to draft a Science Plan for a field campaign entitled “Controls on Open Ocean Productivity and Export Experiment”. The original COOPEX plan, which required constraining both the production and the fate of fixed organic carbon, was too ambitious and had budgetary limitations. Therefore, it was decided that the field campaign should focus on the fates of fixed carbon and not its production. It was during the COOPEX expert meeting at the University of California Santa Barbara (UCSB) in June 2013, that the EXport Processes in the Ocean from RemoTe Sensing (EXPORTS) field campaign was born.

Science plan for the field campaign was published as a report (EXPORTS Writing Team, 2015) and an overview paper(Siegel et al., 2016). Following the extensive peer and panel reviews of the proposed science plan, in October 2015, NASA OBB selected another team of scientists to
form a Science Definition Team, who was tasked with development of the implementation plan for EXPORTS. After a year of work, and several meetings, team published the final report (EXPORTS Science Definition Team, 2016).

Following the recommendations from the NASA EXPORTS science plan (EXPORTS Writing Team, 2015), EXPORTS science implementation plan (EXPORTS Science Definition Team, 2016), and NSF’s Biology of the Biological Pump workshop report (Burd et al., 2016), EXPORTS is being conducted as a three phase project. EXPORTS commenced with the release of the Request For Proposals (RFP) by NASA Ocean Biology and Biogeochemistry in August 2016 to fund data mining and Observing System Simulation Experiment (OSSE) numerical modeling in support of EXPORTS planning and science activities. Following, NSF’s released a Dear Colleague Letter (DCL) stating they would consider proposals that leveraged the NASA investment with objectives that supported the BoBP workshop report. In 2017, NASA released a general call for funding of the field portion of the EXPORTS, known as the First phase/stage of EXPORTS, that will span over the two field campaigns, starting in the North Pacific, in 2018, and North Atlantic in 2020. The second of EXPORTS, targets the implementation and translation of the knowledge learned during the field campaigns into predictive and forecasting models and satellite-derived NPP algorithms.

**Figure 1-1. Timeline of the EXPORTS field campaign.**

EXPORTS field campaigns combine research vessels, autonomous platforms and remote sensing observations to achieve the overarching goals. The North Pacific field campaign, that took place in 2018, consisted of two UNOLS vessels: (1) a process ship (R/V Roger Revelle)
responsible for the deployment of short-term drifting arrays, sediment traps, most net tows, CTD casts to collect material for shipboard experimentation, and incubation-based biological rate determinations; and (2) a survey ship (R/V Sally Ride) responsible for the physical and geochemical measurements requiring distributed CTD sampling, large-volume in situ pumping, and towed profiler surveys. The autonomous array consisted of 2 biogeochemical (BCG) Argo floats, 6 Neutrally Buoyant Sediment Traps (NBST), single Lagrangian float, a Wire Walker, and a surface-tethered trap. The North Atlantic field Campaign was planned to have an additional vessel (through collaboration with WHOI’s Ocean Twilight Zone project) and additional AUV platforms, however the protocols and sampling regime will remain very similar to the North Pacific field work. As this is being written, global pandemic of Covid-19 has postponed the planned North Atlantic campaign to foreseeable future. In addition to the vessel and autonomous assets, the field campaigns were planned near long-term monitoring stations such as the Ocean Observatories Initiative Station Papa in the North Pacific and the Porcupine Abyssal Plain (PAP) site in the North Atlantic which will provide additional historical and reference data.

In early phase of the EXPORTS field stage, the Project Office and PIs implemented a parameter group organizational structure to facilitate the inter-project communication, planning, and data management. The main goal of the parameter groups was to compile all the methods and protocols used during the EXPORTS field campaigns. The parameter groups were clustered around five main topics (Table 1.1): (1) Stocks, Proxies & Context, (2) Optics, (3) Particle Characterization, (4) Rates, and (5) Exports. Stocks, Proxies and Context parameter group is responsible for all biogeochemical measurements (e.g., dissolved oxygen, DOC, POC), physical observations including those collected using sensors onboard the ship and autonomous platforms, and synthesized observations created by mapping one variable onto another (e.g., POC derived from optical backscattering). The optics parameter group is responsible for all the inherent and apparent optical properties such remote sensing reflectance, water absorption and scattering, concentration of optical water constituents and attenuation. Particle characterization group focused on different methods of characterization of quantity and quality (community composition) of particles in the water column. Rates working group focused on rates of change of different biogeochemical stocks, encompassing different estimates of production and removal. Lastly, Exports working group focused on vertical fluxes of carbon, and other biogeochemical tracers.

The main goal of the parameter groups was to compile all the methods and protocols used during the EXPORTS field campaigns. Such documents, here named as protocol documents, define the method, targeted deliverable parameters, dependencies on other parameters measured or modeled during the field campaigns and uncertainties associated with deliverables. Ultimately, protocol documents have listings of parameter names, as submitted to the databases - SeaBASS and BCO-DMO. Protocols were written prior to the North Pacific experiment and finalized within the year following the cruise. In addition to being the ultimate reference place for all collected measurements during the EXPORTS field campaign, protocols provided a path to post cruise synthesis work and facilitated answering of the EXPORTS science questions (Siegel et al., 2016). This connection to overarching goals of the EXPORTS
was highlighted in the reports that each of the parameter groups produced; reports collated in this technical memorandum.
Organized around the five main topics and some of the protocols may be listed multiple times as they may answer the questions under several topics targeted by EXPORTS.

### Table 1.1 Summary of the parameter groups objectives and measurements

<table>
<thead>
<tr>
<th>Parameter Group</th>
<th>Team objectives</th>
<th>Type of measurements/instruments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stocks, Proxies &amp; Context</td>
<td>Provide documentation to help EXPORTS investigators understand, access, and utilize stock, proxy, and context observations</td>
<td>Stocks are biogeochemical measurements in quantity per volume or mass seawater (e.g., DOC and dissolved oxygen). Proxies are synthesized observations created by mapping one variable onto another (e.g., POC derived from optical backscatter). Context observations include all physical/sensor measurements from the EXPORTS assets (i.e., ships, autonomous vehicles, floats, moorings and remote sensing observations)</td>
</tr>
<tr>
<td>(Chapter 2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optics (Chapter 3)</td>
<td>Provide high quality optical data that can be used to characterize the various ocean constituents and that can be further used to translate the EXPORTS data into satellite-derived algorithms.</td>
<td>Apparent optical properties (e.g., radiance, irradiance and reflectance). Inherent optical properties (e.g., attenuation coefficient, absorption coefficient, volume scattering function, fluorescence). The direct imaging devices, although optical measurements, were included in the Particle Characterization group. Measurements include those from ship underway system, ship-deployed profilers, above water radiometer, gliders, Lagrangian BGC Argo floats and wirewalker.</td>
</tr>
<tr>
<td>Particle Characterization</td>
<td>Provide high quality measurements of the abundance, composition and size distribution of suspended particles (including live organisms) in the water column.</td>
<td>Optical and imaging instruments were used for enumeration, sizing and classification of particles (e.g., plankton taxonomy and functional groups, biomass, biovolume, abundance, size distribution). Measurements were complemented with (1) metagenomics and DNA barcoding, (2) microscopic image analyses of particles, and (3) characterization of particles collected on polyacrylamide gels in sediment traps. Instrumentation includes epifluorescence microscopy, flow cytometry, Imaging FlowCytoBot-IFCB, MOCNESS/Zooscan, Underwater Video Profiler, LISST, Coulter counter, etc.</td>
</tr>
<tr>
<td>(Chapter 4)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rates (Chapter 5)</td>
<td>Provide high quality measurements of primary production, bacterial and community respiration, secondary production, and grazing rates.</td>
<td>Primary productivity (GPP, NPP, NCP): incorporation of isotopes (stable and radioactive) over 6 hr and 24 hr incubations to measure uptake of dissolved inorganic carbon. The production of biogenic gases (e.g., oxygen) to estimate PP. New production estimated through stable isotope NO3 incorporation. Diatom-specific production estimated through uptake of silicon. Respiration: Bacterial and Community respiration measured by O2 drawdown assays and DOC remineralization assays. Zooplankton respiration assessed by O2 drawdown, analysis of Electron Transport System (ETS) enzyme activity, and converting community composition and biomass measurements to community respiration. Measurements also include bacterial production, secondary production and grazing, aggregation and sinking rates, and nutrient uptake rates.</td>
</tr>
<tr>
<td>Exports (Chapter 6)</td>
<td>Understand the mechanisms controlling the magnitude and attenuation of fluxes to depth (biological and physical export pathways out of the euphotic zone and flux pathways below the euphotic zone).</td>
<td>High magnification microscope images of polyacrylamide gel collectors on sediment traps are used to enumerate, characterize and determine the distribution of sinking particles. The presence or absence of particles collected with the Marine Snow Catcher. The biological composition determined by genetic sequencing and proportional contributions to bulk traps of amino acids. Respiration rates on sinking particles, determined using RESPIRE traps and experimentally with particles from the Marine Snow Catcher. Profiled Underwater Vision Profiler (UVP) images and LISST particle size distributions are analyzed to identify aggregates and their water column size distribution. Aggregate dynamics and coagulation modeling. Depth- and time-dependent distribution of zooplankton biomass are determined using the MOCNESS and UVP, with semi-automated image analysis to identify taxa. Vertical and horizontal variability in zooplankton migratory distributions are characterized qualitatively with acoustics systems on the survey ship, and on the autonomous platforms. Fecal pellet production rates determined experimentally.</td>
</tr>
</tbody>
</table>

Following the protocols developed by the working groups. Please note that the protocols are organized around the five main topics and some of the protocols may be listed multiple times as they may answer the questions under several topics targeted by EXPORTS.
Figure 1-2 EXPORTS data tree
(https://coggle.it/diagram/Wq85Q8GZi/y3bf0xO/t/export/37265e74477f7634f76c10dc391269b84fbc01c36d487976f4b3155e4e742)
Overview

The goal of the Stocks, Proxies and Context Working Group (SPC WG) is to provide documentation to help EXPORTS investigators understand, access, and utilize Stocks, Proxy and Context observations from the NE Pacific EXPORTS Field Campaign. The other Parameter WGs are focused on specific aspects of the field program’s measurement suite; Optics, Rates, Export Pathways, and Particle Characterization. Hence, one of the foci for the SPC WG is ensuring that there are no gaps in measurements among the five working groups.

Definitions and parameter tables

Definitions for Stocks, Proxy and Context measurements are as follows:

**Stocks** - Stocks are defined here as any measurement of biogeochemical significance measured in quantity per volume or mass seawater. For example, DOC and dissolved oxygen are Stocks, while salinity is not (salinity will be defined as a Context measurement). Hence, Stocks are the measurements that contribute to the boxes in the EXPORTS wiring diagram (Fig. 2.1). For convenience, all taxa-specific information can be found under the report of the Particle Characterization Working Group (Chapter 4), while chemical composition information (either on particles or dissolved constituents) is covered in this chapter. Similarly, genomics-based determinations are not Stocks, so measurements such as OTUs/volume are covered by the Particle Characterization Working Group (Chapter 4). All optics measurements are considered in the Optics WG report (Chapter 3). Stocks measurements are primarily ship-based observations from the cruise and as such will be largely static in time once an analysis is run (compared with Proxies whose implementation will be more fluid, based on the updates provided by post-cruise analyses).
Stocks protocol documents describe analyses performed and provide measures of uncertainty for each parameter. It is likely that many, if not most, of the protocol documents will be “owned” by another Parameter WG. Finally, when appropriate, protocols will include intercalibration information amongst variables (assessing variations among various POC or Chl-a determinations made).

Proxies - Proxies are defined here as synthesized observations created by mapping one variable onto another. Development of proxies usually serves the purpose of increasing the spatial / temporal resolution of a discrete measurement using a sensor-based measurement and typically a simple algorithm linking them. For example, using optical backscatter to represent POC is a proxy using a derived empirical relationship between the two parameters. Furthermore, to ensure consistency across the platforms, pre-cruise instrument intercalibration was performed (Chapter 3), and during the cruise numerous planed and serendipitous in field intercalibration exercises were performed. Due to their derived nature, it is unlikely that the process creating these proxies will be locked down early in the analysis phase. The Proxy Measurement Table (Table 2.1) is included here to illustrate our intent in creating the table. It is likely that this activity will not be complete by the time these WG reports are completed. Again, protocol documents will describe the methods used to derive the proxy measurements and provide measures of uncertainties for each.

Context - Context observations are useful for understanding the context of the observations made during EXPORTS. These include physical / sensor measurements from the EXPORTS ships’ sampling systems (i.e., CTD/rosettes, ADCP, UW, echo sounders), all autonomous assets (SeaGlider, BioARGO floats, Lagrangian float, etc.) as well as measurements made by nearby (PMEL, OOI moorings, Line P, etc.) and remote observations (MODIS and VIIRS imagery, merged altimetry, etc.). When appropriate, short documents will describe analyses used to process the observations into data sets and how to access them.
## Stock, Proxy and Context Measurement Tables

Table 2.1 Stocks as defined in EXPORTS, with associated information. Platforms are P - process ship (R/V Revelle), S - survey (R/V Sally Ride), _uw - underway, _CTD - rosette, _pumps - pumps, _exp - measurement for experiment. Complete table can be found at: https://drive.google.com/open?id=1Kzvna3bDEU9DyD9EJeR9EFsQI1polzG8SG0M35RjTs

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Parameter name in database (SeaBASS)</th>
<th>Units (SeaBASS)</th>
<th>Platform</th>
<th>Parameter document</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxygen (discrete, continuous)</td>
<td>Oxygen, oxygen_kg</td>
<td>mL L^-1, umol/kg</td>
<td>S_CTD, P_CTD, WW, LF, SG</td>
<td>Oxygen</td>
</tr>
<tr>
<td>Macronutrients (NO3, NO2, SiO4, PO4)</td>
<td>NO3, NO2, SiO4, PO4</td>
<td>mmol/m^3</td>
<td>P_CTD, S_CTD</td>
<td>Inorganic_nutrients_protocol_UCSB</td>
</tr>
<tr>
<td>Nitrate (continuous)</td>
<td>NO3</td>
<td>mmol/m^3</td>
<td>S_uw, S_CTD</td>
<td>BGC SUMO + ISUS</td>
</tr>
<tr>
<td>Ammonia [NH4+]</td>
<td>NH4</td>
<td>mmol/m^3</td>
<td>P_CTD</td>
<td>Santoro_ammoniumOPA</td>
</tr>
<tr>
<td>Particulate Organic Carbon and Particulate Organic Nitrogen</td>
<td>POC, PON</td>
<td>mg/m^3</td>
<td>P_CTD, S_CTD, S_uw, P_uw</td>
<td>Three or 4 methods</td>
</tr>
<tr>
<td>Particulate Inorganic Carbon</td>
<td>PIC</td>
<td>mol/m^3</td>
<td>P_CTD, S_CTD</td>
<td>Particulate_Inorganic_Carbon</td>
</tr>
<tr>
<td>Biogenic Silica</td>
<td>Bsi</td>
<td>mmol/m^3</td>
<td>P_CTD, S_CTD</td>
<td>Psi protocol Teflon Tubes_BrzezinskiLab</td>
</tr>
<tr>
<td>Dissolved Organic Carbon</td>
<td>DOC_L, DOC</td>
<td>umol L^-1, umol/kg</td>
<td>P_CTD, S_CTD</td>
<td>DOC</td>
</tr>
<tr>
<td>Size fractionated (1, 20, 50 um) Particulate organic Carbon, Particulate Nitrogen, Total Particulate Phosphates</td>
<td>POC, PN, PTP</td>
<td>mg/m^3, mmol/m^3</td>
<td>S_pump</td>
<td>size fractionated particles</td>
</tr>
<tr>
<td>Size fractionated (1, 20, 50 um) Particulate Inorganic Carbon</td>
<td>PIC</td>
<td>mol/m^3</td>
<td>S_pump</td>
<td>size fractionated particles</td>
</tr>
<tr>
<td>Size fractionated (1, 20, 50 um) Biogenic Silica</td>
<td>Bsi</td>
<td>mmol/m^3</td>
<td>S_pump</td>
<td>size fractionated particles</td>
</tr>
<tr>
<td>HPLC pigments (including Chl a)</td>
<td>Chl_a and many other</td>
<td>mg/m^3</td>
<td>P_CTD, S_CTD, S_uw, P_uw, S_pumps</td>
<td>Final_method_HPLC</td>
</tr>
<tr>
<td>Fluorometric Chl a (acetone extraction)</td>
<td>Chl</td>
<td>mg/m^3</td>
<td>P_CTD, S_CTD</td>
<td>Chlorophyll_analysis_Roesler</td>
</tr>
<tr>
<td>Fluorometric Chl a (methanol extraction)</td>
<td>Chl_experiment</td>
<td>mg/m^3</td>
<td>P_exp</td>
<td>Chlorophyll_extraction_Menden-Deuer</td>
</tr>
<tr>
<td>Phytoplankton abundance</td>
<td>Abun_phyto</td>
<td>Cells/L</td>
<td>P_CTD, S_CTD, S_uw, P_uw</td>
<td>yes</td>
</tr>
<tr>
<td>Bacterioplankton abundances</td>
<td>Abun_bacterioplankton</td>
<td>Cells/L</td>
<td>P_CTD</td>
<td>Bacterial abundance</td>
</tr>
<tr>
<td>Small Zooplankton (&lt;63 um)</td>
<td>Conc_particles_id</td>
<td>Particles/L</td>
<td>P_other</td>
<td>63 micron net</td>
</tr>
<tr>
<td>Zooplankton</td>
<td>Conc_particles_id</td>
<td>Particles/L</td>
<td>P_other</td>
<td>Zooplankton + MOCNESS</td>
</tr>
<tr>
<td>Aggregates</td>
<td>Conc_particles_id</td>
<td>Particles/L</td>
<td>P_other</td>
<td>Snowcatcher, density</td>
</tr>
<tr>
<td>TEP</td>
<td>TEP_bottle, TEP_MSC</td>
<td>ug_Gxan EQUIV/L</td>
<td>P_CTD, P_other</td>
<td>Snowcatcher</td>
</tr>
<tr>
<td>DIC/TA</td>
<td>DIC, total_alkalinity</td>
<td>umol/kg, none</td>
<td>S_CTD, S_uw</td>
<td>Discrete TA + Discrete DIC</td>
</tr>
<tr>
<td>pH</td>
<td>pH</td>
<td>None</td>
<td>S_CTD, S_uw</td>
<td>BGC SUMO + Discrete pH</td>
</tr>
<tr>
<td>Lipids</td>
<td></td>
<td></td>
<td>P_CTD</td>
<td>Lipidomics_method_for_EXPORTS</td>
</tr>
<tr>
<td>234Th</td>
<td>Conc_Th_234</td>
<td>Dpm/L</td>
<td>P_CTD</td>
<td>Buesseler Thorium export pathways protocol briefs</td>
</tr>
</tbody>
</table>
Table 2.2 Proxy measurements developed during North Pacific EXPORTS. Complete table can be found at https://docs.google.com/spreadsheets/d/16dqU0WHnOwMyYbZx7Z1jCNoa2OhSICgbUhy6BKTrQ/edit#gid=0

<table>
<thead>
<tr>
<th>‘Parameter’</th>
<th>Ship-based water sample measurement</th>
<th>Simple sensor</th>
<th>AUV measurement</th>
<th>Protocol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phytoplankton Chl</td>
<td>Chl, HPLC Chl</td>
<td>Chl F; a(676)</td>
<td>Chl F</td>
<td>Chlorophyll proxies for AUVs</td>
</tr>
<tr>
<td>Phytoplankton Chl</td>
<td>Chl, HPLC Chl</td>
<td>Ed(412, 442, 553)</td>
<td>Ed(412, 442, 553)</td>
<td>Chlorophyll proxies for AUVs</td>
</tr>
<tr>
<td>Phytoplankton C</td>
<td>plankton C from imaging, FCM, and/or sorting</td>
<td>bbp</td>
<td>bbp</td>
<td>POC proxies for AUVs</td>
</tr>
<tr>
<td>Plankton community composition</td>
<td>FCM&amp; imaging (1º) HPLC (2º)</td>
<td>Chl F/ bbp</td>
<td>Chl F/bbp</td>
<td></td>
</tr>
<tr>
<td>POC</td>
<td>POC</td>
<td>cp, bbp</td>
<td>cp, bbp</td>
<td></td>
</tr>
<tr>
<td>Nitrate</td>
<td>Chemical nitrate</td>
<td>N/A</td>
<td>Suna</td>
<td>Fassbender_unnderway_nitrate_pH</td>
</tr>
<tr>
<td>Oxygen</td>
<td>Winkler</td>
<td>SBE 43</td>
<td>Optode</td>
<td></td>
</tr>
<tr>
<td>Aggregate abundance</td>
<td>N/A</td>
<td>optical spikes cp, bbp; LISST &amp; UVP on CTD</td>
<td>optical spikes cp, bbp; OmandPerry_OpticalSpikes_1page</td>
<td></td>
</tr>
<tr>
<td>Zooplankton stock</td>
<td>Zooplankton from MOCNESS</td>
<td>MOCNESS with ADCP</td>
<td>ADCP</td>
<td></td>
</tr>
</tbody>
</table>

Table 2.3 Context Measurement Summary

<table>
<thead>
<tr>
<th>EXPORTS NP</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTD-SIO (Revelle &amp; Ride), CTD-TNC (Revelle), CTD-MOCNESS (Revelle)</td>
</tr>
<tr>
<td>UW-SIO (Revelle &amp; Ride)</td>
</tr>
<tr>
<td>ADCP (Revelle &amp; Ride)</td>
</tr>
<tr>
<td>EK80 echo sounder (Ride)</td>
</tr>
<tr>
<td>Mets (Revelle &amp; Ride)</td>
</tr>
<tr>
<td>SeaGlider (AUV team)</td>
</tr>
<tr>
<td>Lagrangian Float (AUV team)</td>
</tr>
<tr>
<td>BioARGO floats (AUV team)</td>
</tr>
<tr>
<td>OOI UW measurements</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Line P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bottle Files &amp; CTD profiles (recent and historical)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Station P Infrastructure</th>
</tr>
</thead>
<tbody>
<tr>
<td>PMEL Mooring Mets, Fluxes and University of Washington Observations</td>
</tr>
<tr>
<td>OOI Subsurface Moorings &amp; Gliders</td>
</tr>
<tr>
<td>APL Waverider Mooring</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Satellite</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ocean color imagery</td>
</tr>
<tr>
<td>SST (thermal &amp; microwave)</td>
</tr>
<tr>
<td>Sea Level (merged sea level)</td>
</tr>
</tbody>
</table>
Relationship to EXPORTS science questions

Stock concentrations play into all EXPORTS Science Questions. In particular, changes in biogeochemical stocks relate to changes in process. Some of the stocks can be derived, by using proxies from in situ optical measurements, or with algorithms from ocean color measurements. Hence, these fundamental measurements need to be monitored continuously.
Overview

The underlying hypothesis of the EXPORTS is that changes in community composition in the surface ocean observed using satellite remote sensing can be used to quantify the export and fate of upper ocean net primary production (NPP). Fundamentally, ocean optics investigate a) the way the light field is changing in the ocean due to the nature and geometry of the ambient light field and ocean itself, also known as apparent optical properties (AOPs); and b) the way that oceanic constituents interact with light which depends on the nature of the medium solely, also known as inherent optical properties (IOPs). While both offer insight into the physical nature of the light in the ocean and provide an insight about the quantity and quality of light available for NPP, EXPORTs optical measurements, either collected from the satellites, in-situ platforms, or used on discrete samples collected in field give an opportunity to measure different pools of biogeochemical parameters and their changes on frequencies and scales not accessible by other means. This is achieved by developing relationships between AOPs and IOPs and targeted parameters, either through the development of Proxies (see Chapter 2) or ocean color algorithms.

Measured parameters and associated methods

Main goal of EXPORTS field campaign is to provide the critical information for quantifying the export and fate of upper ocean net primary production (NPP) using satellite observations and state of the art ocean technologies. Optical measurements, collected as part of EXPORTS, offer
a direct connection to the satellite observations, and a way to extrapolate measurements of stocks and rates to the larger spatial and temporal scales covered by the AUV-based observations. Optical instrumentation was present on almost all of the observational platforms deployed during EXPORTS, making it (following the contextual measurements of the physical properties) most data abundant subsection of EXPORTS observations.

The main goal of this report is to summarize the EXPORTS Science Team’s optical observations made during the 2018 North Pacific Ocean cruise onboard R/V Roger Revelle and Sally Ride, as well as the AUVs deployed as part of this experiment, and catalogize the protocols used to obtain these observations. The optical observations covered in this document include those that measure the AOPs - apparent optical properties (e.g., radiance, irradiance, and reflectance), and the inherent optical properties (e.g., attenuation coefficient, absorption coefficient, volume scattering function, fluorescence), and are shown in Table 3.1 and 3.2. While each of the methods used has an outlined parameter write up (linked in the Tables 3.1 and 3.2, and collated in the end of this technical memorandum), it is important to note that methods used were followed available community standards (e.g. in-line systems on both vessels, operated by PIs Roesler and Boss followed recommendations by Boss et al. (2019)).

Table 3.1 List of instrumentation used, during the EXPORTS NP, to collect Apparent Optical Properties (AOPs). Platforms are P – Process ship, S – Survey ship, LF – Lagrangian Float, WW – Wirewalker, SG – Seaglider.

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Parameter(s) measured</th>
<th>Parameter name in database (SeaBASS)</th>
<th>Units (SeaBASS)</th>
<th>Platform</th>
<th>Parameter document</th>
</tr>
</thead>
<tbody>
<tr>
<td>C-OPS (Compact-Optical Profiling System) at 19 wavelengths</td>
<td>In-water radiometry (Upwelling radiance, downwelling irradiance, downwelling surface irradiance, diffuse attenuation coefficient)</td>
<td>Lu</td>
<td>uW/cm^2/nm/sr</td>
<td>P, S, SG (Ed only)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Es</td>
<td>uW/cm^2/nm</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ed</td>
<td>uW/cm^2/nm</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Kd</td>
<td>uW/cm^2/nm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HyperSAS</td>
<td>Above-water radiometry (water-leaving radiance)</td>
<td>Lu, Lu, Es</td>
<td>uW/cm^2/nm/sr, uW/cm^2/nm</td>
<td>P, S</td>
<td></td>
</tr>
<tr>
<td>HTSRB</td>
<td>Above-water radiometry in floating mode</td>
<td>Lu</td>
<td>W/cm^2/nm/sr,</td>
<td>P, S</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ed</td>
<td>uW/cm^2/nm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C-OPS</td>
<td>Photosynthetic available radiation</td>
<td>PAR</td>
<td>uE/cm^2/s</td>
<td>P, S</td>
<td></td>
</tr>
<tr>
<td>PAR instrument on CTD/glider</td>
<td>Photosynthetic available radiation</td>
<td>PAR</td>
<td>uE/cm^2/s</td>
<td>P, S, SG</td>
<td></td>
</tr>
<tr>
<td>Surface PAR</td>
<td>Photosynthetic available radiation</td>
<td>PAR</td>
<td>uE/cm^2/s</td>
<td>P</td>
<td></td>
</tr>
<tr>
<td>PAR on wirewalker/Seaglider</td>
<td>Photosynthetic available radiation</td>
<td>PAR</td>
<td>uE/cm^2/s</td>
<td>WW, SG, P</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter(s) measured</th>
<th>Instrument</th>
<th>Parameter name in database (SeaBASS)</th>
<th>Units (SeaBASS)</th>
<th>Platform (mode)</th>
<th>Parameter document</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Hyper)Spectral absorption (particulate, dissolved total), Particulate attenuation</td>
<td>Ac-S/Ac-9</td>
<td>ap</td>
<td>1/m</td>
<td>P(I,v),S(I,v)</td>
<td>Boss and Roesler</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ad</td>
<td>1/m</td>
<td>S(I,v), P(v)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>cp</td>
<td>1/m</td>
<td>P(I,v),S(I,v)</td>
<td></td>
</tr>
<tr>
<td>Absorption (particulate, phytoplankton, dissolved)</td>
<td>Discrete</td>
<td>ap, aph, ad</td>
<td>1/m</td>
<td>S(d), P(d)</td>
<td>Roesler, Nelson</td>
</tr>
<tr>
<td>Particulate attenuation (single λ)</td>
<td>LISST-Deep</td>
<td>cp</td>
<td>1/m</td>
<td>S(v), P(v)</td>
<td>McDonnel</td>
</tr>
<tr>
<td></td>
<td>C-star</td>
<td>cp</td>
<td>1/m</td>
<td>S(v), P(v)</td>
<td>Hydro</td>
</tr>
<tr>
<td></td>
<td>C-Star</td>
<td>cp</td>
<td>1/m</td>
<td>WW</td>
<td>Omand/hydro</td>
</tr>
<tr>
<td></td>
<td>LISST-VSF</td>
<td>cp</td>
<td>1/m</td>
<td>S(d)</td>
<td>Zhang</td>
</tr>
<tr>
<td>Spectral backscattering (λ,φ), as derived from Volume Scattering Function</td>
<td>Eco-Triplet (BB3)</td>
<td>bbp</td>
<td>1/m</td>
<td>S(i), P(i)</td>
<td>Boss, Roesler</td>
</tr>
<tr>
<td></td>
<td>FLBBRTD</td>
<td>bbp</td>
<td>1/m</td>
<td>S(i), P(v)</td>
<td>CTD</td>
</tr>
<tr>
<td></td>
<td>BBFL2b</td>
<td>bbp</td>
<td>1/m</td>
<td>S(i)</td>
<td>Roesler</td>
</tr>
<tr>
<td></td>
<td>HS-6</td>
<td>bbp</td>
<td>1/m</td>
<td>P(v)</td>
<td>Boss</td>
</tr>
<tr>
<td></td>
<td>ECO-BB9</td>
<td>bbp</td>
<td>1/m</td>
<td>P(v)</td>
<td>Boss</td>
</tr>
<tr>
<td></td>
<td>BBFL2SSC</td>
<td>bbp</td>
<td>1/m</td>
<td>WW</td>
<td>Omand</td>
</tr>
<tr>
<td></td>
<td>FLBBCDLC</td>
<td>bbp</td>
<td>1/m</td>
<td>OOI G</td>
<td>Roo</td>
</tr>
<tr>
<td></td>
<td>FLNTU</td>
<td>bbp</td>
<td>1/m</td>
<td>LF</td>
<td>D’Asaro</td>
</tr>
<tr>
<td></td>
<td>Hydrosocat 6</td>
<td>bbp</td>
<td>1/m</td>
<td>P(v)</td>
<td>Boss</td>
</tr>
<tr>
<td></td>
<td>MCOMSC</td>
<td>bbp</td>
<td>1/m</td>
<td>BCG-A</td>
<td>Fassbender</td>
</tr>
<tr>
<td></td>
<td>ECO-BB9</td>
<td>bbp</td>
<td>1/m</td>
<td>P(v), S(v)</td>
<td>Boss, Scott</td>
</tr>
<tr>
<td>Chlorophyll Fluorescence</td>
<td>ALFA</td>
<td>Chl_stimf</td>
<td>mg/m^3</td>
<td>P(i)</td>
<td>Boss</td>
</tr>
<tr>
<td></td>
<td>Eco-Triplet</td>
<td>Chl_stimf</td>
<td>mg/m^3</td>
<td>S(i)</td>
<td>Roesler</td>
</tr>
<tr>
<td></td>
<td>FLBBRTD</td>
<td>Chl_stimf</td>
<td>mg/m^3</td>
<td>S(v), P(v)</td>
<td>Hydro team</td>
</tr>
<tr>
<td></td>
<td>MCOMSC</td>
<td>Chl_stimf</td>
<td>mg/m^3</td>
<td>BCG-A</td>
<td>Fassbender</td>
</tr>
<tr>
<td></td>
<td>BBFL2SSC</td>
<td>Chl_stimf</td>
<td>mg/m^3</td>
<td>WW</td>
<td>Omand</td>
</tr>
<tr>
<td></td>
<td>FLBBCDLC</td>
<td>Chl_stimf</td>
<td>mg/m^3</td>
<td>OOI G</td>
<td>Roo</td>
</tr>
<tr>
<td></td>
<td>FLNTU</td>
<td>Chl_stimf</td>
<td>mg/m^3</td>
<td>LF</td>
<td>D’Asaro</td>
</tr>
<tr>
<td></td>
<td>FLBBRTD</td>
<td>Chl_stimf</td>
<td>mg/m^3</td>
<td>S(v), P(v)</td>
<td>CTD</td>
</tr>
<tr>
<td>CDOM fluorescence</td>
<td>ALFA</td>
<td>cdfm</td>
<td>mg/m^3</td>
<td>P(v)</td>
<td>Boss</td>
</tr>
<tr>
<td></td>
<td>Eco-Triplet</td>
<td>cdfm</td>
<td>mg/m^3</td>
<td>S(i)</td>
<td>Roesler</td>
</tr>
<tr>
<td></td>
<td>HORIBA_JY_Fluoromax4 cdfm</td>
<td>ppb</td>
<td>S(d), P(d)</td>
<td>Nelson</td>
<td></td>
</tr>
<tr>
<td>Volume Scattering Function (λ,φ).</td>
<td>FLNTU</td>
<td>VSF</td>
<td>1/m/sr</td>
<td>LF</td>
<td>D’Asaro</td>
</tr>
<tr>
<td></td>
<td>VSF – 9</td>
<td>VSF</td>
<td>1/m/sr</td>
<td>S(v)</td>
<td>Freeman</td>
</tr>
<tr>
<td></td>
<td>LISST - Deep</td>
<td>VSF</td>
<td>1/m/sr</td>
<td>S(v)</td>
<td>McDonnel</td>
</tr>
<tr>
<td></td>
<td>LISST-100X(B)</td>
<td>VSF</td>
<td>1/m/sr</td>
<td>S(d), P(i)</td>
<td>Gray, Boss</td>
</tr>
<tr>
<td></td>
<td>MVSM</td>
<td>VSF</td>
<td>1/m/sr</td>
<td>S(d)</td>
<td>Gray</td>
</tr>
<tr>
<td></td>
<td>LISST-VSF</td>
<td>VSF</td>
<td>1/m/sr</td>
<td>S(d)</td>
<td>Zhang</td>
</tr>
<tr>
<td></td>
<td>ECO-BB9</td>
<td>VSF</td>
<td>1/m/sr</td>
<td>P(v), S(v)</td>
<td>Boss, Scott</td>
</tr>
<tr>
<td>Particle size distribution</td>
<td>LISST-100X(B)</td>
<td>PSD_DNSD</td>
<td>number/m^3/um</td>
<td>S(d), P(i)</td>
<td>Gray, Boss</td>
</tr>
<tr>
<td></td>
<td>LISST-Deep</td>
<td>PSD_DNSD</td>
<td>number/m^3/um</td>
<td>S(v), P(v)</td>
<td>McDonnel</td>
</tr>
<tr>
<td></td>
<td>Coulter Counter</td>
<td>PSD_DNSD</td>
<td>number/m^3/um</td>
<td>S(d), P(d)</td>
<td></td>
</tr>
<tr>
<td>Variable fluorescence</td>
<td>ALFA</td>
<td>FV_Fm</td>
<td>Unitless</td>
<td>P(i)</td>
<td>Boss</td>
</tr>
</tbody>
</table>
The direct imaging devices, such as those taking images or videos, though in general belong to optical observation and play an important role in achieving the EXPORTS goal, are NOT covered in this Chapter, and can be found in report from the EXPORTS Particle Characterization working group (Chapter 4). As mentioned before, certain optical observations will be used, in conjunction with discrete measurements of biogeochemical stocks, to develop a suite of biogeochemical proxies (Table 2.2, Chapter 2). Ultimate goal is that some of the optical parameters collected here will be used for the development of the ocean color algorithms that can support biological carbon pump research, targeting specifically new generation of hyperspectral satellites (PACE, Werdell et al. (2019)).

Inter-instrument calibration

As visible from the Table 3.2, there was a large number of small instruments measuring backscattering ($b_{bp}$) and Chlorophyll Fluorescence ($Chl\ F$). As these will be the primary instruments used to develop and extrapolate stock measurements of Particulate Organic Carbon and Chlorophyll a, prior to cruise Optics working group conducted a laboratory intercomparison. First, all sensors were taped with electrical tape and immersed in filtered seawater to estimate their dark counts/dark voltage. These dark counts measurements, together with the in-situ profiles with taped sensors (for some instruments) will be ultimately compared with the laboratory (factory) calibrations. Following, $b_{bp}$ sensors were immersed in serial dilution with beads, and $Chl\ F$ sensors were immersed in the serial dilution of algal culture and seawater. Actual bead and phytoplankton concentrations are irrelevant, as the comparison was used to compare the linearity of the instrument response in after-cruise intercalibration.
Chapter 4 EXPORTS Particle Characterization working group

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Overview

High quality measurements of the abundance, composition and size distribution of suspended particles in the water column are central for addressing all three science questions (and associated sub-questions) of the EXPORTS program. An impressive suite of optical and imaging instruments, covering the full spectrum of particle size in the ocean, were deployed for enumeration, sizing and classification of particles (e.g., according to taxonomy or functional groups) during the North Pacific cruise in August-September 2018. These measurements were augmented with (1) metagenomics and DNA barcoding of whole seawater samples (to increase resolution depth of taxonomic composition and functional genes), (2) microscopy/Zooprocess analyses of particles recovered from plankton nets, and (3) characterization of particles collected on polyacrylamide gels in sediment traps.

The instruments and methods that were used to characterize particles differ in their fundamental measurement principles, as well as their size ranges, sensitivities, and selectivity of particle
detection. The comparability of data provided by different instruments and methodologies (e.g., different particle sizers, imaging/microscopy vs. metagenomics, etc.) and the integration of data across many technological platforms have not been fully evaluated yet (but see Boss et al., 2018) and will be examined during the data processing phase. The goal of this document is to provide an overview of the different measurements, to facilitate their integration with emerging research questions, and to facilitate validation of remote sensing algorithms and ecosystem models. Some of the products are in formats easily handled by SeaBASS while data from other sensors/methodologies require integration with other platforms (e.g., imaging, metagenomics).

Summary of measurements to characterize particles:

I. Concentrations (abundances) of particles

<table>
<thead>
<tr>
<th>Particle type</th>
<th>Platform</th>
<th>Units</th>
<th>Contact</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacterioplankton</td>
<td>Epifluorescence microscopy</td>
<td>cellsx10^9 L^{-1}</td>
<td>Carlson, Hansell</td>
</tr>
<tr>
<td>Synechococcus</td>
<td>Influx/Guava flow cytometer</td>
<td>Cells L^{-1}</td>
<td>Graff, Menden Deuer</td>
</tr>
<tr>
<td>Prochlorococcus</td>
<td>Influx/Guava flow cytometer</td>
<td>Cells L^{-1}</td>
<td>Graff, Menden Deuer</td>
</tr>
<tr>
<td>Picoeukaryotes</td>
<td>Influx/Guava flow cytometer</td>
<td>Cells L^{-1}</td>
<td>Graff, Menden Deuer</td>
</tr>
<tr>
<td>Nano- and microphytoplankton (total and by taxon)</td>
<td>IFCB</td>
<td>Cells mL^{-1}</td>
<td>Sosik/Roesler, Karp-Boss/Boss</td>
</tr>
<tr>
<td>Zooplankton abundance (by size and taxon)</td>
<td>MOCNESS net/ ZOOSCAN</td>
<td>Individuals m^{-3}</td>
<td>Maas/ Steinber</td>
</tr>
<tr>
<td>Non-living particles</td>
<td>UVP</td>
<td>Individuals m^{-3}</td>
<td>McDonnell, Boss/Karp-Boss</td>
</tr>
<tr>
<td>Zooplankton (by taxon)</td>
<td>UVP</td>
<td>Individuals m^{-3}</td>
<td>McDonnell, Boss/Karp-Boss</td>
</tr>
</tbody>
</table>

Other biomass measurements:

Cphyto (Influx flow cytometer; Graff)
Nphyto (Influx flow cytometer; Graff)
POC (Nelson, Graff, Roesler)
PON (Nelson, Graff, Roesler)

II. Particle Size Distribution (PSD)

Suspended particles in the ocean span 5 orders of magnitude in size, ranging from a few nm to large aggregates that reach a few cm in dimension. Particle size is often expressed in terms of an equivalent spherical diameter which is derived from either the particle’s volume or its cross-sectional area. PSD is usually derived by counting the number of particles within a given size class and normalizing that number by the width of the size class and the volume sampled; thus, accurate measurements of both size and concentrations are needed. Different instruments for
detecting and sizing particles in seawater are based on different principles and sensitivities to size descriptors (Table 4.2) and encompass different parts of the particle size spectrum (Figure 4.1).

### Table 4.2 Commercially available particle sizing instruments and their measurement principles, leading to potential discrepancies.

<table>
<thead>
<tr>
<th>Approach</th>
<th>Measurement</th>
<th>Principle associated particle characteristic</th>
<th>Example instruments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrical impedance</td>
<td>resistivity</td>
<td>Volume</td>
<td>Coulter Counter, Elzone Counter</td>
</tr>
<tr>
<td>Optical</td>
<td>light scattering</td>
<td>axis dimension (e.g. diameter)</td>
<td>MVSM, LISST, flow cytometers</td>
</tr>
<tr>
<td>Imaging</td>
<td>image</td>
<td>cross-sectional dimensions</td>
<td>IFCB, FlowCAM, UVP, ZooScan</td>
</tr>
</tbody>
</table>

Figure 4-1 Particle size distribution (modified from Stemmann and Boss 2012) and the different PSD measurements during Exports. Identification of overlapping regions between instruments and evaluation of associated uncertainties will allow to reconstruct PSD across a broad range of particle size.

Measurements of PSD will be obtained at different spatial and temporal resolutions, depending on the instrument and its mode of deployment (Table 4.3).
Table 4.3 A summary of PSD measurements, sampling frequency, and mode of deployment.

<table>
<thead>
<tr>
<th>PSD measurement</th>
<th>Platform</th>
<th>Mode of operation/frequency of sampling</th>
<th>Size range</th>
<th>Contact for data products</th>
</tr>
</thead>
<tbody>
<tr>
<td>ViewSizer</td>
<td>Survey ship</td>
<td>Discrete samples from Niskin bottles</td>
<td>~ 100-700 nm</td>
<td>Zhang</td>
</tr>
<tr>
<td>Multispectral Volume Scattering Meter (MVSM)</td>
<td>Survey ship</td>
<td>Discrete samples from Niskin bottles</td>
<td>~ 0.02 - 200 μm</td>
<td>Zhang</td>
</tr>
<tr>
<td>Flow cytometer (Heidi?)</td>
<td>Survey ship</td>
<td>discrete samples from in line and Niskin bottles</td>
<td>0.5 - 80 μm</td>
<td>Sosik/Roesler</td>
</tr>
<tr>
<td>Flow cytometer (Influx)</td>
<td>Process ship</td>
<td>Discrete samples from Niskin bottles</td>
<td>~ 0.5- 80 μm</td>
<td>Graff</td>
</tr>
<tr>
<td>Flow cytometer (Guava)</td>
<td>Process ship</td>
<td>Discrete samples from Niskin bottles</td>
<td>~ 1- 30 μm</td>
<td>Menden-Deuer/Rynearson</td>
</tr>
<tr>
<td>FlowCAM</td>
<td>Process ship</td>
<td>-Subsamples from marine snow catcher deployments -Subsamples from zooplankton net tows</td>
<td>~ 10-200 μm</td>
<td>Menden-Deuer/Rynearson/Passow</td>
</tr>
<tr>
<td>IFCB</td>
<td>Survey ship &amp; Process ship</td>
<td>Inline: surface (5mL/20 min) and discrete sample from Niskin bottles</td>
<td>~5-150 μm (minimum dimension; elongated targets up to ~300 μm in length)</td>
<td>Sosik/ Roesler/Huot</td>
</tr>
<tr>
<td>LISST 100X (B)</td>
<td>Process ship</td>
<td>Inline: surface; 1 sample per ~ 10 s (with 0.2-μm FSW every hour to account for drift and blank)</td>
<td>1.25-250 μm (based on manufacturer)</td>
<td>Boss</td>
</tr>
<tr>
<td>LISST DEEP (B)</td>
<td>Survey ship &amp; Process ship</td>
<td>Profiling mode</td>
<td>1.25-250 μm (based on manufacturer)</td>
<td>McDonnell/Siegel</td>
</tr>
<tr>
<td>UVP</td>
<td>Survey ship</td>
<td>CTD rosette. Volume imaged: ~ 1 L</td>
<td>&gt;50 μm</td>
<td>McDonnell/Karp-Boss</td>
</tr>
<tr>
<td>Gel traps</td>
<td>Process ship</td>
<td>NBST and STT, 5 discrete depths, 3-5 day flux integrations</td>
<td>~10-10000 μm</td>
<td>Durkin</td>
</tr>
<tr>
<td>Zooscan (MOCNESS)</td>
<td>Process ship</td>
<td>Discrete samples from MOCNESS</td>
<td>&gt; 200 μm for MOCNESS Whatever we image! (UVP)</td>
<td>Maas/Steinberg</td>
</tr>
</tbody>
</table>

Other size proxies:
Spectral slope of beam attenuation and scattering (links to optics working group)
Size groups derived from HPLC (pico-, nano-, micro-)
Size fractionated optical properties (in line ACS, $b_{bp}$, fluorometry)
### III. Composition: morphology and genomics

**Table 4.4 Approaches used to study the community/particle composition during EXPORTS.**

<table>
<thead>
<tr>
<th>Approach</th>
<th>Platform</th>
<th>Mode of operation/frequency of sampling and sampling volume</th>
<th>Target organisms/particles</th>
<th>Information</th>
<th>Contact</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNA sequencing</td>
<td>Process &amp; Survey</td>
<td>filtered biomass from 0.2-10 L water from Niskin bottles, flow through system &amp; marine snow catchers.</td>
<td>Whole community (&gt;0.2 µm) and size fractionated (0.2-5 µm &amp; &gt;5 µm)</td>
<td>Taxonomy, prokaryotes &amp; eukaryotes</td>
<td>Gifford (16S, metagenomics), Cassar (18S), Rynearson (18S), Durkin (18S), Siegel (18S), Jenkins (18S), Santoro (16S), Carlson (16S)</td>
</tr>
<tr>
<td>IFCB</td>
<td>Process &amp; Survey</td>
<td>Inline: surface (5mL/20 min) and discrete sample from Niskin bottles</td>
<td>Phytoplankton and associated chlorophyll containing organisms and particles</td>
<td>Taxonomy, functional groups/trait, Abundance Biovolume (and other size measures)</td>
<td>Sosik Karp-Boss</td>
</tr>
<tr>
<td>FlowCAM</td>
<td>Process</td>
<td>Subsamples of marine snow catcher deployments. Zooplankton:63 micron net tows - 2 day/night pairs per epoch</td>
<td>Chl-a containing organisms/particle; limited zooplankton composition 63-200 µm</td>
<td>Taxonomy, functional groups/trait, Abundance Biovolume (and other size measures)</td>
<td>Menden-Deuer, Rynearson, Passow, Maas, Steinberg</td>
</tr>
<tr>
<td>UVP</td>
<td>Process &amp; Survey</td>
<td>CTD rosette. Volume imaged: ~1 L</td>
<td>Zooplankton, particularly gelatinous zooplankton, Rhizaria, large aggregates, marine snow, and fecal pellets</td>
<td>Taxonomy (course resolution), functional groups/trait, Abundance Biovolume (and other size measures)</td>
<td>McDonnell Karp-Boss</td>
</tr>
<tr>
<td>Microscopy</td>
<td>Process</td>
<td>200ml from Niskin bottles (focus on grazing except T0 and TF)</td>
<td>Phytoplankton (&gt;5 µm) and microzooplankton</td>
<td>Taxonomy, functional groups/trait, Abundance, Biovolume</td>
<td>Menden-Deuer</td>
</tr>
<tr>
<td>Microscopy (MOCNESS, other nets)</td>
<td>Process</td>
<td>MOCNESS: 6 day/night paired tows Other nets: variable-animals used in experiments</td>
<td>Zooplankton (metazoans, primarily &gt;200 um)</td>
<td>Taxonomy, functional groups/trait, Abundance, Biovolume</td>
<td>Steinberg</td>
</tr>
<tr>
<td>Zooscan (MOCNESS)</td>
<td>Process</td>
<td>Variable</td>
<td>Zooplankton (metazoans, primarily &gt;200 um)</td>
<td>Taxonomy, functional groups/trait, Abundance Biovolume (and other size measures)</td>
<td>Maas/ Steinberg</td>
</tr>
<tr>
<td>RNA later traps</td>
<td>Process</td>
<td>Flux reaching traps</td>
<td>eukaryotes, bacteria, archaea in sinking particles</td>
<td>Taxonomy (18S and 16S)</td>
<td>Durkin (18S) and Santoro (16S)</td>
</tr>
<tr>
<td>Gel traps (microscopy)</td>
<td>Process</td>
<td>Flux reaching traps</td>
<td>phytoplankton and protists</td>
<td>Taxonomy</td>
<td>Durkin</td>
</tr>
<tr>
<td>Gel traps (DNA)</td>
<td>Process</td>
<td>individual particles picked out from gel traps</td>
<td>eukaryotes, bacteria, archaea in aggregates, fecal pellets</td>
<td>Taxonomy (18S and 16S)</td>
<td>Durkin (18S) and Santoro (16S)</td>
</tr>
</tbody>
</table>
A combination of imaging tools (IFCB, FlowCAM, UVP, Zooprocess), microscopy, and DNA sequence analysis provides information on community composition for both eukaryotes and prokaryotes. The taxonomic depth that can be achieved with any one of the approaches depends on the size range of the target organisms, the degree of morphological variation among species, and the level of detail available in databases or taxonomic identification guides. The ability to capture and detect rare or delicate organisms depends on the volume analyzed and the mode of sampling (e.g., niskin vs. flow through). The integration of different approaches, with their different strengths, will likely yield a thorough understanding of plankton composition. Details are described in Table 4.4.

**Other sources of information for particle characterization**

HPLC - functional groups (pico, nano, micro phytoplankton)

Flow cytometry - functional groups (pico-eukaryotes, *Synechococcus* and *Prochlorococcus*)
The overall goal of the EXPORTS program is to develop a predictive understanding of the export and fate of global ocean primary production (PP) and its implications for present and future climates. To accomplish this, a coordinated, process-oriented approach will be taken that includes a robust field campaign where the majority of parameters known to influence the formation, transformation and fate of organic matter synthesized via net primary production (NPP) will be measured. Although quantification of the stock of PP and the amount of flux in various pathways provides an instantaneous snapshot of the system’s state, to move on to predictive modeling, it is necessary to determine the rates at which PP is accumulated, transformed, remineralized and sinks to depth as well as the chemical and hydrographic parameters modulating these rates.

The objective of this document is to describe the EXPORTS Science Team’s conceptual framework and methods for determining these rates within an ecosystem/carbon cycle state. This will serve as a resource for identifying transformations and modulating factors that are not well characterized by the EXPORTS Science Team, and will hopefully serve as a resource for future and international programs exploring flux. In section A we describe the major processes.
that contribute to the transfer of carbon in the upper water column. These have been summarized in a figure that details the rate based processes depicted in the EXPORTS “wire diagram” (Figure 5.1). In section B, descriptions of rate measurements determined in the EXPORTS program are provided in the context of whether they are involved in the formation, alteration or remineralization of organic matter. These have been divided into five categories, which are: 1) Primary Production, 2) Respiration, 3) Organic carbon transformations and 4) Aggregation/Sinking and 5) Nutrient Uptake. The protocol briefs then provide detailed descriptions of each method, their uncertainties, key data products, and related references to facilitate data management, coordination and transparency. Rate measurements are given as a change in standing stock of a particular parameter (e.g., phytoplankton biomass, DOC, etc.) over time and can be used to estimate turnover and/or fluxes.

Overview

Primary production by phytoplankton fixes approximately 50 Pg carbon (C) annually (Field et al., 1998). Rapid consumption and remineralization of this food source removes most of this carbon, leaving only an estimated 5 to 12 Pg (10 to 24%) for export out of the euphotic zone (Li and Cassar, 2016; Siegel et al., 2016). Export of the remaining carbon from the surface waters is mediated by aggregation and sinking processes in both the euphotic zone (EZ) and the deeper water column, which is referred to as the twilight zone (TZ) in the EXPORTS paradigm. In this midwater region repackaging and consumption of surface flux by midwater organisms modify both the total carbon export as well as the aggregation and sinking rates. This net reduction in vertical transfer of C is referred to as attenuation.

Understanding and predicting central paradigms of ocean ecosystem function, including export production and responses to environmental change requires recognition of grazing and inclusion of this dominant loss factor in global biogeochemical, ecosystem and cross-biome comparison models (e.g. Stock and Dunne (2010)). In all ocean ecosystems, grazing by herbivorous protists (i.e. microzooplankton) constitutes the single largest loss factor of marine primary production (PP) and alters the abundance and size spectra of particles (Banse, 2013; Worden et al., 2015; Steinberg and Landry, 2017). Global estimates of herbivorous protist grazing across major biogeochemical provinces in the ocean range from 49% to 77% primary production removed, with an overall average of 62% (Schmoker et al., 2013). Currently the trophic linkages and transfer efficiencies among microzooplankton, mesozooplankton and macrozooplankton remain poorly characterized, making analyses of grazing rate and secondary production of the mesozooplankton a large source of uncertainty (Steinberg and Landry, 2017). The transfer to higher trophic levels is even more poorly constrained (Burd et al., 2016).

Once grazed, around half of the C consumed by the zooplankton is respired as CO₂ while the remainder contributes to growth and secondary production. In surface waters respiration acts as remineralization of PP, while midwater zooplankton respiration after consumption of particle flux is a source of flux attenuation. In contrast, the respiration of diel vertical migratory zooplankton, which consume organic particles in the surface waters at night and metabolize (i.e., respiring, excreting, egesting) surface-ingested POM in the mesopelagic zone during the day, is one of the major pathways of export flux. This component of the biological pump is referred to as “active” transport, as opposed to the passive sinking from surface waters of particles such as
fecal pellets. The largest component of the active transport of C by vertical migrators is the respiratory flux (respiration of CO₂ at depth), which scales positively with migrant animal biomass. Current estimates suggest that 31-40% of the PP in the euphotic zone is respired by microzooplankton, 21% by mesozooplankton and 50-90% is respired by bacteria (Anderson and Ducklow, 2001; Rivkin and Legendre, 2001). Mesozooplankton respiration through the full water column accounts for an estimated 17–32% of global PP (Steinberg and Landry, 2017).


Subgroup Primary Production

Primary production in marine pelagic environments is the amount of dissolved carbon dioxide fixed by photosynthesis over time. In the euphotic zone, the vast amount of primary production is carried out by autotrophic single-celled prokaryotes and eukaryotes (collectively termed phytoplankton). Rates of primary production in the ocean vary spatially and temporally and are largely a function of the physical and chemical environment that influences phytoplankton composition, abundance and physiology.

A. Gross primary productivity (GPP): GPP is the total quantity of organic carbon that is fixed by photosynthesis.

B. Net primary productivity (NPP): NPP is equal to gross primary productivity minus losses due to respiration by autotrophs.

C. Net community productivity (NCP): NCP is equal to GPP minus community respiration (CR), or NPP minus heterotrophic respiration (HR). Negative NCP (i.e. net heterotrophic conditions) is associated with remineralization of POC and DOC. Conversely, a positive NCP (i.e. net autotrophic conditions) leads to a net production of POC and DOC at the ocean surface, which can either accumulate or be exported. Accordingly, NCP is equal to export production and new production (see below) when the POC and DOC inventories are at steady-state.

\[
gpp = \frac{\text{GPP}}{\text{NPP}} \cdot \text{AR} \cdot \text{Organic matter} + O_2
\]

\[
\text{Export production} = \text{NCP} - \text{MLD} \times \frac{d(\text{POC} + \text{DOC})}{dt}
\]

D. New production: New production refers to primary production fueled by allochthonous (external) nutrient supply (e.g., vertical mixing of nutrient to the ocean surface, atmospheric nutrient deposition, and N_2 fixation). Under steady-state conditions, new production can also be used as a proxy for export production (Eppley and Peterson 1979).

E. Regenerated production: Regenerated production refers to primary production fueled by the remineralization of nutrients at the ocean surface. Taken together, the ratio of new production divided by the total production (i.e., new plus regenerated production) yields the f-ratio, which is often used as a way to describe the export flux of organic matter from the surface ocean by way of the carbon biological pump.

F. Diatom production: Diatom production is the amount of dissolved inorganic carbon fixed specifically by diatom photosynthesis.

Primary productivity will be measured using multiple approaches that will estimate gross and net primary productivity as well as net community productivity. Both the incorporation of isotopes
(stable and radioactive) over short (6 hr) and long (24 hr) incubations will be used to measure uptake of dissolved inorganic carbon. In addition, the production of biogenic gases (e.g., oxygen) will also be measured and used to estimate PP. New production is estimated through stable isotope NO$_3$ incorporation. Diatom-specific production will be estimated through uptake of silicon.

**Subgroup Respiration**

Respiration is the cellular metabolic processes used to generate energy via the oxidation of organic matter and the release of waste products. In the context of EXPORTS the term is used to characterize the aerobic conversion of organically fixed C (PP) to energy (in the form of ATP) and CO$_2$. It is one of the major transformation pathways of PP and, depending on where it occurs in the water column, can contribute to flux or to the attenuation of flux.

Here we divide respiration into three categories: bacterial, community, and zooplankton.

A. **Bacterial Respiration (BR):** BR is the remineralization of organic matter by free-living microbes and microbes associated with particles passing through a 5 µm pore size filter.

B. **Community Respiration (CR):** CR is the remineralization of organic matter by all members of the bacterioplankton, phytoplankton, and zooplankton communities. This measurement provides a holistic approximation of total heterotrophic respiration (HR). This measure underestimates mesozooplankton, fish and nekton contributions to community respiration due to methodological constraints.

C. **Zooplankton Respiration (ZR):** Estimates of total ZR are made of the >200 µm zooplankton community sampled comparing daytime and nighttime vertically stratified net tows (MOCNESS). Smaller metazoans that are not retained in the nets (<200 µm) are sampled with a 63 µm mesh net and the community composition and biovolume will be used to calculate the ZR of this size-fraction. Very large mobile zooplankton, fish and nekton respiration are underestimated by current EXPORTS protocols.

Bacterial and Community respiration will be measured in two complimentary ways: O$_2$ drawdown assays and DOC remineralization assays. Zooplankton respiration will also be assessed in two ways 1) analysis of the Electron Transport System (ETS) enzyme activity and 2) converting community composition (from ZooScan analysis) and biomass measurements to community respiration. The latter will be accomplished using experimentally measured individual respiration rates as well as the published equations based on temperature and scaling coefficients (Ikeda 2014).

**Subgroup Organic Carbon transformations**

Carbon transformations are the conversion of NPP into the biomass of other prokaryotic or eukaryotic organisms via heterotrophy. This conversion can have different efficiencies depending on the carbon source (i.e. food quality)–characterizing the rate of growth based on photosynthetic community and DOM pool is critical for understanding how community composition influences export versus recycling.

A. Bacterial production
We will determine the flux of the most labile fraction of DOC to bacterioplankton (i.e., bacterial carbon demand; BCD) from independent measurements of heterotrophic bacterial production (BP) and estimates of bacterial growth efficiency (BGE). The most labile fraction of DOC (LDOC) is rapidly consumed, supporting the metabolic energy and nutrient demands of heterotrophic prokaryotes, with turnover rates on time scales of hours to days. Bacterial carbon demand (BCD) best represents the flux of LDOC, with greater than 50% of net PP flowing through the labile DOM pool on a daily basis (Ducklow, 1999; Williams, 2000). Determining the flux of LDOC will constrain the respiratory fate of a large fraction of NPP.

We will also conduct DOC remineralization experiments to directly assess the fraction of the accumulated DOC that is available to bacterial production vs respiration. Simultaneous measurements of the change in bacterial biomass and DOC in microbial dilution cultures will allow us to estimate bacterial growth efficiency (BGE) of the natural microbial assemblages (Carlson et al., 2004) as follows:

\[
\text{BGE} = \frac{\Delta BB}{\Delta DOC} \quad (2)
\]

where \(\Delta BB\) and \(\Delta DOC\) represent the change in bacterial carbon biomass and DOC removal through stationary growth in the microbial remineralization experiments. Estimates of water column BCD (0-500 m), or the gross flux of carbon to heterotrophic bacterioplankton, will be determined from BP profiles (\(^3\)H-Leu incorp) and BGE estimates from the EZ and MZ determined on the Lagrangian cruise: \(\text{BCD} = \text{BP} / \text{BGE} \quad (3)\)

B. Secondary production/Grazing

To quantify predation and secondary production effects on export production, particle abundance and size distribution will be measured as a function of co-occurring environmental and biological conditions. Herbivorous grazing rates on specific groups of phytoplankton in the euphotic zone will be quantified using a two-point dilution method (Morison and Menden-Deuer, 2017) and flowcytometry. To achieve high-resolution grazing rate measurements, on time scales similar to the capacity of autonomous or remote sensors, we are developing novel ecogenomic tools that will open an entirely new field for remotely sensed products that assess zooplankton grazing (see proposed genomic methods by Menden-Deuer). Moreover, long-term incubations where predators have been concentrated to increased grazing signal will be used to resolve feeding potential and transformations of organic matter below the euphotic zone. Secondary production and feeding of higher trophic levels (mesozooplankton, nekton and fish) are not measured by EXPORTS protocols.

Subgroup Aggregation/Sinking rates

As particles sink through the water column they can coagulate (aggregate) to form (larger) aggregates, which may be fragmented (disaggregated) due to both physical and biological processes. The sinking of aggregates serve as one of the main export flux pathways, while disaggregation is a major driver of attenuation processes in the midwater. Aggregates are complex particles made up of phytodetritus, resident eukaryotic and prokaryotic communities, fragmented fecal pellets, organic polymers, detritus, mineral particles, and other compounds that become connected as they sink through the water column. The EXPORTS interest in these processes are to determine the size class and composition of various particle types throughout
the water column, and to estimate their sinking, aggregation and disaggregation rates as well as other attenuation mechanisms (grazing). Aggregation will be studied both by comparing particles captured at various depths (trap, bottle, Marine Snow Catcher) as well process studies on various particle types. Sinking and respiration rates of particular particle types will be measured, while disaggregation and loss due to feeding will be assessed using grazing studies.

Subgroup Nutrient (Si, Fe, N) Uptake rates

It is well known that environmental factors can modify the physiological rates of all of the processes of interest to the EXPORTS program. The N Pacific is known to be nutrient (iron) limited, so it is valuable to determine how changes in the nutrient field will influence the physiology and associated rates of the primary producers. The primary hypothesis to be tested during EXPORTS is that differences in the type (Si, Fe, N) and degree of nutrient stress experienced by co-occurring phytoplankton (diatom) taxa can predict the export pathways that each will follow through the food web. Short term incubations using isotopically traced nutrients, will be conducted to measure changes in the rate of primary productivity and assess stoichiometry of nutrient use and a second set of longer duration incubation experiments will be carried out to find transcriptomic markers of nutrient stress. These two process studies will be linked with analyses of the vertical structure of the natural community composition and physiology to connect measured rates and transcriptomic tracers of nutrient stress with observed changes in natural assemblages as a function of depth and nutrient fields.
Table 5.1 Summary of Protocol Briefs. This table provides a cross reference between the rate measured, the rate depicted on the rates wiring diagram (Figure 5.1) and the protocol descriptions associated with this working group and associated subgroups (SGs) (P = production, R = respiration, T = transformation (Corg->Corg), A = aggregation, N = Nutrient Rate Processes).

<table>
<thead>
<tr>
<th>Rate</th>
<th>Wire Diagram Abbreviation</th>
<th>SG:</th>
<th>Lead PI</th>
<th>Contact Person</th>
<th>Method protocol</th>
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<td>NCP</td>
<td>P,R</td>
<td></td>
<td>Lee</td>
<td>Nicholson</td>
<td>O2, NO3, rate of change of C from backscatter</td>
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<tr>
<td>NCP</td>
<td>P,R</td>
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<td>Estapa</td>
<td>Omand</td>
<td>diel cp and oxygen</td>
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<td>NCP</td>
<td>NCP1</td>
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<td>Cassar</td>
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<td>Marchetti</td>
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<td>Marchetti</td>
<td>Marchetti</td>
<td>mRNA sequencing of poly-A selected genes</td>
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<td>Metatranscriptomics of prokaryotic plankton</td>
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<td>Marchetti</td>
<td>Gifford</td>
<td>mRNA sequencing of genes</td>
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<td>GPP</td>
<td>P</td>
<td>Marchetti</td>
<td>Marchetti</td>
<td>H13CO3, 6 hr incubation</td>
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<tr>
<td>GPP/CR</td>
<td>P</td>
<td></td>
<td>Lee</td>
<td>Nicholson</td>
<td>diel oxygen</td>
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<td>Halsey</td>
<td>H14CO3 - 24 hr incubation</td>
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<td>Marchetti</td>
<td>Marchetti</td>
<td>H13CO3, 24 hr incubation</td>
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<td></td>
<td>Lee</td>
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<td>Menden-Deuer</td>
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<td>Chlorophyll and fluorescence changes 24 hour</td>
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<td>Nicholson</td>
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<td>function of size and temperature</td>
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<td>Zooplankton respiration and excretion</td>
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<td>R</td>
<td>Steinberg</td>
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<td>Carlson</td>
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<td>Passow/Carlson</td>
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<td>O2 change incubation + gifford</td>
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<td>Process</td>
<td>Code</td>
<td>Type</td>
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<td>Methods</td>
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<td>T</td>
<td>Siegel, Passow/Steinberg</td>
<td>Incubations for MS grazing and fragmentation rate of zooplankton on marine snow</td>
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<tr>
<td>Bacterial Production</td>
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<td>Carlson</td>
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<td>Phytoplankton loss processes, via microzooplankton/protist grazing</td>
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<td>Incubation vs. in situ Chl a accumulation rates</td>
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<td>Grazing rate marker</td>
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<td>Mesopelagic grazing rate potential</td>
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<td>Mesozooplankton grazing</td>
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<td>T</td>
<td>Steinberg/Menden-Deuer, Menden-Deuer/Steinberg/Mass</td>
<td>Incubation</td>
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<td>NPP</td>
<td>VSi/DPP</td>
<td>P,N</td>
<td>Jenkins, Brzezinski</td>
<td>32Si uptake 0.6-5.0 &amp; 5.0 um size fractions 24h</td>
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<td>NPP</td>
<td>VSi/DPP; VFe (NPP9); nPP; GPP/NPP 1-8</td>
<td>P,N</td>
<td>Jenkins, Brzezinski</td>
<td>Nuri ent stress assessments. Effects of Fe and Si on rates of C, Si and N use (dawn to dusk)</td>
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<td>Nit</td>
<td>R,T</td>
<td>Santoro</td>
<td>15NH4 addition</td>
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<td>NCP (POC + DOC)</td>
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<td>Fassbender</td>
<td>Fassbender</td>
<td>BGC tracer budgets (DIC, TA, NO3, O2)</td>
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<tr>
<td>CaCO3 Production (PIC)</td>
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<td>Fassbender</td>
<td>Fassbender</td>
<td>BGC tracer budgets (DIC, TA, NO3, O2)</td>
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<tr>
<td>NPP</td>
<td>VSi/DPP</td>
<td>N</td>
<td>Jenkins, Brzezinski</td>
<td>32Si uptake 0.6-5.0 &amp; 5.0 um size fractions 24h</td>
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<td>Jenkins</td>
<td>57FeCl3 uptake into diatom size fraction</td>
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<tr>
<td>particle aggregation</td>
<td>Sum of A1, A2, A3, A4 and A6</td>
<td>A</td>
<td>Lam</td>
<td>Lithogenic particle concentration analysis followed by inverse modeling</td>
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<tr>
<td>particle disaggregation</td>
<td>Sum of A5 and physical &amp; bacterial diagggregation (not in the diagram)</td>
<td>A</td>
<td>Lam</td>
<td>Lithogenic particle concentration analysis followed by inverse modeling</td>
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<td>R,T</td>
<td>Lam</td>
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<tr>
<td>sinking velocity of aggregates</td>
<td>Sum of S1, S2, and S3</td>
<td>A</td>
<td>Lam</td>
<td>Lithogenic particle concentration analysis followed by inverse modeling</td>
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Overview

Critical to the EXPORTS program goal of estimating biological carbon fluxes from satellite observations is understanding of mechanisms controlling the magnitude and attenuation of those fluxes to depth in the ocean. The main goal of this document is to describe the EXPORTS Science Team’s observational methods that are used to distinguish between (i) biological and physical export pathways out of the euphotic zone and (ii) flux pathways below the euphotic zone. It is our intention that this document will serve as a resource not only for Science Team members, but also for other scientists contributing to other programs with similar goals. This report also outlines a plan for integrating and performing internal consistency checks (i.e., achieving closure) among the various measurements of biological carbon fluxes, and gives an overview of observational and model uncertainties. Thus we hope it will also be a useful resource for data managers. The remainder of this overview defines key concepts and their
relationships to the main EXPORTS science questions. The document then summarizes measurement techniques contributing to determination of each of the five export pathways, and describes the plan for achieving measurement-measurement closure. Finally, a series of protocol briefs describe each method, its uncertainties, key data products, and related references.

The biological and physical pathways that govern the export of carbon from the euphotic zone and its attenuation with depth vary across space and time in the ocean. The EXPORTS program’s overarching scientific goal is to determine how satellite-observable surface properties of both the ocean ecosystem and ocean circulation control the relative importance of these pathways. The EXPORTS Science Plan defines five export pathways:

1. Gravitational settling of intact phytoplankton cells
2. Gravitational settling of aggregates
3. Gravitational settling of zooplankton products
4. Active transport of carbon by diurnally or ontogenetically migrating zooplankton
5. Net vertical transport of suspended particulate or dissolved carbon by physical and microbial processes

The EXPORTS “wiring diagram” (Figure 6.1) illustrates the relationships among these pathways, and the expected variations in the relative importance of the pathways in different locations and times.
The goal of the EXPORTS field campaigns is to characterize export via each of these pathways over operationally-defined time periods – referred to as “epochs” or “ecosystem and carbon cycling (ECC) states”, equivalent to the time necessary for sinking particles to exit the euphotic zone and enter sediment traps in the upper 500 m. In the North Pacific field campaign, each of the measurement techniques described below was repeated during three 8-day states. Work was conducted aboard two ships: a “Process Ship” operating in a water parcel-following Lagrangian mode, and a “Survey Ship” in a gridded survey mode. Additionally, autonomous
platforms (gliders and floats or “AUVs”) were used to increase the spatiotemporal coverage of some measurements.

Flux measurements are made within a vertical reference frame where the base of the “well-lit surface ocean” (as defined in the EXPORTS Science Questions, below), or “export reference depth”, is theoretically defined as the depth where new production equals export production (Buesseler and Boyd, 2009). Prior to obtaining direct observations of production, we operationally estimate this as the depth where in vivo chlorophyll fluorescence drops to 10% of its subsurface maximum (Owens et al., 2015). Flux measurements are made at the export reference depth as well as 50 m and 100 m below it, in order to capture processes that typically result in rapid flux attenuation at the shallowest depths. Diel vertical migration of zooplankton influences multiple export pathways and so a fourth measurement depth targets the deep biomass maximum. The fifth and deepest flux measurement may be constrained by the operational limits of sediment traps but targets seasonal zooplankton migrators where possible.

The main EXPORTS Science Questions are:

1. How do upper-ocean ecosystem characteristics determine the vertical transfer of carbon from the well-lit surface ocean?
2. What controls the efficiency of vertical transfer of carbon below the well-lit surface ocean?
3. How can the knowledge gained be used to reduce uncertainties in contemporary and future estimates of the export and fates of global ocean net primary production (NPP)?

Observations of export pathways at the base of the euphotic zone or mixed layer contribute directly to EXPORTS Science Question 1 by establishing the magnitude of the vertical transfer of carbon from the well-lit surface ocean. By further establishing which pathways are responsible for the vertical transfer, these observations allow connections to be made to characteristics of the upper ocean ecosystem. Similarly, measurements at depths below the euphotic zone and mixed layer directly address Science Question 2 by quantifying the transfer efficiency of different pathways. Observations of the relative importance of the different export pathways under different surface ecosystem states will be useful in addressing Science Question 3.

Summary of measurements by export pathway

Pathway 1 (sinking, single phytoplankton cells) is characterized using high magnification microscope images of polyacrylamide gel collectors on sediment traps, and estimating C per cell using literature values and shipboard measurements (see “Stocks and Proxies” and methods for determination of phytoplankton carbon). The presence or absence of phytoplankton cells in the non-sinking, slow sinking or fast sinking category of particles (e.g. the partitioning of cells between these pools) as collected with the Marine snow catcher addresses the importance of this pathway.

Pathway 2 (sinking aggregates) is characterized by synthesizing four methods. As with Pathway 1, microscope images of gel traps are used to enumerate, visually characterize, and determine the size distribution of sinking aggregates. The biological composition of the aggregates in gel traps are determined by genetic sequencing of representative aggregate types and by the
proportional contributions to bulk traps of amino acids due to phytodetritus, microbes, and fecal pellets. Rapidly-sinking aggregates are also sampled using Marine Snow Catchers and, if present, characterized for size distribution, carbon content, biomineral content, porosity, density, sinking velocity and respiration loss. Respiration rates on sinking particles, determined using RESPIRE traps and experimentally with particles from the Marine Snow Catcher, provide information on the mechanisms driving attenuation of Pathway 2 flux with depth. Profiled Underwater Vision Profiler (UVP) images and LISST particle size distributions are analyzed to identify aggregates and their water column size distribution. Finally, aggregate dynamics and coagulation modeling are used to synthesize the size distributions of suspended and sinking aggregates, their compositions, and physical properties to provide a size-resolved estimate of aggregate-mediated C fluxes as a function of depth.

Pathway 3 (sinking zooplankton products) is determined in three ways. First, zooplankton products are identified in gel traps and their C and N contents are estimated using published and on-board measurements. In addition, zooplankton excretion rate experiments are carried out on board to measure dissolved organic C and N (protocols are documented under Biological Rates working group). The rates determined by these experiments will be applied to measures of zooplankton biomass and community composition to generate estimates of total export via sinking zooplankton products. Partitioning of fecal pellets between non-sinking, slow sinking or fast sinking category of particles, as determined using the marine snow catcher, assesses the relative importance of fecal pellet sinking. Genetic characterization of sinking fecal pellets collected in the gel traps, as well as compound-specific amino acid composition of zooplankton and their food sources, provide qualitative evidence of ecosystem interactions contributing to Pathway 3.

Pathway 4 (zooplankton active transport) is determined by applying analyses of zooplankton metabolic rates to estimates of migratory biomass. First, the depth- and time-dependent distribution of zooplankton biomass are determined using the MOCNESS and UVP, with semi-automated image analysis to identify taxa. Vertical and horizontal variability in zooplankton migratory distributions are characterized qualitatively with acoustics systems on the survey ship, and on the autonomous platforms. These biomass calculations are converted to active transport in three ways, via application of 1) metabolic respiration rates (respiration of CO₂, excretion of DOC, egestion of POC as fecal pellets) determined experimentally on board the ship 2) estimates of metabolic respiration rate calculated from organism size and temperature coefficients and 3) electron transport system enzymatic activity measurements on the MOCNESS samples that are used to estimate the depth-resolved CO₂ production by zooplankton respiration. Both the ETS and metabolic respiration rate protocols are documented by Biological Rates working group.

Pathway 5 (physical transfer of POC and DOC) is comprised of advective and turbulent diffusive transfer of POC and DOC to depth, along and across density surfaces, as well as microbial processes acting upon POC and DOC. Characterization of this pathway requires quantifying POC and DOC stocks and lability, determining rates of microbial utilization, and describing the physical transport and mixing of these pools over a range of spatiotemporal scales, from submesoscale motions up to annual vertical mixing. Certain smaller-scale circulation processes
are parameterized in terms of larger-scale hydrographic and biogeochemical properties
captured by both satellite observations and EXPORTS measurements.

In addition to measurements of single export pathways described above, the EXPORTS
program also utilizes methods characterizing the sum of two or more pathways. For instance,
spikes in profiles of optical scattering and fluorescence serve as a proxy for the abundance of
particles large enough to occlude optical sensing volumes (large aggregates and zooplankton
products) and provide an estimate of fluxes due to the sum of Pathways 2 and 3. Several bulk
methods provide estimates of fluxes due to all sinking particles (the sum of Pathways 1-3),
including bulk fluxes to sediment traps and marine snow catchers, and optical attenuation fluxes
to optical sediment traps and gel traps. Measurements of flux derived from $^{234}\text{Th}$ and $^{210}\text{Po}$
disequilibria represent the sum of Pathways 1-4 -- all sinking-particle and zooplankton-mediated
pathways -- over the preceding weeks ($^{234}\text{Th}$) and months ($^{210}\text{Po}$). Particle stocks collected with
size-fractionated filtration through large volume pumps provide estimates of the composition of
large particles that are likely sinking via pathways 1-4. Inversion of $^{234}\text{Th}$ activities, and organic
and lithogenic particle stocks will provide estimates of aggregation, disaggregation, and
remineralization rates of particulate material present in two operational size classes that
together represent pathways 1-4. Two qualitative but information-rich methods, compound-
specific isotopic analysis of amino acids, and environmental lipidomics, are used to qualitatively
describe sources and biological drivers of particle fluxes in pathways 1-4. $^{210}\text{Po}$ disequilibrium
from $^{210}\text{Pb}$ in seawater will be used in a similar way to $^{234}\text{Th}$, but at a lower spatial resolution, to
estimate fluxes on a seasonal scale. And finally, geochemical budgets measured through long-
term monitoring of $\text{O}_2$, $\text{NO}_3$, and carbon pools from moorings and profiling floats will
independently constrain all 5 pathways over longer seasonal and annual scales.

**Closure of C flux budgets**

By measuring each of the export pathways using multiple methods, as well as measuring
combinations of more than one pathway, we will rigorously constrain uncertainties in flux
estimates. Even in cases where independent methods do not give the same result for a given
pathway, we can use relationships among different techniques to inform our understanding of
driving processes. These measurements and flux estimates will need to be evaluated in the
context of larger-scale characteristics of the ecosystem and circulation as observed from *in situ*
autonomous array and remote sensing platforms during the EXPORTS field program. Table 6.1
summarizes sets of measurements that we can use to constrain uncertainties in this way.
Table 6.1 Possible closure schemes (aka things that should add up), ignoring issues of time/space averaging scales

<table>
<thead>
<tr>
<th>Pathway(s)</th>
<th>Method(s) 1</th>
<th>Method(s) 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 (sinking zooplankton products)</td>
<td>Gel trap fecal pellet/C contents</td>
<td>Biomass distribution/egestion excretion rate experiments</td>
</tr>
<tr>
<td>4 (zooplankton active transport)</td>
<td>Biomass distribution/ETS activity</td>
<td>Biomass distribution/respiration rate experiments</td>
</tr>
<tr>
<td>Sum of 1, 2, 3 (sinking particles)</td>
<td>Bulk fluxes to sediment traps and in marine snow catchers</td>
<td>Sum of independent measurements of pathways 1-3</td>
</tr>
<tr>
<td>Sum of 1, 2, 3 (sinking particles)</td>
<td>Attenuation with depth of sinking particle fluxes in traps and marine snow catchers</td>
<td>Respiration rates on sinking particles from RESPIRE traps and marine snow catcher experiments</td>
</tr>
<tr>
<td>Sum of 1, 2, 3 (sinking particles)</td>
<td>Inversion of bulk fluxes through reconstruction modeling approaches to give particle size, composition, settling velocity -- for bulk fluxes and possibly for pathways 1-3 independently</td>
<td></td>
</tr>
<tr>
<td>Sum of 1, 2, 3 (sinking particles)</td>
<td>Source partitioning from compound-specific stable isotopes, lipidomics</td>
<td></td>
</tr>
<tr>
<td>Sum of 2, 3 (large, sinking particles)</td>
<td>Optical spike fluxes and fluxes of large particles sinking &gt;18 m/d in marine snow catchers</td>
<td>Sum of independent measurements of pathways 2 and 3</td>
</tr>
<tr>
<td>Sum of 1, 2, 3, 4 (sinking particles + zooplankton)</td>
<td>$^{234}$Th flux/C:Th ratios and $^{210}$Po flux/Po:C ratios of large particles</td>
<td>Sum of independent measurements of pathways 1-4</td>
</tr>
<tr>
<td>Sum of 1,2,3,4 (sinking particles+zooplankton)</td>
<td>Inversion of size-fractionated POC, $^{234}$Th, lithogenic particle concentrations and bulk fluxes to estimate bulk aggregation, disaggregation, and remineralization rates</td>
<td>Sum of independent measurements of pathways 1-4</td>
</tr>
<tr>
<td>Sum of all pathways (1-5)</td>
<td>Geochemical budget estimates of NCP (O$_2$/Ar, DIC, NO$_3$ others?)</td>
<td>Sum of independent measurements of pathways 1-5</td>
</tr>
<tr>
<td>Sum of all pathways (1-5)</td>
<td>Biological rate estimates of NCP (GPP-R)</td>
<td>Sum of independent measurements of pathways 1-5</td>
</tr>
</tbody>
</table>

Complementing the pathway-specific estimates of flux described in this document, net community production will be independently determined using geochemical budgets and biological rate measurements. These methods are detailed in the Biological Rates Working Group report. If the overarching hypothesis is correct, that the total carbon flux is the sum of carbon fluxes via the five pathways defined in the EXPORTS Science Plan, then estimates of total export from the surface ocean should be in agreement with NCP measurements when averaged over appropriate scales of space and time. With closure of flux budgets achieved, the importance of each export pathway to the total can be related to the ecosystem characteristics and remote sensing observables of each ECC state.
## Table 6.2 Protocol briefs (This list of protocol briefs may contain some overlaps with other working groups.)

<table>
<thead>
<tr>
<th>Pathway 1</th>
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<tbody>
<tr>
<td>1. Visual cell ID in gel traps</td>
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<table>
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<tr>
<th>Pathway 2</th>
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<tbody>
<tr>
<td>2. Characterization of Sinking Particles from Marine Snow Catcher samples</td>
<td></td>
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<tr>
<td>3. Visual characterization of sinking aggregates in gel traps</td>
<td></td>
</tr>
<tr>
<td>4. Genetic characterization of sinking aggregates in gel traps</td>
<td></td>
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<tr>
<td>5. Particle characterization from UVP profiles (merged with #15, below)</td>
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<tr>
<td>6. Coagulation modeling</td>
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<thead>
<tr>
<th>Pathway 3</th>
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<tbody>
<tr>
<td>7. Visual characterization of zooplankton products in gel traps</td>
<td></td>
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<tr>
<td>8. Genetic characterization of zooplankton products in gel traps</td>
<td></td>
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<tr>
<td>9. Characterization of feces in the different categories collected with the Marine snow catcher</td>
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<tr>
<td>10. Zooplankton fecal pellet production</td>
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<tr>
<td>11. Compound-specific stable isotope analysis of zooplankton</td>
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<tr>
<td>12. Reconstruction of zooplankton product export from gel traps</td>
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<tr>
<th>Pathway 4</th>
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<tbody>
<tr>
<td>13. Day-night MOCNESS tows</td>
<td></td>
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<tr>
<td>14. Zooplankton biomass, abundance</td>
<td></td>
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<tr>
<td>15. UVP characterization of zooplankton abundance (merged with #5, above)</td>
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<tr>
<td>16. Bio-acoustic measurement of zooplankton biomass</td>
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<tr>
<th>Pathway 5</th>
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<tbody>
<tr>
<td>17. Bulk DOC/TDN stocks</td>
<td></td>
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<tr>
<td>18. Bulk POC stocks - reference to BGC stocks WG doc</td>
<td></td>
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<tr>
<td>19. Dissolved combined amino acids</td>
<td></td>
</tr>
<tr>
<td>20. Subduction of bulk DOC</td>
<td></td>
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<tr>
<td>21. Subduction of bulk POC</td>
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<tr>
<td>22. Physical modeling protocol</td>
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<table>
<thead>
<tr>
<th>Large sinking particle pathways (2,3)</th>
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<tbody>
<tr>
<td>23. Flux of large sinking particles from profiles of optical spikes (bb, cp) - reference to Stocks &amp; Proxies WG doc</td>
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<thead>
<tr>
<th>Sinking particle pathways (1,2,3)</th>
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<tbody>
<tr>
<td>24. Bulk fluxes of particles (&amp; constituent elements) to sediment traps</td>
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<tr>
<td>25. Bulk fluxes of particles to optical sediment traps</td>
<td></td>
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<tr>
<td>26. Genetic characterization of bulk sinking particles in sediment traps</td>
<td></td>
</tr>
<tr>
<td>27. Bulk fluxes of rapidly-sinking particles in marine snow catcher samples</td>
<td></td>
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<tr>
<td>28. Reconstruction of export from particle size distributions</td>
<td></td>
</tr>
<tr>
<td>29. Lipidomics markers for particle sources and biological drivers</td>
<td></td>
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<tr>
<td>30. Compound-specific isotopic markers for particle sources and biological drivers</td>
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<thead>
<tr>
<th>Particle and zooplankton pathways (1,2,3,4)</th>
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<tbody>
<tr>
<td>31. Compositions of large particles collected with in situ pumps</td>
<td></td>
</tr>
</tbody>
</table>
32. $^{234}\text{Th}$ survey combined with C(etc).$^{234}\text{Th}$ from large volume pumps

33. $^{210}\text{Po}$ and C(etc).$^{210}\text{Po}$ from large volume pumps

34. Rates from inversion of size-fractionated particle stocks and $^{234}\text{Th}$ activities

Relevant biological rate measurements (see Bio. Rates Working Group Report)

- Total fluxes from geochemical budgets
- Total fluxes from biological rate measurements
References:


EXPORTS Science Definition Team (2016). "EXPORTS Implementation Plan". https://oceanexports.org/docs_implementation_plan.html


Chapter 7 Protocols

Following is the collection of the “one page” protocols, as developed by the EXPORTS Science team during the preparations for the North Pacific Field Campaign. Some of the protocols here were submitted as part of the SeaBASS dataset submission. Final version of the protocols, as well as the update status of the submitted data is can be found at: https://sites.google.com/view/oceanexports/home
**Method:** Derive hyperspectral reflectance measuring downwelling irradiance, and upwelling radiance, in buoy mode.

**Document author and contact info:** Emmanuel Boss ([emmanuel.boss@maine.edu](mailto:emmanuel.boss@maine.edu))

NB: This document is largely based on Rudick (2017).

**Brief description of protocol:**
This buoy approach includes two methods:
1. Above water radiometry with Skylight-Blocked Approach (Fig. 1)

![Fig. 1. Measuring reflectance by measuring above water leaving radiance and downwelling irradiance. Upwelled radiance needs to be corrected for self-shading (by cone and by buoy) and insure that sensor is not touching the water surface nor is it covered with drops. Reflectance is computed from:](#)

\[ R_{rs}(\lambda) = \frac{L_{w}(\lambda, 0^+)}{E_{d}(\lambda, 0^+)} \]

2. Above and below water radiometry (Fig. 2)

![Fig. 2. Measuring reflectance by measuring below water upwelling radiance and downwelling irradiance and propagating the measurement above the surface.](#)
Upwelled radiance needs to be propagated across the interface and corrected for self-shading (by sensor and buoy). This is done using IOPs measured locally or estimated from the upwelled radiance (e.g. Zibordi et al., 2012).

Reflectance is computed from:

\[ R_{rs}(\lambda) = \frac{L_{w}(\lambda,0)}{E_d(\lambda,0)} \]

**Deployment methodologies:**
Buoy is tethered to vessel but needs to be sufficiently far to not be affected by boat shadow.

**Derived parameters:**
Remote sensing reflectance (ocean color) from which additional parameters can be obtained (IOPs as well as biogeochemical parameters).

**Uncertainties and quality control concerns:**

**For both methods:**
Sensor calibration uncertainty.
Long enough averaging to obtain sufficient number of realization (usually about a minute) as well as average wave focusing. If uncertainties due to spatial heterogeneity are of interest, leave system in water for longer.
Need to get far away from large research vessel to avoid effects due to its shadow. Ideally towards the direction of the sun. Avoid data taken with instrument tilted (need high frequency measurements from which tilted measurements can be removed).

**For above water radiometry with Skylight-Blocked Approach:**
The uncertainties and correction associated with self-shading have been estimated by (Shang et al. 2017), who propose also a correction scheme.

**For above and below water radiometry:**

**Key method references**


**Method:** Derive hyperspectral reflectance measuring downwelling irradiance, downwelling radiance and upwelling radiance, removing sky contribution from upwelling radiance.

**Document author and contact info:** Emmanuel Boss (emmanuel.boss@maine.edu)

NB: This document is largely based on Rudick (2017).

**Brief description of protocol:**
Three measurements of radiance are made including downwelling irradiance, downwelling radiance and upwelling radiance (Fig. 1).

![Figure 1. Schematic of above water radiometry with sky radiance measurement and skyglint removal (from Rudick, 2017).](image)

$L_d$ represent the sky contribution to upwelling radiance in the direction of the downlooking radiometer, $L_u$ the downwelling radiance in the symmetric direction to the viewing angle as will contribute if the ocean was perfectly flat. $E_d$ is the downwelling irradiance.

The spectrum of the water leaving radiance is estimated from:

\[
L_w(\lambda; \theta_v, \phi) = L_u(\lambda; \theta_v, \phi) - \rho_d(\lambda; \theta_v, \phi)
\]

and the remote-sensing reflectance ($R_{rs}$) from:

\[
R_{rs}(\lambda) = \frac{L_u(\lambda; \theta_v, \phi)}{E_d(\lambda)}.
\]
\( \rho \) denotes the fraction of skylight that is reflected into the down-looking sensor, \( \varphi \) the azimuth angle \( \theta_{\nu-} \), the downward zenith angle and \( \theta_{\nu+} \). Mobley (1999) recommends \( \varphi = \mp 135^\circ \) and \( \theta_{\nu\pm} = 40^\circ, 120^\circ \).

**Deployment methodologies:**
On board the deck of the ship with compass and tilt sensors and automatic mechanism to insure correct pointing directions.

**Derived parameters:**
Remote sensing reflectance (ocean color) from which additional parameters can be obtained (IOPs as well as biogeochemical parameters).

**Uncertainties and quality control concerns:**
Sensor calibration uncertainty.
The most critical aspect of above water measurements of \( L_w \) lies in the removal of skylight reflected at the air-sea interface, represented by the coefficient \( \rho \). It is well understood that any modulation of the sea surface will result in from contributions from the sky in directions not viewed with the \( L_d \) sensor (hence patchy clouds can bias the measurements). Approach includes using values from Mobley’s (1999, 2015) published tables, choosing \( \rho \) such that there is zero reflectance at the NIR (Morel, 1980) or choosing \( \rho \) such as to minimize atmospheric signals (as observed in atmospheric absorption bands) in the resulting reflectance (Simis and Olsson, 2013). Various attempt to model \( \rho \) which are not conclusive (See: Rudick, 2017). One method for estimation of uncertainties associated with this method is to consider the spectral consistency of \( R_{rs} (\lambda) \) in the near infrared. For clear waters and at sufficiently long wavelength \( R_{rs} (\lambda) \) can be assumed zero and any offset in measurements can be used as an estimator of total measurement uncertainty, provided this information has not already been used to perform a "residual correction" of data (Hooker and Morel, 2003).

Need to insure not to view boat’s wake or shadow and that irradiance sensor is not shaded by vessel.

**Key method references**


LISST-100X setting when in-line:

**Method:** Near forward particulate VSF at 670nm is derived from measurements of near-forward scattering of a laser. Measurements are inverted to obtain a PSD.

**Document author and contact info:** Emmanuel Boss (emmanuel.boss@maine.edu)

**Deployment methodology:**
Sensors is calibrated at the manufacturer and checked with NIST-traceable beads (Slade and Boss, 2006. Sensors is deployed in-line and is cleaned daily (including measurements of DIW that are not used except for tracking. ). Z-scat (blank) is determined based on VSF at the last 2 min of the 10 min long 0.2 μm filtered period at the end of each hour and interpolated in-between (Boss et al., 2018).

**Processing data:**
Z-scat (blank) is determined based on VSF at the last 2 min of the 10 min long 0.2 μm filtered period. Median-average total scattered data over 10 min to obtain one ‘robust’ near forward scattering spectra and variance around it. Blank for particulate VSF is the interpolated value between two consecutive Z-scats. Compute particulate scattering subtracting the Z-scat to the total data. Invert particulate scattering to PSD using Mie inversion (using the ‘natural particle’ we get significantly bigger deviations from Coulter Counter data). Beam attenuation is also computed.

All the raw data is saved in case we need to reprocess in the future.
Processing code is available at: [https://github.com/OceanOptics/InLineAnalysis](https://github.com/OceanOptics/InLineAnalysis)

**Derived parameters:**
Near forward VSF at 670nm.
Beam attenuation at 670nm and 0.0269° acceptance angle.
Particulate size distribution

**Uncertainties and quality control concerns:**
Uncertainty in VSF stems from:
1. Uncertainties in calibration – these are assessed with beads and corrected (in particular the assumed ring area).
2. Potential for changes in particles within the flow-through system.

Uncertainty in PSD stems from:
1. Assumption regarding particle’s shape. We found that in the open ocean the Mie theory (Agrawal and Pottsmith, 2000) based inversion matched best data from Coulter counter.
2. ‘Contamination’ by particles outside the inverted region – dealt with by removing first and last 3 size bins (Traykovsky et al., 1999).

**SeabASS fields and units:**
/fields=PSD_DNSD_###umsize or PSD_DVSD_###umsize, VSF_670_###ang
/units=TBD
References


**Method:** Particulate backscattering coefficient at one wavelength derived from VSF measurement in one wavelength in the back direction.

**Document author and contact info:** Emmanuel Boss (emmanuel.boss@maine.edu)

**Brief description of protocol:**
Sensors are calibrated with NIST-traceable beads (Sullivan et al., 2013). Salt water contribution is removed using salinity measurements (Zhang and al., 2009). Dark reading are measured on platform instrument is deployed on. Conversion from one angle VSF to the backscattering coefficient is based on literature values (Sullivan et al., 2013)

**Deployment methodologies:**
Sensors are deployed on profiling packages as well as using a special container in the flow-through system. In flow-through system contribution of box to signal needs to be measured (with DIW) and, by running 0.2 μm filtered water, the dissolved contribution can be assessed.

**Derived parameters:**
Particulate backscattering coefficient at wavelength of measurement.
POC from b_{bp} (e.g. Cetinic et al., 2012).
Phytoplankton carbon from b_{bp} (e.g. Graf et al., 2015).

**Uncertainties and quality control concerns:**
Uncertainty in b_{bp} stem from:
1. Uncertainties in calibration (actual angular response of sensor and inhomogeneity of illumination (hot spot of LED).
2. Uncertainties in conversion factor from one angle VSF in the back-direction to backscattering. Based on comparison with instrument using different calibration.
Comparison between instrument calibrated differently result in <10% uncertainty (Boss et al., 2004). Recent comparison between instrument of different angles in back direction showed differences on the O(50%) (Poteau et al., 2017). This was traced back to a problem with calibration coefficient provided by the manufacturer. After correction the uncertainties are O(10%) (Barnard, personal communication, 2017). Uncertainties in POC and phytoplankton carbon will be established based on regression with samples from project.

**SeaBASS fields and units:**
/fields=bbp,POC_cp,phyto_carbo
/units=1/m,mg/m^3,ug/L

**Key method references**
This document describes how we processed the CDOM absorption spectra from the Slow
descent rate optical package (SLOW-DROP) for the EXPORTS 01 cruise (RR1318).

Emmanuel Boss (emmanuel.boss@maine.edu) and Bentley Simpson, 2019-12-05

Sensors:
The CDOM absorptions were measured using different AC-S sensors:
From start to 8-17: used 301.
From 8-18 to 8-19: used 24.
From 8-20 until the end used 298.

Delay:
Based of plotting graphs and comparing temperature sensitive wavelengths we determined
that AC-S were, on average, 6.5 second lagging compared to the temperature (and pressure) of
the CTD (measured outside the water stream). This delay was applied to the AC-S data. We only
use the a-side from the AC-S as it was less sensitive to bubbles and provided more stable values
compared to the c-side.

Salinity:
The Temperature-Salinity relationship is very tight for EXPORTS and the top 110m (where we
deployed the IOP package). Because we had problems with the salinity sensor of our CTD we
decided to use the relationship established from the ship’s CTD to compute salinity. Hence the
salinity we report and use for correction is:
S=32.3+(14-T)/32 if T>6
S= 32.55 +0.9*(6-T) if T<6
Where T is the temperature measured with our CTD

Pressure:
We tarred the pressure at the surface (lowest pressure recorded was taken as offset).

Decent rates:
Varied from 15-30 cm/s.

Spike removal:
A 5pt median filter was applied to remove spikes.

Binning:
We median-binned the data into 1m bins.

Calibration:
We calibrated the sensors daily on the package. We chose the calibration file within two days of
each cast, that when applied to our data, resulted in spectra that were least different from an
exponential function, over all the wavelengths (least absolute difference). We added a function
for salinity and temperature so those were corrected before the absolute difference was
measured. Calibration spectra used are provided in their own files.
Quality control:
1. We expected spectra to be exponential within uncertainty of ~0.005m$^{-1}$. Spectra that were widely different were removed.
2. We compared (by eye) the trace of $a_g(440)$ to the voltage recorded by an analogue flow-through CDOM fluorometer (WETLabs, Wetstar). The traces were very similar in all cases.

Uncertainty estimate:
We estimate the uncertainty in CDOM to be smaller than 0.005m$^{-1}$ based on the manufacturer uncertainty (0.01m$^{-1}$), our binning (typically ~15 scans binned per m), and the variability in successive calibrations (based on the above procedure).
Measurements: Size fractionated (0.6-5.0 & >5.0 um) assessment of Si and Fe limitation of silica and primary production

Document author and contact information: Mark Brzezinski, mark.brzeinski@lifesci.ucsb.edu

Brief description of protocol and relation to export pathways: Seawater is collect from 2 depths from within the euphotic zone. At each depth the rate of silica production and primary production are measured in unaltered seawater (controls) and in the same water that has been augmented with 20 µm silicic acid, 1 nM Fe or both 20 µm silicic acid and 1nM Fe. Rates of silica production and primary production in all controls and treatments are measured in the 0.6-5.0 and the >5.0 µm size fractions following protocols Size fractionated 14C carbon uptake Brzezinski and Size fractionated 32Si silicic acid uptake Brzezinski protocol one pagers.

Other contributing protocols: Determination of 0.6 -5.0 & > 5 µm POC (combustion elemental analysis) and BSi (sodium hydroxide digestion), Par profiles.

Uncertainties and quality control concerns: The light field in deck incubators only approximates the light experience of captured diatom cells in situ. Extrapolation of rates measured from dawn to dusk to daily rates. Costs prohibit routine replication.

Data product originating from this measurement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rate+Si/control</td>
<td>unitless</td>
</tr>
<tr>
<td>Rate +F3/control</td>
<td>unitless</td>
</tr>
<tr>
<td>Rate +Fe+Si/control</td>
<td>unitless</td>
</tr>
</tbody>
</table>

BCO-DMO Fields:

Cruise,Date_Zulu,Time_Zulu,Event_num,Activity,Station,Cast,Latitude,Longitude,Rosette_Bottle,Target_Depth,pcnt_lo,TRMT,PO4,PO4_flag,SiO4,SiO4_flag,NO2,NO2_flag,NO2_NO3,NO2_NO3_flag,POC,POC_flag,PON,PON_flag,BSi_0_6umfilt_5umprefilt,BSi_5umfilt,rate_32Si_uptake_24hr_0_6umfilt_5umprefilt,rate_32Si_uptake_specific_24hr_0_6umfilt_5umprefilt,rate_32Si_uptake_24hr_5umfilt,rate_32Si_uptake_specific_24hr_5umfilt,rate_14C_uptake_24hr_0_6umfilt_5umprefilt,rate_14C_uptake_24hr_5umfilt,ISO_DateTime.UTC

Key method references

Measurements: Size fractionated primary production (0.6-5.0 & > 5.0 µm)

Document author and contact information: Mark Brzezinski, mark.brzeinski@lifesci.ucsb.edu

Brief description of protocol and relation to export pathways: The rate of inorganic carbon fixation into the > 5 µm particle size fraction is measured in seawater samples from the euphotic zone. The radioisotope 14C (bicarbonate) is added to seawater and each sample incubated in deck incubators (24h) screened with neutral density screening to simulate the light intensity at the depth of collection. Particles are recovered by filtration onto 5 µm polycarbonate filters, fumed with HCl to remove carbonates and analyzed for 14C content by liquid scintillation counting. The spiked seawater is sampled at the end of the incubation to determine total tracer added. The method is used alongside measures of 32Si uptake to determine the degree to which silicic acid and iron limit silica production and the rate of primary production.

Other contributing protocols: Determination of 0.6 -5.0 & > 5 µm POC (combustion elemental analysis), Par profiles.

Uncertainties and quality control concerns: The light field in deck incubators only approximates the light experience of captured diatom cells in situ. Extrapolation of rates measured from dawn to dusk to daily rates. Costs prohibit routine replication.

Data product originating from this measurement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Daily 0.6-5.0 &amp; &gt;5 µm C uptake</td>
<td>mol C L⁻¹ d⁻¹</td>
</tr>
</tbody>
</table>

BCO_DMO fields:

Cruise, Date_Zulu, Time_Zulu, Event_num, Activity, Station, Cast, Latitude, Longitude, Rosette_Bottle, Target_Depth, pcnt_lo, PO4, PO4_flag, SiO4, SiO4_flag, NO2, NO2_flag, NO2_NO3, NO2_NO3_flag, BSi_0_6umfilt_5umprefilt, BSi_5umfilt, rate_32Si_uptake_24hr_0_6umfilt_5umprefilt, rate_32Si_uptake_specific_24hr_0_6umfilt_5umprefilt, rate_32Si_uptake_24hr_5umfilt, rate_32Si_uptake_specific_24hr_5umfilt, ISO_DateTime.UTC

Key method references

Measurements: Size fractionated silicic acid uptake

Document author and contact information: Mark Brzezinski, mark.brzeinski@lifesci.ucsb.edu

Brief description of protocol and relation to export pathways: The rates of silicic acid uptake into the > 5 µm and in the 0.6 - 5.0 particle size fraction are measured in seawater samples from multiple depths spanning the euphotic zone in profile mode. The radioisotope $^{32}$Si(OH)$_4$ is added to seawater and each sample incubated in deck incubators (dawn to dusk) screened with neutral density screening to simulate the light intensity at the depth of collection. Particles are recovered by serial filtration through 5 and 0.6 µm polycarbonate filters, each filter is mounted on a planchette and the $^{32}$Si activity measured by low level beta counting. The rates obtained will be combined with parallel measures of the rate of C, N and Fe incorporation to 1) understand the stoichiometry of particle production in the diatom size fraction, 2) provide a baseline rate for assessing the level of nutrient stress in diatoms. The ultimate goal is to relate the nutrient status of diatoms to the specific ‘sinking particle’ pathways (pathways 1, 2, 3) that different diatom taxa follow.

Other contributing protocols: Determination of > 5 and 0.6-5.0 µm biogenic silica concentration (NaOH digestion), silicic acid concentration.

Uncertainties and quality control concerns: The light field in deck incubators only approximates the light experience of captured diatom cells in situ. Extrapolation of rates measured from dawn to dusk to daily rates. Costs prohibit routine replication.

Data product originating from this measurement

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Daily &gt;5 &amp; 0.6-5.0 µm Si uptake</td>
<td>mol Si L$^{-1}$ d$^{-1}$</td>
</tr>
<tr>
<td>Specific Si uptake (&gt;5 &amp; 0.6-5.0 µm)</td>
<td>d$^{-1}$</td>
</tr>
<tr>
<td>Integrated Si production (&gt;5 &amp; 0.6-5.0 µm, euphotic zone)</td>
<td>mmol Si m$^{-2}$ d$^{-1}$</td>
</tr>
</tbody>
</table>

BCO_DMO fields:

Cruise,Date_Zulu,Time_Zulu,Event_num,Activity,Station,Cast,Latitude,Longitude,Rosette_Bottle,Target_Depth,pcnt_lo,PO4,PO4_flag,SiO4,SiO4_flag,NO2,NO2_flag,NO2_NO3,NO2_NO3_flag,BSi_0_6umfilt_5umprefilt,BSi_5umfilt,rate_32Si_uptake_24hr_0_6umfilt_5umprefilt,rate_32Si_uptake_specific_24hr_0_6umfilt_5umprefilt,rate_32Si_uptake_24hr_5umfilt,rate_32Si_uptake_specific_24hr_5umfilt,ISO_DateTime_UTC

Key method references

Method: Size fractionated particles

Document author and contact info: Ken Buesseler kbuesseler@whoi.edu

Brief description of protocol and relation to export pathways: To understand the sources and fate of carbon associated with the biological pump, sampling of particles is needed. To collect enough material to obtain relevant data on the concentrations of C and associated elements on the rarer, larger suspended particles, in-situ pumping is commonly used to filter upwards of 1000 L. We will be deploying battery powered in-situ pumps (McLane Industries) whereby water entering the pump passes first through two screens (51 micron followed by 5 micron nominal pore size) followed by a 1 µm nominal pore-size QMA (quartz based) filter (all are 142 mm diameter). The QMA can be readily subsampled with “punches” of varying size for different analytes as particle distribution is even across the filter (Lam et al. 2015). For the screens, we have considerable experience in rinsing particles gently off these screens on to a 25 mm diameter 1 µm pore size silver filter that is dried and beta counted at sea for $^{234}$Th, and subsequently recounted on shore for $^{234}$Th prior to splitting by weight into subfractions for CHN, PIC and bSi (Lamborg et al. 2008). As in GEOTRACES, the screens will be cut into “pizza” wedges for $^{234}$Th, $^{210}$Po-$^{210}$Pb and Ba analysis (only at 3 stations). Given the sampling of 6 depths (5 trap depths plus 50 m) at 12 stations, this generates 72 samples for each of the three size classes. Moreover, we will have three dip blanks for each filter type, i.e. screens (51 and 5 µm) and QMA filters loaded in a filter holder that will be deployed with the deepest pump down to 500 m three times during the cruise. Samples will be generally dried except for other protocols (HPLC, organic isotopes, and phosphorous for a subset of QMA punches).

Other contributing protocols: Determine C or other element to $^{234}$Th ratios to convert water column-derived $^{234}$Th fluxes into C and other elemental flux estimates, as well as element to $^{210}$Po ratios; comparisons of filtration derived particle size and stocks results to other optical PSD data, such as from UVP and other camera systems; comparison to direct estimates of sinking particle fluxes using sediment traps and optical flux traps; look for links between spatial variability in particle abundance and size to community structure and other variables.

Uncertainties and quality control concerns: Particle data will have uncertainties derived from sample processing, blanks and other steps determined by standards and variability among replicate analyses. Each $^{234}$Th and $^{210}$Po measurement includes an associated analytical counting uncertainty.

Sample Analyses Summary and Units

<table>
<thead>
<tr>
<th>Analyses type</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Size fractionated particulates</td>
<td>&gt;1-5; 5-51; &gt;51 micron pore size</td>
</tr>
<tr>
<td>PC, PN, PIC, bSi</td>
<td>µmol/L</td>
</tr>
<tr>
<td>$^{234}$Th</td>
<td>dpm/L</td>
</tr>
<tr>
<td>$^{210}$Pb, $^{210}$Po</td>
<td>dpm/100L</td>
</tr>
<tr>
<td>Ba</td>
<td>pM</td>
</tr>
<tr>
<td>$^{138/134}$Ba$_{NIST}$</td>
<td>Unitless (ratio quantity)</td>
</tr>
</tbody>
</table>

* POC is determined by PC-PIC

SeaBASS submission fields and units:

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Key analytical methods references


Export Pathways Working Group “protocol briefs”

1. $^{234}$Th survey combined with particulate Element: $^{234}$Th ratios from large volume pumps

2. Ken Buesseler, kbuesseler@whoi.edu

3. Brief protocol description and how it relates to export pathways.
   We will use the particle-reactive tracer, thorium-234 (half-life = 24.1 d) to quantify the spatio-temporal variability in particle flux from the well-lit surface layer, and its attenuation with depth below. The disequilibrium in $^{234}$Th from its soluble parent, uranium-238, provides quantitative information on where particle export and remineralization occurs. The link from $^{234}$Th to C or other elemental fluxes is based upon determining the ratio of particulate $^{234}$Th to C (will be done here also for particulate nitrogen, biogenic silica and particulate inorganic C) measured on depth resolved profiles of size-fractionated particles (>1 to 50-100 µm range) collected using in-situ pumps (subsamples will be shared). The fluxes derived from this approach include all of the sinking particle pathways, as well as net removal of surface ocean particles (EXPORTS pathways 1, 2, 3), if removed by diel migrating zooplankton and released after they return to depth (pathway 4).

4. Other contributing protocols: Direct estimate of sinking particle fluxes using sediment traps and optical flux traps; comparisons to fluxes derived from particle stocks, such as from UVP and other camera systems; comparisons of $^{234}$Th flux to zooplankton and DOM mixing pathways; link between spatial variability from $^{234}$Th compared to community structure

5. Uncertainties and quality control concerns: Each $^{234}$Th measurement includes an associated analytical counting uncertainty. Particle data will have uncertainties derived from sample processing, blanks and other steps determined by variability among replicate analyses. The $^{234}$Th flux is derived from a model, and here we will estimate these terms by measuring local and time-varying transport terms, and by sampling in a Lagrangian mode, non-steady state terms can be assessed. Propagated model and analytical errors will be included in derived flux fields.

6. Data products originating with this method

   A. Sample Analyses

<table>
<thead>
<tr>
<th>Analyses type</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total $^{234}$Th</td>
<td>dpm/L</td>
</tr>
<tr>
<td>Size fractionated particulates</td>
<td>&gt;1-20; 20-53; &gt;53 micron pore size</td>
</tr>
<tr>
<td>PC, PN, PIC, bSi</td>
<td>µM/L</td>
</tr>
<tr>
<td>$^{234}$Th</td>
<td>dpm/L</td>
</tr>
</tbody>
</table>

   *note- size classes of particulates tbd and may vary from 53 micron.

   ** POC is determined by PC-PIC

   B. Derived products

   We will combine $^{234}$Th data with satellite products and results from a 3D high-resolution coupled physical-biogeochemical model with $^{234}$Th dynamics to produce synthesized flux data products.

   7. Key analytical methods references

   Buesseler, K.O., Benitez-Nelson, C., Rutgers Van Der Loeff, M., Andrews, J., Ball, L., Crossin,


Export Pathways Working Group “protocol briefs”

1. **210Pb and 210Po survey combined with particulate Element: 210Po ratios from large volume pumps**

2. Ken Buesseler, kbuesseler@whoi.edu; Montserrat Roca-Martí, mrocamarti@whoi.edu

3. **Brief protocol description and how it relates to export pathways**

Both polonium-210 (half-life = 138.4 d) and its parent lead-210 (half-life = 22.3 y) have a strong affinity for particle surfaces. However, 210Po is also incorporated into the cytoplasm of bacteria and phytoplankton (Cherrier et al., 1995; Fisher et al., 1983) and is preferentially assimilated by zooplankton with respect to 210Pb (Stewart and Fisher, 2003). This results in a disequilibrium in 210Po from its parent in seawater that can be used to quantify biogenic particle flux from the well-lit surface layer and its attenuation with depth below, in a similar way to 234Th. Due to its half-life, 210Po integrates a time scale of several months prior to the sampling. Export estimates from 210Po will be compared to other methods with shorter time scales, 234Th (weeks) and sediment traps (days), with the aim to provide more insights into the export and attenuation of sinking particle fluxes below the euphotic zone. The link from 210Po to particulate organic carbon (POC) and other elemental fluxes (particulate nitrogen, particulate inorganic carbon and biogenic silica) is based upon determining the ratio of particulate 210Po to the corresponding element measured on depth resolved profiles of size-fractionated particles (>1 to 50-100 µm range) collected using in-situ pumps. The fluxes derived from this approach include all the sinking particle pathways (EXPORTS pathways 1, 2, 3), as well as net removal of surface ocean particles by diel migrating zooplankton (pathway 5).

4. **Other contributing protocols:** Comparison to direct and other indirect estimates of sinking particle fluxes, including sediment traps, 234Th, and fluxes derived from particle stocks, such as UVP and other optical systems; link between spatial variability from 210Po compared to community structure.

5. **Uncertainties and quality control concerns:** The uncertainties associated with 210Pb and 210Po measurements are mainly due to the counting of polonium isotopes by alpha spectrometry and the determination of both the 209Po activity and the recovery of stable Pb (209Po and Pb are used as internal tracers to monitor losses of Po and Pb during sample processing). These uncertainties will be quantified appropriately following Rigaud et al. (2013). 210Po-derived fluxes will have uncertainties derived from both data and modeling assumptions.

6. **Data products originating with this method**

A. **Sample Analyses**

<table>
<thead>
<tr>
<th>Analyses type</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total 210Pb; Total 210Po</td>
<td>dpm 100L⁻¹</td>
</tr>
<tr>
<td>Size fractionated particulates</td>
<td>&gt;1-5; 5-51; &gt;51-micron pore size</td>
</tr>
<tr>
<td>PC, PN, PIC, bSi</td>
<td>µM L⁻¹</td>
</tr>
<tr>
<td>210Pb; 210Po</td>
<td>dpm 100L⁻¹</td>
</tr>
</tbody>
</table>

* POC is determined by PC-PIC

B. **Derived products**

Export flux estimates of POC (mmol C m⁻² d⁻¹) and other elements at different depths in the upper 500 m of the water column.

7. **Key analytical methods references**


Export Pathways Working Group “protocol briefs”

1. Abundance and isotopic composition of barium

2. Ken Buesseler, kbuesseler@whoi.edu; Montserrat Roca-Martí, mrocamarti@whoi.edu; Tristan Horner, tristan.horner@whoi.edu

3. Brief protocol description and how it relates to export pathways

How do upper ocean ecosystems characteristics determine the vertical transfer of organic matter from the well-lit surface ocean? This question will be tackled by measuring the abundance and isotopic composition of barium - an emerging geochemical proxy for organic carbon oxidation – in seawater and particulate samples from the upper mesopelagic (≤ 500 m). Earlier work has shown that barite (BaSO₄) micro-crystals precipitate inside biogenic aggregates in association with heterotrophic bacterial production, reflecting remineralization of organic matter (e.g., Dehairs et al., 2008; Martinez-Ruiz et al., 2018). However, our knowledge about barite cycling in seawater is limited. We are primarily interested in quantifying the relationship between organic matter remineralization and barite precipitation (and the Ba-isotopic composition thereof) in the context of the quantitative rate information derived from radionuclide proxies (²³⁴Th/²³⁸U and ²¹⁰Po/²¹⁰Pb). Such a study has not been conducted before and can be extremely beneficial in terms of understanding how Ba cycling relates to carbon remineralization. Ba-isotopic fractionation in seawater and marine particles (δ¹³⁸/¹³⁴Ba_NIST) and a new tracer, Ba* (defined as the difference between expected and observed Ba; e.g., Horner et al., 2015; Bates et al., 2017) will be used to investigate particle cycling and the biogeochemical processes that govern Ba distributions in seawater. Ba-derived data will be compared to heterotrophic respiration rates and flux attenuation estimates from sediment traps, besides radionuclide-derived data.

4. Other contributing protocols: Comparison to direct and indirect estimates of particle flux attenuation, including sediment traps, ²³⁴Th and ²¹⁰Po, as well as carbon respiration rates by bacteria and zooplankton.

5. Uncertainties and quality control concerns:

Long-term uncertainties for total and particulate Ba concentration measurements are estimated at ± 2 % (RSD; relative standard deviation; e.g., Horner et al., 2015). Similarly, uncertainties for Ba-isotopic measurements of samples with unknown composition have been estimated as ± 0.03 ‰ (Horner et al., 2015; Bates et al., 2017). Data accuracy is monitored via processing and analysis of international reference materials as part of each batch of samples (e.g., GEOTRACES GSP, GSC, D1). Lastly, blanks—extraneous Ba added during sample processing—are monitored via processing and analyzing of sub-ng aliquots of an isotopic double spike (¹³⁵Ba–¹³⁶Ba) through all sample purification procedures. Analyses of samples processed in batches with unreasonably high blank Ba concentrations will be discarded and the offending samples re-processed.

6. Data products originating with this method

A. Sample Analyses

<table>
<thead>
<tr>
<th>Analyses type</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Ba] (total Ba concentrations)</td>
<td>nM</td>
</tr>
<tr>
<td>δ¹³⁸/¹³⁴Ba_NIST (total Ba-isotopic compositions)</td>
<td>Unitless (ratio quantity)</td>
</tr>
<tr>
<td>p[Ba] (particulate Ba concentrations)</td>
<td>pM; determined for &gt;1–5, 5–51, and &gt;51 μm size fractions</td>
</tr>
<tr>
<td>δ¹³⁸/¹³⁴Ba_NIST (particulate Ba-isotopic compositions)</td>
<td>Unitless (ratio quantity); determined for &gt;1–5, 5–51, and &gt;51 μm size fractions</td>
</tr>
</tbody>
</table>

B. Derived products

Total: Ba concentrations, Ωбаріte, Ba*, and Ba-isotopic compositions. Particulate: Ba concentrations and Ba-isotopic compositions for three size fractions.

7. Key analytical methods references


Pelagic barite precipitation at micromolar ambient sulfate. Nat. Commun. 8: 1342. doi:10.1038/s41467-017-01229-5
The aim of the coagulation model is to model bio-physical processes affecting the particle size distribution with depth in the water column. The model is primarily an inverse-model that will be used to obtain rate parameters that are difficult or impossible to obtain from field observations or experiments (e.g. aggregation and disaggregation rates etc.). Two types of models will be used to minimize the effects of model uncertainty. The first model has been used in multiple studies before and represents the size spectrum with a high-resolution in particle size. This model assumes that particles are homogeneous and so cannot differentiate between particle types or particles of different composition. However, the model has the advantage of being well-established. The second model is under development and will be able to accommodate particles of multiple compositions. One version of this model has already been developed and used to successfully predict the amounts of oil, organic carbon, and mineral ballast in sedimenting particles. A second, more computationally efficient version of this model is currently being developed.

**Model Input Data:**
UVP and optically derived particle size distributions will be used as data for the inverse model, along with relevant zooplankton and fecal pellet concentrations, and microbial rates. Particle concentration by type will be used as data for determining particle production rates. Model output will also be compared with export flux measurements from the field.

**Model Output:**
Model output will consist primarily of particle size spectra, particle composition, and rate parameters including aggregation, disaggregation and consumption rates, particle stickiness, aggregate fractal dimension, and particle settling velocity.

**Model Quality Control and Parameter Uncertainties:**
The model code will be developed, documented and made publicly available using current best practices (e.g. unit and regression testing etc.)

Parameter uncertainties will be determined using, for example, Monte Carlo simulations and using inter-model comparisons.

Units of model input/output variables are:

- stickiness — unitless
- Aggregate fractal dimension — unitless
- Settling velocity of aggregates — m d$^{-1}$
- Particle size spectra — m$^{-4}$
- Particle degradation/consumption/dissolution rates — mg C d$^{-1}$
- Particle aggregation/disaggregation rates — d$^{-1}$
**Method:** Bulk DOC and DON stocks

**Document author and contact info:** Craig Carlson, carlson@lifesci.ucsb.edu

**Brief description of protocol and relation to export pathways:** Samples for the determination of bulk DOC and DON concentrations were collected over the surface 1000 m to capture DOM variability through the euphotic and mesopelagic zones on both survey (each station; spatial resolution) and process cruises (at least once per day; temporal resolution). Additional DOM profiles were collected and will continue to be collected from cruises deploying and recovering autonomous assets, or through collaborations with Line P PIs providing higher resolution of DOM seasonality. Water was passed through in-line GF/F filters into combusted EPA vials, then acidified to pH 3 with 4N HCL and stored. DOC concentrations will be determined by the high temperature combustions method ashore using a Shimadzu TOC-V or TOC-L (Carlson et al., 2010). Total dissolved nitrogen (TDN) is determined on the same instruments but with a detector for NO by chemiluminescence (Walsh, 1989). DON is calculated as the difference between TDN and DIN (the latter measured on samples collected by the EXPORTS hydro team). We hypothesized that net DOC production is a fairly regular fraction of NCP. Bulk DOC stock analysis will provide the mechanistic foundations to constrain DOC net production efficiency as well as its persistence.

**Other contributing protocols:** DIN (nutrient analysis by hydro team)

**Uncertainties and quality control concerns:** The systems’ responses are standardized daily with a four-point calibration curve of glucose or nitrate solution in Nanopure water. Each analytical run includes a set of deep and surface seawater “working” reference materials calibrated with DOC and TDN consensus reference material (CRM) (Hansell 2005). All samples are systematically referenced every 6 – 8 analyses throughout a run and generally have a coefficient of variation ranging between 1-3% over the 3-7 independent analyses (number of references depends on size of the run).

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>DOC</td>
<td>µmol L⁻¹, µmol kg⁻¹</td>
</tr>
<tr>
<td>DON</td>
<td>µmol L⁻¹, µmol kg⁻¹</td>
</tr>
</tbody>
</table>

**SeaBASS submission fields and units:**

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Key method references:


Method: DOC Remineralization Experiments

Document author and contact info: Craig Carlson, carlson@lifesci.ucsb.edu

Brief description of protocol and relation to export pathways: Microbial remineralization culture experiments were set up as seawater dilution cultures in which a source microbial assemblage (EZ or MZ) was inoculated into a naturally occurring seawater media i.e. 0.2 μm filtrate (EZ or MZ; DOM and nutrients) (Carlson et al. 2004). The culture was incubated at in situ temperatures (maintained in upright incubators) in the dark for days to weeks. Samples monitoring changes in bacterioplankton abundance, DOM concentrations and dissolved combined amino acid or dissolved combined neutral concentrations were collected from experiments at regular intervals for periods of days to weeks. Bacterial growth efficiency (BGE) will be obtained from concomitant changes in DOC concentration and bacterial carbon (Wear et al, 2015). DNA and RNA samples were also collected to assess changes in microbial community and function (in conjunction with S. Gifford).

Data generated from microbial remineralization experiments are relevant to Question SQ1a and SQ1b and essential to quantify the fraction of seasonally accumulated DOM that is bioavailable to microbial remineralization at surface versus how much survives degradation and is available for export during physical mixing (i.e. export potential). In addition, microbial remineralization experiments with mesopelagic microbes and surface accumulated DOM will simulate/investigate microbial responses to export events.

Other contributing protocols: O₂ respiration measurements (Scott Gifford), 16S rRNA gene phylogeny and transcriptomics (Scott Gifford), DNA sample collection, bulk DOC stocks, DOM composition and bacterial abundances.

Uncertainties and quality control concerns: All experimental treatments for microbial remineralization experiments will be replicated and each experiment will be conducted at least three times per Lagrangian cruise to assess variability among ECC states.

Data products originating with this method:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>DOC Removal Rate (over incubation)</td>
<td>μmol C L⁻¹ day⁻¹</td>
</tr>
<tr>
<td>Bioavailable DOC (over incubation)</td>
<td>μmol C L⁻¹, and % of total DOC</td>
</tr>
<tr>
<td>DCAA or DCNS (Change / diagenetic index)</td>
<td>nmol L⁻¹, relative contribution</td>
</tr>
<tr>
<td>Bacterioplankton counts</td>
<td>Cells L⁻¹</td>
</tr>
<tr>
<td>BGE</td>
<td>%</td>
</tr>
<tr>
<td>Bacterial community structure</td>
<td>Relative contribution of OTUs</td>
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Key method references:

**Method:** Dissolved organic matter composition

**Document author and contact info:** Craig Carlson, carlson@lifesci.ucsb.edu

*Note: Whether either DCAA or DCNS product is used for the full suite of EXPORTS samples is currently under evaluation.*

**Dissolved Combined Amino Acid (DCAA) Analysis:** Seawater was passed through in-line GF/F filters and stored at -20°C. Dissolved free amino acids (DFAA) will be analyzed as o-phthaldialdehyde (OPA) derivatives in high performance liquid chromatography (Lindroth and Mopper, 1989). Total dissolved amino acids (TDAA) will be analyzed in the same way as DFAA but after hydrolysis in 6 M HCl in sealed ampoules for 20 h at 110 °C (Henrichs, 1991; Kaiser and Benner 2009). DCAA will be calculated as the difference between TDAA and DFAA. Concentrations of DCAA will be measured and normalized to bulk DOC concentrations to produce the DCAA yield index. The DCAA yield and the mole ratio of individual amino acids reveal diagenetic patterns of DOM production and subsequent consumption in the epipelagic and mesopelagic zones.

**Dissolved Combined Neutral Sugars (DCNS):** The same principle of a diagenetic status index of DOM as identified for DCAA yield has also been applied to DCNS (Goldberg et al., 2009) with slightly differing isolation methods for DCNS as follows. Polymers of DOM are acid hydrolyzed (0.85M H2SO4) for 20 h at 100 °C to produce monomers that are detectable by pulsed amperometric detection high performance liquid chromatography (PAD-HPLC). DCNS yields are then normalized to DOC concentrations to produce a diagenetic index for DOM.

**Other contributing protocols:** DOC analysis

**Uncertainties and quality control concerns:** All of the field measurements will have had their measurement uncertainties quantified and standard errors reported for all variables. Certified standards are utilized to calibrate amino acids and neutral sugars with each sample being run in duplicate to triplicate to assess analytical error.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
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<td>nmol L⁻¹</td>
</tr>
<tr>
<td>DCAA yield</td>
<td>%</td>
</tr>
<tr>
<td>Or</td>
<td></td>
</tr>
<tr>
<td>DCNS</td>
<td>nmol L⁻¹</td>
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<tr>
<td>DCNS yield</td>
<td>%</td>
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Key method references:


**Method:** $^3$H- Leucine incorporation rate as a proxy for net heterotrophic bacterial production

**Document author and contact info:** Craig Carlson, carlson@lifesci.ucsb.edu

**Brief description of protocol and relation to export pathways:** The flux of the most labile fraction of DOC to bacterioplankton requires independent measurements of net heterotrophic bacterial production (BP) and estimates of bacterial growth efficiency (BGE). Net BP was estimated by $^3$H-Leucine incorporation (20nM addition; SA 60 Ci/mmol) using a modified version of the microcentrifuge method (Smith and Azam, 1992; Halewood et al 2012). Samples were collected on all SIO Cast profiles for depths spanning the surface to 500 m and were incubated in the dark for 2-4 hr at in situ temperatures. The $^3$H-Leu incorporation rates can be converted to bacterial carbon production (BP) using common conversion factors as described in Simon and Azam (1989; i.e. 1.5 – 3 kg C (mol leucine)$^{-1}$). To convert net BP to bacterial carbon demand (BCD) requires estimates of BGE from the literature or derived from independent remineralization experiments (see DOC Remineralization protocol) in which simultaneous measurements of the change in bacterial biomass and DOC in microbial dilution cultures are used to estimate BGE. BCD is determined as net BP / BGE and best represents the net flux of labile DOC (LDOC) through heterotrophic bacterioplankton. Determining the flux of LDOC will constrain the respiratory fate of a large fraction of NPP.

**Other contributing protocols:** Bacterial growth efficiency (BGE) from microbial remineralization experiments (see DOC remineralization experiments).

**Uncertainties and quality control concerns:** $^3$H- Leu incorporation rates were determined for 12 depths from euphotic to mesopelagic (1- 500 m) waters using replicate incubations with killed controls. BGE will be obtained from duplicate treatment incubations of microbial remineralization experiments.

**Data products originating with this method:**

<table>
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<th>Parameter</th>
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</tr>
</thead>
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<td>$^3$H- Leu incorporation rate</td>
<td>pmol L$^{-1}$ h$^{-1}$</td>
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**SeaBASS submission fields and units:**

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```

74
Key method references:


**Method:** Bacterioplankton Cell Enumeration

**Document author and contact info:** Craig Carlson, carlson@lifesci.ucsb.edu

**Epifluorescence microscopy protocol:** The epifluorescence microscopy method described below enumerates total prokaryotic abundance but is not able to differentiate between bacterial and archaeal domains. The cell density of the combined bacterial and archaeal domains is often referred to as bacterioplankton abundance (BA) in the literature (Glockner et al. 1999). The principle of analysis is that bacterioplankton are preserved with particle free formalin (at final concentration of 3.7%), stained with a 4'-6'-diamidino-2-phenylidole (DAPI) concentrated onto a blacked (acid black dye) polycarbonate 0.2 µm filter and enumerated with an epifluorescence microscope (60X; Porter and Feig 1980). Individual cells were identified on 10 images captured from the microscope and were counted using automated algorithms developed using ImageJ software (Bankhead, 2014). ImageJ software was used to determine BA as well as cell maximum and minimum length dimension to estimate cell biovolume (Baldwin and Bankston, 1988) of DAPI stained samples. The epifluorescence microscopy assay is appropriate for measuring bacterioplankton abundance within the dynamic range of $10^7$ – $10^9$ cells L$^{-1}$. These data will be used to estimate the variability of bacterioplankton stocks over specific depth horizons as well as throughout dissolved organic carbon (DOC) remineralization experiments. Cell abundance can be converted to C, N and P units using conversion factors from the literature or by independent analyses.

**Flow cytometry protocol:** Flow cytometric analyses of bacterioplankton stained with nucleic acid binding fluorochrome (SYBR-Green; Fisher Scientific) (Noble and Fuhrman, 1998) were used to enumerate bacterioplankton in remineralization experiments. SYBR-Green stain was added to formalin-preserved samples at a 1000-fold final dilution and allowed to rest for 10 minutes before analysis by a Guava easyCyte 5HT HPL Flow Cytometer (Blue 488 nm 150 mW laser; EMD Millipore). Stained bacterioplankton cells were enumerated using gating that excluded regions known to contain background noise previously identified by SYBR-stained HPLC-grade water.

**Uncertainties and quality control concerns:** Slides were spot checked at sea to ensure loading density and slide preparation quality and a subset of samples were prepared in triplicate as a check for reproducibility. Flow cytometry and epifluorescence microscopy output will be intercalibrated for direct comparison.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
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</thead>
<tbody>
<tr>
<td>Bacterioplankton Abundance</td>
<td>Cells E9 L$^{-1}$</td>
</tr>
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</table>
Mean cell volume  \( \mu m^3 \)

**SeaBASS submission fields and units:**

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/associated_archives=EXPORTSNP_Bact_Abund_Profiles.tar.gz
/associated_archive_types=imaging_epifluorescence

**Key method references:**


Method: DOC export

Document author and contact info: Craig Carlson, carlson@lifesci.ucsb.edu

Brief description of protocol and relation to export pathways: Estimates of the annual DOM export out of the surface 100 m and into the mesopelagic zone will be constrained by assessing changes in mesopelagic DOC inventories associated with physical measurements of subduction and deep convective mixing (Carlson et al. 1994, Hansell and Carlson 2002). For annual estimates of export flux we will leverage DOM collection from EXPORTS asset deployment/recovery cruises and time series cruises to Station Papa. High resolution sampling on survey ship will provide coincident measure of submesoscale physical variability along with DOM variability. Sensors on autonomous assets remaining after cruises will provide additional temporal variability of T, S, density used evaluated physical subduction of surface water to depth as well as changes in mixed layer depth (MLD). In summary, measurements of changes in mesopelagic DOM inventories from cruises and extended time series samples along with and estimates of subduction and maximal vertical mixing, observed with Bio-Argo floats, together provide insights on DOM export from the euphotic zone.

Other contributing protocols: autonomous sensor data for Temperature, salinity, density and MLD estimates.

Uncertainties and quality control concerns: All of the field measurements for DOC will have had their measurement uncertainties quantified and standard errors reported for all variables. Our laboratories use consensus and calibrated reference materials 3-4 times in every analytical run to ensure comparability between analytical runs within and between UCSB and U Miami laboratories. We will draw replicate 10% of DOC samples for our measurements enabling precision to be quantified. The glider and float sensors will be calibrated (by other groups) with discrete bottle samples collected upon deployment and upon occupation during “spatial” and “Lagrangian” campaigns.

Data products originating with this method: Parameter | Units
---|---
DOC export | µmol L\(^{-1}\) y\(^{-1}\)

Key method references:
Method: Winkler titrations for total dissolved O$_2$ concentration and saturation

Document authors and contact info: Nicolas Cassar, nicolas.cassar@duke.edu, David Nicholson, dnicholson@whoi.edu.

Brief description of protocol and relation to biogeochemical stocks: The O$_2$ concentration and saturation in seawater reflect a multitude of biological and physical processes, including photosynthesis, respiration, bubble injection through breaking waves, mixing of water masses, temperature and atmospheric pressure changes. The method for determining O$_2$ concentration in seawater was originally presented by Winkler (1888). We use the modified version of Carpenter (1965) as outlined in the WOCE Standard Operating Procedures Manual (Dickson, 1996). Briefly, O$_2$ in a known volume of sample oxidizes iodide ions (I$^-$) to iodine (I$_2$). The amount of iodine generated through this reaction is quantitatively titrated with a standard of thiosulfate (S$_2$O$_3^{-2}$) using an automated potentiometric end-point titrator. The conversion stoichiometry is four moles of thiosulfate for every mole of O$_2$. This assay is appropriate for measuring O$_2$ concentration in seawater under most oceanic conditions when hydrogen sulfide is not present. These data will be used to measure O$_2$ concentration and saturation and most importantly to calibrate the O$_2$ sensors (optodes).

Other contributing protocols: This is complementary to the Equilibrator Inlet Mass Spectrometer measurements which specifically measures the biological O$_2$ saturation and concentration. From these two methods, the physical O$_2$ supersaturation can be derived.

Uncertainties and quality control concerns: Samples for O$_2$ analyses will be collected at multiple stations. The main purpose is to calibrate and validate the O$_2$ sensors data on the ships and the autonomous platforms. Samples will be collected in triplicates to check replication error and precision. Precision and accuracy of dissolved O$_2$ concentration measurements should be 0.2% of air saturation or better (coefficient of variation). Air saturation values are calculated relative to the salinity and temperature dependent solubility of oxygen in seawater (Garcia and Gordon, 1992).

Data products originating with this method:

<table>
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<th>Parameter</th>
<th>Units</th>
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<td>Dissolved O$_2$</td>
<td>µmol kg$^{-1}$</td>
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<tr>
<td>concentration</td>
<td></td>
</tr>
<tr>
<td>Total O$_2$ saturation</td>
<td>%</td>
</tr>
</tbody>
</table>

During the 2018 EXPORTS North Pacific cruise, Winkler titrations were conducted on the R/V Sally Ride by Weiyi Tang, Duke University with assistance from Alex Niebergall, Duke University using reagents and equipment provided by the Nicholson...
Lab at WHOI. Over the course of the cruise, 287 individual Winkler bottle samples from calibration casts were titrated. Titrations on five calibration casts were conducted for the Seaglider, four for the Lagrangian Float, three for the BGC Argo float and one for the Wirewalker. Most were sampled as sets of triplicates with each replicate drawn from a different Niskin bottle fired at the same depth. Of the 287 samples, 3 were flagged as unexplained outliers and 6 were flagged due to identified problems that included Niskin misfires, a broken flask, and a software crash. Subsequent results are reported for the remaining 278 samples. Oxygen concentration ranged from 16.5 μmol kg⁻¹ to 300.9 μmol kg⁻¹. The median standard deviation for triplicate samples was 0.52 μmol kg⁻¹ (0.2% of saturation).

Oxygen concentration were determined in units of μmol kg⁻¹ and subsequently converted to units of ml L⁻¹ as required by SeaBASS using the equation:

\[
O_2(\text{ml/L}) = O_2(\text{μmol/kg}) \times 22.392 \times (\sigma_0 + 1000) \times 10^{-6}
\]

where 22.392 is the virial molar volume of oxygen (L mol⁻¹) and \(\sigma_0\) is potential density. The equilibrium dissolved oxygen concentration (\(O_2\)eq) was calculated using the TEOS Gibbs Seawater Toolbox function gsw_O2sol_SP_pt.m which returns units of μmol kg⁻¹. Dissolved oxygen saturation is then calculated as:

\[
O_2 \text{ saturation} = 100 \times \frac{O_2(\text{μmol/kg})}{O_2\text{eq}}
\]

**SeaBASS fields and units:**

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**Key method references:**

**Method:** O₂/Ar Net Community Production Estimates

**Document author and contact info:** Nicolas Cassar, nicolas.cassar@duke.edu

Brief description of protocol and relation to export pathways: Export production is believed to be a function of NCP which is defined as the balance between net primary production and heterotrophic respiration, or the difference between gross primary production and community respiration. If the organic matter pool (POC+DOC) in the ML is at steady state, export production is equal to NCP. Export may lag production, in which case NCP is expected to be greater than export. Conversely, without allochthonous sources of organic matter, if the organic matter inventory in the ML decreases, export production will be predicted to be transiently greater than NCP. We can account for the lack of steady-state by measuring changes in the O₂ and POC and DOC pools over time. We derive NCP from the ratio of O₂ to the inert gas argon (Ar). The biological O₂ supersaturation can be estimated from O₂/Ar because O₂ and Ar have similar solubility properties (Craig and Hayward 1987). ΔO₂/Ar is measured underway from the ship’s flow-through seawater line by Equilibrator Inlet Mass Spectrometry (EIMS) as described in Cassar et al. (2009). NCP in units of mmol O₂ m⁻² day⁻¹ is calculated from ΔO₂/Ar and other factors, including a gas transfer velocity for O₂ (m day⁻¹) and a wind speed parameterization and a gas exchange weighting. NCP for O₂ is then converted to carbon (mmol C m⁻² day⁻¹).

Other contributing protocols: DNA sequencing for 16S and 18S.

Uncertainties and quality control concerns: Lack of steady-state (can be accounted for in Lagrangian mode), vertical mixing/exchange with other water masses, gas exchange parameterization, conversion factor O₂/C. See Cassar et al. (2014) for a thorough description of the uncertainties.

Data products originating with this method:

<table>
<thead>
<tr>
<th>Parameter*</th>
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<tbody>
<tr>
<td>Biological O₂ saturation</td>
<td>%</td>
</tr>
<tr>
<td>Physical O₂ saturation</td>
<td>%</td>
</tr>
<tr>
<td>NCP (*)</td>
<td>mmol-C m⁻² d⁻¹</td>
</tr>
</tbody>
</table>

* Converted from biological O₂ flux to the atmosphere.

SeaBASS submission fields and units:

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**Key method references**


Method: Compound-specific isotope analysis of amino acids (AA-CSIA) isolated from sinking particles, size-fractionated particles, and zooplankton fecal pellets

Document author and contact info: Hilary G. Close (hclose@rsmas.miami.edu)

Brief description of protocol and relation to export pathways: Sinking particles will be collected via sediment traps. Two size fractions of particles (0.8–51 μm, >51 μm) and subsamples of 1–5 μm particles will be collected via in situ pumps. Four size fractions of particles (0.3–1 μm, 1–6 μm, 6–51 μm, >51 μm) will be collected at a subset of depths. Fecal pellets will be harvested from zooplankton incubations. Filters may be subsampled for size-fractionated total particulate carbon, particulate organic carbon, and total particulate nitrogen concentrations and isotope ratios. Fourteen protein-forming amino acids will be analyzed for compound-specific C and N isotope analysis and quantitation and enantiomer ratios. From this data, several parameters will be calculated that can distinguish the extent to which microbial and metazoan heterotrophy have acted upon detrital particles and/or contributed secondary biomass. We will use a multivariate statistical framework to estimate phytoplankton, heterotrophic microbe, and fecal pellet contributions to particles and thus to size-fractionated particle flux. We anticipate coordinating with several projects to refine our statistical framework and eventually to construct a dynamical model of organic matter sources and transformations.

Other contributing protocols: AA-CSIA of size-fractionated zooplankton, microbial and zooplankton community characterization/quantification, microbial and zooplankton respiration, fecal pellet production and flux, visual particle characterization, particle size distributions, thorium-derived size-fractionated carbon and nitrogen flux, particle fluxes from sediment traps, particle aggregation/disaggregation rates.

Uncertainties and quality control concerns: Blank “dipped” filters as full-process blanks. Synthetic amino acids added to samples as internal isotope and recovery standards. Suites of amino acid standards prepared and analyzed with sample batches as external standards for quantitation, mass-balance correction for derivative carbon, and racemization correction. Cross-lab standard material prepared and analyzed with sample batches. Triplicate analysis, propagation of analytical uncertainty in calculated parameters. Propagated uncertainty in δ¹³C from analysis of derivatization standards. Estimated uncertainty in filtered volumes and thus concentrations.

Data products originating with this method

<table>
<thead>
<tr>
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<tr>
<td>Natural-abundance nitrogen isotope ratios of individual amino acids, size fractionated particles and incubated fecal pellets</td>
<td>Values of δ¹⁵N (air)</td>
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<tr>
<td>Natural-abundance carbon isotope ratios of individual amino acids, size fractionated particles and incubated fecal pellets</td>
<td>Values of δ¹³C (VPDB)</td>
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<td>D:L enantiomer ratio of alanine, size fractionated particles</td>
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<td>*Bulk nitrogen concentration, size fractionated particles ([PN])</td>
<td>μmol L⁻¹</td>
</tr>
<tr>
<td>*Natural-abundance bulk nitrogen isotope ratios, size</td>
<td>Values of δ¹⁵N (air)</td>
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</table>
fractionated particles (δ¹⁵N<sub>PN</sub>)

| *Bulk total carbon concentration, size fractionated particles ([PC]) | μmol L⁻¹ |
| *Natural-abundance bulk carbon isotope ratios, size fractionated particles (δ¹³C<sub>PC</sub>) | Values of δ¹³C (VPDB) |
| *Bulk organic carbon concentration, size fractionated particles ([POC]) | μmol L⁻¹ |
| *Natural-abundance bulk organic carbon isotope ratios, size fractionated particles (δ¹³C<sub>POC</sub>) | Values of δ¹³C (VPDB) |
| *PC:PN and POC:PN ratios, size fractionated particles | mol/mol |

*As sample size permits and where not redundant

**Key method references**


Method: Genetic characterization of bulk sinking particles in sediment traps

Document author and contact info: Colleen Durkin, cdurkin@mlml.calstate.edu

Brief description of protocol and relation to export pathways: Each sediment trap will include 1 tube containing RNAlater preservative. Particles in the RNAlater will be collected onto a filter and frozen. DNA will be extracted and the V4 hyper-variable region of the 18S and 16S rRNA DNA markers will be amplified using primer sets modified by Apprill et al. 2015 and Penna et al. 2017. Amplified PCR products will be sequenced on an Illumina MiSeq with a 300 bp paired end run. DNA sequence identities will be assigned by comparing to sequence databases. This analysis contributes to export pathways 1, 2, and 3 by specific organisms with particle export. When combined with particle-specific DNA sequencing data, the organisms only exported in small particles can be inferred.

Other contributing protocols: Genetic characterization of zooplankton products in sediment traps, genetic characterization of surface plankton communities, genetic characterization of aggregates in sediment traps.

Uncertainties and quality control concerns: While the presence of a DNA sequence confirms the link between an organism or its remains with sinking particles, the absence of a DNA sequence does not confirm the absence of this link. It is possible for organic matter to be exported without any DNA evidence of its organismal source. To relate these data to surface phytoplankton communities, it is critical that the same PCR primers are used to amplify 16S and 18S rRNA of surface plankton communities.

Data products originating with this method:
Data table of species detected in bulk sinking material – csv file
DNA sequences – fasta file

SeaBASS submission fields and units:
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/units=%,none,TBD

Key method references:

Method: Genetic characterization of sinking aggregates in gel traps

Document author and contact info: Colleen Durkin, cdurkin@mlml.calstate.edu

Brief description of protocol and relation to export pathways: Jars containing polyacrylamide gel layers will be deployed in 1 tube of each deployed sediment trap. Sinking particles and organisms that settle into the gel layer remain distinctly separated, preserving original characteristics of size and quantity and constituents. Individual aggregates will be pipetted out of the gel layer and frozen in cryovials immediately after trap recovery. DNA will be extracted and the V4 hyper-variable region of the 18S and 16S rRNA DNA markers will be amplified using primer sets modified by Apprill et al. 2015 and Penna et al. 2017. Amplified PCR products will be sequenced on an Illumina MiSeq with a 300 bp paired end run. DNA sequence identities will be assigned by comparing to sequence databases. This analysis contributes to export pathway 2 by connecting specific organisms with transport by sinking aggregates.

Other contributing protocols: Genetic characterization of bulk particles in sediment traps, genetic characterization of surface plankton communities.

Uncertainties and quality control concerns: This analysis is performed on the largest particles in the gel but sequences that are specific to the small sinking particles can be inferred by identifying differences between individual particles and the bulk trap material sequence compositions. While the presence of a DNA sequence confirms the link between an organism or its remains with aggregates, the absence of a DNA sequence does not confirm the absence of this link. It is possible for organic matter to be exported without any DNA evidence of its organismal source. Additionally, particles will degrade over the course of the deployment period, and may alter the DNA sequences detected. The extent of these changes is currently being examined. To relate these data to surface phytoplankton communities, it is critical that the same PCR primers are used to amplify 16S and 18S rRNA of surface plankton communities.

Data products originating with this method:
Data table of species per aggregate – csv file
DNA sequences – fasta file

SeaBASS submission fields and units:
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Key method references:

**Method:** Genetic characterization of zooplankton products in gel traps

**Document author and contact info:** Colleen Durkin, cdurkin@mlml.calstate.edu

**Brief description of protocol and relation to export pathways:** Jars containing polyacrylamide gel layers will be deployed in 1 tube of each deployed sediment trap. Sinking particles and organisms that settle into the gel layer remain distinctly separated, preserving original characteristics of size and quantity and constituents. Individual fecal pellets will be pipetted out of the gel layer and frozen in cryovials immediately after trap recovery. DNA will be extracted and the V4 hyper-variable region of the 18S and 16S rRNA DNA markers will be amplified using primer sets modified by Apprill et al. 2015 and Penna et al. 2017. Amplified PCR products will be sequenced on an Illumina MiSeq with a 300 bp paired end run. DNA sequence identities will be assigned by comparing to sequence databases. This analysis contributes to export pathway 3 by connecting specific organisms with transport by sinking fecal pellets.

**Other contributing protocols:** Genetic characterization of bulk particles in sediment traps, genetic characterization of surface plankton communities.

**Uncertainties and quality control concerns:** This analysis is performed on the largest fecal pellets in the gel but sequences that are specific to the small sinking particles can be inferred by identifying differences between individual particles and the bulk trap material sequence compositions. While the presence of a DNA sequence confirms the link between an organism or its remains with fecal pellets, the absence of a DNA sequence does not confirm the absence of this link. It is possible for organic matter to be exported without any DNA evidence of its organismal source. Additionally, particles will degrade over the course of the deployment period, and may alter the DNA sequences detected. The extent of these changes is currently being examined. To relate these data to surface phytoplankton communities, it is critical that the same PCR primers are used to amplify 16S and 18S rRNA of surface plankton communities.

**Data products originating with this method:**
Data table of species per fecal pellet – csv file
DNA sequences – fasta file

**SeaBASS submission fields and units:**
/fields=abun, namespace_manual, identification_manual, biotic_group, DNA_counts, associated_files, associated_file_type
/units=%, none, TBD

**Key method references:**

**Method:** Visual cell ID in gel traps

**Document author and contact info:** Colleen Durkin, cdurkin@mlml.calstate.edu

**Brief description of protocol and relation to export pathways:** Jars containing polyacrylamide gel layers will be deployed in 1 tube of each deployed sediment trap. Sinking particles and organisms that settle into the gel layer remain distinctly separated, preserving original characteristics of size and quantity and constituents. Gel jars will be visually examined by microscopy to quantify the flux and identity of individually-sinking organisms. To detect the large and potentially rare organisms (phytoplankton and zooplankton), the entire gel layer will be surveyed at low magnification (30x-50x) with a stereomicroscope and all visible organisms will be identified and counted. To quantify smaller cells, a gridded transparency will be placed underneath the gel and all cells within ≥20 grid squares (0.25 cm² area per square) will be identified and counted at 115x under darkfield illumination. Cell fluxes will be calculated by dividing the total number of cells counted by the area examined and further divided by the deployment time (number m⁻² d⁻¹). Approximate cell volumes will be calculated and number fluxes will be converted to carbon flux using the equations of Menden-Deuer and Lessard 2000.

**Other contributing protocols:**

**Uncertainties and quality control concerns:** The uncertainty of fluxes can be estimated by the square root of the number of counts. The carbon per cell is an estimate based on an empirically derived equation.

**Data products originating with this method:**
Cells fluxes: cells m⁻² d⁻¹
Zooplankton fluxes or swimmer contamination: organisms m⁻² d⁻¹
Phytoplankton carbon flux: mmol C m⁻² d⁻¹

**SeaBASS submission fields and units:**

```
/id_fields_definitions=1id:passive,2id:aggregate,3id:long_fp,4id:dense_detritus,5id:large_loose,6id:short_fp,7id:mini_pellet,8id:salp_pellet,9id:phyto,10id:foraminifera,11id:rhizaria,12id:fiber,13id:copepod,14id:pteron,15id:amphipod,16id:other_zooplankton,17id:zooplankton_part,18id:unidentifiable
```

```
/fields=R2R_Event,sample,SN,station,depth,date,time,date_start,time_start,date_end,time_end,date_resurface,time_recovery,elapsed_time,lat,lon,lat_start,lon_start,lat_end,lon_end,lat_resurface,lat_recovery,lon_recovery,bin_diameter_center,bin_diameter_upper,bin_diameter_lower,flux_particles_1id,flux_particles_1id_unc,bincount_1id,flux_particles_2id,flux_particles_2id_unc,bincount_2id,flux_particles_3id,flux_particles_3id_unc,bincount_3id,flux_particles_4id,flux_particles_4id_unc,bincount_4id,flux_particles_5id,flux_particles_5id_unc,bincount_5id,flux_particles_6id,flux_particles_6id_unc,bincount_6id,flux_particles_7id,flux_particles_7id_unc,bincount_7id,flux_particles_8id,flux_particles_8id_unc,bincount_8id,flux_particles_9id,flux_particles_9id_unc,bincount_9id,flux_particles_10id,flux_particles_10id_unc,bincount_10id,flux_particles_11id,flux_particles_11id_unc,bincount_11id
```
Key method references:


**Method:** Visual characterization of zooplankton products in gel traps

**Document author and contact info:** Colleen Durkin, cdurkin@mlml.calstate.edu

**Brief description of protocol and relation to export pathways:** Jars containing polyacrylamide gel layers will be deployed in 1 tube of each deployed sediment trap. Sinking particles and organisms that settle into the gel layer remain distinctly separated, preserving original characteristics of size, quantity, and constituents. A process blank gel is prepared and remains on the ship to quantify background particle contamination. Gel layers are imaged under a stereomicroscope at multiple magnifications (e.g. 7x, 20x, 50x, 115x), in multiple focal planes, and under brightfield and oblique lighting sources. Image data are processed using a python script that isolates, counts, measures, and saves images of individual particles. Particle size distribution (PSD) (number m\(^{-2}\)) is calculated for log-transformed size bins spanning optimal magnification ranges. Each sample PSD is blank-corrected by subtracting the average of all process blank PSDs and converted to flux by dividing by deployment time (number m\(^{-2}\) d\(^{-1}\)). Every detected particle is manually assigned an identity (e.g. fecal pellet, aggregate) using a python-based graphical user interface that allows rapid identity assignment of thousands of images. The percent of fecal pellets in each size bin is multiplied by the total particle number flux to determine fecal pellet number fluxes across size bins. Fecal pellet number fluxes will be converted to fecal pellet carbon fluxes using conversions measured directly from concurrent zooplankton grazing experiments (mmol C m\(^{-2}\) d\(^{-1}\)). This analysis contributes to export path 3 by calculating the quantity of fecal pellets that are contributing the carbon flux at each depth.

**Other contributing protocols:** Zooplankton fecal pellet production, Visual characterization of aggregates in gel traps

**Uncertainties and quality control concerns:** The uncertainty of fluxes can be estimated by the square root of the number of counts. Oblique lighting enables detection of translucent particles and can resolve smaller size bins, whereas brightfield lighting is more comparable to optical instruments like the Optical Sediment Trap.

**Data products originating with this method:**
- Fecal pellet fluxes across size bins (10-1000 um diameter): pellets m\(^{-2}\) d\(^{-1}\)
- Estimated carbon flux from fecal pellets at each depth: mmol C m\(^{-2}\) d\(^{-1}\)

**SeaBASS submission fields and units:**
- /fields= flux_carbon_fecalpellet
- /units=mmol C m-2 d-1

**Key method references:**
Method: Bulk particle flux to sediment traps

Document author and contact info: Meg Estapa, mestapa@skidmore.edu

Brief description of protocol and relation to export pathways: Sediment traps are used to directly collect sinking particles at discrete, sub-mixed layer depths. Collected particles are analyzed for particulate carbon (PC), particulate inorganic carbon (PIC), biogenic silica (bSi), $^{234}$Th, $^{210}$Po, $^{210}$Pb, Ba, and mass and converted to fluxes by normalizing to the trap collection area and length of deployment. POC flux is determined as the difference between PC and PIC fluxes. Bulk compositional analysis does not discriminate among sinking particles from different export pathways (single cells, aggregates, zooplankton products) so this method provides an estimate of the sum of all “sinking particle” pathways (1, 2, and 3). EXPORTS field measurements use two different sediment trap platform designs (one neutrally-buoyant, the other surface-tethered) both carrying cylindrical trap tubes with closing lids. Other analytes not described here may be analyzed by other contributing groups.

Other contributing protocols: Analytical determination of PC (combustion elemental analysis), PIC (coulometry), bSi (alkaline digestion and spectrophotometric determination), $^{234}$Th (low level β emission), $^{210}$Po (alpha spectrometry), $^{210}$Pb, Ba, and particle mass (gravimetry).

Uncertainties and quality control concerns: Influence of hydrodynamic biases which depend upon trap design; zooplankton “swimmer” presence in samples; sample solubilization during deployment; intra-platform (i.e. “tube-to-tube”) variability; inter-platform (i.e. “trap-to-trap”) variability; handling (process) blanks for all analytes; mechanical issues with traps during deployments (e.g. depth variability, lid closures)

Data products originating with this method:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>PC flux</td>
<td>mmol-C m$^{-2}$ d$^{-1}$</td>
</tr>
<tr>
<td>POC flux</td>
<td>mmol-C m$^{-2}$ d$^{-1}$</td>
</tr>
<tr>
<td>PIC flux</td>
<td>mmol-C m$^{-2}$ d$^{-1}$</td>
</tr>
<tr>
<td>bSi flux</td>
<td>mmol-Si m$^{-2}$ d$^{-1}$</td>
</tr>
<tr>
<td>$^{234}$Th flux</td>
<td>dpm m$^{-2}$ d$^{-1}$</td>
</tr>
<tr>
<td>$^{210}$Po flux</td>
<td>dpm m$^{-2}$ d$^{-1}$</td>
</tr>
<tr>
<td>$^{210}$Pb flux</td>
<td>dpm m$^{-2}$ d$^{-1}$</td>
</tr>
<tr>
<td>Ba flux</td>
<td>mmol-Ba m$^{-2}$ d$^{-1}$</td>
</tr>
<tr>
<td>mass flux</td>
<td>mg m$^{-2}$ d$^{-1}$</td>
</tr>
</tbody>
</table>

* Each to be reported separately for surface-tethered and neutrally-buoyant sediment traps, and as a function of depth

SeaBASS submission fields and units:
/fields= flux_PC, flux_PN, flux_POC, flux_PIC, flux_bSi, flux_Th_234, flux_mass,flux_pb_210,flux_Po_210,flux_ba, flux_P

/units= mmol m-2 d-1, mmol m-2 d-1, mmol m-2 d-1, mmol m-2 d-1, mmol m-2 d-1, dpm m-2 d-1, mg m-2 d-1, dpm m-2 d-1, dpm m-2 d-1, mmol m-2 d-1, mmol m-2 d-1
Key method references
**Method:** Particle flux to optical sediment traps  
**Document author and contact info:** Meg Estapa, mestapa@skidmore.edu

**Brief description of protocol and relation to export pathways:** An upward-looking optical attenuance sensor (either bulk or imaging) or optical sediment trap (OST) is deployed on a quasi-Lagrangian platform and used to measure the accumulation rate of sinking particles while the platform drifts on a neutral surface. The rate of increase in accumulated particles’ attenuance (ATNOST flux) is converted to a particulate carbon flux (PCATN flux) using an empirical calibration function. These calibration data are generated by co-deploying OSTs on the same platforms as bulk sediment traps, and measuring ATNOST flux and PC flux simultaneously. Independent estimates of attenuance flux will also be determined from brightfield microscopy on polyacrylamide gel sediment traps carried on these platforms. In EXPORTS, C-Rover 2000 beam transmissometers (WETlabs, specs here: [http://www.seabird.com/c-rover-2000](http://www.seabird.com/c-rover-2000)) will be used as bulk OSTs in the first (N. Pacific) field campaign, and imaging sensors will be added in the second (N. Atlantic) campaign. PCATN flux is an estimate of the sum of all “sinking particle” pathways (1, 2, and 3).

**Other contributing protocols:** Bulk PC fluxes to sediment traps, visual characterization of sinking particles in gel traps.

**Uncertainties and quality control concerns:** Platform-flow interactions biasing collection of particles by size; integration of FATN measurement over sufficient area and time to achieve statistical confidence; inter-platform variability

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>ATNOST flux</td>
<td>m² m⁻² d⁻¹</td>
</tr>
<tr>
<td>ATNgel flux</td>
<td>m² m⁻² d⁻¹</td>
</tr>
<tr>
<td>PCATN flux</td>
<td>mmol-C m⁻² d⁻¹</td>
</tr>
</tbody>
</table>

**SeaBASS submission fields and units:**

/fields= flux_ATN  
/units= m2 m-2 d-1

**Key method references**


Method: Net community production, net primary production, and particulate inorganic carbon export from chemical tracer budgets using biogeochemical profiling floats

Document author and contact info: Andrea Fassbender, fassbender@mbari.org

Brief description of protocol and relation to export pathways: Float observations are used to evaluate bulk nitrate, dissolved oxygen (DO), dissolved inorganic carbon (DIC), and total alkalinity (TA) transformations throughout the water column and close chemical tracer budgets to estimate carbon export. Nitrate, DO, and pH are directly measured by the floats while TA is estimated from salinity, temperature, and oxygen measurements and used with pH to calculate DIC. Additionally, float pH is used to estimate the partial pressure of carbon dioxide ($pCO_2$) at the ocean surface [Williams et al., 2017] to calculate air-sea CO$_2$ exchange that contributes to the DIC budget. By closing multiple tracer budgets, it is possible to quantify the export of particulate and dissolved organic carbon (POC & DOC) and particulate inorganic carbon (PIC) when integrated over the annual cycle, assuming steady state [Fassbender et al., 2016; Plant et al., 2016]. Net primary production (NPP) will also be estimated from float bio-optical sensors and regional chlorophyll fluorescence-particle backscattering-NPP relationships determined during the EXPORTS cruise.

Other contributing protocols: Analytical determination of: POC (combustion elemental analysis) and PIC (by difference using acidified filter); DOC and TOC (combustion catalytic oxidation), achieving POC by difference; absorption coefficient of CDOM (UV/Vis spectrophotometry); DIC (NDIR spectrophotometry), TA (open cell titration), and pH (UV/Vis spectrophotometry); empirical backscatter-POC relationships; and empirical chlorophyll fluorescence-particle backscattering-NPP relationships.

Uncertainties and quality control concerns: Unresolved physics, application of a TA algorithm in a calcium carbonate production region, C:N and C:O ratios applied in chemical tracer budgets, POC-backscatter relationship, NPP-chlorophyll relationship.

Data products originating with this method:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>POC export</td>
<td>mol-C m$^2$ yr$^{-1}$ (or day)</td>
</tr>
<tr>
<td>PIC export</td>
<td>mol-C m$^2$ yr$^{-1}$ (or day)</td>
</tr>
<tr>
<td>DOC export</td>
<td>mol-C m$^2$ yr$^{-1}$ (or day)</td>
</tr>
<tr>
<td>NCP (POC + DOC export)</td>
<td>mol-C m$^2$ yr$^{-1}$ (or day)</td>
</tr>
<tr>
<td>NPP</td>
<td>mol-C m$^2$ yr$^{-1}$ (or day)</td>
</tr>
</tbody>
</table>

* These parameters will be estimated every ~3 days during the EXPORTS cruise, and every ~10 days otherwise for an anticipated 3-5 years; however, the most robust results may come from integration over the seasonal/annual cycle.

Data quality control and access: Float sensor data quality control procedures are outlined in Johnson et al., 2017. Float data are available in near real time from the MBARI FloatViz and Argo websites:
Float 0949 (Dory) WMO #: 5905988
Float 0948 (Nemo) WMO #: TBD. The float has exhibited serious problems since the December 2018 redeployment. As a result, we may report data from the EXPORTS deployment, at which time the float will be issued a WMO number.

**Key method references**


**Method:** Discrete dissolved inorganic carbon (DIC) – (µmol kg⁻¹)

**Document author and contact info:** Andrea Fassbender, fassbender@mbari.org

**Brief description of protocol and relation to export pathways:** Discrete DIC samples were collected into 500 mL and 250 mL borosilicate bottles and preserved with 200 µL and 100 µL, respectively, of saturated mercuric chloride for later analysis at the Monterey Bay Aquarium Research Institute (MBARI). A custom analysis system was used in which a Kloehn V6 syringe pump (5 mL syringe) handles fluid control, delivering 1.75 mL of sample to a custom designed CO₂ stripping chamber. 200 µL of 5% phosphoric acid (H₃PO₄) is subsequently added to the CO₂ stripping chamber for acidification of the sample. CO₂-free gas is then bubbled through the acidified sample and the evolved CO₂(g) is delivered to a LiCOR 7000 Nondispersive Infrared gas analyzer for measurement. Carrier gas flow rate is controlled using a mass flow controller. The DIC concentration of the sample is proportional to the integral of the LiCOR CO₂(g) sample peak. All samples were run in triplicate, and the results were averaged. Instrument performance was monitored by measuring Certified Reference Material (CRM; provided by Andrew Dickson at SIO) approximately every hour. The average accuracy of the instrument relative to CRMs is better than 1 µmol kg⁻¹.

The following systematic, concentration-dependent correction for samples analyzed on our laboratory instrumentation, was applied: DICCorrected = DICSample – (DICSample – CRM) × -0.023. The standard error of the slope (-0.023 ± 0.002) results in correction uncertainties ranging from ± 0.2 to ± 0.75 µmol kg⁻¹.

Discrete DIC samples are used to validate DIC values estimated from profiling float pH measurements and TA estimates. A biogeochemical tracer budget for DIC is used to quantify carbon export over the float lifetimes.

**Other contributing protocols:**

**Uncertainties and quality control concerns:** Measurement precision was estimated by analyzing sets of replicate samples drawn from the same Niskin bottle during rosette casts. The average standard deviation between replicate sets was found to be ± 0.02 µmol kg⁻¹ (n = 2). Sampling reproducibility was estimated by analyzing sets of replicate samples drawn from different Niskin bottles during rosette casts. The average standard deviation between replicate pairs and was found to be ± 1.3 µmol kg⁻¹ (n = 22). Combining instrument accuracy, sampling reproducibility, and the concentration-dependent correction uncertainty, the estimate of overall measurement uncertainty is ± 1.8 µmol kg⁻¹.

**Key method references:**
**Method:** Discrete pH – (no units)

**Document author and contact info:** Andrea Fassbender, fassbender@mbari.org

**Brief description of protocol and relation to export pathways:** Discrete samples for pH were collected into 250 mL borosilicate bottles and analyzed aboard the *R/V Sally Ride* spectrophotometrically using an automated system designed after the one described in *Carter et al.*, [2013]. The temperature of the sample was held constant at 20 °C using a 10-cm jacketed cell, and every sample was immersed in a 20 °C water bath for at least 25 minutes before analysis. An indicator dye (purified m-cresol purple from Dr. Robert Byrne’s lab) solution (2 mM) was used to assess sample pH. The sample pH perturbation caused by dye addition was quantified by adding both the normal amount and twice the amount of dye to seawater solutions of ~pH 7.4, 7.8, and 8.1. The dye perturbation was conducted prior to the cruise, during the cruise, and after the cruise yielding an average perturbation (relative to the isosbestic point, Aiso) of: \( \Delta pH/\Delta A_{iso} = -0.0462 \times pH + 0.344 \).

Multiple laboratory comparisons between the ZEISS MMS spectrophotometer used on the cruise and the standard, state-of-the-art Agilent 8453 spectrophotometer used in the laboratory yielded the following pH dependent correction: \( \Delta pH_{Agilent-MMS} = 6.33 \times 10^{-3} \times pH_{Agilent} - 0.0406 \).

Discrete pH samples were used to validate independent, algorithm approaches (e.g., *Juranek et al.*, 2011; *Carter et al.*, 2016, 2017; *Sauzède et al.*, 2017; *Bittig et al.*, 2018) for estimating pH to correct pH sensor drift on profiling floats throughout their lifetimes. Float pH observations are used with TA estimates to calculate dissolved inorganic carbon (DIC) for biogeochemical tracer budget assessment of carbon export. Additionally, float pH measurements and TA estimates are used to calculate sea surface \( pCO_2 \) to quantify the air-sea exchange of carbon dioxide, which is relevant for the DIC tracer budget.

**Other contributing protocols:** N/A.

**Uncertainties and quality control concerns:** Measurement precision was estimated by analyzing sets of duplicate samples (same Niskin bottle) from each rosette cast \( (n = 45) \). The average standard deviation between duplicate pairs and was found to be \( \pm 0.0005 \). Sampling reproducibility was estimated by analyzing sets of replicate samples (different Niskin bottle) from some rosette casts \( (n = 21) \). The average standard deviation between replicate pairs and was found to be \( \pm 0.0009 \).

**Key method references:**


Method: Discrete total alkalinity (TA) – (μmol kg⁻¹)

Document author and contact info: Andrea Fassbender, fassbender@mbari.org

Brief description of protocol and relation to export pathways: Discrete TA samples were collected into 500 mL and 250 mL borosilicate bottles and preserved with 200 μL and 100 μL, respectively, of saturated mercuric chloride for later analysis at the Monterey Bay Aquarium Research institute (MBARI). TA was analyzed using a Metrohm 855 automated titrator following standard open cell alkalinity titration procedures. The titrant was comprised of 0.1 M hydrochloric acid (HCl) in a 0.7 M sodium chloride (NaCl) background solution. The titration temperature was held constant at 20.0 ± 0.2 °C throughout the titration. The temperature of the sample is measured immediately upon delivery to the jacketed cell for later sample mass determination using density and volume. All samples were run in triplicate, and the results were averaged. Certified Reference Material (CRM; provided by Andrew Dickson at SIO) were run every 10 samples (approximately once every hour) to ensure accuracy. The average accuracy of the instrument relative to CRMs is better than 4 μmol kg⁻¹.

Discrete TA samples were used to validate TA estimates derived from algorithms [Carter et al., 2016, 2017; Sauzède et al., 2017; Bittig et al., 2018] applied to profiling float observations that are used to close biogeochemical tracer budgets and quantify carbon export.

Other contributing protocols:

Uncertainties and quality control concerns: Measurement precision was estimated by analyzing sets of replicate samples drawn from the same Niskin bottle during rosette casts. The average standard deviation between replicate sets was found to be ± 0.04 (n = 2). Sampling reproducibility was estimated by analyzing sets of replicate samples drawn from different Niskin bottles during rosette casts. The average standard deviation between replicate pairs and was found to be ± 2.0 (n = 23).

Key method references:
**Method:** High-Resolution Nitrate Profiles on CTD Rosette – (µmol kg⁻¹)

**Document author and contact info:** Andrea Fassbender, fassbender@mbari.org

**Brief description of protocol and relation to export pathways:** Vertical water column profiles of nitrate at ~1 m resolution were obtained at each sampling station using an In Situ Ultraviolet Sensor (ISUS) for nitrate [Johnson and Coletti, 2002]. The instrument was custom built in the Chemical Sensor Laboratory at the Monterey Bay Aquarium Research Institute and calibrated in the laboratory prior to the cruise. The ISUS was mounted vertically on the rosette frame approximately 50 cm above the SeaBird Electronics 911 conductivity-temperature-depth sensor (CTD). Nitrate concentration was calculated from the ISUS ultraviolet absorbance spectra (200-240 nm) using an updated algorithm, where the bromide spectra is calculated from salinity [Sakamoto et al., 2009]. Corrections for the pressure dependence of bromide ultraviolet absorption were also included [Sakamoto et al., 2017]. The sensor collected data at approximately 1 Hz. Raw ISUS nitrate data and contemporaneous, quality-controlled CTD data (time stamp matched) were used to compute the final nitrate values. Results from the downcast were binned at a 1 m interval to comprise the final profile data.

High-resolution nitrate profiles from ISUS sensor measurements on the CTD rosette were compared to nitrate profiles from two biogeochemical profiling floats in the region. The high-resolution nitrate profiles provide context about vertical nutrient gradients, which can be challenging to capture from lower-resolution (e.g., 5-10 m) observations, such as on the floats.

**Other contributing protocols:**

**Uncertainties and quality control concerns:** To be completed.

**Data products originating with this method:** NO3 (mmol/m³)

**Key method references**


**Method:** Underway Nitrate (µmol kg⁻¹) and pH (unitless)

**Document author and contact info:** Andrea Fassbender, fassbender@mbari.org

**Brief description of protocol and relation to export pathways:** A prototype, dual nitrate-pH system was integrated into the ship’s underway seawater line. Nitrate was measured using an In Situ Ultraviolet Sensor (ISUS) (1), and pH was measured using a Deep-Sea-Durafet (DSD) (2). A SeaBird Electronics 45 thermosalinograph was located directly downstream (< 10 cm) of the system flowcell to make underway temperature and salinity measurements near the sensors. The instruments were powered through an isolation transformer to prevent ground loop issues. The system was polled using a LabView interface, and measurements were made every 15 to 20 seconds. The pH sensor was calibrated by taking discrete samples from the underway line (n = 9) throughout the cruise.

Underway nitrate and pH measurements were used to identify fronts. Additionally, the underway observations were compared to those on two biogeochemical profiling floats deployed in the region.

**Other contributing protocols:** High-Resolution Nitrate Profiles on CTD Rosette

**Uncertainties and quality control concerns:** To be completed.

**Data products originating with this method:** NO3 (mmol/m³), pH (none)

**Key method references**


Introduction

This dataset contains Fast Repetition Rate (FRR) fluorescence data, incident Photosynthetically active radiation (PAR) data, and sea surface temperature data for NASA EXPORTS north pacific field campaign.

FRR and SST data were collected continuously during the cruise using sample water drawn from the ship’s flow through seawater system.

PAR data reported here were collected with a Licor cosine collector positioned on the top rail of the port side aerosol van on the upper forward weather deck of the Atlantis, providing measurements relatively free of any ship shading. PAR data are reported in units of uM quanta/cm^2/s.

The FRR was characterized by the manufacturer, Zbignew Kolber. The Licor sensor was calibrated by Licor shortly before the cruise. Data submitted to SeaBASS from the FRR include initial fluorescence (F0), Maximum Fluorescence (Fm), Variable Fluorescence (Fv/Fm), and the functional cross section of photosystem II (Sigma-PSII). These properties were derived from the single turnover flash sequence from the FRR. 16 individual flash sequence results were averaged into each reported value. Seawater sample analyzed by the FRR was exposed to darkness from the time the water was drawn into the ship to the time of measurement (estimated as a few minutes).
Method: Volume Scattering Function at 9 angles (VSF-9)

Document author and contact info: Scott Freeman (scott.freeman@nasa.gov)

Brief description of protocol: The ECO volume scattering function meter measures scattering at nine angles at 532 nm. The vsf-9 uses LEDs modulated at 1 kHz for source light. The source light enters the water volume and scattered light is detected by a detector positioned such that the acceptance angle forms a specific intersection with the source beam.

Calibration

The calibration applied will be conducted by M. Twardowski and company using NIST-traceable microsphere beads. We will also have an intercalibration exercise before and after the cruise. The angles, in degrees, are: 62, 76, 80, 90, 110, 120, 140, 160, 170.

Deployment Methodology

The package is lowered to 5-10 meters to thermally equilibrate and de-gas. The instrument is switched on after a time delay, then the package is brought to the surface and a slow descent (~0.3 m$^{-1}$) is started. Near the seafloor or approximately 100 meters, the package is held for a minute before being raised at ~0.5 m$^{-1}$ or faster. Only downcast data are used.

The VSF-9 is mounted to the FSG Wetlabs IOP cage, facing downward and positioned such that the cage reflects none of the light emitted by the LEDs. Data are stored in a Wetlabs DH-4 and offloaded after each deployment.

4) Derived Parameters

Particulate VSF at 9 angles, 532 nm ($\beta(\theta, 532)_p$) [1/m/sr], after subtraction of VSF$_{water}$ (Zhang et al, 2009); and integrated particulate backscatter at 532 nm ($b_{bp532}$) [1/m].

Uncertainties and quality control concerns:
Uncertainty is reported as standard deviation in a one-meter bin.

5) References


Method: Microbial metagenomics and metatranscriptomics

Document author and contact info: Scott Gifford, sgifford@email.unc.edu

Brief description of protocol and relation to export pathways: The primary objective is to characterize prokaryote community composition via metagenomics and microbes’ physiological and metabolic responses to the carbon pool via community transcriptome sequencing (metatranscriptomics). Linkages between microbial community composition and metabolic activities with respiration, primary productivity, and net community production rate measurements will illuminate the underlying factors controlling carbon availability for export. Seawater samples (4L) will be fractioned between particle-associated and free-living bacterial communities by filtration through 5 μm and 0.22 μm filters, respectively, and then immediately flash freezing both filters in liquid nitrogen. DNA will be extracted using MoBio’s DNA isolation kit. RNA extractions will use Ambion’s mirVana kit. For both DNA and RNA samples, internal standards will be added just prior to extraction. Internal standard recovery efficiencies after sequencing will enable reads of gene or transcript abundances to be placed on a per cell or liter basis (Gifford et al., 2011, 2013). After extraction, DNA libraries will be prepared using KAPA Biosystem’s Hyperplus kit with Roche barcodes. RNA samples will be rRNA depleted using the custom rRNA subtraction protocol of Stewart et al. (2010) and libraries prepared using EpiCentre’s ScriptSeq v2 kit. After pooling, libraries will be sequenced via the Illumina HiSeq 4000 platform at UNC’s High-Throughput Sequencing Facility. Metagenomes will be sequenced to a target depth of 50 -100 million reads per sample. Metatranscriptomes will be sequenced to a target depth of 10-20 million reads per sample. After sequencing, reads will undergo quality control trimming and pairing using custom workflows on the Gifford lab’s slipstream appliance. Metagenomic reads will assembled into contigs using MetaBAT (bitbucket.org/berkeleylab/metabat). Taxonomic identification and putative protein function will be annotated via Diamond homology searches against NCBI’s nr and RefSeq database. Metatranscriptomic reads will be mapped to metagenomic contigs using Bowtie.

Other contributing protocols: DNA sequencing for 16S and 18S, metagenomics and metatranscriptomics, bacterial abundance, DOC concentrations, marine snow characterization.

Uncertainties and quality control concerns: RNAs have short half-lives (minutes) and are highly susceptible to degradation by RNAases, which are ubiquitous in the environment. Per liter and per cell gene or transcript abundance estimates from meta-omic datasets requires good measurements of filtration volumes and stringent extraction protocols.

Data products originating with this method:

Parameter | Units
--- | ---
Taxa abundances | Genome equivalents L⁻¹
Transcript abundances | Transcripts L⁻¹ or Transcripts genome-equivalent⁻¹
Gene abundances | Genes L⁻¹

Key method references
**Method:** O\(_2\) drawdown community and bacterial respiration rates

**Document author and contact info:** Scott Gifford, sgifford@email.unc.edu

**Brief description of protocol and relation to export pathways:** The primary objective is to measure carbon remineralization rates of the community and bacterioplankton in both the free-living and particulate associated size fractions. These rates will help constrain the remineralization component of NCP and reduction in carbon export potential. Biological Oxygen Demand (BOD) bottles (60 ml) containing PreSens oxygen optode sensors spots will be acid washed and triple rinsed with sample seawater before each incubation. The bottles will be filled with either whole (unfiltered) or 5 µm filtered seawater directly from the Niskin bottles. After filling, the BOD bottles are capped with a glass stopper and submerged in a dark water bath set at in situ temperature. O\(_2\) concentrations will be measured every 4 to 6 hours for 48 hours using the PreSens Fibox Fiberoptic sensor to determine oxygen drawdown (Edwards et al. 2011). Using this technique, we have previously measured rates over a range of productivity gradients, from highly productive upwelling systems to open ocean oligotrophic systems, where respiration rates are often 1 µM O\(_2\) d\(^{-1}\) or less.

**Other contributing protocols:** DNA sequencing for 16S and 18S, metagenomics and metatranscriptomics, bacterial abundance, DOC concentrations, marine snow characterization.

**Uncertainties and quality control concerns:** Low DOC standing stocks can introduce two issues: 1) Respiration rates < 1 µM per day can be difficult to resolve. We have previously overcome this limitation by increasing the incubation time to 48 hours. However, one must ensure that drawdown is linear over this period and bottle effects are not biasing results. 2) DOC contamination: The low ambient DOC concentrations mean that even small amounts of carbon contamination can artificially increase rates.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Community respiration</td>
<td>mmol-C m(^{-3}) d(^{-1})</td>
</tr>
<tr>
<td>Free-Living bacterial respiration</td>
<td>mmol-C m(^{-3}) d(^{-1})</td>
</tr>
<tr>
<td>Non-sinking particle respiration</td>
<td>mmol-C m(^{-3}) d(^{-1})</td>
</tr>
<tr>
<td>Small-sinking particle respiration</td>
<td>mmol-C m(^{-3}) d(^{-1})</td>
</tr>
</tbody>
</table>

**Key method references**


Method: Phytoplankton concentrations and elemental stocks

Document author(s) and contact info: Jason Graff, jrgraff@science.oregonstate.edu, Michael Behrenfeld, mjb@science.oregonstate.edu

Brief description of protocol and relation to export pathways:

Phytoplankton cell concentrations for four major groups (Prochlorococcus, Synechococcus, picoeukaryotes, and nanoeukaryotes) will be determined using a Becton Dickinson Influx Cells Sorter (BD ICS) flow cytometer. Whole seawater samples will be analyzed for multiple depth profiles each day. At each depth, 4 ml of seawater are to be collected into sterile 5 ml polypropylene tubes (3x rinsed) and immediately stored at ~40°F in the dark until analysis on the BD ICS. Groups specific identifications are determined based on fluorescence and scattering properties. Our ICS is equipped with a blue (488 nm) laser and four detectors; forward scatter (FSC) with enhanced small particle detection, side scatter (SSC), fluorescence at 692 +/- 40 nm (FL692) and fluorescence at 530 +/- 40 nm (FL530). Sample flow rates are required for normalizing cell counts collected over time to volume and are calculated from volumetric changes in a 1 ml water sample over a known time (60 s or greater) using a pipettor to determine the volume of water lost. The ICS is calibrated with fluorescent beads following standard operating protocols (Spherotech, SPHEROTM 3.0 µm Ultra Rainbow Calibration Particles and Drop Delay beads are used to calibrate instrument timing for cells sorting).

Our new method for measuring phytoplankton carbon and nitrogen (C_phyto, N_phyto) involves separating phytoplankton from natural assemblages using the BD ICS, which is a particle sorting flow cytometer, and then performing elemental analysis on the sorted sample using a Shimadzu TOCN analyzer (full details are published in Graff et al. 2012 and Graff et al. 2015). Briefly, prior to sample analysis and sorting, whole seawater is passed through a 64µm screen to eliminate large, but generally rare, cells and particles that cannot pass through the 100 µm nozzle that we typically use for this method. Sorted cells, and the associated sheath fluid in which the cells are passed to for analysis, are collected and then stored in LN or at -80 C until analysis. A ‘blank’ or correction sample of the carrier sheath fluid is collected immediately following sorting of each sample to account for non-target carbon and nitrogen. These samples are also frozen with the sorted cell samples until elemental analysis. Samples from the surface mixed layer and from specific depths will be targeted for direct phytoplankton elemental analysis. The contribution of larger cells to biomass can be made from size fractionated chlorophyll and carbon estimates of phytoplankton from the Imaging Flow Cytobot (IFCB). Net primary production and phytoplankton biomass (C_phyto - in units of carbon per volume) occupy the first step in the transfer of inorganic carbon to the diverse ecological pathways of organic carbon. Traditional approaches for quantifying phytoplankton biomass from retrievals of surface chlorophyll are compromised because of drastic seasonal changes in cellular chlorophyll:carbon (Chl:C) ratios (Westberry et al. 2016). This variability is driven by seasonality in mixed layer light conditions, severity of iron stress, and species compositional shifts, and it has significantly impacted our understanding of carbon cycling in the region. Our direct measurements of the community structure and biomass, coupled with optical proxies of carbon stocks, are the important first steps to understanding carbon pathways leading to export.
Other contributing protocols: $^{14}$C NPP and dilution experiment derived $\mu$; NPP/$\mu = C_{\text{phyto}}$, optical proxies for $C_{\text{phyto}}$ – specifically particulate backscattering ($b_{bp}$), IFCB analysis of overlapping and larger phytoplankton

Uncertainties and quality control concerns: Multiple samples collected each day for cell counts and elemental analysis will constrain daily values and the characterization of changes occurring throughout each 8-day epoch planned for the EXPORTS field program.

Data products originating with this method: None – all currently exist within the SeaBASS framework.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concentration of Synechococcus</td>
<td>cells L-1</td>
</tr>
<tr>
<td>Concentration of Prochlorococcus</td>
<td>cells L-1</td>
</tr>
<tr>
<td>Concentration of picoeukaryotes</td>
<td>cells L-1</td>
</tr>
<tr>
<td>Concentration of nanoeukaryotes</td>
<td>cells L-1</td>
</tr>
<tr>
<td>$C_{\text{phyto}}$</td>
<td>$\mu g$ L-1</td>
</tr>
<tr>
<td>$N_{\text{phyto}}$</td>
<td>$\mu g$ L-1</td>
</tr>
</tbody>
</table>

Key method references:


Method: Photoacclimation

Document author and contact info:
Jason Graff
jrgraff@science.oregonstate.edu
541-737-4090

Brief description of protocol and relation to export pathways:
Growth, grazing, and accumulation rates are targeted parameters in the EXPORTS program for relating surface plankton properties to carbon export to the deep ocean. Phytoplankton are able to rapidly respond to changes in mixed layer growth irradiance through a process known as photoacclimation and can skew rate estimates based on chlorophyll (Chl) if not properly taken into account. Our primary objective is to measure phytoplankton photoacclimation by observing changes in phytoplankton specific Chl to carbon (C_{phyto}) ratios or their proxies of fluorescence (FL) and forward scatter (FSC) from in-situ and incubated whole seawater samples. In-situ sampling and on-deck experiments, with and without iron additions, will be performed to track changes in Chl:C_{phyto} and FL:FSC ratios in response to different light treatments and in-situ changes in growth irradiance. Multiple light treatments in on-deck incubations allow us to determine the growth irradiance to which cells were acclimated at the time of collection and the change due to experimental light conditions. This is critical for correctly assessing the balance between Chl based phytoplankton growth rates and zooplankton grazing rates in on-deck incubations, e.g. dilution experiments. In-situ tracking of Chl:C_{phyto} can provide corrections for net accumulation rates based on Chl or FL alone. Specific measurements that will be made include high performance liquid chromatography (HPLC) Chl, C_{phyto} using cell sorting approaches, and phytoplankton FL and FSC parameters from flow cytometry.

Other contributing protocols: Establishing active mixing and/or mixed layer depths will be critical for evaluating the light environment experienced by cells. Contributing assets that do not include discrete measurements include optical parameters measured via in-line flow through seawater, water column profiles, and autonomous vehicles.

Uncertainties and quality control concerns:
TBD – these have yet to be addressed using this protocol and will be determined from recently completed experiments.

Data products originating with this method
Percent correction (+/-) for Chl based rates of phytoplankton accumulation in-situ and on-deck experiments.

7. Key method references.
Graff, J.R. and M.J. Behrenfeld. (in prep) Deep mixing and re-stratification events drive vertical differentiation in phytoplankton physiology (and accumulation) in the North Atlantic ocean
**Method:** Volume Scattering Function measurements and particle size distributions.

**Document Author and Contact Info:** Deric Gray  (deric.gray@nrl.navy.mil)

**Brief Description of Instrument Protocol:** The volume scattering function (VSF) of seawater will be measured with two instruments: the Multi-Spectral Volume Scattering Meter (MVSM), a prototype instrument developed at the Marine Hydrophysical Institute in Sevastopol, Crimea, and the LISST-100X (Sequoia Scientific). The MVSM measures the VSF in 0.25° increments over the angles from 0.5 – 179°, and at eight wavelengths: 443, 488, 510, 532, 555, 565, 590, and 620 nm. The LISST (operating at 532 nm) measures the VSF from 0.1 – 13° in 32 angular intervals. The data from both instruments are combined to produce the VSF from 0.1 to 179°. The resulting VSF is then inverted to derive particle size distributions (PSD) from 0.02 – 200 µm.

**Deployment Methodology:** Discrete water samples will be collected from the CTD rosette and measured with the instruments in the lab. Approximately 10L of water is needed from each depth, and measurements will take 20 minutes per depth. Water filtered through a 0.2 µm filter from the deepest depth will be measured to establish a baseline for the LISST instruments, and additional baselines measured as needed. Surfaces measurements from the inline flow-through system will also be measured periodically throughout the cruise.

**Data Products:** The directly measured data are the volume scattering functions. Data from the LISST is inverted by the standard processing software (LISST-SOP 5.1) to provide particle size distributions from 1-250 µm. The complete VSF from the MVSM+LISST is also inverted to provide PSDs from 0.02 – 200 µm.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>VSF</td>
<td>0.1 – 179° (443, 490, 510, 532, 555, 565, 590, 620 nm)</td>
<td>m$^{-1}$ Str$^{-1}$</td>
</tr>
<tr>
<td>PSD</td>
<td>0.02 – 200 µm</td>
<td>m$^3$ µm$^{-3}$</td>
</tr>
</tbody>
</table>

**Uncertainties:**
Instrument: The largest instrument uncertainties result from calibration errors and instrument drifts during long deployments. Instrument drifts will be monitored by periodic measurements of pure water during the deployment. The instruments will be calibrated pre- and post-cruise in laboratory, and also mid-cruise on the vessel. Measurement: Repeated measurements will be made of each water sample during the cruise to assess measurement uncertainties. Sample uncertainty will be minimized by mixing water from all Niskin bottles collected at each depth.

**Key Method References:**
Cruise name: EXPORTS 1
Cruise id: RR1813
Ship: R/V Roger Revelle
Location: Station Papa, North East Pacific
Dates at sea: 2018/08/11 to 2018/09/12 01:00
   Epoch 1:2018/08/14 4:30 to 2018/08/23 9:00
   Epoch 2:2018/08/23 9:00 to 2018/08/31 9:00
   Epoch 3:2018/08/31 9:00 to 2018/09/08 9:00
At Station P: 2018/08/14 0:00 to 2018/08/23 9:00
Operators: Nils Haëntjens and Emmanuel Boss
Group Leaders: Emmanuel Boss and Lee Karp-Boss
ACS serial numbers: 298 and 301

Revision 2:
Gamma in the file EXPORTS-EXPORTSNP_InLine-ACS-Products_20180811-20180912_R1 was not processed for the ACS 301 and was populated with -9999 (NaN) values. The bug was corrected and a beautiful diel cycle can now be observed for the entire campaign. Only the file EXPORTS-EXPORTSNP_InLine-ACS-Products_20180811-20180912_R2 is concerned by this revision. The processing stays identical.

We use a calibration independent technique (Slade et al., 2010) to obtain particulate absorption (ap) and attenuation (cp) by differencing measurements with a 0.2um filter from measurements made with no filter. Dissolved absorption and attenuation are obtained by subtracting daily MilliQ run from .2um filtered measurements. Filters are exchanged weekly and flow-tubes are cleaned every day. Switching between filtered and unfiltered measurements is done every 60min (50min total, 10min dissolved). In addition, MilliQ water was run every day after the instrument cleaning through the instrument to obtain ag and cg spectrums.

Two ACS, serial number 298 and 301, are used during the campaign, ACS 298 was used from the beginning of the campaign to 2018/08/20 18:10 and ACS 301 was used from 2018/08/20 20:17 to the end of the expedition. The change in ACS is not related to any dysfunction of instrument on the InLine system but rather an issue with the ACS-301 when profiling, not affecting underway observations. Bad spectrums are removed manually and arise, generally due to bubbles going through the instrument.

For each minute, the remaining data between 15th and 75th percentiles are binned-averaged and their standard deviation is kept for reporting. The particulate bins are processed by subtracting the filtered measurements from the unfiltered measurements. Filtered values needed to obtain the particulate values are interpolated to the time of particulate measurements linearly. Dissolved bins are computed by differencing MilliQ and filtered data. The MilliQ values are linearly interpolated to match the filtered periods.
The mismatch in spectral band positions between absorption and attenuation are corrected using interpolation. We use the 3rd method of Zaneveld et al., 1994 to correct for scattering with 730nm as the null wavelengths simultaneously performing a residual temperature correction (Slade et al., 2010). Attenuation is also corrected for residual temperature effect. Then, we perform a spectral unsmoothing based on the method in Chase, A., et al., 2013. We have left spectra with negative absorption in the blue regions, as these values are not significantly different from zero.

The device files are used for wavelength registration, to convert binary counts from the instruments into scientific units (1/m) independent of instruments temperature, and to indicate the last service of the instrument.

While the dissolved absorption and attenuation spectrums are available they must be used with caution and only a couple of hours a day might be valid due to bio-fueling of the instrument which can’t be assessed with the current method. For more information please contact us (emmanuel.boss@maine.edu, nils.haentjens@maine.edu).

Additional products derived directly from the ap or cp spectrums are provided.

- Chlorophyll a (chl) is computed using the particulate absorption line height at 676 nm and the global relationship from Tara Ocean (Boss et al. 2013):
  - line_height = a_p(676) - (39/65 × a_p(650) + 26/65 × a_p(715))
  - chl = 157 × line_height^1.22 (relationship NOT applied here, from Tara Ocean)
- The particulate organic carbon (POC) is computed using the particulate attenuation at 660 nm Using the global relationship from Gardner et al. (2006):
  - POC = 380 × c_p(660)
- Gamma is computed using the method of Boss et al. 2001.

**Figure 1.** Particulate absorption spectrums during the entire expedition measured with ACS 298 (left) and ACS 301 (right). The higher absorptions at the beginning and the end of the expedition are in to coastal waters.
References:

Emmanuel Boss, Marc Picheral, Thomas Leeuw, Alison Chase, Eric Karsenti, Gabriel Gorsky, Lisa Taylor, Wayne Slade, Josephine Ras, Herve Claustre, 2013. The characteristics of particulate absorption, scattering and attenuation coefficients in the surface ocean; Contribution of the Tara Oceans expedition, Methods in Oceanography.


Cruise name: EXPORTS 1
Cruise id: RR1813
Ship: R/V Roger Revelle
Location: Station Papa, North East Pacific
Dates at sea: 2018/08/11 to 2018/09/12 01:00
   Epoch 1: 2018/08/14 4:30 to 2018/08/23 9:00
   Epoch 2: 2018/08/23 9:00 to 2018/08/31 9:00
   Epoch 3: 2018/08/31 9:00 to 2018/09/08 9:00
   At Station P: 2018/08/14 0:00 to 2018/08/23 9:00
Operators: Nils Haëntjens and Emmanuel Boss
Group Leaders: Emmanuel Boss and Lee Karp-Boss
ALFA Serial Number: 011

The WETLabs Aquatic Laser Fluorescence Analyzer, (ALFA) spectrofluorometer was mounted on the flow through system of the ship, after a vortex debubbler and switching system. The switching system automatically ran filtered seawater (0.2 um) the first 10 minutes of every hour, unfiltered/total seawater was running through the instruments the rest of the time. The instrument was cleaned daily at night time with bleach and laboratory grade soap. A peristatic pump was used to pump the water through the instruments of the underway.

The data presented here was collected and preprocessed by the manufacturers software, manually quality checked (un-realistic values mainly being due to bubbles running through the system), and minute binned. Data collected when the water was filtered was removed from the dataset and not used in the processing but can be distributed per request to the data owner.

TSG, PAR, and the chlorophyll fluorometer (Wetlabs WS3S) data from the R/V Roger Revelle are added to the SeaBASS file to help for data analysis.

Parameters collected by the ALFA are:
- Chl_stimf_ex405: chlorophyll a fluorescence excited at 405 nm
- Chl_stimf_ex514: chlorophyll a fluorescence excited at 514 nm
- Fv_Fm_ex405: Fv/Fm value with 405 nm excitation
- Fv_Fm_ex514: Fv/Fm value with 514 nm excitation
Cruise name: EXPORTS 1
Cruise id: RR1813
Ship: R/V Roger Revelle
Location: Station Papa, North East Pacific
Dates at sea: 2018/08/11 to 2018/09/12 01:00
  Epoch 1:2018/08/14 4:30 to 2018/08/23 9:00
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  Epoch 3:2018/08/31 9:00 to 2018/09/08 9:00
  At Station P: 2018/08/14 0:00 to 2018/08/23 9:00

Operators: Nils Haëntjens and Emmanuel Boss
Group Leaders: Emmanuel Boss and Lee Karp-Boss
ECO-BB3 serial numbers: 349

WetLabs ECO-BB3 serial number 349 was measuring the angular scatterance at 1 Hz at the three wavelength (470, 532, 660 nm). The data was logged with a home-grown data-logger (Inlinino, http://inlinino.readthedocs.io/). The system is run on an hourly schedule of whole water for 50min followed with 10min of filtered measurements (using a 0.2 µm filter). In processing, the first 400 seconds data collected after switching from total filtered and vice-versa are ignored corresponding to the time it takes to renew the water in the BB-Box. 20-40 L of MilliQ water are run daily through the system to estimate the dissolved backscattering coefficient.

Period with obvious bad measurements are removed manually (most likely due to large clouds of bubbles or accumulation of big particles in coastal waters).

For each minute, the measurements between the 15th and 75th percentiles are averaged and their standard deviation is kept for reporting. Both the dissolved ($\beta_g$) and particulate ($\beta_p$) VSF are computed depending on switch position. The dissolved VSF is obtained by subtracting the MilliQ measurements from filtered measurements (interpolating in time between successive daily MilliQ values). The particulate VSF is obtained by subtracting the filtered from the total values (filtered values are linearly interpolated). Those differences take care of the dark and wall effects of the BB box. The slope coefficient used (table 1) comes from the latest calibration done by Jim Sullivan of FAU on 6/29/16. A temperature and salinity correction is performed on the dissolved using Zhang et al. 2009.

The particulate backscattering coefficient ($b_{bp}$) is computed using $\chi=1.076$ (nominal angle 124, Sullivan et al, 2013). Note: the reported value for particulate backscattering does not include the contribution of the fraction below 0.2um.

<table>
<thead>
<tr>
<th>Nominal Wavelength [nm]</th>
<th>Slope [sr$^{-1}$ count$^{-1}$]</th>
<th>Dark [counts]</th>
<th>Uncertainties</th>
</tr>
</thead>
<tbody>
<tr>
<td>468.7</td>
<td>8.407E-6</td>
<td>55.5</td>
<td>Max(11%,6E-5)</td>
</tr>
<tr>
<td>527</td>
<td>4.624E-6</td>
<td>50.8</td>
<td>Max(10%,3E-5)</td>
</tr>
<tr>
<td>652</td>
<td>4.090E-6</td>
<td>43.7</td>
<td>Max(17%,5-5)</td>
</tr>
</tbody>
</table>
References
**Method:** NPP, 24 h and 2 h H${}^{14}$CO$_3$ incubations

**Document author and contact info:** Kim Halsey, halseyk@science.oregonstate.edu, 541-737-1831; James Fox, james.fox@oregonstate.edu

**Brief description of protocol and relation to export pathways:** Our primary objective is to measure $^{14}$carbon assimilation in seawater samples. These (A) 24 hour and (B) short-term (2 h) incubations will allow us to calculate net organic carbon production (NPP) and will yield information about the physiology of the phytoplankton community. NPP describes the rate of CO$_2$ conversion into organic matter; the first step of the carbon cycle. A key goal of EXPORTS is to determine the fraction and rate of NPP that is transferred to the dark ocean. We will apply the commonly used $^{14}$C-uptake method (Steeman Nielsen, 1952). Seawater samples will be spiked with $^{14}$C-labeled bicarbonate, incubated at a range of light levels, filtered, acidified, and then measured by scintillation counter to determine the amount of $^{14}$C incorporated into biomass. 24 h incubations will be conducted from dawn-to-dawn in on-deck temperature controlled incubators with and without screening to estimate NPP throughout the euphotic zone. Short term (2 h) incubations will be done in a photosynthetron.

**Other contributing protocols:** NPP can be normalized to a variety of different parameters (e.g., Chla, total absorption, C$_{phyto}$) especially for use in linking to satellite retrievals. Estimates of light intensities with depth will be needed to match $I_g$ to incubation light exposures.

**Uncertainties and quality control concerns:** Duplicate measurements help constrain the uncertainty.

**Data products originating with this method:** Short term incubations yield $P_{max}$ and alpha ($\alpha$) which can be used to determine the maximum quantum yield $\phi_m$ and possibly growth rate ($\mu$).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>$P_{max}$</td>
<td>mol C (m$^3$ d)$^{-1}$</td>
</tr>
<tr>
<td>alpha ($\alpha$)</td>
<td>mol C m$^2$ s (mol photons d)$^{-1}$</td>
</tr>
<tr>
<td>Maximum photosynthetic quantum yield ($\phi_m$)</td>
<td>mol C (mol photons)</td>
</tr>
<tr>
<td>Growth rate ($\mu$)</td>
<td>d$^{-1}$</td>
</tr>
</tbody>
</table>

**SeaBASS submission fields and units:**

(fields=depth,lightlevel,rate_14C_uptake_bottle_24hr_0.2umfilt,R2R_event,date,time
(units=m,%,mol/l/d,none,yyyyymmdd,hh:mm:ss)
Key method references


<table>
<thead>
<tr>
<th>Variable</th>
<th>Biogenic Silica</th>
</tr>
</thead>
<tbody>
<tr>
<td>SeaBASS Name</td>
<td>Bsi</td>
</tr>
<tr>
<td>Units</td>
<td></td>
</tr>
<tr>
<td>Sampling</td>
<td>Niskin bottle on rosette</td>
</tr>
</tbody>
</table>

Document author and contact information: Janice Jones, janice.jones@lifesci.ucsb.edu

## Particulate Si Determination

### At Sea:
1. Filter sample through a 0.6um polycarbonate membrane filter at <10mm Hg, record volume filtered. Nalgene clamp towers work well with the thin PC filters.
2. Fold filter into quarters, place in a screw cap cryovial, cap loosely and dry at 65\(^\circ\)C. Once dry (in ~2 days) store at room temperature. Or cap tightly and freeze at -20\(^\circ\)C and keep frozen until samples can be dried in a drying oven as above.
3. For every box of filters used, make 3 filter blanks – take a clean filter, fold in quarters, place in a cryovial and label BLANK. These blanks will go through the same procedure as the samples.

### In the Lab:

#### Reagents:
- 0.2N NaOH
- 1N HCl
- 0.2 N HF (LSi)
- Saturated aqueous boric acid (LSi)
- Deionized distilled water (Milli-Q or Nanopure)
- Regent for colorimetric seawater silicate

4. 15 ml Teflon centrifuge tubes are stored in 0.5 M HF. Dump the rinse HF from a set of Teflon tubes into the HF reservoir (2L bottle). Rinse the tubes with Nanopure 3x. Transfer a dry filter from a microcentrifuge tube to the bottom of a 15 ml tube keeping filter as open as possible. If necessary, use long forceps to open filter, exposing the surface for NaOH digestion. Rinse forceps well with Nanopure between samples.
5. Make 3 tube blanks – empty Teflon tubes. Treat blanks as you do all other vials with filters.
6. Cover filter with 4ml of 0.2N NaOH. Plug and vortex. Be sure filter remains submerged. Cover each tube with a loose cap to keep condensation drips out of sample.
7. Place in water bath at 95\(^\circ\)C for:
   - RoMP samples = 1 hr
   - BATS samples = 2hr
   - JGOFS = 40 min
   - Plumes and Blooms = 30 min
   - LTER = 40 min
   - EXPORTS = TBD

Updated Apr/2014
8. Cool in ice water bath immediately. Remove foil while cooling. This should take about 3-5 minutes, then samples should be cool enough for acid addition.

9. Add 1.0 ml of 1 N HCl and vortex after each addition. This neutralizes the NaOH and along with the cooling stops the digestion. Work quickly to minimize the time difference between the first and last sample.

10. With a clean spatula or long forceps, gently crunch filter into bottom of tube. Rinse the spatula/forceps with Nanopure between samples. Scrunching the filter makes it easier to withdraw your sample later, but you will need to remove the filter eventually so be gentle.

11. Centrifuge for 10 min at setting 6 to drive the lithogenic Si particles to the bottom of the tube.

12. Withdraw 4ml of the 5ml in the Teflon tube and place in 30ml PP (polypropylene) bottle – be sure to take the sample from the top of the liquid, don't push the pipette tip down into the filter. Do this in front of the light – you can see the liquid level with the backlight. Add 18ml of Nanopure to the PP bottle for a total of 21ml.

13. From the PP bottle containing 21ml of sample, transfer 10ml to another 30ml PP bottle for DSi analysis. Your dilution factor for this 10ml sample will be \((5/4)*(21/10)=2.625\). If you need to dilute the sample, take an aliquot less than 10ml and make up to 10ml with Nanopure. Reflect the change in your dilution factor: i.e. if you only use 5ml of 21ml then \((5/4)*(21/5)=5.25\).

14. If lithogenic silica concentrations are NOT to be determined skip to step 22.

15. To rinse the LSi sample, add 7ml Nano to the remaining 1ml of sample + filter in the Teflon tube. Plug tube, vortex and centrifuge for 10 min at setting 6 on clinical centrifuge (program 15 on new centrifuge). If filter does NOT spin to the bottom of tube, push it down with clean poker and spin again for 10 minutes (remember we are trying to rid ourselves of any remaining dissolved silica from the BSi digestion not the particulate LSi). Aspirate to 1ml (BE SURE to rinse slurper tip before you use it!!)

16. Repeat rinse/aspirate step. Two rinses are necessary but do more if there was a lot of BSi.

17. Remove the plug from the Teflon tubes, cover with large clean Petri dishes (to allow drying while keeping dust out of tubes), and place in drying oven at 65°C until dry (takes approximately 48hours). Once dry, these filters can be stored at room temperature until you have time to continue – cover the tubes with plugs and caps then cover entire rack with plastic wrap if you're going to let them sit around.

18. When dry, remove filters from oven and let cool. Wear gloves when working with HF, and you may want to work in the hood but it's not necessary. Completely cover each filter with 0.2ml of 2.5M HF.

19. Crunch the filter down into the bottom of the tube with the HF poker (teflon or plastic stirring rod or spatula), remove all air bubbles and completely submerge the filter ball under the HF. Rinse the poker well with Nanopure between samples. The filter will have to be removed from the tube later so you'll want to try and flip it over or lift it slightly off the bottom to make that removal easier!!.

20. Use plugs to tightly caps the tubes. Be sure to treat the tube and filter blanks with 0.2ml of 2.5M HF and the HF poker as well. Let samples and blanks sit covered with HF for 48 hours.

21. A couple of hours before the 48hour waiting period is over, the saturated boric acid solution (~1M, 65g/L) needs to be filtered. The \(\text{H}_3\text{BO}_3\) MUST be freshly filtered prior to use. A total volume of 14.8ml per sample is required, plus 110ml for the standard curve plus enough to make dilutions if necessary. The saturated boric acid should be filtered through a 0.6um 47mm PC membrane filter and collected in

**ONCE THE PMP TUBES FOR LSi ARE IN THE OVEN AND THE BSi SAMPLES ARE READY TO ENTER THE DSi PART OF THE ANALYSIS YOU CAN STOP FOR THE DAY. THE BSI SAMPLES SHOULD BE PROCESSED WITHIN 48 HOURS.**

22. Use plugs to tightly caps the tubes. Be sure to treat the tube and filter blanks with 0.2ml of 2.5M HF and the HF poker as well. Let samples and blanks sit covered with HF for 48 hours.

23. A couple of hours before the 48hour waiting period is over, the saturated boric acid solution (~1M, 65g/L) needs to be filtered. The \(\text{H}_3\text{BO}_3\) MUST be freshly filtered prior to use. A total volume of 14.8ml per sample is required, plus 110ml for the standard curve plus enough to make dilutions if necessary. The saturated boric acid should be filtered through a 0.6um 47mm PC membrane filter and collected in
a clean bottle. Filter enough boric acid for all samples, standard curves and dilutions you might need – it doesn’t hurt to filter too much.

22. Vortex the tube to release the HF inside the crunched filter. Set the dispensette on the filtered boric acid to 7.4ml (CHECK VOLUME!!). Dispense 7.4ml of filtered boric acid into the tube, vortex to resuspend filter and transfer the boric acid, HF and crunched filter to a 30ml PP bottle. Dispense a second 7.4ml aliquot of boric acid into the tube, vortex to rinse the tube and transfer volume to same 30ml bottle (you can do this in 3 aliquots totaling 14.8ml for better tube rinsing if you want). The total volume in the bottle is 15ml – 14.8ml of filtered boric acid and 0.2ml of 2.5M HF.

23. Withdraw 10ml of the 15ml in the PP bottle and transfer to a clean 30ml PP bottle for the DSi reaction. The dilution factor here is (15/10)=1.5.

24. The standard curve is prepared in a 0.2ml:14.8ml ratio of 2.5M HF:filtered saturated boric acid. Mix 3ml 2.5M HF and 222ml filtered boric acid in a beaker, transfer 10ml of this solution to each of the standard curve bottles. **DO NOT USE NANOPURE FOR THE LSi STANDARD CURVE.** Use this solution for dilutions as well. If you'll have dilutions, make a larger volume - be sure to maintain the ratio!

25. Follow same protocol as for Dissolved Si Analysis - 4 ml of the Acid/Moly reagent, wait 10 minutes to form silicomolybdic acid, add 6 ml reducing reagent.

26. Tube blanks correct for signal generated by the effect of the NaOH & HF digestions on the tube. Filter blanks correct for signal generated by the effect of the digestions on the filter. When the signal from the tube blank gets too high (<90%T for 1cm cell, <80% for 10cm cell) the tubes need to be discarded – usually they can only be used for HF approximately 10 times. Since the filter blank incorporates the tube blank, the signal from the filter blank is subtracted from the signal on all samples.

27. Calculate the LSi concentration in the original seawater sample (LSi). Correct the value based on the number of nanopure rinses in steps 14/15. The calculation is as follows:

$$\text{LSi umol/filter}_{\text{corrected}} = \text{LSi umol/filter} - ((1/5)*(1/8)^2)*\text{BSi umol/filter} = \text{LSi} - (0.003125*\text{BSi umol/filter})$$
**Method:** Estimation of particle (dis)aggregation and sinking rates based on size-fractionated lithogenic particle concentrations

**Document author and contact info:** Phoebe J. Lam (pjlam@ucsc.edu); Jong-Mi Lee (jm_lee@ucsc.edu); Olivier Marchal (omarchal@whoi.edu)

**Brief description of protocol and relation to export pathways:**
Size-fractionated particle samples will be collected in the upper water column at stations occupied during EXPORTS from in-situ filtration pumps (McLane Industries) whereby water entering the pump passes through two screens (51 micron followed by 0.8 micron nominal pore size). Subsamples of the 0.8-51 micron and >51 micron particles will be digested using HNO$_3$ and HF (Cullen and Sherrell, 1999). The digested solutions will be diluted and run on the Inductively Coupled Plasma Mass Spectroscopy (ICPMS) to measure concentrations of various trace elements and REEs, including Ti which represents lithogenic particles in the samples.
In addition to this effort of sample collection and analysis, we will develop and apply a two-particle size class model of particle and thorium cycling in the oceanic water column. In this model, the particle size classes will coincide with those sampled at sea. Rates of particle aggregation, disaggregation, remineralization, and sinking at stations occupied during the EXPORTS and GEOTRACES programs will be estimated from a model fit to measurements of size-fractionated lithogenic particle and particulate organic carbon (POC) concentrations and $^{234}$Th activity, provided by this and other research groups. A time-dependent version of the model will be applied to infer temporal variations in these rates from EXPORTS data (time series data), and a steady-state version of the model will be applied to infer horizontal variations in these rates from GEOTRACES data (transect data). For both model versions, the model fit to the data will be obtained from the application of inverse methods: methods of optimal estimation theory for the time-dependent version, and methods of total inversion or nonlinear programming for the steady version. Through this approach, we will be able to document variations in particle (dis)aggregation, remineralization, and sinking rates (for export pathways 1, 2, and 3) over the course of a bloom and across different pelagic environments in the open ocean, with due consideration for the uncertainties in the data and the model.

**Other contributing protocols:** Analytical determination of POC and $^{234}$Th, comparison to direct estimates of sinking particle fluxes using sediment traps and optical flux traps, comparison to particle size distribution data from UVP and other optical systems.

**Uncertainties and quality control concerns:**
Particle data will have uncertainties derived from blanks and other steps determined by standards. Rate constants will have uncertainties derived from both data and modeling errors.

**Data products originating with this method**

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<tbody>
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<td>nmol/kg</td>
</tr>
<tr>
<td>aggregation (0.8-51μm to &gt;51 μm) rate constant</td>
<td>d$^{-1}$</td>
</tr>
<tr>
<td>disaggregation (&gt; 51μm to 0.8-51μm) rate constant</td>
<td>d$^{-1}$</td>
</tr>
<tr>
<td>sinking rate of &gt; 51μm particles</td>
<td>m d$^{-1}$</td>
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</table>
remineralization rate constant \( \text{d}^{-1} \)

**Key method references**


**Method:** Estimation of particle (dis)aggregation and sinking rates based on size-fractionated lithogenic particle concentrations

**Document author and contact info:** Phoebe J. Lam (pjlam@ucsc.edu); Jong-Mi Lee (jm_lee@ucsc.edu); Olivier Marchal (omarchal@whoi.edu)

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**Other contributing protocols:** Analytical determination of POC and $^{234}$Th, comparison to direct estimates of sinking particle fluxes using sediment traps and optical flux traps, comparison to particle size distribution data from UVP and other optical systems.

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</tr>
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<td>d⁻¹</td>
</tr>
<tr>
<td>sinking rate of &gt; 51μm particles</td>
<td>m d⁻¹</td>
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<tr>
<td>remineralization rate constant</td>
<td>d⁻¹</td>
</tr>
</tbody>
</table>

### Key method references

Method: Bio-acoustic measurement of zooplankton biomass

Document author and contact info:
Amy Maas, Amy.Maas@bios.edu, Eric D’Asaro dasaro@apl.washington.edu

Brief description of protocol and relation to export pathways:
The day/night abundance, distribution and species composition of the mesozoooplankton control two of the main exports pathways: active transport and fecal pellet production. These communities can be quite patchy, and some members (particularly larger individuals) are capable of net avoidance. To better constrain the distribution and relative abundance of the zooplankton, and to examine both meso- and finer scale variability in migration behavior, analyses of the acoustic sound scattering layer (SSL) will be conducted. These will be used to inform the active flux pathway by providing information about the variability in the behavior.

During the cruise period, raw acoustic backscatter data will be collected by the Simrad EK80 on the R/V Ride at five frequencies (18, 38, 70, 120 and 200 kHz). Raw data will be integrated over 0.1 min intervals and 1 m depth strata per frequency, then concatenated into daily sections (with Matlab code) to produce echograms from depths of a few meters to at least 400 m. These data will be used to compute relative (qualitative) differences in mesoscale variability, timing of migration, migration depths, etc. based on visual scrutiny of the rate, timing, and amplitude of diel vertical migrations evident in the data. Additional, more quantitative analysis may be possible given the dominant zooplankton species as determined from MOCNESS and UVP data combined with acoustic backscattering models.

During the autonomous mission (July – December), a glider will carry a 1 MHz Nortek Signature ADCP operated as a narrow band (2.9°) echosounder with a large number of narrow (3 mm) bins over a range of a few meters away from the glider. Each bin will sample ~1 ml of water so that each bin is likely to contain only one individual. The system will thus i) count animals, ii) measure the scattering cross-section of each animal and iii) roughly measure the size of animals bigger than 1 mm. These measurements will be made many times per day along profiles extending from the surface to 1000m and will thus be capable of observing diel migrations. Comparisons with more established methods will be made during the cruise period using the same model of ADCP operating as an internally recording, battery powered sampler. During MOCNESS tows, it will be mounted with a view of the net opening, so as to make direct measurements of the acoustic signatures of animals captured. During CTD measurements, it will be mounted with a view of the camera sampling volume so as to obtain simultaneous optical and acoustic measurements of the same animals.

Other contributing protocols: MOCNESS sampling and UVP sampling.

Uncertainties and quality control concerns: It is difficult to go from single or multi-beam echosounders to direct quantification of biomass as some species produce a louder backscatter relative to size and abundance (i.e. fish with swim bladders, pteropods, etc.). The use of a high-frequency, high resolution echo sounder to measure zooplankton has not been previously attempted, so its accuracy is not known.

Data products originating with this method:

<table>
<thead>
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<th>Parameter</th>
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<tr>
<td></td>
<td></td>
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<tr>
<td>depth of migration</td>
<td>m</td>
</tr>
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<td>-------------------</td>
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</tr>
<tr>
<td>% community migration</td>
<td>%</td>
</tr>
</tbody>
</table>

**Key method references**


Method: Zooplankton electron transport system

Document author and contact info: Amy Maas, Amy.Maas@bios.edu

Brief description of protocol and relation to export pathways: Respiration of diel vertical migratory (DVM) zooplankton is one of the five export pathways. Although individual direct measurements of respiration and associated scaling via abundance and biomass provide one measure of this active flux they rely upon the assumption that experiments with a limited number of individual and species can characterize the fully community respiratory rate. Analysis of the Electron Transport System (ETS) captures diel changes in whole mesozooplankton community metabolism in discrete depth intervals throughout the water column. The ETS analysis measures the enzymatic capacity of a sample to transfer electrons to a terminal receptor. When performed on samples of mesozooplankton taken with net tows it is generally interpreted as the “potential respiration” of the organisms within a discrete vertical community (Gómez et al., 1996; Packard, 1971). Using a flash frozen fraction of the MOCNESS samples, enzymatic activity is measured on each vertical strata of all MOCNESS samples using the standard ETS methods (Gómez et al., 1996; Packard, 1971). The enzyme activity will be used to directly calculate the respiratory CO\(_2\) production (active transport; µmol CO\(_2\) m\(^{-3}\) h\(^{-1}\)) in each vertical strata using the equations detailed in Packard and Gómez (2013). These measures will capture both diel and ontogenetic variations in active flux.

Other contributing protocols: MOCNESS abundance and biomass sampling to provide biomass calculations.

Uncertainties and quality control concerns: This assay is a measure of “potential respiration” and is based on an assumption that the abundance of ETS relates directly to the respiration of a community (i.e. that an individual organism is always producing the maximum of ATP possible). This is an oversimplification of organismal physiology, but provides a good first approximation of whole community respiration rate. The calculation of active flux via the ETS method for respiration has been calculated to have an uncertainty of 31-38% based on all sources of error (Packard and Gómez, 2013; Packard et al., 1988), while calculations of respiratory C demands of the mesopelagic organisms have a 40% error (Packard and Gómez, 2013).

Data products originating with this method:

<table>
<thead>
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<th>Parameter</th>
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<td>respiratory flux</td>
<td>mg-C m(^{-2}) d(^{-1})</td>
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SeaBASS submission fields and units:
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Key method references


**Method:** Zooplankton biomass/abundance

**Document author and contact info:** Amy Maas, Amy.Maas@bios.edu

**Brief description of protocol and relation to export pathways:**
The day/night abundance, distribution and species composition of the mesozoooplankton control two of the main exports pathways: active transport and fecal pellet production. To characterize these communities biomass and abundance analyses of the water column will be conducted.

**Biomass:** Biomass analysis will be conducted following established protocols and will provide cross calibration of our results with other datasets (Madin et al. 2001; Steinberg et al. 2000, 2008), with the sample split poured through nested sieves (0.2-, 0.5-, 1-, 2-, and 5-mm mesh), and contents of each concentrated onto pre-weighed, 0.2-mm Nitex mesh filters and frozen on board at -20 °C. Filters will be subsequently thawed, weighed to obtain wet weight, then dried for 24 h at 60°C and weighed again to obtain dry weight. Each dried sample will then be homogenized using a mortar and pestle, and a weighed subsample analyzed for C&N content using an elemental analyzer (Madin et al., 2001). From these measurements wet, dry, and C&N biomass (mg m⁻³) will be calculated by dividing the biomass by the volume filtered through the net. In the event of a salp, doliolid, or other gelatinous zooplankton bloom, measured abundance, size, and biovolume from fresh samples will be converted to wet/dry weight and C&N (Madin et al., 2001; Stone and Steinberg, 2014).

**Abundance:** To characterize mesozooplankton size distribution and taxonomic composition within each depth strata, we will use both the MOCNESS and UVP. The MOCNESS sample split will be preserved in buffered formaldehyde and an aliquot of each preserved sample will be imaged with a ZooSCAN optical imaging system (Hydroptic) using two size classes (> 2 mm, < 2 mm) following established procedures (Gorsky et al., 2010; Picheral et al., 2017; Vandromme et al., 2012). Image analysis of at least 1000 particles per sample will be conducted via the widely used ZooProcess and EcoTaxa pipeline. UVP automated image analyses will be also done using ZooProcess and EcoTaxa. From both the ZooSCAN and UVP analyses, counts and biovolumes will be obtained by major taxon (copepods, euphausiids, doliolids, ostracods, etc). Certain taxa of interest (e.g., known ontogenetic migrators of the calanoid copepod families) will be identified to species using ZooSCAN or microscopy. Biovolumes from the ZooSCAN will be calculated by dividing the biovolumes per group by the split and volume filtered through the net. They will then be added together by animal size to directly compare to the size fractionated biomass measurements to provide an estimation of uncertainty using the equations of Davis & Wiebe (1985).

**Other contributing protocols:** MOCNESS sampling.

**Uncertainties and quality control concerns:** Although the ZooScan and EcoTaxa pipeline are semi-automated, the quality of organismal identification via machine learning algorithms is variable. The learning set will be carefully cultivated and images quality-checked to optimize identification.

**Data products originating with this method:**

<table>
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<th>Parameter</th>
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<tr>
<td>zooplankton dry biomass*</td>
<td>mg m⁻³ d⁻¹</td>
</tr>
<tr>
<td>zooplankton C&amp;N biomass*</td>
<td>mg-C or N m⁻³ d⁻¹</td>
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</tbody>
</table>
Zooplankton abundance** # individuals taxonomic group$^{-1}$ m$^{-3}$

Reported for each of 5 size fractions for each vertical strata of the MOCNESS during both day and night.

**Reported for each vertical strata of the MOCNESS during both day and night.

SeaBASS fields and units:
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Key method references


Method: Zooplankton respiration, excretion, and egestion experiments

Document author and contact info: Amy Maas, Amy.Maas@bios.edu

Brief description of protocol and relation to export pathways:
Respiration of diel vertical migratory (DVM) zooplankton is one of the five export pathways, while excretion of DOC and egestion of POC as fecal pellets at depth may also contribute to “active flux”. We will perform live respiration and excretion experiments on dominant migratory species to provide measurements of key species’ contribution to community active C flux during each ecosystem state sampled. Oxygen consumption will be measured and will serve as a proxy for respiratory CO\textsubscript{2} excretion while DOC excretion and POC egestion will be measured directly. These will be scaled to community level export using the biomass measures and DVM species identification made with the MOCNESS.

Incubations will be carried out for 12 h, in the dark, in filtered seawater, and at daytime residence depth temperature to mimic time spent and conditions (non-feeding, cold) at depth. Prior to incubation, 20 L of water will be collected, filtered (0.2 µm), and stored in an incubator at \textit{in situ} mesopelagic temperature and in darkness for < 8 h prior to use in respiration experiments.

Migrators are collected from the EZ at night. Individuals of each of the dominant migratory groups (e.g., euphausiids, \textit{Metridia} spp, \textit{Calanus}/\textit{Neocalanus} spp., and hyperiid amphipods) will be gently selected using a wide-bore pipette or small ladle. Effort is also made to obtain more rare migratory organisms (gelatinous species, decapods). Individuals are selected to provide a large within-species range of sizes, with at least 10 replicates for each dominant species sampled over the course of each ecosystem state.

Immediately following collection, individuals are placed into custom 20-250 mL chambers (dependent upon organism size) containing 0.2 µm filtered seawater (Saba et al., 2011; Schnetzer and Steinberg, 2002). Each chamber contains an optical sensing spot to allow for semi-continuous measurements of oxygen using a multi-channel FireSting optical oxygen meter (PyroScience, Aachen Germany; Maas et al., 2016). Chambers are maintained in the dark in onboard incubators at \textit{in situ} mesopelagic temperature. Chambers without animals serve as controls.

Separate incubations will also be set up to determine the contribution of bacterial respiration associated with any egested fecal pellets. Incubations are monitored semi-continuously, with measurements of oxygen concentration taken at 2 h intervals for a total of 12 h. At the end of the incubation individuals are removed and frozen in liquid N. Upon return to land these individuals are imaged for biovolume cross-calibration, dried, and weighed on a Mettler Toledo Microbalance. A sample of incubation water will then be collected for DOC and analyzed post-cruise using a Shimadzu TOC analyzer 5000A (Peltzer et al., 1996; Sharp et al., 2002). Both individual and weight-specific respiration, excretion, and egestion are calculated for each dominant migratory species. Regressions for each species, along with 95% confidence intervals are reported for allometric scaling results. Species counts from net tows are used to scale up individual rates to community respiration, excretion, and egestion, and active transport for dominant species (Steinberg et al., 2000).

Other contributing protocols: MOCNESS abundance and biomass sampling to scale individual respiration and excreta experiments to community export.

Uncertainties and quality control concerns: Physiological effect of net capture and incubation; circadian patterns in physiology; effect of pressure (which is not controlled for). When scaling – the effects of zooplankton patchiness and variation in species physiology.
Data products originating with this method:

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<th>Parameter (DOC flux vertical strata)</th>
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<tbody>
<tr>
<td>active</td>
<td>mg-C m(^{-2}) d(^{-1})</td>
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</table>

Each to be reported separately for each dominant species

SeaBASS submission fields and units:

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Key method references


vertically migrating zooplankton in the Sargasso Sea. Marine Ecology Progress Series 234, 71-84.

Zooplankton vertical migration and the active transport of dissolved organic and inorganic
Method: Zooplankton respiration, excretion, and egestion as a function of size and temperature

Document author and contact info: Amy Maas, Amy.Maas@bios.edu

Brief description of protocol and relation to export pathways:
Respiration of diel vertical migratory (DVM) zooplankton at depth is one of the five export pathways, while excretion of DOC and egestion of POC as fecal pellets at depth may also contribute to “active flux”.

The biomass and abundance measurements from the MOCNESS and UVP sampling are used to calculate downward active flux of respiratory CO$_2$ by migrant zooplankton (mg C m$^{-2}$ d$^{-1}$) as by Al-Mutairi & Landry (2001) using respiration rates calculated from published empirical allometric relationships based on zooplankton taxon, weight, and temperature (Ikeda, 1985; Ikeda, 2014). Migrants are assumed to reside below the EZ 12 h during the day, with equal time spent in the surface waters at night, and the average temperature experienced by migrants at depth during the day for each tow is applied (Dam et al. 1995, Steinberg et al. 2000, 2008a or b, 2012, Al-Mutairi & Landry 2001, Hannides et al. 2009).

Downward active flux of DOC excreted, and POC egested as fecal pellets, by migrant zooplankton (mg C m$^{-2}$ d$^{-1}$) are each calculated as 31% of downward active flux of CO$_2$, based on experimental results from previous studies (Steinberg et al. 2000, Schnetzer & Steinberg 2002, Goldthwait & Steinberg 2008). Weight-specific respiration, excretion, and egestion rates are scaled up (for each size fraction) to the entire migrating biomass (night minus day biomass in EZ or mixed layer).

Other contributing protocols: MOCNESS abundance and biomass sampling and environmental parameters. UVP sampling of the gelatinous community. Ground truth of respiratory demand, DOC and POC production from experiments.

Uncertainties and quality control concerns: Application of scaling factors assumes that biomass and temperature are the main constraints on respiratory and excreta rate, if there are species specific differences or variations in physiology due to some other factor (food availability, oxygen, circadian rhythm, etc), they will not be accounted for. The factors used in allometric equations to calculate metabolic rates (temperature, taxonomic group, size and depth habitat) characterize ~93% of the variance in respiration data (Ikeda, 2014).

Data products originating with this method:

<table>
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<tr>
<td>zooplankton active POC flux</td>
<td>mg-C m$^{-2}$ d$^{-1}$</td>
</tr>
<tr>
<td>zooplankton active DOC flux</td>
<td>mg-C m$^{-2}$ d$^{-1}$</td>
</tr>
<tr>
<td>zooplankton active respiratory flux</td>
<td>mg-C m$^{-2}$ d$^{-1}$</td>
</tr>
</tbody>
</table>

Key method references
Measurement and instrument: Analysis of zooplankton particles from MOCNESS net sampling with ZooScan and associated EcoTaxa Pipeline

A brief description of the method: A sample split from each net of all MOCNESS tows was preserved in buffered formaldehyde and an aliquot from each was imaged with a ZooSCAN optical imaging system (Hydroptic) using two size classes (> 2 mm, < 2 mm) following established procedures (Gorsky et al. 2010, Bachiller et al. 2012, Vandromme et al. 2012, Lebourges-Dhaussy et al. 2014). Briefly the sample was passed through a sieve to create the two size fractions and then split to attain ~1000-1500 particles. The subsample was poured into the ZooSCAN then particles were manually separated. An image was then taken at 4800 dpi. All images were processed using the pipeline described below.

Data processing: Image analysis of at least 1000 particles per MOCNESS sample was conducted via the widely used ZooProcess and EcoTaxa pipeline. The software automatically separates particles and measures their major dimensions. They images were then loaded into a database where machine learning algorithms compare the images to other validated zooplankton images, assigning a taxonomic group to the image. This was validated manually. Counts were then be obtained for major taxon (copepods, euphausiids, doliolids, ostracods, etc) and biovolumes calculated using taxon specific equations and measurements. Biovolumes from the ZooSCAN were calculated by dividing the biovolumes per group by the split and volume filtered through the net for the MOCNESS.

Data products originating from the method: Images of individual zooplankton; estimates of taxonomic composition of imaged zooplankton, abundance and biomass (biovolume) by size and taxon.

Key method references:
Vandromme, P., L. Stemmann, C. Garcia-Comas, L. Berline, X. Sun, and G. Gorsky. 2012. Assessing biases in computing size spectra of automatically classified zooplankton from
imaging systems: A case study with the ZooScan integrated system. Methods in Oceanography 1–2:3-21.
**Method:** Particulate inorganic carbon (PIC) by coulometry

**Document author and contact info:** Antonio Mannino (antonio.mannino@nasa.gov)

**Brief description of protocol:**
Large volumes of seawater (4 to 10 L) are filtered onto plastic (polypropylene or polycarbonate) 47 mm diameter filters in duplicate (daily triplicates), which do not adsorb appreciable amounts of salts. Residual salts are removed from the filter by rinsing with a borate buffer. Filter sample blanks should also be collected daily by filtering a similar volume of pre-filtered (<0.2 μm) seawater as the samples. Filters are packaged individually in foil packets and stored frozen (-20°C should suffice; -80°C even better). The coulometer is set up according to manufacturer recommendations. Samples should be permitted to reach room temperature prior to analysis. A standard calibration curve of calcium carbonate should be performed that spans the full mass range of expected PIC on the filters. The filter or carbonate standard is inserted into the bottom of a glass analysis flask and connected to the digestion system on the Coulometer and sealed. Phosphoric acid is introduced into sample flask and allowed to digest the sample until no additional CO₂ is measurable inside the Coulometer cell. The inorganic carbon content from each filter analyzed is computed based on the calcium carbonate calibration curve. Finally, the seawater filter blank value is subtracted from total PIC measured on each sample filter. Instrument performance is validated daily with the DIC reference material available from the Dickson lab at Scripps.

**Measured Parameter and Units:**
Particulate Inorganic Carbon – mg m⁻³ or micromoles kg⁻¹

**Uncertainties and quality control concerns:**
Uncertainty in PIC stem from:
1. Seawater filter sample blanks with measurable carbon content; blanks with <0.5 μg C are achievable.
2. Uncertainties in the calcium carbonate calibration curve, including the purity of the standard, weighing of the standard, and performance of the Coulometer.
3. DIC reference material correction factor.
4. Accuracy in volume filtered and other filtration handling issues.
5. Low signal due to low concentration of PIC in the absence of high abundances of coccolithophores and other calcium carbonate bearing plankton.

**Key method references**
Method: 18S rDNA amplicon analysis of eukaryotic plankton

Document author and contact info: Adrian Marchetti, amarchet@email.unc.edu

Brief description of protocol and relation to export pathways: Duplicate seawater samples (4L) are collected in low density polyethylene (LDPE) cubitainers and immediately filtered onto Millipore Supor filters (0.8 μm pore size, 47 mm). Filters are immediately flash frozen in liquid nitrogen and stored at -80 °C. Onshore, DNA is extracted from individual cut-up filters using the Qiagen DNeasy Plant Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s protocols with an initial bead beating step. Genomic DNA of *Thermus Thermophilus*, which is not expected to be present in the water samples, is added as internal standards to quantitatively characterize community composition. The amount of internal standards is determined with qPCR to have internal standards only comprise around 1% of the total sequenced reads. The V4 hypervariable region is amplified with barcoded custom 18S V4 primers F (5′-CCAGCASCYGCGGTAATTCC-3′) and R (5′-ACTTTCGTTCTTGAT-3′) (Wang *et al.*, 2018). PCR products are quality checked on a gel and pooled to have similar amounts of DNA from each sample. The pooled library is submitted for sequencing on an Illumina Miseq platform (San Diego, CA, USA). DNA sequences are analyzed in QIIME 2 to profile eukaryotic plankton community composition (Caporaso *et al.*, 2010).

Other contributing protocols: Metagenomics analysis of surface water eukaryotic phytoplankton (Marchetti).

Uncertainties and quality control concerns: 18S rRNA gene copy variation, PCR bias and sequencing errors.

Data products originating with this method:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>OTU abundance</td>
<td>Counts</td>
</tr>
<tr>
<td>DNA sequences</td>
<td>fastq</td>
</tr>
</tbody>
</table>

Key method references:


**Method:** mRNA sequencing of poly-A selected (eukaryotic) genes

**Document author and contact info:** Adrian Marchetti, amarchet@email.unc.edu; Weida gong (wdgong@live.unc.edu)

**Brief description of protocol and relation to export pathways:** Triplicate seawater samples (10L) will be collected in LDPE cubitainers and immediately filtered onto Millipore Supor filters (0.45 μm pore size, 142 mm) by way of a peristaltic pump. Filter times will be limited to a maximum of 30 minutes and conducted under dim light. Filters will be immediately flash frozen in liquid nitrogen and then stored at -80 °C. In the laboratory, RNA will be extracted from individual thawed filters using TRIzol reagent (Invitrogen, Carlsbad, CA, USA) according to manufacturer protocols with an initial bead beating process. Sequence library preparation and sequencing will be performed at the UNC High Throughput Sequencing Facility in Chapel Hill using standard Illumina library prep protocols (TruSeq Stranded Total RNA kit, San Diego, CA, USA) and the Illumina HiSeq 4000 Platform. Sequences will be assembled using Trinity (v 2.4.0) and annotated with tBLASTx (v 2.6.0) against MarineRefII and Kyoto Encyclopedia of Genes and Genomes (KEGG) for taxonomic and functional information. Sequence abundance will be normalized with R package DESeq2 to assess expression levels of genes. Genes that have significant relationships with rate processes and other physiological parameters will be identified and used to infer the physiological status of both autotrophic and heterotrophic protists. See Marchetti et al., (2012) for an overview of the metatranscriptomic approach and Gong et al., (2016) for more details on RNA extraction, library preparation and bioinformatics pipeline.

**Other contributing protocols:** RNA extraction, cDNA synthesis.

**Uncertainties and quality control concerns:** RNA extraction efficiency and library preparation quality.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transcript abundance</td>
<td>Counts per Million (CPM)</td>
</tr>
<tr>
<td>Gene expression</td>
<td>Fold change</td>
</tr>
</tbody>
</table>

**Key method references:**


Method: Experimental chlorophyll - 90% ethanol extraction

Adrian Marchetti (amarchet@email.unc.edu), Weida Gong
University of North Carolina

Triplicate seawater samples (400 ml) were filtered onto polycarbonate filters (via gravity) and GF/F filters (via gentle vacuum) arranged in a series cascade. Filters were then immersed in 6 ml 96% ethanol in scintillation vials for 12 hours before the start of extraction (Graff and Rynearson, 2011). The extracted chl a was quantified using a Turner Designs 10-AU fluorometer using the acidification method (Parsons et al., 1984). Water-column integrated chl a was calculated down to the depth of 1% incident irradiance level.
Method: New production/Gross primary productivity/Net primary productivity

Document author and contact info: Adrian Marchetti, amarchet@email.unc.edu; Weida Gong wdgong@live.unc.edu

Brief description of protocol and relation to export pathways: Seawater samples (1L) are collected prior to dawn in triplicate using a Trace Metal Clean Rosette (TMC) system at five depths corresponding to 65%, 38%, 20%, 10% and 1% of the incident irradiance ($I_0$) and inoculated with pre-determined amounts of NaH$^{13}$CO$_3$ and Na$^{15}$NO$_3$ stable isotopes (i.e., non-radioactive isotopes). From Epoch 2 Day 6 to Epoch 3 Day 2, seawater samples are collected at depths corresponding to 40%, 20%, 10%, 5% and 1% of the incident irradiance to capture particulate maximum at depth of 5% light level. For C, isotope inoculations are performed at concentrations of 180 µM, assuming an approximate DIC concentration of 1800-2000 µM. For N, isotope inoculations are adjusted to achieve ~10% of the in situ NO$_3$ concentrations, as premeasured via a submersible ultraviolet nitrate analyzer (SUNA) located on the Survey ship. Inoculated samples are then incubated within on-deck surface seawater flow-through incubators screened to mimic in situ light levels of the sampled depths. Seawater samples from mixed layer (65%, 38% and 20% of irradiance level) are incubated for 6 hours to estimate mixed layer integrated Gross Primary Productivity (GPP) and another set of seawater samples from all five depth are incubated for 24 hours to provide depth profile of Net Primary Productivity (NPP). After incubation, one sample is filtered directly onto a pre-combusted (450 °C for 4 h) GF/F filter. The other 2 samples are first filtered onto a 5 μm polycarbonate (via gentle vacuum pressure) that contains a collection flask for the filtrate. This filtrate is then filtered onto a pre-combusted GF/F filter (via gentle vacuum pressure). Cells collected on the 5 μm polycarbonate filter are washed onto a pre-combusted GF/F using an 0.2 µm filtered seawater. All GF/F filters are placed into acid-cleaned 30 mm petri dishes, sealed with vinyl tape and frozen at -20 °C until onshore analysis. At each Epoch, a time zero is performed on a single sample at a single depth where the seawater sample is filtered immediately following isotope addition onto a pre-combusted GF/F filter. Onshore, filters are dried at 60 °C overnight and pelletized in tinfoil squares (Elemental Analysis). Mass spectrometry analysis is performed at the UC Davis Stable Isotope Facility. Particulate nitrogen (PN), particulate carbon (PC), atom % $^{15}$N and atom % $^{13}$C are obtained for each sample along with filter blanks. Estimates of $^{13}$C and $^{15}$N incorporation into cells are performed using equations outlined in Slawyk et al., (1977) and Dauchez et al., (1995) and will provide measurements of uptake of DIC, NO$_3$ along with PC and PN concentrations. DIC/NO$_3$ uptake rates are normalized by POC/PON, or Chl $a$ concentrations to obtain biomass-specific uptake rates. To calculate rates, measurements of dissolved NO$_3$ and DIC concentrations are required to be measured on the same samples. DIC uptake rate from 6 hours and 24 hours incubation are depth integrated to estimate depth-integrated GPP and NPP, respectively. New production estimates are calculated from NO$_3$ uptake rates multiplied by the Redfield ratio and normalized to DIC uptake rates as outlined in Aufdenkampe et al. 2002.

Other contributing protocols: mRNA sequencing of poly-A selected (eukaryotic) genes (Marchetti)
Uncertainties and quality control concerns: Amount of $^{13}\text{C}/^{15}\text{N}$ incorporated into organic matter that gets respired and/or excreted from phytoplankton during the 6 hour (short-term) and 24 hour (long-term) incubations.

Data products originating with this method:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrate uptake rate</td>
<td>mol N m$^{-3}$ d$^{-1}$</td>
</tr>
<tr>
<td>f-ratio</td>
<td>no units</td>
</tr>
<tr>
<td>Gross primary production rate</td>
<td>mol C m$^{-3}$ d$^{-1}$</td>
</tr>
<tr>
<td>Net primary production rate</td>
<td>mol C m$^{-3}$ d$^{-1}$</td>
</tr>
</tbody>
</table>

SeaBASS submission fields and units:

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/units=yyyymmdd,m,mol/L/d,mol/L/d,mol/L/d,mol/L/d,mol/L/d,mol/L/d,mol/L/d

Key method references:


Optics Working Group
Protocol Brief: Particle Size Distributions from LISST-DEEP

Primary Author: Andrew McDonnell, amcdonnell@alaska.edu

Brief Description of Protocol: The LISST (Laser In-Situ Scattering Transmissometer) DEEP is an in situ instrument that is equipped with a collimated laser diode and a ring detector that measures intensity of scattering of the laser through the water as a function of scattering angle. The resulting small angle Volume Scattering Function (VSF) is used in a mathematical inversion in order to derive the particle size distribution for particles ranging in size between 1.25 and 250 µm.

Deployment methodologies: The LISST DEEP is a profiling instrument that will be attached to the CTD rosettes of both the EXPORTS Process and Survey Ships. The instrument measures the scattering that results from particles contained within the water as the rosette descends through the water column. Measurements are conducted at a 1 Hz sampling rate and this data is used to construct a vertical profile of the size distribution for every CTD rosette cast conducted. The instrument is powered by the CTD and rated to 3000 m, enabling deep profiles from the surface, through the mesopelagic zone, and into the

Contributing Approaches: (Analysis - approach - potential collaborators)
In situ measurements of the particle size distribution throughout the water column are a core component of EXPORTS and have many potential synergies with other observations. Together, the LISST DEEP and the UVP5 produce information on the in situ particle size distribution across a wide and overlapping range of size classes, enabling the estimation of a unified size distribution estimate. Bottle-collected measurements of the full VSF will also be conducted, and compared to the in situ measurements from the LISST DEEP (which only measures small angle scattering). Size distribution data will be compared to particles collected and analyzed via other methods such as the marine snow catcher (MSC), in situ pumps, bulk sediment traps, polyacrylamide gels. Together, these measurements will be used to derive estimates of particle composition, sinking velocity, and flux as a function of particle size.

Uncertainties and Quality Control concerns:
Filtered sea water is used to regularly measure scattering blanks in order to determine the background scattering, and this will be used to correct the VSF. Each calibration and computational step does involve unavoidable assumptions and uncertainties especially with regard to particles with differing optical properties. Uncertainties also arise at low particle concentrations.

Data products:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optical transmission @ 670 nm</td>
<td>m⁻¹</td>
</tr>
<tr>
<td>Volume Scattering Function</td>
<td>m$^{-1}$ sr$^{-1}$</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>--------------------</td>
</tr>
<tr>
<td>Particle concentration size distribution</td>
<td>$\mu$L L$^{-1}$ µm$^{-1}$</td>
</tr>
</tbody>
</table>

**References:**


Protocol Brief:
Particle and zooplankton characterization from the Underwater Vision Profiler (UVP)

Document authors
Andrew McDonnell, amcdonnell@alaska.edu,
Lee Karp-Boss, lee.karp-boss@maine.edu

Contacts regarding data:
Revelle: Emmanuel Boss, Emmanuel.boss@maine.edu
Lee Karp-Boss, lee.karp-boss@maine.edu
Sally Ride: Andrew McDonnell, amcdonnell@alaska.edu

A brief description of the method: The Underwater Vision Profiler (UVP5) is an underwater camera system that was designed to record the vertical distributions of large aggregates and zooplankton (> 50 µm) down to 6,000 m. Two units that consist of red-light emitting diodes (LEDs) illuminate (in 100 µm flashes) an area of 4 X 20 cm which provides a volume sampling of ~1L per frame. The UVP5-HD was mounted on the bottom of each CTD rosette and collected data on each CTD cast (data are collected during down cast).

Data processing: The UVP5 software acquires and process images in real time. The gain, shutter and LED pulses are controlled and the background image is removed. Images are acquired and processed to get size and grey level for each image. Size information on all detected particles is stored but only Images of particles and plankton larger than 500 µm in equivalent spherical diameter are segmented and saved for later identification. Image post processing and metadata acquisition is accomplished with the Zooprocess software. Tabulated particle data are used to sum the number and volume of particles within predefined size bins, allowing for the computation of the Datasets. Data and images have been uploaded to the Ecotaxa website (http://ecotaxa.obs-vlfr.fr/) which serves as a tool for particle and zooplankton identification with machine learning and human verification, as well as a repository for all globally collected UVP data. Data files for particle and zooplankton abundances

Calibrations: Calibration of the UVP5 are done by the manufacturer and include quantifying the illuminated volume and determining the appropriate conversion between particle area in pixels and mm. The latter is done by measuring the size of particles with a microscope and their corresponding area in the UVP (dropping particles one at a time in the field of view of the camera). The two instruments were sent together to the manufacturer for pre-cruise calibration, where inter-calibration against the same ‘standard’ UVP was done at the same time. Inter-calibration between UVP
instruments and a ‘standard’ is done in situ (Bay of Villefranche, France) and allows for the comparability of all UVP5 data within the global database.

**Uncertainties and Quality Control concerns:**
Uncertainties in concentrations increase with increasing particle size as these particles are less abundant. Small image volumes (~1L) and low abundances of zooplankton often lead to high uncertainties in concentration due to a low number of identifiable zooplankton. Additionally, some zooplankton have avoidance swimming behavior and avoid being sampled. Thus, quantitative measures of zooplankton may not be possible at high resolutions in space, depth and time. Low image resolution only allows taxonomic identification at the rank of subclass or higher.

**Data products:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total particle concentration (includes zooplankton)</td>
<td># m^3</td>
</tr>
<tr>
<td>Total particle concentration size distribution</td>
<td># m^4</td>
</tr>
<tr>
<td>Total particle volume concentration (includes</td>
<td>unitless</td>
</tr>
<tr>
<td>zooplankton)</td>
<td></td>
</tr>
<tr>
<td>Total particle volume concentration size distribution (includes zooplankton)</td>
<td>m^3</td>
</tr>
<tr>
<td>Zooplankton concentration</td>
<td># m^3</td>
</tr>
<tr>
<td>Zooplankton concentration size distribution</td>
<td># m^4</td>
</tr>
<tr>
<td>Zooplankton particle volume concentration</td>
<td>unitless</td>
</tr>
<tr>
<td>Zooplankton volume concentration size distribution</td>
<td>m^3</td>
</tr>
<tr>
<td>Zooplankton concentration by taxonomic group</td>
<td># m^3</td>
</tr>
<tr>
<td>Non-zooplankton particle concentration</td>
<td># m^3</td>
</tr>
<tr>
<td>Non-zooplankton particle concentration size</td>
<td># m^4</td>
</tr>
<tr>
<td>distribution</td>
<td></td>
</tr>
<tr>
<td>Non-zooplankton particle volume concentration</td>
<td>unitless</td>
</tr>
<tr>
<td>Non-zooplankton particle volume concentration size distribution</td>
<td>m^3</td>
</tr>
<tr>
<td>Particle concentration by type</td>
<td># m$^{-3}$</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>----------</td>
</tr>
</tbody>
</table>

**Key References:**


**Document author and contact info:** Susanne Menden-Deuer (URI), smenden@uri.edu

**Measurement and instrument:** Size and abundance measurements of suspended particles with a Beckman Coulter Counter, Multisizer III (version 3.53; Beckman Coulter, Inc, Brea, CA)

**A brief description of the method:** The measurement of particle sizes and concentrations proceeds through the Coulter principle, which quantifies changes in electrical impedance within a sensing zone surrounding a specified aperture (20-500 µm) produced by particles suspended in an electrolyte. The instrument measures only the impedance and does not provide characteristics of particles. As such, particles are not characterized as living or dead. Measurements are made within the 3 to 65% range of the aperture opening, which is typically 100 µm for applications within EXPORTS. Measurements are acquired from discrete samples, from volumes of 1 ml within <30 seconds. Thus, high replication is achievable.

**Data processing:** Size bin and abundance data are stored along with specified standard operating protocol files that help to identify specifics of the run settings. Analysis includes the identification of abundance peaks, their means and standard deviations. The slope of the abundance size spectra is calculated as a means to identify shifts in particle composition.

**Calibration:** The instrument is regularly calibrated, and calibration is verified with manufacturer supplied microbeads.

**Uncertainties and quality control concerns:** Abundance of particles naturally decays with increasing size and statistically reliable quantification of particles >10 µm can be difficult. Particle size is reported as ‘equivalent spherical diameter’. True particle dimensions (e.g. aspect ratios) are not known, as the electrical impedance within the sensing zone is converted to a volume and then to the ESD. High particle concentrations can lead to coincidence within the sensing zone and result in underestimates of total particle counts and/or volume.

**Data products originating from the method:** Particle size distributions, abundances and size abundance spectra of discrete samples (from surface flow through and vertical profiles) in the size range of 3 to 60 µm.

**Key method references:**
1. **Method:**  *Vertical profiles of protistan grazing capacity, including mesopelagic*

2. **Contact Info:** Susanne Menden-Deuer, smenden@uri.edu; Heather McNair, hmcnair@uri.edu

3. **Brief description:** To overcome the significant limitation in our ability to quantify grazing rates below the euphotic zone, we will use fluorescent stains and flow cytometry in a series of incubations to measure rate of herbivory, bacterivory and quantify the abundance of active predators from 50-300 m.

   To overcome the significant limitations in our ability to quantify feeding below the euphotic zone, we will use incubation assays with live stain LysoTracker Green (LTG), activated by the predators feeding vacuole. In addition, we will enumerate bacteria in all samples and offer phytoplankton prey stained with CellTracker and LysoSensor (non-diatom and diatom respectively) in short term incubation experiments aboard the ship to derive vertical profiles of relative feeding frequency and capacity, in parallel to empirical measurements of grazing rates in surface waters. Seawater samples (5L each) are collected from discrete depths down to 500 - 1500 m using a CTD rosette with mounted Niskin bottles. The exact depths will depend on water masses and focus of other EXPORTS investigators. To increase signal to noise ratio, samples are concentrated 10-fold and stained with the live stains. Samples are incubated in the dark in the shipboard walk in incubator set to an appropriate temperature (~4°C). Subsamples of 1-5 ml are removed at discrete intervals for up to 24 hours. Three discrete sample types will be taken for 1) immediate analysis using a flow cytometer, 2) archive samples fixed with paraformaldehyde and glutaraldehyde and 3) an archive sample filtered onto dark, 0.2 µm polycarbonate filters, mounted on microscope slides using immersion oil and frozen. The data acquired will deliver estimates to what degree protists consume biomass in the twilight zone and might reduce remineralization rates through bactivory. Together, these rate estimates provide assessments of the transfer rates of organic matter through the twilight zone across ECC states, which are key to building a predictive and global model of carbon export rates.

4. **Other Contributing Protocols:** Flowcytometry (Guava), vertical profiling, Dilution method, marine snow abundance and characteristics

5. **Uncertainty and quality control:** The development of these methods is relatively recent, so we will quantify uncertainties through replication at every step (triplicate incubation, replicate sample analysis) to yield coefficients of variation.

6. **Data products** will be depth (m), predator abundance (cells L-1), ingestion frequency (%), possibly relative fluorescence (RFU).

6. **SeaBASS submission fields and units:**
   /id_fields_definitions=1id:all_phytoplankton,2id:Synechococcus,3id:heterotrophic_prokaryotes
7. Key Method references


Method: *Plankton population dynamics via deckboard incubations (dilution method)*

Contact Info: Susanne Menden-Deuer, smenden@uri.edu; Heather McNair, hmcnair@uri.edu

**Brief description:** Seawater samples are collected from 1-5 discrete depths corresponding to a range of incident irradiance using a CTD rosette with mounted Niskin bottles. Water is screened through a 200 µm mesh to remove large grazers. Water from each light-depth is partitioned into triplicates of three treatments: whole sea water (<200 µm), diluted, and nutrient amended treatments. Diluted treatments are prepared with whole seawater diluted with 0.2 µm filtered seawater from the corresponding depth. All bottles are incubated for 24 hours in deckboard incubators with screen-manipulated light levels (dark, 10-20%, 40%, 65% and 100%). Light-depths above the thermocline were maintained at surface seawater temperature; light-depths below the thermocline were chilled to the corresponding in situ temperature. Grazing, net phytoplankton growth, and gross phytoplankton growth rate estimates are based on differential changes in chlorophyll a between the diluted and whole seawater treatments. Samples for size fractionated chl (GFF and 5 µm), flow cytometry, and microscopy are collected from each bottle following the 24 h incubation. Chl a is extracted in ethanol and read on a Turner 10AU Fluorometer. Herbivorous protist grazing rate is calculated as the difference in phytoplankton growth rate in the diluted vs undiluted seawater. Gross phytoplankton growth rate is calculated as the sum of the grazing rate and the net phytoplankton growth rate (i.e. the rate of change in chlorophyll a) in the undiluted bottles. The ratio of the grazing vs gross phytoplankton growth rate provides an estimate of primary production consumed and allows predation of a phytoplankton biomass accumulation rate. Flow cytometry and microscopy will facilitate a more detailed view of plankton dynamics with group specific growth and grazing rates and estimates of changes in chlorophyll due to photoacclimation. Experimental design will target euphotic zone integrated rates as well as the effect of light on surface rates. Predation rates are crucial to parameterize carbon flows at the base of the marine food web. These data provide key algorithms to relate plankton community composition, growth and mortality to export through all 5 pathways and across ECC states, which will be essential for building a diagnostic modeling framework to predict export of global PP, now and in future climate scenarios.

**Other Contributing Protocols:** Chl a extraction (ethanol), Flowcytometry (Guava), Nutrient measurements, vertical profiling, photoacclimation, microscopy to measure species composition and biomass

**Uncertainty and quality control:** Based on prior analyses, the uncertainty in the Chl a concentration extraction is between 1-10%. Rate estimates of phytoplankton growth and grazer induced mortality are good within +/- 0.1 d⁻¹ (Morison & Menden-Deuer 2017).

**6. Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phytoplankton growth rate (µ)</td>
<td>d⁻¹</td>
</tr>
<tr>
<td>Heterotrophic protist grazing rate (g)</td>
<td>d⁻¹</td>
</tr>
<tr>
<td>Phytoplankton accumulation rate (r)</td>
<td>d⁻¹</td>
</tr>
<tr>
<td>Chlorophyll a (GFF and 5 µm)</td>
<td>µg L⁻¹</td>
</tr>
</tbody>
</table>
SeaBASS submission fields and units
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Key Method references
The method was first described in:

A recent assessment of uncertainty is here:
Measurement and instrument: Characterization of species composition and size distribution of microphytoplankton with a FlowCAM Benchtop B3 Series.

A brief description of the method: The FlowCAM® enumerates and classifies plankton particles through a combination of flow cytometry, microscopy, and image analysis--simultaneously delivering 30 different properties (including various metrics of size) for each particle. These properties can be used in particle identification. The instrument counts and images particles contained in a seawater sample that is drawn through a glass chamber, the flow cell, by means of a syringe pump. The camera images particles passing within the camera’s field of view. Particles contained within each image frame are then extracted and separated into individual images. The analysis can be performed in either of two modes: in auto-image mode the instrument images all particles flowing in front of the camera field of view at a fixed, user-specified flow rate and frame rate; in trigger mode, the instrument only images particles that trigger the camera when they exceed either a scatter or a fluorescence set threshold. The size range of the particles that can be analyzed is determined by the size of the flow cell and the recommended corresponding magnification: the depth of the flow cell sets the upper size limit of particles to be analyzed, while the lower size limit depends on the smallest size that the magnification can resolve and can be specified by the operator. Flow cells with depths of 50, 100, and/or 300 µm are appropriate for plankton analysis, and are used in combination with 20x, 10X and 4X objectives respectively.

Data processing: Size bin, abundance data, and properties values for each particles are stored along side the operating context files. Size is reported as either ‘equivalent spherical diameter’ derived from dimensions of a sphere obtained from a series of ferret measurements around the particle, or it is reported as ‘Area Based Diameter’, i.e. the diameter of a circle of the same number of pixels as the particle. Biovolume calculation assume all particles to be spherical. The software also automatically stores a summary of the run settings and provides summary statistics of each measured property. Classification into taxa is possible after establishing a training set or can be done manually.

Calibration: The instrument can be calibrated using manufacturer supplied microbeads.

Uncertainties and quality control concerns: Abundance of particles naturally decays with increasing size and the analysis of a large number of particles and thus of a large volume of sample is needed to effectively resolve the size spectrum. Standard flow cells are wider than the camera field of view, and thus only a small percentage of the volume aspirated is processed and imaged, and particles flowing outside the field of view are not imaged. In trigger mode however, particles flowing outside the field of view can trigger the camera, which then captures any particle contained in the frame. The software has no way to determine which particle produced the trigger signal. In trigger mode, samples must be dilute enough to allow for only one particle per image.

For quantitative purposes, Field of View flow cells are recommended. Since the size of the flow cell and the magnification used determine the size range of particles to be analyzed, samples may need to be fractionated, which not only requires manipulation of the sample but also switching flow cells and objectives.

Data products originating from the method: Particle images, species composition, particle size distributions, abundances, and size spectra of discrete samples (from surface flow through and vertical profiles) in the size range of 3 to 300 µm.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Particle images</td>
<td>unitless</td>
</tr>
<tr>
<td>Species composition</td>
<td>taxa (unitless), species diversity indices (unitless)</td>
</tr>
<tr>
<td>Particle size distribution</td>
<td>frequency per size bin (numbers/µm), slope of</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>-----------------------------</td>
<td>----------------------------------------------------------------</td>
</tr>
<tr>
<td>Abundance</td>
<td>biomass (µg C L⁻¹) and cells L⁻¹</td>
</tr>
<tr>
<td>Size spectra</td>
<td>same as particle size distribution but for living organisms, as opposed to all particles</td>
</tr>
</tbody>
</table>

**Key method references:**
Method: Particle characterization using a Guava flow cytometer

Document author and contact info: Heather McNair, hmcnair@uri.edu

A brief description of the method: Samples to be characterized using the Guava flow cytometer will be pipetted into 96-well plates (~200 µL). Particles will be characterized based on the recorded forward scatter, side scatter, red fluorescence (695/50 nm) and yellow fluorescence (583/26 nm) after exciting with a blue laser (488 nm). To avoid clogging the instrument, samples will be screened through 40 µm mesh prior to being run. Gain settings on the photomultipliers will be determined using an initial sample and kept constant for all following samples. A range of different size beads will be used to create an equation to convert forward scatter to size. Data will be processed using InCyte flow cytometry software that allows particles in a sample to be subset into different groups.

Calibration: The instrument may not be calibrated by the user, but the use of microbeads may provide reference to size and an internal standard for calculating concentration (Olson, Vaulot, and Chisholm 1985), and biological controls may aide in establishing settings and interpreting results.

Uncertainties and quality control concerns: The resolution, range, and observation values per particle are highly dependent on collection settings and machine specifications. To minimize complications settings will be kept constant throughout the cruise. Detection limits will be further explored using filtered sea water blanks.

Data products originating from the method:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scattering per particle</td>
<td>Relative units</td>
</tr>
<tr>
<td>Fluorescence per particle</td>
<td>Relative units</td>
</tr>
<tr>
<td>Abundance</td>
<td>Relative units</td>
</tr>
<tr>
<td>Scatter statistics per group</td>
<td>Relative units</td>
</tr>
<tr>
<td>Fluorescence statistics per group</td>
<td>Relative units</td>
</tr>
</tbody>
</table>

SeaBASS fields and units:
/id_fields_definitions=1id:nanoeukaryote,2id:picoeukaryote,3id:Synechococcus

/fields=r_event,lon,lat,depth,lightlevel,date_start,time_start,date_end,time_end,abun_1id,abun_1id_sd,a bun_2id,abun_2id_sd,abun_3id,abun_3id_sd,u_ph_1id,u_ph_1id_sd,u_ph_2id,u_ph_2id_sd,u_ph_3id,u_ph_3id_sd,u_ph_3id_se,g_1id_se,g_2id_se,g_3id_se,FSC-H_1id,FSC-H_1id_sd,FSC-H_2id,FSC-H_2id_sd,FSC-H_3id,FSC-H_3id_sd,FL-H_ex488_em695_1id,FL-H_ex488_em695_1id_sd,FL-H_ex488_em695_2id,FL-H_ex488_em695_2id_sd,FL-H_ex488_em695_3id,FL-H_ex488_em695_3id_sd,FL-H_ex488_em583_3id,FL-H_ex488_em583_3id_sd

/units=none,degrees,degrees,m,%/yyyyymmdd,yy/mm/dd,mm:ss,cells/L,cells/L,cells/L,cells/L,L,L,L,L,L,L,L,L,L,L,L,L,L,L,L,L,L,L,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,d^-1,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits,arbunits

Key method references:
**Variable**  | Light Absorption Coefficient Spectrum of CDOM
---|---
**SeaBASS Name**  | ag
**Units**  | 1/m
**Sampling**  | Niskin bottle on rosette

Document author and contact info: Norman Nelson, normannelson@ucsb.edu

Updated 7/21/16

Light absorption by CDOM will be measured from samples collected and prepared according to the UCSB CDOM Laboratory protocol (Nelson et al., 2007). Samples are analyzed in an UltraPath spectrophotometer according to the same method with recent modifications as specified by the draft NASA CDOM Group UltraPath analysis protocol (reference TBA).

Samples are collected from Niskin bottles using silicone tubes by analysts wearing nitrile gloves. The samples are collected into combusted 60ml brown borosilicate glass EPA vials with Teflon lid liners.

To remove particles the samples are filtered through 0.2 micron pore 25mm Nuclepore polycarbonate filters that have been pre-extracted with 60ml ultrapure water (Barnstead Nanopure, 18MΩ-cm, low carbon filter cartridge).

Samples are stored at 4C in the dark until analysis, generally within 6 months to 1 year. 2-year stability of samples stored in this manner has been documented by Swan et al. (2009).

On analysis day sample vials are allowed to equilibrate to room temperature and are analyzed for UV and visible absorbance in an UltraPath single-beam long path (200cm cell) spectrophotometer. Samples are referenced against ultrapure water (Barnstead Nanopure, 18MΩ-cm, low carbon filter cartridge) and are corrected for refractive index effects using concurrently measured apparent absorption spectra of 30 g/l and 40 g/l solutions of sodium chloride (SigmaUltra > 99.97% NaCl), interpolated by the salinity of the sample. Corrected spectra are converted to naperian absorption coefficient by the formula $a = 2.303A/l$ where A is the decadal absorbance of the sample, l is the pathlength (m), and 2.303 converts decadal to natural log scale.

Reference spectra of a water solution of Suwanee River Fulvic Acid Standard 1, IHSS #1S101F (http://www.humicsubstances.org/elements.html) vs ultrapure water at approximately 0.25 mg/l is also measured at the time of sample analysis, and is converted to specific absorption ($m^2/mg$).

All spectra are null corrected at long wavelength (690-710 nm avg), in accordance with the current draft protocol.

Reported data include the CDOM absorption spectrum ($a_g$), the estimated error (based on the average of ~6 scans over 30 s, propagated through the analysis arithmetic), the raw absorbance average and its standard deviation, and the SRFA and NaCl reference spectra.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>CDOM absorption, $a_g$</td>
<td>m$^{-1}$</td>
</tr>
</tbody>
</table>
References:


At-sea operations: The Biospherical C-OPS is a light weight, hand deployed package to measure the
downwelled irradiance and upwelled radiance sunlight field (Morrow et al 2010). Nineteen discrete
wavelengths in the UV, visible, and near IR are measured and there is a surface mounted deck irradiance
sensor with matching wavelengths. The package also has pressure, tilt and roll, water temperature. The
package weighs about 30 lbs in air and just a few lbs in water. The package is attached by an electrical
cable that is held on a cable bucket (35 gal poly trash can) on the fantail. The instrument is allowed to
free-fall to approximately 100m, which takes about 6 minutes. The instrument is then hand hauled back to
the surface by 1 or 2 persons and the cable is spooled back into the cable. Between casts, the profiler is
kept in the lab to prevent overheating of the sensors.

Constraints: The following parameters control the quality of the light profiles (Mueller et al 2003):
1) Ship shadow. Our deployment procedure attempts to keep the profiler away from the ship’s
underwater shadow.
2) Ship bearing. If the sun direction is anywhere aft of amidships, we should have no problems.
Worst bearing is with the sun directly ahead.
3) Ship movement through the water. We have 300m of cable that we spool out as the instrument
sinks, so if the ship is also moving through the water, much of the cable is laid out horizontally and
we don’t reach our desired depth before reaching the end of our cable. Ideal forward speed is a
one half knot.

Before the cast: Controlling computer is started up, and the Biospherical software is used to collect dark
current data on both profiler and surface reference instruments.

Start of cast: We will require radio or squawk-box comms between the person deploying the light
profiler, the bridge, and the computer operator in the lab. A minimum of 2 persons on deck, with one
primarily for communications since the deployer will have both hands occupied. There is no need for the
ship to return to the Station location for the light cast (we’re close enough.)

We attempt to get the profiler about 20m behind the ship before we start the free-fall. Getting the
probe clear of the stern is the most demanding part of the cast. Often a 30 sec mild bump ahead
with the main will both move the ship forward and sweep the instrument aft with the prop wash.

Once the light profiler is about 20m aft and at the surface, the instrument is allowed to free-fall to the
desired depth. During the free-fall, the cable is kept slack by paying out the wire just fast enough to keep
the wire smoothly pulling over the stern railing near the port corner. The angle that the cable enters the
water is constantly observed to make sure it continues to lead aft and away from the stern.

As the light profiler nears the target depth, the computer operator calls out every 10 m pressure mark. At
the target depth, or when the maximum safe amount of cable to let out is reached, cable deployment is
stopped and retrieval begun.

Data processing: We use standard ocean color radiometry protocols (Mueller et al 2003) carried out through
a Matlab implementation of the BBOP data processing system (Siegel et al. 2005). Tilt criteria and the depth
of extrapolation for determination of surface properties are determined for each cast depending on the
conditions.
References:


This README file details the CTD processing for the EXPORTS 2018 North Pacific campaign. Four CTDs were deployed on the campaign that we present here, all SeaBird 911+ units. One was on the rosette sampler on the R/V Sally Ride (SR 1812, or "survey" in the SeaBASS files), one on the main rosette on the R/V Roger Revelle (RR 1813 SIO, "process"), one on the UHawaii trace metal rosette (RR 1813 TM, "TM") and one on the MOCNESS system (RR 1813 MOCNESS, "MOCNESS"). Other CTDs present on the IOP packages, the wirewalker, etc. are not covered in this data release. This release should be still considered preliminary, as we refine post-cruise corrections to the data new releases will be made.

Notes on SeaBASS preparation for this release (20191002)

There are two SeaBASS files for every cast. One is the "unbinned" processed data (see below for processing pipeline) which features minimal filtering, and flagging by loopedit and wildedit for data quality. One is the 1 m binned downcast only data. The SeaBASS files do not contain the 'secondary' T and S sensor data.

Flags and Bins

We ran the SeaBird programs WILDEDIT and LOOPEDIT on the unbinned data files to identify outliers and data contaminated by ship heave. LOOPEDIT identifies samples collected on upward heaves during a cast and increments the flag column. WILDEDIT uses a two pass process to identify data points lying 2 or 20 standard deviations outside the mean, respectively. Data scans flagged in this way are not used to compute bins, but the data in the "Unbinned" data files are not altered by flagging.

SeaBird's software allows the recovery of the numbers of samples used in each bin, but no statistics. There is a "bincount" column in each SeaBASS file which gives the number of valid samples used to compute the binned value.

SeaBird's binning algorithm is as follows (as applied using depth)

\[
X_i = \frac{(X_c - X_p) \cdot (z_i - z_p)}{z_c - z_p} + X_p
\]  

(1)

where \(X_i\) is the value variable \(x\) at depth \(i\), \(z_p\) is the average depth of the previous bin, \(z_c\) is the average pressure of bin i, \(X_c\) is the average value of the X data in bin i, \(z_i\) is the center value for depth in bin i, and \(X_p\) is the average value of variable X in the previous bin (SeaBird Scientific University, Module 13, Advanced Data Processing).
Calibration Data

Configuration and calibration data for each sensor package / cast are included in the .XMLCON files that correspond to each cast. This XML format file contains calibration coefficients and dates for each sensor, as deployed. It is important to examine these files because of configuration problems with the SIO rosette CTD package in the first week of the cruise caused significant changes in the configuration.

Post-Cruise Corrections

This release (R1) of the SeaBASS files contains corrections to selected variables as computed by the members of the EXPORTS Synthesis Working Group 4 (lead authors A. Thompson (T&S), D. Nicholson (O₂), chlorophyll fluorescence (M. Feen) and optical backscatter (X. Zhang). These preliminary reports are attached to this file. What follows here is a brief summary and the reports include more details.

Temperature and salinity corrections were found by using the Sally Ride primary C & T sensors as the "gold standard." No corrections were deemed necessary to apply between the Ride and the Revelle sensors, so no changes were made. Rough corrections to the process cruise TM and MOCNESS sensors were made by comparison to adjacent Revelle CTD rosette cast data by N. Nelson. These corrections may be revised in the future.

Winkler titration measurements of dissolved oxygen collected on the survey cruise were used to correct the Ride and Revelle CTD oxygen sensors, and corrections were applied to both for the SeaBASS release.

In situ stimulated chlorophyll a fluorescence measurements were calibrated to in situ chlorophyll using total chlorophyll a as determined by the GSFC HPLC lab, from bottle samples on both survey and process ships during night-time (PAR values < 20 umol/m²/s) casts only (to avoid nonphotochemical quenching of fluorescence during the day). M. Feen used the raw voltage signals from the fluorometers compared to the discrete HPLC measurements to arrive at a distinct computation for each of the main rosette sensors. N. Nelson used the Revelle values to prepare a rough correction of the fluorometers on the MOCNESS and TM packages.

At present we are using factory calibrations for the backscatter sensors on the rosette packages. X. Zhang evaluated the sensor on the survey ship CTD and determined the 05/18 calibration (not the most recent) was valid.

Post-cruise correction of beam transmissometer data is planned for Release 2, once POC and transmissometer data from lowered packages are available.

Original processing notes from version 20190306 / 20190916
The 20190916 version of the data should be identical to the 20190306 version of the data, which was apparently deleted from the Google Drive by a user who was not aware of their actions.

We used identical processing pipelines for each data file from the different instruments. All data were processed from raw (.HEX) format files (and .XMLCON configuration and .BL bottle trip data files) using SeaBird Data Processing version 7.26.7 on a Windows 10 system at UCSB following the cruise. This data release includes all the raw data, Program Setup (.psa) files, batch processing configuration scripts, and batch processing meta-scripts that can be used to reprocess the entire data set quickly. Tau corrections were used for oxygen sensors.

Within each instrument directory there is one subdirectory for each final product, as well as subdirectories for the raw data (including the .xmlcon and .bl files) and for the .psa files. A scratch directory is present but no relevant files are there. The instrument directory includes the SeaBird and Windows batch files used to process the data.

The file formats for each product are SeaBird .cnv or .btl ascii files. I've provided a simple loadcnv.m script to load the .cnv files and selected metadata into memory in MATLAB.

There is also a directory containing WHPO exchange format (ascii .csv) files with the bottle data, and the scripts used to create them from the .btl files. The loadwhpobot.m script will load these files into memory in MATLAB. These files do not have the statistics (min/max/nscans) that are included with the .btl files.

Further information on sensor serial numbers and calibrations and so forth is available in the xml sections of the data files.

**Important notes on this release (20190306):**

Post-cruise instrument corrections are **not** applied to the data found in the EXPORTS L0 CTD DATA folders on the Google Drive. Post cruise instrument corrections **ARE** applied to the SeaBASS files.

**Fixes in this release (20190306):**

MOCNESS and RR SIO CTD data streams had the incorrect raw voltage out channels. Minor changes to the filter settings and wildedit settings to ensure consistency between all instruments.

The WHPO Exchange Format bottle files are not included in this release (broader fixes are necessary and they will be added at a later date)

**Processing pipeline schematic:**
DatCnv.exe
RawData --------------------> RawConverted
(.hex, .xmlcon, .bl)   (.cnv, .ros)
AlignCTD.exe
RawConverted ---------------> ScratchData
(.cnv)                        (.cnv)
CellTM.exe
ScratchData --------------------> UnfilteredData
(.cnv)                        (.cnv)
Filter.exe
UnfilteredData --------------> ScratchData
(.cnv)                        (.cnv)
WildEdit.exe
ScratchData --------------------> ScratchData
(.cnv)                        (.cnv)
LoopEdit.exe
ScratchData --------------------> UnbinnedData
(.cnv)                        (.cnv)
BinAvg.exe
UnbinnedData --------------> Bin1mData
(.cnv)                        (.bin)
BinAvg.exe
RawConverted --------------> Bin2HzData
(.cnv)                        (.bin)
BottleSum.exe
RawConverted --------------> BottleData
(.ros)                        (.btl)

Explanation of the processing pipeline by step:

DatCnv.exe takes raw data from the .hex files, configuration data from the .xmlcon files, and bottle trip data from the .bl files in the RawData and creates an ascii raw data file (.cnv) and rosette summary (.ros) file in the RawConverted directory.

AlignCTD.exe synchronizes the pressure, conductivity, and temperature data in the RawConverted file to account for the different position of the temperature+pressure and conductivity sensors in the package. The aligned file is placed in ScratchData.

CellTM.exe corrects the conductivity data for cell thermal mass effects in the file in ScratchData and places the corrected file in UnfilteredData. This is a product. Standard SeaBird settings for the 911 sensor package were used.

Filter.exe applies low-pass filters to the pressure and conductivity sensors. We used the default SeaBird settings for low-pass filtration. UnfilteredData .cnv files back to ScratchData.
WildEdit.exe looks for outliers in the data using a $2\sigma$ criterion (first pass) and a $20\sigma$ criterion (second pass). Note this does not remove the data from the file but it increments the flag column in the data. ScratchData .cnv files back to ScratchData.

LoopEdit.exe attempts to flag 'loop' features in the profile caused by ship roll and gradients. This also increments the flag column in the data. We used the SeaBird default settings. ScratchData .cnv files were edited and saved to UnbinnedData. This is a product.

BinAvg.exe was used to make 1 m downcast only files from the unbinned data .cnv files saved as .bin (still ascii files in cnv format) in Bin1mData. This is a product. BinAvg.exe was used to make 2Hz files from the (complete cast) unbinned data .cnv files saved as .bin (still ascii files in cnv format) in Bin2HzData. This is a product.

BottleSum.exe was used to make bottle files (in SeaBird ascii .btl format) from the .ros files in RawConverted and the .xmlcon files in RawData. Bottle files were saved in the BottleFiles subdirectory. This is a product. We included statistics (min/max/number of scans averaged) as well as the averages.

Directory structure of this release:
https://drive.google.com/open?id=1j02wGDwVYBe0H3dDhXVKqgh0jnLPv-8H

```text
EXPORTS L0 CTD DATA
loadcnv.m
20181023 (obsolescent)
20190116 (obsolescent)
20190306 (this edition) (== 20190916)
SR 1812 20190306
Bin1mData
UnfilteredData
PSA
UnbinnedData
BottleFiles
RawConverted
ScratchData
Bin2HzData
RawData
exports_batch_v3.txt
exports_batch_batch.bat
SeaBASS
RR1813 MOCNESS CTD 20190306
Bin1mData
UnfilteredData
PSA
UnbinnedData
```
BottleFiles
RawConverted
ScratchData
Bin2HzData
RawData
exports_batch_MOCNESS.txt
exports_batch_batch_MOCNESS.bat
SeaBASS
RR1813 TM CTD 20190306
Bin1mData
UnfilteredData
PSA
UnbinnedData
BottleFiles
RawConverted
ScratchData
Bin2HzData
RawData
exports_batch_batchTM.bat
exports_batchRR1813TM10.txt
exports_batchRR1813TM.txt
SeaBASS
RR1813 SIO CTD 20190306
Bin1mData
UnfilteredData
PSA
UnbinnedData
BottleFiles
RawConverted
ScratchData
Bin2HzData
RawData
exports_batch_batchRR1813.bat
exports_batch_RR1813S.txt
exports_batch_RR1813.txt
SeaBASS

Key to the voltage channels in each data set

RR1813 MOCNESS
V8: WETLabs C-star
V4: Fluorometer

RR1813 SIO
Casts 1-4, 9-10
V0: WETLabs ECO NTU
V1: Seatech/WETLabs FLF Fluorometer (Chl)
V4: LISST
V6: WETLabs C-star transmissometer
Casts 5-8, 11-84
V0: WETLabs ECO Fluorometer (chl)
V1: WETLabs ECO NTU
V4: LISST
V6L WETLabs C-star transmissometer

RR1813 TM
V2: WETLabs WETStar Fluorometer (chl)

SR1812 CTD
V2: WETLabs ECO Fluorometer (chl)
V3: WETLabs ECO BB
V4: WETLabs C-star transmissometer
V5: PAR

EXPORTS Synthesis WG 4 Preliminary Instrument Intercomparison Reports

Chlorophyll Fluorescence Calibration_10_7_19_Final_.pdf
EXPORTS2018_02_Cal_Oct8.pdf
inter-calibration of bb sensors_Oct8.pdf
Method: HPLC Pigment analysis

Document authors and contact info: Norm Nelson, norm.nelson@ucsb.edu; Collin Roesler, croesler@bowdoin.edu; Ivona Cetinic, ivona.cetinic@nasa.gov; Sasha Kramer, sasha.kramer@lifesci.ucsb.edu

Brief description of protocol and relation to export pathways: Algal pigments will be analyzed with High Pressure Liquid Chromatography on samples obtained from both the survey and process vessels. Water samples will be collected with a CTD-Niskin bottle rosette at approximately 8 depths, 3-4 times a day. In order to ensure compatibility between pigment, carbon and optical samples, Niskin bottles from each depth will be collected into a large volume carboy using a funnel with ½ inch tubing. Carboys will be kept cold and dark until subsampling. Subsamples for each analysis will be collected from the carboy after gentle mixing via the Perry method; swirling three times in one direction, three time in the reverse direction, three times in the first direction. Additional samples will be collected from the inline flow-through system once or twice a day, at the intake depth of approximately 3m. The inline samples on the survey vessel will be size fractionated with 3 µm and 20 µm cartridge filters (Pall®) in addition to the unfiltered samples. Additional pigment samples will be collected during the “calibration profiles” that will be conducted either during pick up, deployment or encounter with the autonomous assets.

Sample particulates will be collected via low pressure (less than 5 mm Hg) vacuum filtration on a pre-combusted (450C for 4 hours) 25 mm 0.7 μm Whatman® GF/F filter. Exact volume and volume uncertainty are recorded for each sample. Filters will be folded in half, sample side inward, transferred into labeled aluminum foil packages, and stored in liquid nitrogen from the time of collection until the time of delivery to NASA GSFC. Analysis will be conducted at Ocean Ecology Lab at NASA GSFC following the methods described in SIMBIOS intercalibration exercise, and in the SeaWIFS HPLC round-robins.

Other contributing protocols: None

Uncertainties and quality control concerns: Duplicate samples will be collected at a rate of approximately one every ten samples (i.e., once every other CTD cast or every other day for the inline system). GSFC’s sample processing methods include an assessment of the duplicate samples, sample precision by reinjection of samples, and an effective limit of quantitation (lower detection limit) which are all provided with the data.

Data products originating with this method:

<table>
<thead>
<tr>
<th>Pigment Name</th>
<th>Parameter Symbol</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alloxanthin</td>
<td>allo</td>
<td>mg m⁻³</td>
</tr>
<tr>
<td>Alpha beta carotene</td>
<td>alpha_beta_car</td>
<td>mg m⁻³</td>
</tr>
<tr>
<td>Name</td>
<td>Abbreviation</td>
<td>Unit</td>
</tr>
<tr>
<td>---------------------------------------------</td>
<td>--------------</td>
<td>------------</td>
</tr>
<tr>
<td>19’-Butanoyloxyfucoxanthin</td>
<td>but_fuco</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Chlorophyll c3</td>
<td>chl_c3</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Chlorophyllide a</td>
<td>chlide_a</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Diadinoxanthin</td>
<td>diadino</td>
<td>mg m$^{-3}$</td>
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<tr>
<td>Diatoxanthin</td>
<td>diato</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Divinyl Chlorophyll a</td>
<td>dv_chl_a</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Divinyl Chlorophyll b</td>
<td>dv_chl_b</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Gyroxanthin diester</td>
<td>gyro</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Fucoxanthin</td>
<td>fuco</td>
<td>mg m$^{-3}$</td>
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<tr>
<td>19’-Hexanoyloxyfucoxanthin</td>
<td>hex_fuco</td>
<td>mg m$^{-3}$</td>
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<tr>
<td>Lutein</td>
<td>lut</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Monovinyl chl-a</td>
<td>mv_chl_a</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Monovinyl chl b</td>
<td>mv_chl_b</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Neoxanthin</td>
<td>neo</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Peridinin</td>
<td>perid</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Pheophorbide a</td>
<td>phide_a</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Pheophytin a</td>
<td>phytin_a</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Prasinoxanthin</td>
<td>pras</td>
<td>mg m$^{-3}$</td>
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<tr>
<td>Total Chlorophyll a</td>
<td>tot_chl_a</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Total Chlorophyll b</td>
<td>tot_chl_b</td>
<td>mg m$^{-3}$</td>
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<tr>
<td>Total Chlorophyll c</td>
<td>tot_chl_c</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Violaxanthin</td>
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<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Zeaxanthin</td>
<td>zea</td>
<td>mg m$^{-3}$</td>
</tr>
</tbody>
</table>

**Key method references:**
SIMBIOS intercalibration exercise and the SeaWIFS HPLC round-robs.
Inorganic nutrients are analyzed using flow injection analysis, from rosette samples collected and frozen at sea.

Samples will be collected in clean, pre-rinsed plastic HDPE 20ml scintillation vials. Samples being analyzed for Si(OH)$_4$ must be collected in plastic HDPE vials. For all other analytes, glass or plastic HDPE containers are acceptable. Some adsorption of PO$_4$ during storage in plastic containers has been noted, so glass may be preferred if PO$_4$ is to be determined.

Caps should have a plastic liner. All foil lined lids should be avoided.

Storage blanks are required if NH$_4$ is to be measured. Storage blanks are recommended for all other analytes to detect possible contamination.

A minimum of 5mL of sample is required for the determination of a single nutrient species. 15-17mL of sample is necessary for simultaneous determination of all analytes. It is entirely unnecessary to submit more than 20mL of sample. If you must submit more than 20mL of sample or have other volume concerns, please contact the Analytical Lab.

Mark all samples clearly and directly on the sample container, as well as the lid. If you plan on shipping samples, any masking tape or colored lab-style tape placed on sample vials will fall off during shipment if in the presence of dry ice. **Do not use these types of tape to label samples if you plan on shipping samples on dry ice.**

Samples should be frozen immediately after collection and stored at -20°C or below. Please allow sufficient headspace when freezing for expansion of the sample without breaking the container.

Do not acidify samples for nutrient analysis.

Please ship samples overnight express using insulated shipping containers and dry ice to keep the samples frozen during transport.

**Flow Injection Analysis – Nutrients**

- **Manufacturer:** Lachat Instruments Div., Zellweger Analytics, Inc.
- **Model:** QuikChem 8500
- **Capabilities**
Simultaneous determination of up to 5 analytes per sample
Random-access autosampler with racks for up to 120 samples per batch
Analytical manifolds available:
  - Nitrite
  - Nitrate plus Nitrite
  - ortho-Phosphate
  - Silicic Acid
  - Ammonium

**General Description**

Flow injection analysis (FIA) is a continuous-flow technique for automated wet-chemical analysis. The methodology used by the flow injection analyzer is similar to that used by AutoAnalyzers, with continuously flowing reagent streams, reaction 'manifolds', and flow-through detectors. However, FIA does not use air-bubble segmentation to separate samples and promote mixing. Instead, small diameter tubing is used in the manifolds, resulting in laminar flow conditions in which mixing takes place by axial and radial diffusion, and the manifolds are self-cleaning. Diffusion (dilution and mixing) is controlled by manifold design. A major practical advantage of this technology over that of air-segmentation, is that analytical results are usually available within a minute or so from the time the sample is aspirated, so any problems in the system can be spotted quickly and corrected, with little wasted time. The overall analysis times also tend to be shorter with FIA, so more samples can typically be analyzed in a given period of time. Precision and detection limits are generally comparable between the two technologies.

The software that controls our instrument is Windows 7 based, and displays real-time graphical output of any or all of the active detectors. This is extremely useful for during-run monitoring of the instrument's performance. Data is stored on disk, and is available for post-run processing in a variety of formats. The software also provides a wide range of quality control options to ensure accurate and reliable results. These include check-standards, control samples, same-vial or different-vial replicates, and spikes.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Concentration Range, μM</th>
<th>Precision</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrate (NO₃) + Nitrite (NO₂)</td>
<td>0.2 - 300</td>
<td>+/-5%</td>
</tr>
<tr>
<td>Nitrite (NO₂)</td>
<td>0.1 - 100</td>
<td>+/-5%</td>
</tr>
<tr>
<td>Phosphate (PO₄)</td>
<td>0.1 - 200</td>
<td>+/-5%</td>
</tr>
<tr>
<td>Silicic Acid (Si(OH)₄)</td>
<td>1.0 - 600</td>
<td>+/-5%</td>
</tr>
<tr>
<td>Ammonium (NH₄)</td>
<td>0.1 - 200</td>
<td>+/-5%</td>
</tr>
</tbody>
</table>
**Method:** Net community production from mass balance

**Document author and contact info:** David Nicholson, dnicholson@whoi.edu

**Brief description of protocol and relation to export pathways:** Mass balance of dissolved oxygen (O$_2$), dissolved inorganic nitrogen (DIN) and total organic carbon (C$_{org}$) in the euphotic will be used to solve for net community production (NCP), which is the whole ecosystem balance between photosynthesis and respiration and provides an important constraint representing the sum of all export pathways. Within a euphotic zone of thickness, $h$, the column integrated mass balance equations for C$_{org}$, O$_2$ and DIN (mol m$^{-2}$ d$^{-1}$) are:

$$ h \frac{d(C_{org})}{dt} = \int_0^{EZ} \left( P^C - R^C \right) dz - F_{EZ}^C + F_{phys}^C $$

$$ h \frac{d(O_2)}{dt} = \int_0^{EZ} \left( P^O - R^O \right) dz - F_{as}^O + F_{phys}^O $$

$$ h \frac{d(NO_3)}{dt} = -P^N + R^N + F_{phys}^N $$

where $P$ is photosynthesis, $R$ is respiration, $F_{EZ}$ is sinking flux across the base of the euphotic zone, $F_{phys}$ is physical transport by advection and mixing, $F_{as}$ is air-sea gas exchange and superscripts ‘$C$’, ‘$O$’ and ‘$N$’ refer to whether the term is in carbon, oxygen or nitrogen. Sign conventions for $F_{as}$ is positive out of the ocean, $F_{EZ}$ is positive for downward flux. For each mass balance equation NCP is equal to the integral

$$ \int_0^{EZ} (P - R) dz $$

where NCP in units of C, N and O are stoichiometrically related by the photosynthetic quotient and modified Redfield ratios.

**Platforms:** Proxy for POC, itself to be determined by the hydro group, will be developed from backscatter ($b_{bp}$) on the Lagrangian float and Seagliders and beam transmission of the Lagrangian float ($c_p$). Oxygen will be measured by O$_2$ optodes on the float and gliders. Nitrate will be measured by an ultraviolet nitrate sensor on the float and possibly on a glider as well. NH4 will be determined in water collected from the rosette as part of the nutrient suite by the hydro group. Similarly, dissolved organic carbon (DOC) (the complement to suspended POC, together comprising total organic carbon in the euphotic zone) will be determined on water collected during both survey and process cruises from the rosette samples.

**Uncertainties and quality control concerns:** The magnitude and uncertainty of each term in 1–3 varies depending on state of the upper ocean system. For example, for O$_2$ the primary balance is often between $(NCP)^{O_2}$ and $F_{as}^{O_2}$. Seasonally in the subarctic, NO$_3$ drawdown can dominate such that $(NCP)^N$ and $h \frac{d(hN)}{dt}$ are the leading order terms. The C$_{org}$ budget will be limited by the lack of means to autonomously measure DOC. Quantifying $NCP$ using multiple approaches will add confidence to our estimates, reduce uncertainty and provide a check on assumed values for Redfield ratio and photosynthetic quotient.

**Other contributing protocols:** Bottle POC, DOC, Winkler O$_2$, Underway O$_2$/Ar, Bottle nitrate and ammonium

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>NCP</td>
<td>mol C m$^{-3}$ d$^{-1}$</td>
</tr>
</tbody>
</table>

**Key method references:**


**Method:** Mesopelagic Oxygen Utilization Rate from Autonomous Platform Sensors

**Document author and contact info:** David Nicholson, dnicholson@whoi.edu

Below the euphotic zone, respiration consumes O2 (and produces nitrate via remineralization). The rate of oxygen consumption is termed Oxygen Utilization Rate (OUR). OUR can be estimated from the observed time rate of change in dissolved oxygen on an isopycnal in the mesopelagic. A vertical profile of OUR is a means to characterize a ‘Martin Curve’ or the attenuation of sinking flux with depth. The Lagrangian float will drift directly below the euphotic zone where remineralization and flux attenuation is most rapid, thereby minimizing advection and optimizing the POC, O2 and NO3 budgets. In this Lagrangian frame, the time-rate of change of O2 and NO3 yield independent estimates of OUR and thus the attenuation of export flux. Our Lagrangian approach is ideally suited to quantifying OUR at depth, because effects of lateral advection are removed which, in the Eularian frame, can be of similar magnitude to OUR in the mesopelagic (Pelland, 2015). Profiles from the gliders, and potentially Bio-Argo can potentially extend OUR estimates through the mesopelagic down to 1000m and yield full depth profile of OUR.

**Platforms:** Oxygen optode and UV nitrate sensor on Lagrangian float.

**Uncertainties and quality control concerns:** A primary source of uncertainty is the quantification of advective fluxes. In the purely Lagrangian frame, advection is zero, but current shear will result in all Pacific mesopelagic has been estimated as approximately 4 μmol kg⁻¹ y⁻¹ on average. Directly below the well calibrated O2 optode can be controlled to ±0.5 μmol kg⁻¹ y⁻¹ using air calibration (Bushinsky et al., 2016), indicating that we will be able to resolve expected OUR rates on about weekly to monthly scales for the upper mesopelagic and monthly to seasonal scales for the lower mesopelagic.

**Other contributing protocols:** NBST traps, ²³⁴Th flux, Bottle respiration rates, Seaglider O2 and NO3.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>O₂ derived OUR</td>
<td>mol O m⁻³ d⁻¹</td>
</tr>
</tbody>
</table>

**Key method references:**


(Bushinsky et al., 2016; Martz et al., 2008; Pelland, 2015)
**Method:** 16S/18S amplicon sequencing of surface ocean microbial community

**Document author and contact information:** Alex Niebergall, alex.niebergall@duke.edu

**Brief description of protocol and relation to export pathways:** Duplicate 4L water samples were taken from the 5m Niskin bottle in the dawn cast and filtered through a 0.2um Sterivex filter. Samples were filtered within 1 hour of collection. Sterivex were sealed with clay and flash frozen in liquid nitrogen immediately after filtering. Samples were stored in -80°C until DNA extraction. The cells on filter were lysed by bead-beating for 1 min using 0.2 g of zirconium beads in 400 µl of lysis buffer AP1 (Qiagen). DNA was then extracted following protocols from Qiagen DNeasy Plant Mini Kit. V4 and V4-V5 hyper-variable regions of the 16S and 18S rRNA gene will be amplified using primers outlined in Parada et al. 2016 and Walters et al. 2015. Individual 6bp barcodes, designed following Bystrykh 2012, will be added to each sample primer for sample identification after sequencing. Following Fadrosh et al. 2014, a 0 to 7bp ‘heterogeneity spacer’ will be added to the primers to allow the samples to be sequenced out of phase. Internal spikes of a known quantity of *Thermus thermophilus* and *Schizosaccharomyces pombe* will be used for quantitative sequencing following methods outlined in Lin et al (submitted) and Wang et al. 2018. Amplified PCR products will be sequenced on an Illumina MiSeq with a 300 bp paired end run. DNA sequences will be assigned into amplicon sequence variants following protocol outlined in Callahan et al. 2017.

**Other contributing protocols:**
NCP O2/Ar Protocol

**Uncertainties and quality control concerns:**
To relate these data to sinking aggregates, it is critical that the same PCR primers are used to amplify 16S and 18S rDNA of sinking aggregates.

**Data products originating with this method:**
Data table of amplicon sequence variants (ASVs) - csv file
DNA sequences- Fasta files

**Key method references:**

Callahan BJ, McMurdie PJ, Holmes SP. Exact sequence variants should replace operational taxonomic units in marker-gene data analysis. The ISME journal. 2017 Dec;11(12):2639.


Flux of large sinking particles from profiles of optical spikes (Pathways 2 and 3)
Melissa Omand; momand@uri.edu, Mary Jane Perry; perrymj@maine.edu
Last updated: December 27, 2017

This document describes protocols for estimating the flux associated with large sinking particles that appear in bio-optical measurements of backscatter (bbp). Spikes that appear in beam attenuation (cp) profiles may also be associated with large sinking aggregates, and this could easily become a methodological contribution from EXPORTS. Presently however, Briggs et al (2011, hereafter B2011) is the only paper to explicitly quantify the spikes (from bbp) and directly link them to a sinking flux. Therefore, this document focuses upon the methodology described in this manuscript.

Step 1: Create a uniform ‘data point density’ over a profile and across platforms.
The analysis presented in B2011 is primarily derived from glider profiles, which generally sample more slowly (intervals of 5 to 90 seconds), and move vertically more gradually (~10 cm/s) than CTD casts or Wirewalker profiles. It is recommended that the various platforms be sub-sampled, or adjusted in some appropriate manner to maintain a constant sample rate over depth and time. Since raw bbp typically has a skewed distribution with spikes representing rare, high bbp events, in order to adequately quantify these as a proxy for large particle concentration, we must statistically sample the spikes in a manner that creates some consistency between profiles and across platforms. It may be most useful to convert the raw data into a ‘data point density’ for a particular sample strategy (the number of data points collected per meter profiled) and make this quantity consistent within a single platform profile and across platforms.

Step 2: Convert raw data to bbp.
Convert raw voltages (for FLNTU) or digital counts (for ECO Pucks) to volume scattering functions $\beta(\theta, \lambda)$ using scale factors from manufacturer’s calibrations, modified to account for the centroid angle $\theta$ of each instrument (see Sullivan et al. 2013). Next, subtract the volume scattering function of seawater, $\beta_{sw}(\theta, \lambda)$, (Zhang et al. 2009) from $\beta(\theta, \lambda)$ to get the scattering due to particles, $\beta_p(\theta, \lambda)$ and convert this to bbp (integrated particulate backscattering) using the equation: $\text{bbp}(\lambda) = 2\pi \chi \beta_p(\theta, \lambda)$, where $\chi$ is 1.132 for FLNTU and 1.077 for ECO Pucks.

Step 3: Filter for spikes in bbp.
There are a variety of options for spike filtering. The basic idea is to apply some low-pass filter to the bbp data, and then subtract this from the original signal to get a spike data set bbpsp. B2011 used a 7-point running minimum filter followed by 7-point running maximum filter, and then subtracted a noise threshold based on twice the 90th percentile value prior to a date when large spikes were rare (setting all values below this to zero). Some MatLAB functions for spike filtering created by Nathan Briggs are given at the end of this file.

Step 4: Bin average the remaining spike dataset.
Now bin-average bbp$_p$ in time and depth. Bins should be selected such that they contain at least 100 points (including zeros). In B2011, the bins were 2 days and 50 meters, in a data set that combined 4 glider records.

Step 5: Calculate an aggregate POC mass from the binned data.
The final step is to look for patterns - particularly any descending features in the bin-averaged dataset. These features may be indicative of major sinking flux events. A linear regression would then allow the estimation of a sinking rate. The concentration of POC associated with the spikes can be inferred from the slope of the POC vs bbp relationship derived from bottle and CTD comparisons. Finally, the flux attenuation coefficient (ie. the
Martin Curve exponent) can be derived from the decrease in $\text{bbp}_{sp}$ along the sinking feature (if seen) or may be applied from other EXPORTS observations.

**Key method references:**


% Matlab Scripts for Spike Filtering

```matlab
function [baseline,spikes] = separate_spikes(y,n)
% SYNTAX:
%                  [baseline,spikes] = separate_spikes(y,n);
% INUPTS:
%                     y: timeseries to filter (with equal time steps)
%                     n: window size
% OUTPUTS:
%             baseline: filtered timeseries (without spikes)
%               spikes: residuals (only spikes)
% This function calculates a running minimum followed by a running maximum,
% both with window size n, returning the filtered data i
% last modified 15 Oct 2010
% by
% Nathan Briggs - nathan.m.briggs@maine.edu
baseline = nan(size(y));
notnan = ~isnan(y);
baseline(notnan) = maxfilt(minfilt(y(notnan),n),n);
% correction for even windows
if ~rem(floor(n),2)
    warning('Odd window size is recommended')
[rw col] = size(y);
if rw > col
    baseline = [min(y(1:rw/2)); baseline; min(y(end-rw/2+1:end))];
else
    baseline = [min(y(1:rw/2)) baseline min(y(end-rw/2+1:end))];
end
end
spikes = y - baseline;
```

```matlab
function [baseline,spikes] = separate_spikes_median_enhanced(y,n)
% SYNTAX:
%                  [baseline,spikes] = separate_spikes_median_enhanced(y,n);
% INUPTS:
%...
% y: timeseries to filter (with equal time steps)
% n: window size
% OUTPUTS:
% baseline: filtered timeseries (without spikes)
% spikes: residuals (only spikes)
% DESCRIPTION:
% This function calculates a running median, but excludes high outliers
% from the median calculation
% last modified 21 Sep 2011
% by
% Nathan Briggs - nathan.m.briggs@maine.edu
baseline = nan(size(y));
notnan = ~isnan(y);

% calculation:

% correction for even windows
if ~rem(floor(n),2)
    warning('Odd window size is recommended')
end

% spike detection

spikes = y - baseline;

% filter function

function y_filt = minfilt(y,n)
% SYNTAX:
% y_filt = minfilt(y,n);
% INUPTS:
% y: timeseries to filter (with equal time steps)
% n: window size
% OUTPUTS:
% y_filt: filtered timeseries
% This function calculates a running minimum with window size n
% last modified 29 Dec 2010
% by
% Nathan Briggs - nathan.m.briggs@maine.edu

% make sure y is a row vector
col_vector = size(y,2) < size(y,1);
if col_vector
    y = y';
end

% calculation:

% correction for even windows
if ~rem(floor(n),2)
    warning('Odd window size is recommended')
end

% spike detection

spikes = y - baseline;

% filter function

function y_filt = minfilt(y,n)
% SYNTAX:
% y_filt = minfilt(y,n);
% INUPTS:
% y: timeseries to filter (with equal time steps)
% n: window size
% OUTPUTS:
% y_filt: filtered timeseries
% This function calculates a running minimum with window size n
% last modified 29 Dec 2010
% by
% Nathan Briggs - nathan.m.briggs@maine.edu

% make sure y is a row vector
col_vector = size(y,2) < size(y,1);
if col_vector
    y = y';
end

% calculation:
leg = floor((n-1)/2);
y_padded = [nan(1,leg) y nan(1,leg)];

% filter
for ii = 1:n
    y_mat(ii,:) = y_padded(ii:end-n+ii);
end
y_filt = min(y_mat);

if col_vector
    y_filt = y_filt';
end

function y_filt = maxfilt(y,n)
% SYNTAX:
% y_filt = maxfilt(y,n);
% INUPTS:
% y: timeseries to filter (with equal time steps)
% n: window size
% OUTPUTS:
% y_filt: filtered timeseries
% This function calculates a running maximum with window size n
% last modified 29 Dec 2010
% by
% Nathan Briggs - nathan.m.briggs@maine.edu

% make sure y is a row vector
col_vector = size(y,2) < size(y,1);
if col_vector
    y = y';
end

leg = floor((n-1)/2);
y_padded = [nan(1,leg) y nan(1,leg)];

% filter
for ii = 1:n
    y_mat(ii,:) = y_padded(ii:end-n+ii);
end
y_filt = max(y_mat);
if col_vector
    y_filt = y_filt';
end
**Method:** Quantification of *Transparent Exopolymer Particles (TEP)* in the water (collected via Niskin bottles)

**Document author and contact info:** Uta Passow, uta.passow@mun.ca

**Brief description of protocol and relation to export pathways:**

Transparent exopolymer particles, TEP, are essential for aggregation, as TEP form the matrix of aggregates, holding the individual component particles together. TEP thus allow the formation of rapidly sinking aggregates. However, excess density of TEP is negative, so they impact the sinking velocity of aggregates, and if unassociated with ballasting particles, TEP move upwards into the sea surface microlayer.

*Sample collection:* water for TEP measurements was sampled about twice per epoch at 10-12 depths between the surface and 500 m, using the SIO-CTD. Between 3 and 5 liters of water were sampled at each depth, and between 1 and 1.5 liters filtered onto 0.4 PC (25mm) filters in triplicates, stained with Alcian Blue and frozen until later TEP analysis.

*TEP analysis:* TEP concentrations were determined on triplicate filters (PC 0.4 µm, 25 mm diameter) using the colorimetric method (Passow and Alldredge, 1995). Between 100 and 2000 mL were filtered and stained and stained filters stored frozen in facon tubes till analysis. Results are expressed in Gum Xanthan equivalents (GXeq.) and the calibration f-factor was 84.

**Uncertainties and quality concerns:** Only a first quality control was performed, for this reason values need to be used with caution. TEP values at detection limit (DL) have been removed. Values were considered to be at detection limit if measured uncorrected values were ≤ 2 times the blank. If uncorrected values were > 2 times the blank, but ≤ 3 times the blank, they were considered to be near detection and flagged as near DL.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>TEP Bottle</td>
<td>ug_Gxan_equiv/L</td>
</tr>
</tbody>
</table>

**Key methods references:**
**Method:** Quantification of POC, PON, total C, BSi, LSi and TEP in sinking particle fractions collected from the Marine Snow Catcher

**Document author and contact info:** Uta Passow, uta.passow@mun.ca

**Brief description of protocol and relation to export pathways:**

The marine snow catcher allows the collection of particles fractionated according to their sinking velocity (rather than size). POC, PON, total C, BSi, LSi and TEP content in non-sinking, slow sinking and fast sinking particles were measured.

**Sample Collection:** Collection of samples from marine snow catchers (MSC) were conducted ≥ 3 times in each 8-day epoch, each time from 3 depths, ranging from 25 m (within mixed layer depth) to 500 m. Choice of sampling depths depended in part on Fluorescence and UVP profiles, but one MSC was always deployed at 95 m (below euphotic zone). Upon retrieval of the MSC, they were secured in the upright position and a five-liter t=0 sample (T0) collected from the center tap. The MSC were kept in the upright position for exactly 2 hours, and then 10-15 liters of the top (t) fraction was collected from the central tap, and the upper portion of the MSC drained through the center tap. This took 5 minutes. Then the lower tap was opened about 1/3 of the way and the lower portion of the upper section of the MSC drained slowly taking 25-30 minutes, so as not to disturb particles in the base of the MSC. Thereafter the upper section of the MSC was removed using a crane, allowing sampling of the base. Overlying water in the base section was siphoned off (approximately 5 liters) and is called the base (b) fraction, whereas the particles that settled in the tray (500-1000 mL) are considered to be the tray (tr) fraction. The trays were placed at the bottom of the MSC prior to deployment and collected particles sinking rapidly enough to reach the bottom of the MSC. Trays were investigated for the presence of marine snow, and particles > 0.5 mm photographed, but no marine snow sized particles (> 0.5 mm) were observed in any of the samples. Sometimes fibers were observed, but it was assumed that they were contaminants due to handling of the MSC. The total volume of the tray (tr) fraction was determined and all four fractions subsampled for further analysis.

**Analysis method:** Each analysis was done 1-2 times per epoch, POC was always measured.

- **POC/ PON/total_C:** Replicate filters (combusted GF/F, Whatmann 25 mm) were measured in a CEC44OHA elemental analyzer (Control equipment) after fuming with 10% HCL. Duplicate filters were analyzed the same way, but without prior acidification and generated total carbon (total_C). Data Product: Particulate organic carbon, particulate organic nitrogen and total carbon concentration in the specific fraction of the MSC. Unit: mg m⁻³.

- **BSi/LSi:** Filter samples were covered in 4mL of 0.2N NaOH and placed in a 95°C water bath for 40 minutes. The samples were then cooled, where 1.0 mL of 1 N HCl was added and vortexed. Samples were centrifuged for 10 minutes at 2500 rpms to separate LSi from BSi. Using a different tube, 4 mL of the sample solution was added to 6 mL of Nanopure where the reduced molybdosilicic acid spectrophotographic method by Strickland and Parsons 1968 was used. Samples were then run on a spectrophotometer. Filters left over from BSi were used to determine LSi. Filters were rinsed using Nanopure and left to dry. Once filters were completely dried, they were cooled, and 0.2mL of 2.5M HF was added to the filter and left to soak for 48 hours. Filters were then vortexed proceeding with an addition of 9.8mL of boric
acid. In a different tube, 8 mL of the sample solution was added to 2 mL of Nanopure where the reduced molybdosilicic acid spectrophotographic method by Strickland and Parsons 1968 was used. Samples were then measured on a spectrophotometer.

- **TEP analysis**: TEP concentrations were determined on triplicate filters (PC 0.4 µm, 25 mm diameter) using the colorimetric method (Passow and Alldredge, 1995). Between 100 and 2000 mL were filtered and stained and stained filters stored frozen in falcon tubes till analysis. Results are expressed in Gum Xanthan equivalents (GXeq.) and the calibration f-factor was 84.

**Uncertainties and quality concerns**: Only a first quality control was performed, for this reason values need to be used with caution. Values at detection limit (DL) have been removed. TEP values were considered to be at detection limit if measured uncorrected values were ≤ 2 times the blank. If uncorrected values were > 2 times the blank, but ≤ 3 times the blank, they were considered to be near detection and flagged as near DL. The coefficient of variation (standard deviation/average) between triplicates was on average 9%, and always < 20%, with one exception. POC and PON detection limits were 2-7 µg and 1-2 µg, respectively.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>POC</td>
<td>mg/m³</td>
</tr>
<tr>
<td>PON</td>
<td>mg/m³</td>
</tr>
<tr>
<td>total C</td>
<td>mg/m³</td>
</tr>
<tr>
<td>TEP</td>
<td>µg Gxan equiv/L</td>
</tr>
<tr>
<td>BSi</td>
<td>mmol/ m³</td>
</tr>
<tr>
<td>LSi</td>
<td>mmol/ m³</td>
</tr>
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</table>

**MSC dimensions:**

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSC_height</td>
<td>149.9  cm</td>
</tr>
<tr>
<td>MSC_diameter</td>
<td>27.6  cm</td>
</tr>
<tr>
<td>Base_height</td>
<td>12.7  cm</td>
</tr>
<tr>
<td>Tray_height</td>
<td>4.4   cm</td>
</tr>
<tr>
<td>Tray_area</td>
<td>280.0 cm²</td>
</tr>
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</table>

**Legend:**

<table>
<thead>
<tr>
<th>MSC_fraction</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>T0</td>
<td>Total POC/PON/total C/BSi/LSi/TEP contained in the Marine Snow Catcher before partitioning particles according to their sinking velocity (equivalent to T0)</td>
</tr>
<tr>
<td>t</td>
<td>POC/PON/total C/BSi/LSi/TEP contained in the top of the Marine Snow Catcher after 2h of settling</td>
</tr>
<tr>
<td>----</td>
<td>-----------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>b</td>
<td>POC/PON/total C/BSi/LSi/TEP contained in the base of the Marine Snow Catcher after 2h of settling</td>
</tr>
<tr>
<td>tr</td>
<td>POC/PON/total C/BSi/LSi/TEP contained in the tray placed in the Marine Snow Catcher after 2h of settling</td>
</tr>
</tbody>
</table>

**Key methods references:**


Sample Collection: Water for TEP measurements was sampled twice per epic at 10-12 depths between the surface and 500 m, using the SIO-CTD. Between 3 and 5 liters of water were sampled at each depth, and between 1 and 1.5 liters filtered onto 0.4 PC (25mm) filters in triplicates and stained with Alcian Blue for later TEP analysis.

Collection of samples from marine snow catchers (MSC) were conducted ≥ 3 times in each 8-day epic from 3 depths, ranging from 25 m (within mixed layer depth) to 500 m, with a 95 m (below euphotic zone) depth at all times. Choice of depths depended in part on Fluorescence and UVP profiles. Upon retrieval of MSC, they were secured in the upright position and a five-liter t=0 sample collected from the center tap. The MSC were kept in the upright position for exactly 2 hours, and then 10-15 liters of the NSP fraction was collected from the central tap, and the upper portion of the MSC drained through the center tap. This took 5 minutes. Then the lower tap was opened about 1/3 of the way and the lower portion of the upper section of the MSC drained slowly taking 25-30 minutes, so as not to disturb particles in the base of the MSC. Thereafter the upper section of the MSC was removed using a crane, allowing sampling of the base. Overlaying water in the base section was siphoned off (approximately 5 liters) and is called the SSP fraction, whereas the particles that settled in the tray (500-1000 mL) are considered to contain the FSP. Trays were investigated for the presence of marine snow, and particles > 0.5 mm photographed, but no marine snow sized particles (> 0.5 mm) were observed in any of the samples. The volume of the FSP fraction was determined and all four fractions subsampled for further analysis. The collected FSP and the SSP fractions both contain NSP, and the FSP contains SSP. This was accounted for mathematically.

At times a petri-dish with polyacryl gel was positioned into the tray to collect and immobilize sinking particles. However, these gels were never used when activity (bacterial activity, O2 consumption, DOC remineralization) was measured, so not to contaminate with carbon.

Salp fecal pellets (produced from salps kept in buckets by Steinberg group) were individually incubated in 1.2 liter rolling tanks at 41Fahrenheit in the dark to determine their sinking velocity using the orbit method. Before incubation pellets were photographed to allow accurate sizing. After 3-5 hours of incubation, when solid body rotation was established the orbit of the particle in the rolling tank was filmed. Thereafter each pellet was photographed again and then filtered onto a GFF for POC analysis. Respiration rate of salp fecal pellets was determined using the microoptode from Unisense in a 700 uL chamber with a stir bar and a screen separating pellet from stir bar. The instrument was calibrated daily (with zero and 100% oxygenated water) and calibration and sample measurements were submerged in a water bath that held temperature very constant (water bath?). A UPS device helped ensure stable current. Pellets were photographed, inserted into the chamber and stirring speed set to 600 rpm. Oxygen measurements were conducted for three or four 10-minute periods with 10 minute breaks between each measurement period. Each measurement point was averaged over 10 seconds.
Particle types & Analyses of MSC samples:
All particles combined at t=0, POC for reference only

I. Sinking and Non-sinking particles (Non-Sinking (NSP); Slow sinking (SSP: < 18 m d^{-1}); Fast sinking (FSP <0.5-1mm & > 18 m d^{-1}))
   1. Basic Characterization
      a. POC/PON content
      b. Fixed samples
   2. Composition
      a. Size-frequency distribution (Flow cam)
      b. Biochemical composition (POC, BSI, PIC).
      c. Lithogenic – Phoebe Lam
      d. Omics – Alyson Santura, Ewelina
   3. Aggregate formation
      a. TEP
      b. Stickiness/ aggregation potential
   4. Loss processes (profile # 3 of each stage)
      a. Respiration - Scott Grifford
      b. Bacterial production & abundance - Craig Carlson
      c. Remineralization rate – Craig Carlson

II. Marine Snow – no marine snow sized particles were collected in the MSC

Uncertainties and Quality Control concerns: High detail on individual characterization implies low sample numbers and low replication; not all measurements could be done each sampling day, so spatial and temporal heterogeneity make comparisons more difficult.

Data products
Vertical profiles of suspended, slow sinking and fast sinking small particles:
   o Partitioning of biochemical parameters like POC between fast, slow and non-sinking, small particles
   o Size frequency distributions in each category
   o Biogeochemical Characteristics (POC, BSI, PIC) of each category
   o TEP content in mass Gum Xanthan equivalent per volume in each category and in whole water
   o Stickiness/ aggregation potential (% relative number: fraction of particles that aggregate within a fixed time period)
   o Respiration rates of each category
   o Bacterial production rates

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biochemical content of NS, SS, FS, MS*</td>
<td>μg L^{-1} of POC, PIC, BSi</td>
</tr>
<tr>
<td>Parameter</td>
<td>Unit/Description</td>
</tr>
<tr>
<td>---------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Size distribution of NS, SS, FS</td>
<td># size bin(^{-1})</td>
</tr>
<tr>
<td>Salp pellets sinking velocity as a function of size</td>
<td>m s(^{-1})</td>
</tr>
<tr>
<td>TEP content of NS, SS, FS, and whole water</td>
<td>mass gum xanthan equivalent m(^{-3}) or mass gum xanthan equivalent MS particle(^{-1})</td>
</tr>
<tr>
<td>Proxy for stickiness (aggregation potential) of suspended or small sinking particles</td>
<td>fraction of particles that aggregate hr(^{-1})</td>
</tr>
<tr>
<td>Respiration rates of NS, SS, FS</td>
<td>O(_2) consumed per MS hr(^{-1})</td>
</tr>
<tr>
<td>Bacterial production rates</td>
<td>pmol marine snow hr(^{-1})</td>
</tr>
<tr>
<td>Remineralisation Rate</td>
<td>µg C L(^{-1}) hr(^{-1})</td>
</tr>
</tbody>
</table>

*non-sinking, slow-sinking, fast-sinking particles (NS, SS, FS)
**Method:** Determine whether microbially-degraded biomass is important for midwater zooplankton metabolism

**Document author and contact info:** Brian N. Popp (popp@hawaii.edu) and Hilary G. Close (hclose@rsmas.miami.edu)

**Brief description of protocol and relation to export pathways:** Size-fractionated zooplankton (0.2–0.5 mm, 0.5–1 mm and 1–2 mm) will be collected from a paired daytime/nighttime 1m² MOCNESS tow deployed by D. Steinberg. Each tow will be as quantitative as possible, with co-located environmental data (CTD, DO, PAR, fluorometer). Zooplankton from each net will be sieved and the fractions concentrated on pre-weighed Nitex screens, rinsed with isotonic ammonium formate to remove salts, frozen at -20°C, dried, re-weighed for dry biomass, and prepared for bulk and amino acid isotopic analyses. A total of 54 zooplankton samples will be analyzed: 3 sizes, 9 depths, and day versus night. Bulk zooplankton isotope and C:N ratios will be determined via an on-line C-N analyzer coupled with isotope ratio mass spectrometer and will be used to guide our choice of samples for amino acid compound specific isotope analysis (AA-CSIA) and to compare to published bulk data. Zooplankton will be prepared for AA-CSIA by acid hydrolysis and derivatization to produce trifluoroacetic amino acid esters and analyzed by isotope ratio mass spectrometry (IRMS) using procedures that are now routine (Hannides et al. 2009, Dale et al. 2011). The nitrogen isotopic composition of individual amino acids will allow us to calculate the trophic position of zooplankton. The amino acid carbon and nitrogen isotopic compositions of zooplankton will be compared with that of particles to assess the changes in zooplankton dietary sources over depth and organism size to determine which particle size fractions are used by zooplankton to meet their metabolic demand. Carbon isotopic compositions of essential amino acids (EAA) will be used to distinguish between marine producers versus bacterial sources of EAA to midwater zooplankton, which is related to microbial metabolism on the particles. Our results will provide basic information on the depth variability of food web-particle interactions that are needed to develop predictive models of the ocean's biological pump. Evaluating how variations in surface ocean productivity affect mesopelagic food webs will significantly further our understanding of the role of zooplankton in the export or attenuation of C and N flux. Our refined size-class differentiation of particles (see H. Close protocol) will additionally clarify the extent to which zooplankton food webs rely on discrete particle components/size fractions or along the particle size continuum.

**Other contributing protocols:** Compound-specific carbon and nitrogen isotope analyses of amino acids in size-fractionated particles (H. Close protocol).

**Uncertainties and quality control concerns:** All AA-CSIA will be conducted in triplicate; analytical uncertainty will be propagated in the calculation of metavariables, and additional uncertainty in δ¹³C values of amino acids will be propagated from analysis of derivatization standards. Suites of amino acid standards will be prepared and analyzed concurrently with each sample batch as external standards for quantitation and mass-balance correction for derivative carbon. Two synthetic amino acids will be added to samples as internal isotope and recovery standards. In addition, cross-lab standard materials (dried, homogenized fish tissue and homogenized cyanobacterial cells) will be prepared and analyzed concurrently with each sample batch.
Data products originating with this method

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zooplankton bulk carbon isotope analysis</td>
<td>‰, vs. VPDB</td>
</tr>
<tr>
<td>Zooplankton bulk nitrogen isotope analysis</td>
<td>‰, vs. AIR</td>
</tr>
<tr>
<td>Zooplankton bulk carbon content</td>
<td>µg g⁻¹</td>
</tr>
<tr>
<td>Zooplankton bulk nitrogen content</td>
<td>µg g⁻¹</td>
</tr>
<tr>
<td>Zooplankton bulk carbon:nitrogen ratio</td>
<td>mol/mol</td>
</tr>
<tr>
<td>Zooplankton carbon isotope analysis of individual amino acids</td>
<td>‰, vs. VPDB</td>
</tr>
<tr>
<td>Zooplankton nitrogen isotope analysis of individual amino acids</td>
<td>‰, vs. AIR</td>
</tr>
</tbody>
</table>

Key method references

Method: The N and O isotopic composition of nitrate (nitrate $\delta^{15}$N and $\delta^{18}$O) in seawater using the denitrifier method

Document author and contact info: Patrick Rafter, prafter@uci.edu; Alyson Santoro, asantoro@ucsb.edu

Brief description of protocol and relation to export pathways: Nitrate $\delta^{15}$N and $\delta^{18}$O will be determined in seawater samples off-ship by the quantitative reduction of nitrate ($\text{NO}_3^-$) and nitrite ($\text{NO}_2^-$) to $\text{N}_2\text{O}$ using the denitrifier method. Values will be used in support of identifying the degree of surface ocean nitrate consumption and the subsurface source of surface waters.

Other contributing protocols: none

Uncertainties and quality control concerns: Nitrate contamination introduced during sample collection (e.g. collection tubes).

Data products originating with this method:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrate $\delta^{15}$N</td>
<td>per mil [‰]</td>
</tr>
<tr>
<td>Nitrate $\delta^{18}$O</td>
<td>per mil [‰]</td>
</tr>
<tr>
<td>Uncertainty in $\delta^{15}$N</td>
<td>per mil [‰]</td>
</tr>
<tr>
<td>(standard deviation of replicate samples)</td>
<td>per mil [‰]</td>
</tr>
<tr>
<td>Uncertainty in $\delta^{18}$O</td>
<td>per mil [‰]</td>
</tr>
<tr>
<td>(standard deviation of replicate samples)</td>
<td>per mil [‰]</td>
</tr>
</tbody>
</table>

Key method references


Method: Chlorophyll analysis - 90% acetone method.

Document author and contact info: Collin Roesler, croesler@bowdoin.edu;

1. Preparation Steps.
   a. Make up sufficient volume of 90% acetone solution for sample extraction. Each centrifuge tube requires 10 ml; we generally analyze triplicates for each sample.
      i. To make up 500 ml of 90% acetone, pour 450 ml of acetone in a graduated cylinder. This is not a volumetric reaction (i.e. adding 50 ml distilled water will not result in 500 ml of 90% solution). Bring up to 500 ml volume with deionized or distilled water ensuring that the solution is well mixed.
      ii. Work quickly and ensure all containers with acetone are capped, as it is very volatile.
      iii. Work with acetone in the hood only and use gloves.
   b. Make up sufficient volume of 10% HCl solution for the fluorometric readings. Each sample reading requires 3 drops. The dropper bottle holds 30ml but it is important that the acid is freshly made (<1 month old) to ensure complete reaction.
   c. Label sample centrifuge tube (3 per sample).
   d. Using the bottle-top pipettor, pipette once or twice into Actone squirt bottle to clear line. Dispense 10 ml (sometimes we will use 5 ml) of 90% acetone into each tube. Be very accurate in this step, this is the largest source of error in the analysis. Cap tube tightly. Cap the pipettor after usage.

2. Water sample processing.
   a. Place a glass fiber filter (size GF/F is nominal pore size of 0.7 μm) in each filter holder. Attach the filter cups ensuring that they are correctly seated. Don’t over tighten.
   b. Check the level on the filter trap; it should be empty before you begin.
   c. Check the pump pressure, the intake should not exceed 5 mm Hg when blocked, the outtake should not exceed 20 when blocked.
   d. Gently swirl your sample three times in each direction, three times. Allowing sample to settle is one of the other largest sources of random error in the analysis. Shaking or vigorous swirling can break particle aggregates.
   e. Filter a pre-determined volume of sample through each filter. The filtered volume should be sufficient to just see faint color on the filter. The fluorometric method is very sensitive so only a small amount of material is necessary. If too much material is captured on the filter, the extraction efficiency will be reduced and large errors can result.
   f. Collect triplicates for each sample until you are confident that you can obtain <5% coefficient of variation in replicates, then you may do duplicates.
g. Turn the vacuum dial off for each sample as it finishes, preventing air from being drawn through the filter.

h. Fold filter and place each filter in its respective centrifuge tube; ensuring the filter is well below the acetone level. Take care not to scrape material off filter with forceps.

i. Shake the sample vigorously a few times, then, Vortex the sample for about 15 seconds. Again, ensure filter is below acetone level.

j. Store centrifuge tubes in the freezer for 24 to 48 hours. Keep them covered in foil to prevent light damage.

3. Readings with the Turner Fluorometer.
   a. Turn on fluorometer; warm up at least 15 minutes.
   b. Remove sample tubes from the freezer, keeping them protected from light. Process in batches of 6 as possible.
   c. Shake sample vigorously and vortex 5s.
   d. Centrifuge the tubes for five minutes on setting 5. Keep samples vertical to prevent filter material from resuspending. Centrifuge tubes have to arranged in a balance distribution. If all six slots are not filled, place 4 such that the six slots are Y Y N N Y Y. 3 would be arranged as Y N Y N Y N; and 2 would be Y N N Y N N.
   e. Run 90% acetone blanks on each scale of the fluorometer. Read only the 0-10 dial scale, record the sensitivity.
   f. Pour a sample from the centrifuge tube into the glass fluorometer vials, minimizing the amount of filter material in the tube. While sample is stabilizing in fluorometers, dump remaining sample in waste container, remove filter and wipe sample name from tube.
   g. Take a reading on the fluorometer, setting the sensitivity so that the reading is between 2 and 8 if possible. Record both the scale (and the X1 or X100 setting) and the dial reading to the nearest 0.1.
   h. Add 3 drops of 10% HCL, invert with parafilm, set aside.
   i. Repeat f-h for the remaining 5 samples.
   j. Take a reading on the first acidified sample. Record both the scale (and the X1 or X100 setting) and the dial reading to the nearest 0.1, the value will be lower than the original reading and likely on a more sensitive scale. Pour sample into waste container when done.
   k. Process the remaining samples from c to j.
   l. Wash tubes and vials with micro cleaner and rinse well with RO water. Dry upside down in rack or on towels. Regarding tubes on the fluorometers, the assumption is that if they are upside down they are clean, right side up are dirty.

   a. Record the calibration coefficients written on the Turner Fluorometer (M = _______ and A= ________) and note the date of those coefficients.
   b. Chl (mg m$^{-3}$) = $M \times (F_0 - F_a)$ * (1)
   Phaeo (mg m$^{-3}$) = $M \times ((A F_a) - F_0)$ * (2)
Where $M$ and $A$ are the calibration coefficients noted in step 4b. $V_{acOH}$ is the volume of acetone in the centrifuge tube (10 ml) and $V_{filt}$ is the volume of sample filtered in ml. $F_0$ and $F_a$ are calculated as follows:

$F_0 = (3)$

$F_a = (4)$

Where $R_o$ and $R_a$ are the dial reading values for the sample before and after acidification, respectively, scale is the scale that the reading was made at and $R_{blank}$ is the dial reading for pure 90% acetone on the scale that $R_o$ or $R_a$ was made at. For example, if $R_o$ was made on the 100 scale, $R_{blank}$ in equation (3) is the dial reading for acetone on the 100 scale; if $R_a$ was made on the 316 scale, $R_{blank}$ in equation (4) is the dial reading for acetone on the 316 scale.

References for extractive technique and applications:


Reference for field fluorometer technique:

Method: In-line Hyperspectral Size–Fractioned Particulate Absorption

Document author and contact info: Collin Roesler, croesler@bowdoin.edu;

Brief description of protocol and relation to export pathways: Hyperspectral size-fractioned particulate absorption coefficients were obtained from continuous inline observations collected with a WETLabs acs using a differencing method between unfiltered and filtered observations following Slade et al. (2010). We collected continuous inline observations of temperature and salinity using a SeaBird TSG thermsalinograph. Particulate absorption provides an assessment of the total particulate load and a means to assess phytoplankton from non-phytoplankton particulates. It also provides an assessment of absorption line height for the estimation of chlorophyll concentration in the absence of non-photochemical quenching. The hyperspectral beam attenuation provides an additional proxy for total particulate load and additionally is a proxy for particulate organic carbon. The spectral slope of the beam c is a size proxy.

Deployment methodologies: The flow through system is configured so that the inline water system is supplied with a diaphragm pump to minimize particle disaggregation. Water entered the inline system through a vortex debubbler followed by a SeaBird thermsalinograph, then a Sequoia Flow Meter and finally to the acs. The flow meter automatically switched flow from unfiltered to filtered through a large volume 0.2 µm cartridge filter. The automated switch operated continuously through the day such that during each hour 50 minutes is unfiltered and 10 minutes is filtered. Approximately twice per day, an additional filtered configuration was manually triggered such that the hour was parsed into 20 minutes of unfiltered, 15 minutes of 20-µm filtration, 15 minutes of 5-µm filtration and 10 minutes of 0.2-µm filtration. Discrete water samples were collected from each of the size fractions and processed for spectrophotometric particulate absorption, HPLC and POC. An inline CDOM fluorometer, Chl fluorometer and backscattering sensors provide continuous monitoring of dissolved absorption, phytoplankton biomass and particulate biomass, respectively, which are used to interpolate the absorption time series through the fractions.

Data processing
Raw data were processed to remove noisy data associated with bubbles by computing initial 1-minute median bins, then removing raw data that was outside of 1 standard deviation of the median. This was then applied across a range of channels (wavelengths). The raw data were then reprocessed to 1-minute median bins. Standard deviation values were computed for the 1-minute bins.

One-minute binned data was merged with the Ship’s navigation data sets that were also processed to 1-minute bins.

Data time series were then parsed into size bins (unfiltered, <0.2 um, <5 um and <20um). The <0.2um filtered time series was interpolated over the time vector. Particulate absorption was computed by subtracting the interpolated filtered time series from the unfiltered time series for a and c. Uncertainty of the difference was propagated from the 1-minute standard deviation values.
The spectrally flat scattering correction was applied to the particulate absorption (selecting the 725nm to 750 nm median offset). This is in keeping with the observation that particulate absorption measured in the integrating sphere across the 3 epochs yielded negligible NIR absorption. Scattering spectra were computed by difference. Uncertainty was propagated mathematically for each computation.

**Uncertainties and quality control concerns**: Uncertainties associated with natural variations in absorption were determined from the standard deviations of the one-minute bin averaged data. Uncertainties for each computation were computed arithmetically.

There was significant fouling of the inline system. Steps were taken to clean the ship’s lines with bleach approximately every 3 days. All tubing in the instrumented system was replaced every 3-5 days as fouling was observed. Raw data exhibited the signal of fouling (exponentially increasing signals in a and c). However, once the particulate IOPs were computed the biofouling trends have disappeared. The values for the inline system have not been validated against ap from center-mounted integrating sphere method yet.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Particulate absorption coefficient, total</td>
<td>(a_{pT})</td>
<td>(m^{-1})</td>
</tr>
<tr>
<td>Particulate absorption coefficient, &lt;20 µm</td>
<td>(a_{p&lt;20})</td>
<td>(m^{-1})</td>
</tr>
<tr>
<td>Particulate absorption coefficient, &lt;5 µm</td>
<td>(a_{p&lt;5})</td>
<td>(m^{-1})</td>
</tr>
<tr>
<td>Dissolved absorption coefficient, &lt;0.2 µm</td>
<td>(a_{p&lt;0.2})</td>
<td>(m^{-1})</td>
</tr>
<tr>
<td>Particulate absorption coefficient, 0.2-5 µm</td>
<td>(a_{p0.2-5})</td>
<td>(m^{-1})</td>
</tr>
<tr>
<td>Particulate absorption coefficient, 5-20 µm</td>
<td>(a_{p5-20})</td>
<td>(m^{-1})</td>
</tr>
<tr>
<td>Particulate absorption coefficient, &gt;20 µm</td>
<td>(a_{p&gt;20})</td>
<td>(m^{-1})</td>
</tr>
</tbody>
</table>

**Daily files and format**

Daily files for particulate absorption, beam attenuation and scattering (ap, cp and bp, respectively) were constructed for submission to SEABASS. The format is as follows:

- **ap**: Date, time, latitude, longitude, Chl, UChl, ap(1-78 channels), Uap(1-78)
- **cp**: Date, time, latitude, longitude, gamma, RMSEgamma, cp(1-78 channels), Ucp(1-78)
- **bp**: Date, time, latitude, longitude, bp(1-78 channels), Ubp(1-78)

where Chl is the chlorophyll concentration (mg.m3) derived from red peak absorption line height, gamma is the spectral slope of cp, U stands for the propagated uncertainty, and RMSEgamma is a measure of the spectral slope model fit to each spectrum.
Key method references:


Method: In-line Multi-excitation Chlorophyll Fluorescence, WET Labs ECO 3X1M

Document author and contact info: Collin Roesler, croesler@bowdoin.edu

Date: 1 August 2019

Brief description of protocol: Multi-excitation chlorophyll fluorescence was obtained from continuous inline observations collected with a WETLabs ECO 3X1M fluorometer using the differencing method between unfiltered and filtered observations following Slade et al. (2010). The fluorometer excites with 3 LEDs (nominally 440 nm, 470 nm, and 532 nm) and measures chlorophyll fluorescence (695 nm). The instrument is calibrated against a dilution series of the monospecific culture of the diatom Thalassiosira pseudonana to quantify the relative fluorescence response to each of the 3 excitations. Earlier studies have demonstrated that the fluorescence ratios vary as pigmentation varies (Yentsch and Phinney 1985), primarily due to taxonomically-driven variations in pigment composition with secondary dependence on photoacclimation and growth phase (Proctor and Roesler 2010; Thibodeau et al. 2014). The standard method for estimating chlorophyll a concentration is to calibrate in situ observations of chlorophyll fluorescence derived from a 470 nm LED to paired in situ HPLC TChl measurements. The fluorescence at the other channels is scaled according to the T. pseudonana response. Thus if the in situ phytoplankton was T. pseudonana then all three channels would yield the same calibrated chlorophyll values. Variations in the derived calibrated chlorophyll between channels is then interpreted as pigment variations relative to those observed in T. pseudonana, as published in Proctor and Roesler (2010).

Deployment methodologies: A diaphragm pump installed in the sea chest of the R/V Armstrong delivered continuous seawater to the wet lab, with minimal particle disruption. The inline optical system consists of serial flow in the following order: MSRC VDB-1 vortex debubbler, Seabird thermosalinograph, Sequoia Flow Meter, WET Lab acs, WET Labs ECO 3X1M, WET Labs ECO BBFL2 and WET Labs ECO BB3. The ECO 3X1M was configured in a WET Labs cylindrical ECO flow cell. The flow meter automatically switched flow from unfiltered to filtered through a large volume 0.2 μm cartridge filter. The automated switch operated continuously through the day such that during each hour 50 minutes is unfiltered and 10 minutes is filtered. Approximately twice per day, an additional filtered configuration was manually triggered such that the hour was parsed into 20 minutes of unfiltered, 15 minutes of 20-μm filtration, 15 minutes of 5-μm filtration and 10 minutes of 0.2-μm filtration. Discrete water samples were collected from each of the size fractions and processed for spectrophotometric particulate absorption, HPLC and POC.

Data processing: All flow through data were processed into 1 minute bin median and standard deviation values. Observations compromised by bubbles were removed. Data were time-merged with the ship navigation GPS data streams, also processed into 1-minute bin median values. Data stream was parsed into unfiltered and filtered intervals, with transitional data removed. Filtered observations were interpolated to 1–minute intervals and subtracted from unfiltered data, thus yielding blank-subtracted fluorescence values (Slade et al. 2010). Non-negligible non-photochemical quenching was observed within 2 hours of local midnight. Match-ups between
discrete HPLC TChl values and unquenched inline chlorophyll fluorescence values were used to generate an in situ calibration for the 470 nm excitation channel, with \( M \left( \frac{mg}{m^3 dc^{-1}} \right) \), representing the slope of the matchup relationship

\[
Chl \left( \frac{mg}{m^3} \right) = M \left( \frac{mg}{m^3 dc} \right) \times (F_{un\text{filt}} - F_{filt}).
\]

The calibration was scaled to the 440 nm and 532 nm channels using the 440:470 and 532:470 blank-corrected digital count ratios established with the \textit{T. pseudonana} culture. All values are reported in units (mg m\(^{-3}\)); non-photochemically quenched values are provided and should not be interpreted as chlorophyll concentration.

**Uncertainties and quality control concerns:** Uncertainties associated with natural variations in chlorophyll fluorescence were determined from the standard deviations of the one-minute bin median data for filtered, \( \sigma_{filt} \), and unfiltered, \( \sigma_{un\text{filt}} \), observations. Uncertainty in the chlorophyll calculation, \( \sigma_{Chl} \), was propagated (Jcgm 2008) as:

\[
\sigma_{Chl} = Chl \times \sqrt{\sigma_{un\text{filt}}^2 + \sigma_{filt}^2 + \left( \frac{\sigma_M}{M} \right)^2}
\]

where \( \sigma_M \) is the uncertainty in the regression slope of the HPLC and fluorescence match-up.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calibrated Fchl (ex470 nm)</td>
<td>Fchl(470)</td>
<td>mg m(^{-3})</td>
</tr>
<tr>
<td>\textit{Thalassiosira pseudonana}-equivalent Fchl (excitation 440 nm)</td>
<td>Fchl(440)</td>
<td>mg m(^{-3})</td>
</tr>
<tr>
<td>\textit{Thalassiosira pseudonana}-equivalent Fchl (excitation 532 nm)</td>
<td>Fchl(532)</td>
<td>mg m(^{-3})</td>
</tr>
</tbody>
</table>

**Key method references:**


Method: In-line Three-Channel Backscattering, WET Labs ECO BB3

Document author and contact info: Collin Roesler, croesler@bowdoin.edu

Date: 1 August 2019

Brief description of protocol: Three-channel backscattering was obtained from continuous inline observations collected with a WETLabs ECO BB3 sensor using the differencing method between unfiltered and filtered observations following Slade et al. (2010). The backscattering sensor was comprised of 3 LEDs, nominally 412 nm, 595 nm, 715 nm. Sunstone Scientific calibrated the sensor just prior to the EXPORTS North Pacific expedition.

Deployment methodologies: A diaphragm pump installed in the sea chest of the R/V Armstrong delivered continuous seawater to the wet lab, with minimal particle disruption. The inline optical system consists of serial flow in the following order: MSRC VDB-1 vortex debubbler, Seabird thermosalinograph, Sequoia Flow Meter, WET Lab acs, WET Labs ECO 3X1M, WET Labs ECO BBFL2 and WET Labs ECO BB3. The ECO BB3 was configured in a custom-designed casket (Dall'olmo et al. 2009) that minimizes impacts of scattered light within the flow. The flow meter automatically switched flow from unfiltered to filtered through a large volume 0.2 μm cartridge filter. The automated switch operated continuously through the day such that during each hour 50 minutes is unfiltered and 10 minutes is filtered. Approximately twice per day, an additional filtered configuration was manually triggered such that the hour was parsed into 20 minutes of unfiltered, 15 minutes of 20-μm filtration, 15 minutes of 5-μm filtration and 10 minutes of 0.2-μm filtration. Discrete water samples were collected from each of the size fractions and processed for spectrophotometric particulate absorption, HPLC and POC.

Data processing: All flow through data were processed into 1-minute bin median and standard deviation values. Observations compromised by bubbles were removed. Data were time-merged with the ship navigation GPS data streams, also processed into 1-minute bin median values. Data stream was parsed into unfiltered and filtered intervals, with transitional data removed. Filtered observations were interpolated to 1-minute intervals and subtracted from unfiltered data, thus yielding blank-subtracted fluorescence values (Slade et al. 2010).

The particle backscattering coefficient at each wavelength was computed from:

\[ b_{bp} \ (m^{-1}) = M \left( m^{-1} / dc \right) \times \left( DC_{bbunfilt} - DC_{bbfilt} \right) . \]

Where \( M \) is the calibration slope provided by Sunstone Scientific (with a best-case accuracy of 2.1%) and \( DC \) indicates the 1-minute median digital count values for unfiltered or interpolated filtered observations.

Uncertainties and quality control concerns: Uncertainties associated with natural variations in particle backscattering were determined from the standard deviations of the one-minute bin median data for filtered, \( \sigma_{filt} \), and unfiltered, \( \sigma_{unfilt} \), observations. Following the Guide to Uncertainty Measurements (Jcgm 2008), the uncertainty in the particle backscattering coefficient for each channel is computed as:
\[ \sigma_{bbp} = b_{bp} \times \sqrt{\sigma_{unfilt}^2 + \sigma_{filt}^2 + \left(\frac{\sigma_M}{M}\right)^2} \]

where \( \sigma_M \) is the calibration uncertainty.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>bbp (412 nm, 595 nm, 715 nm, )</td>
<td>( b_{bbp} ) (412) ( b_{bbp} ) (595) ( b_{bbp} ) (715)</td>
<td>mg m(^{-3})</td>
</tr>
</tbody>
</table>

**Key method references:**


Method: In-line Single Channel Backscattering and Chlorophyll and CDOM Fluorescence, WET Labs ECO BBFL2

Document author and contact info: Collin Roesler, croesler@bowdoin.edu

Date: 1 August 2019

Brief description of protocol: Single-excitation backscattering, and chlorophyll and CDOM fluorescence was obtained from continuous inline observations collected with a WETLabs ECO BBFL2 sensor using the differencing method between unfiltered and filtered observations following Slade et al. (2010). The backscattering sensor was comprised of a 660 nm LED, the Chl fluorometer excites with a 470 nm LED (emission 695 nm), the CDOM fluorometer excites with a 370 nm LED (emission 460 nm). Calibrations were performed just prior to the EXPORTS North Pacific expedition: the backscattering sensor was calibrated by Sunstone Scientific; the CDOM fluorometer was calibrated by WET Labs; the Chl fluorometer was calibrated by WET Labs and also in-house using a monospecific culture of Thalassiosira pseudonana (Proctor and Roesler 2010; Roesler et al. 2017). The standard method for estimating in situ chlorophyll concentration is to calibrate in situ observations of chlorophyll fluorescence derived from a 470 nm LED to paired in situ HPLC TChl measurements.

Deployment methodologies: A diaphragm pump installed in the sea chest of the R/V Armstrong delivered continuous seawater to the wet lab, with minimal particle disruption. The inline optical system consists of serial flow in the following order: MSRC VDB-1 vortex debubbler, Seabird thermosalinograph, Sequoia Flow Meter, WET Lab 3acs, WET Labs ECO 3X1M, WET Labs ECO BBFL2 and WET Labs ECO BB3. The ECO BBFL2 was configured in a custom-designed casket (Dall'olmo et al. 2009) that minimizes impacts of scattered light within the flow. The flowmeter automatically switched flow from unfiltered to filtered through a large volume 0.2 μm cartridge filter. The automated switch operated continuously through the day such that during each hour 50 minutes is unfiltered and 10 minutes is filtered. Approximately twice per day, an additional filtered configuration was manually triggered such that the hour was parsed into 20 minutes of unfiltered, 15 minutes of 20-μm filtration, 15 minutes of 5-μm filtration and 10 minutes of 0.2-μm filtration. Discrete water samples were collected from each of the size fractions and processed for spectrophotometric particulate absorption, HPLC and POC.

Data processing: All flow through data were processed into 1-minute bin median and standard deviation values. Observations compromised by bubbles were removed. Data were time-merged with the ship navigation GPS data streams, also processed into 1-minute bin median values. Data stream was parsed into unfiltered and filtered intervals, with transitional data removed. Filtered observations were interpolated to 1-minute intervals and subtracted from unfiltered data, thus yielding blank-subtracted fluorescence values (Slade et al. 2010).

The particle backscattering coefficient was computed from:

$$b_{bp} (m^{-1}) = M \left( m^{-1} / d_c \right) \times \left( DC_{bbunfilt} - DC_{bbfilt} \right).$$
Where $M$ is the calibration slope provided by Sunstone Scientific (with a best-case accuracy of 2.1%) and $DC$ indicates the 1-minute median digital count values for unfiltered or interpolated filtered observations.

Non-negligible non-photochemical quenching was observed within 2 hours of local midnight. Match-ups between discrete HPLC TChl values and unquenched inline chlorophyll fluorescence values were used to generate an in situ calibration for the 470 nm excitation channel, with $M \left(\frac{mg}{m^3} dc^{-1}\right)$, representing the slope of the matchup relationship (with a computed uncertainty in the regression slope of $\sigma_M$):

$$Chl \left(\frac{mg}{m^3}\right) = M \left(\frac{mg}{m^3} dc\right) \times (F_{unfilt} - F_{filt}).$$

Non-photochemically quenched values are provided and should not be interpreted as chlorophyll concentration.

**Uncertainties and quality control concerns**: Uncertainties associated with natural variations in particle backscattering and chlorophyll fluorescence were determined from the standard deviations of the one-minute bin median data for filtered, $\sigma_{filt}$, and unfiltered, $\sigma_{unfilt}$, observations. Following the Guide to Uncertainty Measurements (Jcgm 2008), the uncertainty in both backscattering coefficient and chlorophyll concentration are computed as:

$$\sigma_X = X \times \sqrt{\sigma_{unfilt}^2 + \sigma_{filt}^2 + \left(\frac{\sigma_M}{M}\right)^2}$$

where $X$ is either $b_{bp}$ or $Chl$, and $\sigma_M$ is the uncertainty in the calibration.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>bbp (660 nm)</td>
<td>$b_{bp}$ (470)</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>Calibrated Fchl (ex470 nm)</td>
<td>Chl</td>
<td>mg m$^{-3}$</td>
</tr>
</tbody>
</table>

**Key method references:**


Method: Surface Hyperspectral Radiometry

Document author and contact info: Collin Roesler, croesler@bowdoin.edu;

Brief description of protocol and relation to export pathways: Hyperspectral remote sensing reflectance will be obtained at each daytime (1000-1400) CTD station using a Satlantic Hyper Tethered Spectral Radiometer Buoy (H-TSRB). Downwelling irradiance is measured with a cosine sensor in air at 123 wavebands from 398 nm to 800 nm with approximately 3 nm resolution. Upwelling radiance is measured 63 cm below the air-sea interface with a nadir-viewing radiance sensor with 8½ ° FOV. Upwelling radiance is propagated through the interface following the approach of Morel et al. (2007) modified for hyperspectral radiometry to retrieve water leaving radiance, \( L_w(\lambda) \). Remote sensing reflectance is computed from the ratio of hyperspectral water leaving radiance to incident downwelling irradiance. Remote sensing reflectance measured at the sea surface provides the ground truth observation for top-of-the atmosphere remote sensing reflectance obtained from ocean color sensors such as the currently in orbit multi-spectral MODIS and the propose hyperspectral PACE sensors. Semi-analytic inversions of ocean color reflectance yield estimates of inherent optical properties such as component absorption and backscattering coefficients and carbon-based optical proxies such as POC and DOC.

Deployment methodologies: The H-TSRB is deployed by hand from the deck of the ship of the sunny side of the vessel. It remains tethered while it is allowed to float away to greater than three optical depths from the ship to avoid shadow and reflection. At least one minute of observations is collected. The radiometer is temporarily retrieved, caps are placed on the radiometers and it is redeployed in order to obtain temperature and stability dark readings that are removed from sample observations.

Uncertainties and quality control concerns: Uncertainties associated with natural variations in the radiometry observations are computed by standard deviation of the 60-second burst samples. Instrument uncertainty is assessed by mean and standard deviation of the 60-second burst sample of dark readings. These are arithmetically propagated from radiance and irradiance to remote sensing reflectance computations.

Data products originating with this method:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upward radiance at 63 cm depth</td>
<td>( L_u(\lambda, 0.63) )</td>
<td>( \mu \text{W cm}^{-1} \text{nm}^{-1} \text{sr}^{-1} )</td>
</tr>
<tr>
<td>Upward radiance at 0 cm depth</td>
<td>( L_u(\lambda, 0.0) )</td>
<td>( \mu \text{W cm}^{-1} \text{nm}^{-1} \text{sr}^{-1} )</td>
</tr>
<tr>
<td>Water leaving radiance</td>
<td>( L_w(\lambda) )</td>
<td>( \mu \text{W cm}^{-1} \text{nm}^{-1} \text{sr}^{-1} )</td>
</tr>
<tr>
<td>Incident downward irradiance</td>
<td>$E_d(\lambda, 0^+)$</td>
<td>$\mu W \ cm^{-1} \ nm^{-1}$</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>----------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td>Remote sensing reflectance</td>
<td>$R_{rs}(\lambda)$</td>
<td>$sr^{-1}$</td>
</tr>
</tbody>
</table>

**Key method references:**
Brief Overview: Spectral absorption coefficients of particulate matter is determined from spectrophotometric analysis of particles filtered onto glass fiber filters. The particulate matter is partitioned chemically into in vivo pigments (phytoplankton absorption) and non-extractable particulate matter (non-algal particles). The conversion of absorbance to absorption required quantification of the geometric pathlength (m), accounted for by the ratio of the volume filtered (m³) and the effective filter area (m²), and the ratio of the geometric to optical pathlengths, accounted for by experimental comparisons between particulate samples measured in suspension to those on filter pads.

Water sample collection: Discrete water samples were collected from a Niskin bottle rosette sampler from approximately 8 depths, 1-3 times per day. In order to ensure compatibility between pigment, carbon and optical samples, the entirety of each Niskin bottle (from each depth) was collected into a large volume carboy. Carboys were be kept cold and dark until subsampling, within one hour of CTD sampling. Carboys were kept well mixed, by multiple-directional changes in swirling, during subsampling into analysis-specific sampling bottles. Discrete water samples were also collected from an inline water system, pumped from the sea chest with a diaphragm pump, that was coupled to the flow-through optical sensor suite. Samples were collected once or twice a day, in conjunction with a CTD cast. The intake depth was approximately 5m, coincident with the surface Niskin bottle depth. The inline samples were size fractionated with 5 µm and 20 µm cartridge filters (Pall®), in addition to the unfiltered samples. Real time optical data display was used to identify when the outflow water was completely size fractionated (approximately 7 minutes for complete sample flow through and reservoir turn over).

Water sample preparation: Discrete water sub-samples were collected in 2-Liter Nalgene sample bottles and filtered through Whatman® glass fiber filters (25 mm diameter, nominal pore size GF/F) under low pressure (< 5 mm Hg) vacuum with an aspirator pump. The exact volume and volume uncertainty of each sample was recorded. A set of 3-5 blank filter pads for baseline and blank reference scans were prepared in the similar manner but with comparable volumes of MilliQ® purified water filtered for each run of samples. Filter pads were transferred to taped glass petri dishes and laid, in order, on top of moist Kim Wipes® to maintain uniform moisture. Samples were measured immediately after filtration.

Absorbance measurements: Absorbance scans were measured spectrophotometrically in a Cary 300 dual beam spectrophotometer configured with a Labsphere® integrating sphere with center-mounted Plexiglas sample holder. The integrating sphere was aligned before the ship departed and the alignment was checked throughout the cruise. The wavelength range was 350 nm to 800 nm, the slit band width was 2 nm, the wavelength interval was 1 nm, the integration time was 0.2 s for a scan rate of 300 nm/s. One of the set of blank filter pads was selected as the baseline filter. After the baseline scan, the filter, untouched, was run as a sample. This is referred to as the zero absorbance scan. This represents the uncertainty due to instrument noise. The remaining
blank filters were scanned relative to the baseline scan, these are referred to as blank scans. These represent the variability in absorbance due to filter-to-filter variability. Each sample filter was scanned relative to the baseline. The initial scan is referred to as the particulate absorbance scan. The filter is then placed back into the filter cup and extracted with approximately 10 ml of hot methanol for 15 minutes. After the methanol was filtered through, samples were rinsed with approximately 10 ml of MilliQ water and examined for residual color. If necessary, extraction was repeated. The extracted filter was scanned; this is referred to as the non-algal particle (or nap) scan (Kishino et al. 1985). Most filters were extracted twice. By the end of the cruise, there was a shortage of methanol and only a single extraction was performed. Phycobilipigments are not extracted with methanol and those samples were extracted with hot water (Roesler and Perry 1995). In some cases residual pigment absorption was observed and the fully extracted absorbance was estimated by exponential fit (see data processing).

Data Processing: The absorbance scans from the EXPORTS NE Pacific cruise were at the low end of the optimal absorbance range, approximately 0.1 to 0.2 in the blue to green spectral band. However, they were much noisier than is typically found for coastal water samples with similar sample loading on the filter. This was true for samples collected on the Revelle process ship, stored in liquid nitrogen, and processed in the same spectrophotometer in the lab. Thus it is not an instrumental artifact nor a filter loading artifact, but likely a result of the size and composition of the natural particles. The sample absorbance scans were thus smoothed with a 9 nm moving box average filter. They were smoothed twice.

The sample absorbance, $A_{sample}$, was corrected for pathlength amplification following the empirical relationship of Stramski et al. (2015):

$$A = 0.323 \times (A_{sample}^{1.0867}).$$

Spectral absorption coefficients, $a(\lambda)$ ($m^{-1}$), for both the particulate and non-algal particles were computed from the corrected absorbance scans using the following equation:

$$a(\lambda) = 2.303 \times 100 \frac{A}{V_{filt}/\pi r^2}$$

where 2.303 converts the $\log_{10}$ absorbance to $\log_e$ absorption, $A$ is the measured absorbance corrected for pathlength amplification, $V_{filt}$ is the volume filtered (ml or cm$^3$), and $\pi r^2$ is the effective area of the filter (cm$^2$) and the factor of 100 converts cm to m. Absorption by the phytoplankton component was computed by difference between particulate and non-algal particle absorption coefficients. Uncertainty values were computed by mathematical propagation of instrumental and sample uncertainty terms following NASA protocols (2018).

In the case of incomplete extraction, an exponential fit to the non-algal particle absorption spectrum was estimated over the wavelength range 375 nm to 750 nm, the endpoints representing wavelengths for which extractable algal pigment absorption is minimal. The spectral slope of the exponential was assumed to be 0.01 (nm$^{-1}$), based upon the least-square best
fit to well-extracted samples. The revised phytoplankton absorption, $\hat{a}_{\text{phyt}}$, was computed from the difference between the particulate absorption and the modeled non-algal particle absorption, $\hat{a}_{\text{nap}}$.

**SeaBASS data file structure:** The file structure for the SeaBASS data submission is:

wavelength, $A_{\text{zero}}$, median $A_{p,\text{blank}}$, median $A_{\text{nap,blank}}$, $A_{p}$, $A_{\text{nap}}$, $a_{p}$, $\alpha_{\text{nap}}$, $\alpha_{\text{phyt}}$, $\hat{a}_{\text{nap}}$, $\hat{a}_{\text{phyt}}$

where size fractions are ordered total, $<20\mu m$ and $<5 \mu m$ within each data type.

**References:**


Method: Metabarcoding of water column eukaryotic plankton

Document author and contact info: Ewelina Rubin, ewelina_rubin@uri.edu; Tatiana Rynearson, rynearson@uri.edu

Brief description of protocol and relation to export pathways:

This work is being done as part of the particle characterization EXPORTS working group

Biomass on filters was collected daily from depth between 5 and 73m samples onboard the R/V Sally Ride and preserved in -80C. In addition biomass filters were collected from 50-300 m water onboard the the R/V Roger Revelle and preserved Qiagen lysis buffer and stored at 4C. Further, samples from 5-73 m depth were collected, preserved in RNALater and stored at -20C. DNA will be extracted with the Power Water DNA extraction kit (Qiagen). If needed, it will be purified with Zymo Spin IV-µHRC columns (Zymo Research) which removes many environmental PCR inhibitors. The DNA will be quantified with the Qubit DNA broad range assay (Life Technologies). Amplicon libraries targeting the V4 region of SSU rDNA will be generated with the eukaryotic universal primers, designed by Stoeck et al., (2010) and modified by Bradley et al., (2016). These primers will be customized for MiSeq (Illumina) sequencing by adding, in respective order, an adapter for the amplicon binding to the flow cell, a sequencing primer binding site, and a barcode sequence identifying the sample (Kozich et al., 2013). In addition, the forward primer will include a four random-nucleotide sequence to introduce sequence diversity and thereby increase the ability of the MiSeq instrument to discriminate adjacent fluorescent clusters (Nelson et al., 2014). Three 25-µl PCR reactions per sample will be run using Phusion polymerase (Thermo Scientific), with 30 ng of DNA and 0.4 µM of primers. The PCR protocol is: 98°C for 30 s, 30 cycles of 98°C for 10 s, 50°C for 20 s and 72°C for 40 s, and a final step at 72°C for 10 min. Triplicate PCR products will be purified using gel extraction method pooled into one sample and then quantified, and pooled again into a multiplexed library. Paired-end sequencing will be conducted on a MiSeq instrument.

Uncertainties and quality control concerns:

Given the low abundance of plankton in the water column, there are concerns that 1-2 L of filtered seawater may yield low DNA concentrations that are difficult to amplify. If this occurs, samples will be subjected to whole genome amplification prior to amplification of the 18S V4 region. The 18S V4 rRNA region will provide the best resolution at the genus level with only a subset of sequences identified to species. Additional primer sets may be used to enhance resolution of microzooplankton community members.

Data products originating with this method:

Raw data as fastq files (separate for each sample)
Tables and graphs reflecting phylogenetic identification of major eukaryotic groups (mostly likely genera), their relative abundance and the beta diversity for each sampled location.

**Key method references:**


**Method:** Enumeration of phytoplankton and microzooplankton using microscopy

**Document author and contact info:** Tatiana Rynearson, Rynearson@uri.edu

**Brief description of protocol and relation to export pathways:** 20-100ml aliquots of seawater from daily depth profiles will be fixed using 2% final volume Lugol’s solution. Utermoehl settling chambers (1-50ml) and inverted microscopy using 40-400x magnification will be used to identify and enumerate phytoplankton and microzooplankton in the 5-100 micron size range (Lund et al. 1958). Based on the biomass in the fixed samples, settling volumes will be adjusted so that a minimum of 400 cells of each abundant species or genus is counted, to allow for robust downstream statistical analyses (Lund et al. 1958). The settling chambers will initially be enumerated using 40-100x magnification, with the option of examining the sample at higher magnification (200-400x) to identify and count smaller cells and colonies. The length of major axes of the dominant phytoplankton will be measured for 30-50 cells of each species/genus.

**Data Processing:** Cell abundance will be calculated by dividing the total cells of each species/genus counted by the volume settled in the Utermoehl chamber. Approximate cell volumes will be calculated and converted to carbon concentration using the equations of Menden-Deuer and Lessard (2000).

**Calibration:** The major axes of dominant phytoplankton are measured on the inverted microscope using an eyepiece with a ruler etched onto it. The ruler is calibrated using a stage micrometer.

**Uncertainties and quality control concerns:** Phytoplankton can be morphologically identical, especially at the level of light microscopy. The cell counts will be paired with DNA sequencing to resolve species identity of genera known to contain morphologically cryptic species (*Pseudonitzschia*, *Thalassiosira*, *Skeletonema*, *Chaetoceros*). Not all plankton fix equally well in Lugol’s fixative and this will be taken into account when community composition is evaluated using the inverted microscopy counts. The absolute abundance of rare cells will have a large uncertainty. This can be ameliorated up to a point by settling larger volumes.

**Data products originating with this method:** Community composition and carbon concentration of depth profiles for phytoplankton and microzooplankton.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Community composition</td>
<td>cells species(^{-1}), cells genus(^{-1})</td>
</tr>
<tr>
<td>Carbon per cell</td>
<td>pg C cell(^{-1})</td>
</tr>
</tbody>
</table>
Key method references:


1. **Method:** Ecogenomic grazing markers

2. **Contact Info:** Tatiana Rynearson, rynearson@uri.edu, Ewelina Rubin ewelina_rubin@uri.edu

3. **Brief description:** To enhance the resolution of grazing rate measurements, we will be collecting biomass for the identification of genes expressed in actively feeding predators. This work is done in conjunction with laboratory experiments, where biomass from grazing experiments will be harvested and used to validate predator genes whose expression varies depending on feeding status with the long-term goal of using the expression of those genes to determine feeding status in natural communities. Candidate “grazing genes” will be validated using quantitative polymerase chain reaction (qPCR) and/or RNA-seq and compared to traditional methods of determining grazing rate (e.g. dilution experiments).

During EXPORTS, biomass for analysis of grazing genes will be collected from both *in situ* and dilution experiments to broaden the spatial resolution of available grazing rate data. Once to twice a day, on the process and survey ships, seawater samples will be collected from 3-8 discrete depths in the euphotic zone and two depths below using a CTD rosette with mounted Niskin bottles. Water from 3 depths will coincide with those collected for the dilution method. From each depth, 5-10 L will be screened through a 200 µm mesh. Biomass will be collected by filtration, filters will be stored in RNAlater (Life Technologies) and then RNA extracted. The extracted RNA will be treated in two ways depending on the sample. For a subset of the samples, RNA-seq will be used to obtain metatranscriptomes, which will be analyzed for the expression of candidate grazing genes obtained from laboratory experiments. Expression will be benchmarked using grazing rates obtained from deckboard grazing experiments. For another subset of samples, ten of the most differentially regulated genes identified previously in lab experiments and common to a broad taxonomic range of heterotrophic predators will be chosen and their expression quantified using qPCR. Expression levels of each gene will be determined using standards (linearized plasmids with inserts) and reference housekeeping genes (aka. gene for which the expression does not change with treatment – also determined from RNA-Seq grazing experiments) Relative gene expression will be determined using the 2-ΔΔCT method (e.g. Livak and Schmittgen, 2001).

4. **Other Contributing Protocols:** Dilution method, Flow cytometry (Guava), Nutrient measurements, vertical profiling, microscopy to measure species composition and biomass,

5. **Uncertainty and quality control:** Based on previous work, we anticipate the degree of biological uncertainty in the qPCR will reach coefficients of variation on the order of 5% for biological replicates but only 0.5% variation due to replicate technical measurements.

6. **Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relative transcript abundance</td>
<td>No units</td>
</tr>
<tr>
<td>Transcript abundance</td>
<td>Transcripts cell⁻¹</td>
</tr>
</tbody>
</table>
7. Key Method references


Method: 16S rRNA gene amplicon analysis of water column bacterioplankton

Document author and contact info: Alyson Santoro, asantoro@ucsb.edu

Brief description of protocol and relation to export pathways: Seawater samples from the depths of the sediment trap deployments are collected from the Niskin rosette in 2-4L bottles and pressure filtered onto sequential 3 µm pore size polyester membrane filters and 0.2 µm pore size Supor membrane filters. DNA will be extracted from both size fractions and the V4 hyper-variable region of the 16S rRNA gene will be amplified using dual-indexed 515F and 806RB primer sets modified by Apprill et al. 2015 and Parada et al. 2018. Amplified PCR products will be sequenced on an Illumina MiSeq with a 250 bp paired end run. DNA sequence identities will be assigned by comparing to sequence databases and looking for unique sequence variants.

Other contributing protocols: Genetic characterization of bulk sinking particles in sediment traps (Durkin), Genetic characterization of sinking aggregates in sediment traps (Durkin), Genetic characterization of prokaryotes in bulk sinking particles from sediment traps (Santoro), Genetic characterization of prokaryotes in sinking aggregates collected in gel traps (Santoro)

Uncertainties and quality control concerns: While the presence of a DNA sequence confirms the link between an organism or its remains within sinking particles, the absence of a DNA sequence does not confirm the absence of this link. It is possible for organic matter to be exported without any DNA evidence of its organismal source. To relate these data to surface phytoplankton communities, it is critical that the same PCR primers are used to amplify 16S and 18S rRNA of surface plankton communities.

Data products originating with this method:
Data table of amplicon sequence variants (ASVs) detected in bulk sinking material – csv file
DNA sequences – fasta file

Key method references:

Method: Ammonium concentration in seawater using the orthophthalaldehyde (OPA) fluorescence method

Document author and contact info: Alyson Santoro; asantoro@ucsb.edu

Brief description of protocol and relation to export pathways: Dissolved ammonium will be determined in seawater samples on-ship by reaction with OPA and quantification by fluorometry. Deep water (> 1000 m) will be used as an analytical blank and to make up standards. Values will be used in support of nitrification rate measurements and to quantify regenerated nutrient supply to the euphotic zone.

Other contributing protocols: none

Uncertainties and quality control concerns: Ammonium contamination of the rosette from deck operations (e.g. cleaning, smoking), ammonium contamination introduced during sample collection (e.g. collection tubes), high blank values from inappropriate analytical blank

Data products originating with this method:

<table>
<thead>
<tr>
<th>Parameter*</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>[NH₄⁺]</td>
<td>nM</td>
</tr>
<tr>
<td>Uncertainty (standard deviation of replicate samples)</td>
<td>nM</td>
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</table>

Key method references


**Method:** Genetic characterization of prokaryotes in bulk sinking particles from sediment traps

**Document author and contact info:** Alyson Santoro, asantoro@ucsb.edu; Colleen Durkin, cdurkin@mlml.calstate.edu

**Brief description of protocol and relation to export pathways:** Each sediment trap will include 1 tube containing RNAlater preservative. Particles in the RNAlater will be collected onto an 0.2 µm pore-size Supor filter and frozen. DNA will be extracted and the V4 hyper-variable region of the 16S rRNA gene will be amplified using dual-indexed 515F and 806RB primer sets modified by Apprill et al. 2015 and Parada et al. 2018. Amplified PCR products will be sequenced on an Illumina MiSeq with a 250 bp paired end run. DNA sequence identities will be assigned by comparing to sequence databases and looking for unique sequence variants. This analysis contributes to export pathways 1, 2, and 3 by specific organisms with particle export. When combined with particle-specific DNA sequencing data, the organisms only exported in small particles can be inferred.

**Other contributing protocols:** Genetic characterization of bulk sinking particles in sediment traps (Durkin), 16S rRNA gene amplicon analysis of water column bacterioplankton (Santoro), Genetic characterization of sinking particles from the Marine Snow Catcher (Santoro)

**Uncertainties and quality control concerns:** While the presence of a DNA sequence confirms the link between an organism or its remains within sinking particles, the absence of a DNA sequence does not confirm the absence of this link. It is possible for organic matter to be exported without any DNA evidence of its organismal source. To relate these data to surface phytoplankton communities, it is critical that the same PCR primers are used to amplify 16S and 18S rRNA of surface plankton communities.

**Data products originating with this method:**
Data table of amplicon sequence variants (ASVs) detected in bulk sinking material – csv file
DNA sequences – fasta file

**Key method references:**

Method: Genetic characterization of prokaryotes in sinking aggregates collected in gel traps

Document author and contact info: Alyson Santoro, asantoro@ucsb.edu; Colleen Durkin, cdurkin@mlml.calstate.edu

Brief description of protocol and relation to export pathways: Jars containing polyacrylamide gel layers will be deployed in 1 tube of each deployed sediment trap. Sinking particles and organisms that settle into the gel layer remain distinctly separated, preserving original characteristics of size and quantity and constituents. Individual aggregates will be pipetted out of the gel layer and frozen in cryovials immediately after trap recovery. DNA will be extracted and the V4 hyper-variable region of the 16S rRNA gene will be amplified using dual-indexed 515F and 806RB primer sets modified by Apprill et al. 2015 and Parada et al. 2018. Amplified PCR products will be sequenced on an Illumina MiSeq with a 250 bp paired end run. DNA sequence identities will be assigned by comparing to sequence databases and looking for unique sequence variants.

Other contributing protocols: Genetic characterization of bulk sinking particles in sediment traps (Durkin), Genetic characterization of prokaryotes in bulk sinking particles from sediment traps (Santoro), 16S RNA gene amplicon analysis of water column bacterioplankton (Santoro), Genetic characterization of sinking particles from the Marine Snow Catcher (Santoro)

Uncertainties and quality control concerns: While the presence of a DNA sequence confirms the link between an organism or its remains within sinking particles, the absence of a DNA sequence does not confirm the absence of this link. It is possible for organic matter to be exported without any DNA evidence of its organismal source. To relate these data to surface phytoplankton communities, it is critical that the same PCR primers are used to amplify 16S and 18S rRNA of surface plankton communities.

Data products originating with this method:
Data table of amplicon sequence variants (ASVs) detected in bulk sinking material – csv file
DNA sequences – fasta file

Key method references:

**Method:** Water column nitrification rates

**Document author and contact info:** Alyson Santoro; asantoro@ucsb.edu

**Brief description of protocol and relation to export pathways:** Nitrification is the microbial oxidation of reduced nitrogen (ammonia/ammonium) to nitrite and nitrate. The nitrification rate should be related to the water column carbon respiration rate by the C:N stoichiometry of the organic matter being respired. Nitrification rates will be measured in shipboard bottle experiments by the addition $^{15}$N tracer ([$^{15}$N]H$_4$Cl) at approximately 10% of the ambient [NH$_4^+$] or 50 nM, whichever is greater. Incubations are conducted at as close to in situ light and temperature conditions as possible for 24 h. Timecourse measurements are subsampled approximately every 6 hours, frozen, and analyzed for $\delta^{15}$N$_{NO_2^-NO_3^-}$ on return to the laboratory using the 'denitrifier method.' Rates are calculated using a least-squares fitting routine to a model of $^{15}$N/$^{14}$N change in the NO$_3^-$ pool that includes removal by assimilation. Inputs to the model are the starting atom percent ($AP$) $^{15}$N in the NH$_4^+$ pool (calculated from the ambient [NH$_4^+$] and the tracer addition), the starting $AP$ in the NO$_3^-$ pool (measured directly), and the fractionation factor ($\alpha$) for NO$_3^-$ uptake (1.005).

**Other contributing protocols:** [NH$_4^+$] using the OPA-fluorescence method, 16S rRNA amplicon analysis of free-living prokaryotic community

**Uncertainties and quality control concerns:** In addition to the uncertainties associated with any deckboard bottle incubation experiment, key uncertainties in this method are the uncertainty in the [NH$_4^+$] measurement, as the calculated nitrification rate is extremely sensitive to the starting $AP$ NH$_4^+$, isotope dilution of the added tracer by newly remineralized, unlabeled NH$_4^+$, and potential stimulation of the nitrifying community by the added tracer. Uncertainty is also introduced during model fitting.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter*</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrification rate</td>
<td>nM N d$^{-1}$</td>
</tr>
<tr>
<td>Uncertainty</td>
<td>nM N d$^{-1}$</td>
</tr>
</tbody>
</table>

**Key method references**


Method: Particle-associated respiration using an in situ particle capture incubation device (RESPIRE)

Document author and contact info: Alyson Santoro; asantoro@ucsb.edu

Brief description of protocol and relation to export pathways: Microbial respiration on sinking particles will determined using an in situ particle capture incubation device known as a RESPIRE (REspiration of Sinking Particles In the subsuRface ocean) trap. The interceptor is constructed of a titanium cylinder fitted with a dimpled PVC sphere that acts as a valve to the lower portion of the chamber containing an optical oxygen sensor (Aanderaa 4330 optode). The traps operate with an initial ‘collection phase’ in which sinking particles are collected and deposited into the incubation portion of the cylinder by the rotation of the PVC sphere, which successfully excludes zooplankton from the inner chamber. Following the collection phase, the PVC sphere ceases its rotation and oxygen drawdown in the chamber is measured by the optode and recorded by a data logger. Respiration rates are then calculated from the timecourse oxygen data. Upon recovery of the RESPIRE traps, the incubated material will be collected for genetic analysis. Three RESPIRE traps will be deployed at the three shallowest depths of the surface-tethered trap (STT) array. This measurement will contribute to EXPORTS SQ2 to explain what controls the transfer efficiency of carbon in the dark ocean.

Other contributing protocols: Water column respiration (Gifford), Bulk trap fluxes (Estapa), Bacterial production (Carlson), Genetic characterization of bulk sinking particles in traps (Durkin)

Uncertainties and quality control concerns: Bias in particle capture during the particle interception phase, temperature fluctuations influencing optode readings, non-linear drawdown of oxygen during particle incubation phase (i.e. curve fitting), uncertainty in carbon conversion factor going from oxygen to carbon units

Data products originating with this method:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respiration rate</td>
<td>mmol O₂ m⁻² d⁻¹</td>
</tr>
<tr>
<td>Respiration rate</td>
<td>mmol C m⁻² d⁻¹</td>
</tr>
<tr>
<td>Uncertainty</td>
<td>mmol O₂ m⁻² d⁻¹</td>
</tr>
</tbody>
</table>

Key method references


Document History
Measurement and instrument: Imaging of phytoplankton and other particles with Imaging FlowCytobot (IFCB; McLane Research Laboratories, Inc, Falmouth, MA):

A brief description of the method: The IFCB is an imaging-in-flow cytometer. As such, it measures not only individual particle fluorescence and light scattering, but also captures a high resolution (~1 μm) image of each cell or chain in the size range ~5-150 μm width. Controlled flow and illumination conditions ensure a very high rate of images containing in focus, single targets aligned in the flow such that the largest cross-section is imaged. Images can be collected at up to ~10 Hz, depending on particle concentrations encountered. IFCB was operated with chlorophyll fluorescence and scattering triggers enabled and it was configured to sample automatically 5 ml every ~20 minutes from the uncontaminated seawater flow (diaphragm pump source, pre-debubbler to ensure minimal damage to cells). IFCB was also used to analyze discrete samples from Niskin bottles (some with chlorophyll fluorescence triggering only).

Data processing: Full resolution images are stored, though only the portion of the camera field that contains the target of interest (real-time segmentation is done during acquisition). Post-processing includes a variety of automated image processing steps, feature extraction (geometric and other quantities), and machine learning based classification (taxonomic groups and other groupings, such as detrital particles). Processing code and wiki-based documentation is available at: https://github.com/hsosik/ifcb-analysis

Calibration: Main calibration issues are (1) ensuring sample volume is properly quantified (a function design criteria set during manufacture; user verification is good practice, but experience suggests this does not need to be repeated unless there are hardware changes in the instrument); and (2) determination of image scaling (micrometers per pixel; user determined with particles of interest).

Uncertainties and quality control concerns: Sample volume imaged is sensitive to configuration / operating conditions and requires care to ensure that optical alignment and fluidic control are such that the sample core remains within the field of view of the camera. Optical conditions must also be configured to ensure focus is maintained. See procedures to assess status and troubleshoot problems in “IFCB At Sea Guide” referenced below. Additionally, the combination of trigger threshold and fluorescence sensitivity (high voltage on the detector) must be configured to ensure that the smallest particles of interest are consistently captured. Approaches with a combination of standard beads and selected phytoplankton cultures (e.g., Isochrysis, Dunaliella) work well.

Data products originating from the method: Images of phytoplankton (cells, chains, and colonies) and other fluorescent particles, including many protozoa (non-fluorescence particles possible if scattering trigger is utilized) in the size range ~5-150 μm (minimum dimension; elongated targets up to ~300 μm in length); estimates of taxonomic composition of imaged plankton, abundance and biomass (biovolume) by size and taxon.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Images of phytoplankton &amp; other fluorescent particles (~10-200 μm)</td>
<td>unitless</td>
</tr>
<tr>
<td>Biovolume (individual image target)</td>
<td>μm³</td>
</tr>
<tr>
<td>Concentration</td>
<td>Cells mL⁻¹</td>
</tr>
<tr>
<td>Biovolume concentration</td>
<td>μm³ mL⁻¹</td>
</tr>
<tr>
<td>Concentration by taxon</td>
<td>Cells mL^{-1}</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>----------------</td>
</tr>
<tr>
<td>Biovolume concentration by taxon</td>
<td>\mu m^3 mL^{-1}</td>
</tr>
<tr>
<td>Concentration by size class</td>
<td>Cells mL^{-1} \mu m^{-1}</td>
</tr>
<tr>
<td>Biovolume concentration by size class</td>
<td>\mu m^2 mL^{-1}</td>
</tr>
</tbody>
</table>

**Key method references:**


Peacock, E.E., E. T. Crockford, and H.M. Sosik. 2018. IFCB at sea user guide. [https://docs.google.com/document/d/14IfQBriV2AZs1akefM8JYirSAApnVFBdG2XO74kIOI/](https://docs.google.com/document/d/14IfQBriV2AZs1akefM8JYirSAApnVFBdG2XO74kIOI/)
**Method:** Reconstruction of Zooplankton Export from Gel and particle ID Traps

**Document author and contact info:** Deborah Steinberg, debbies@vims.edu

**Brief description of protocol and relation to export pathways:**
Sinking of zooplankton fecal pellets and other products (e.g., mucous feeding webs, carcasses, molts) is one of the five export pathways. We complement the gel trap analyses by Durkin et al. by analyzing in further detail zooplankton fecal pellets and other products in gel traps and in splits from bulk sediment trap samples to determine the relative contribution of different major zooplankton taxa to export. Identified fecal pellets from gel traps, and/or pellets in bulk sediment trap samples, are further classified into fecal pellet type (e.g., cylindrical crustacean pellets, tabular salp pellets, elliptical larvacean pellets) using archived/binned fecal pellet and other particle images (gel traps) and dissecting microscope (bulk trap splits) for each sediment trap depth. The relative contribution of sinking zooplankton fecal pellets and other products vs. other particle classes for each depth is determined using fecal pellet counts and volume to C and N conversions (from the literature, and experimentally-derived from shipboard fecal pellet production experiments with abundant species). The fecal pellet POC/N is subtracted from the total sinking POC/N to determine the relative importance of fecal pellet flux to total flux. Changes with depth in fecal pellet types is analyzed, as an indicator of mesopelagic zone reprocessing.

**Other contributing protocols:** Visual characterization of zooplankton products in gel traps, Visual characterization of aggregates in gel traps, Zooplankton fecal pellet production, MOCNESS abundance to compare zooplankton community structure to zooplankton products in traps

**Uncertainties and quality control concerns:** Uncertainty inherent in particle C to volume conversions. Distinguishing between crustacean molts and carcasses (dead zooplankton that fell into trap), and between carcasses and swimmers (live zooplankton that swam into trap and died), is difficult in trap samples and only possible for some zooplankton species.

**Data products originating with this method:**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fecal pellet fluxes by pellet type* at each depth</td>
<td>number or mg-C m⁻² d⁻¹</td>
</tr>
<tr>
<td>Larvacean house flux at each depth</td>
<td>number or mg-C m⁻² d⁻¹</td>
</tr>
<tr>
<td>Crustacean molt flux at each depth**</td>
<td>number or mg-C m⁻² d⁻¹</td>
</tr>
<tr>
<td>Carcass flux at each depth**</td>
<td>number or mg-C m⁻² d⁻¹</td>
</tr>
</tbody>
</table>

* To be reported separately for each dominant pellet type
**See Uncertainties (above)

**Key method references**


Method: Zooplankton fecal pellet production

Document author and contact info: Deborah Steinberg, debbies@vims.edu

Brief description of protocol and relation to export pathways:
Sinking of zooplankton fecal pellets is one of the five export pathways. We will perform live fecal pellet production experiments on dominant species and whole community size fractions to provide measurements of zooplankton contribution to fecal pellet production during each ecosystem state sampled. These will be scaled to community level export using the biomass measures, as well as species identification and size fractionated biomass made with the MOCNESS.

Animals for live experiments are collected within the epipelagic zone from below the chlorophyll max to the surface, using a 1-m diameter ring net with a non-filtering cod end and slow retrieval rate. Experiments are performed on-board at in situ mixed layer temperature, during night and day, in unfiltered surface seawater using both a mixed size-fractionated zooplankton community (Butler & Dam 1994, Urban-Rich et al. 1999) and numerically important taxa (Urban-Rich et al. 1999, 2001; Wexels-Riser et al. 2001).

Fecal pellet production by each size fraction (0.2-0.5, 0.5-1, 1-2, 2-5, and >5 mm) of the mixed zooplankton community is measured using sets of two large (~3.8 liter) containers with a screen insert in the bottom of the inner container (Butler & Dam 1994). The screen allows pellets to be collected in the outer container, but keeps animals separated from their pellets to prevent pellet consumption. Mixed animals from the tow are size-fractionated live and placed in ambient water in the nested containers and incubated for 4-6 hours. The screen-bottom container is lifted (removing animals) and animals are saved for enumeration and biomass measurements. Fecal pellets collected in the outer container are rinsed, counted, and concentrated onto combusted GF/F filters for POC/PON analysis. For pellet production by abundant species from representative size fractions, a suspended cylindrical insert containing animals is capped at both ends with 200 μm mesh (or larger mesh for larger size classes) and placed into 1-liter experimental bottles containing surface seawater and incubated for 4-6 hours. At the end of the experiment the insert containing animals is removed, and water in the outer jar poured through a 30 μm sieve to collect fecal pellets. Animals and pellets are processed as above. Fecal pellet POC/PON production rates are combined with zooplankton weight measurements from each experimental incubator/bottle to calculate weight-specific fecal pellet POC/PON production rates for the community and within size fractions. Occasionally, fecal pellets of abundant single species of interest (e.g., Neocalanus spp., Salpa fusiformis) are collected in separate ‘bulk’ incubations, to calculate their POC/PON content and better determine their contribution to export.

Other contributing protocols: MOCNESS abundance and biomass sampling to scale individual fecal pellet production experiments to community export.

Uncertainties and quality control concerns: Physiological effect of net capture and incubation. When scaling – the effects of zooplankton patchiness and variation in species physiology.

Data products originating with this method:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fecal pellet production</td>
<td>mg-C (mg body C or dry weight)$^{-1}$ d$^{-1}$</td>
</tr>
<tr>
<td>zooplankton$^{-1}$</td>
<td></td>
</tr>
</tbody>
</table>
Fecal pellet production
zooplankton$^{-1}$

$\text{mg-N (mg body N or dry weight)}^{-1} \text{d}^{-1}$

Zooplankton community
fecal pellet production

$\text{mg-C or mg-N m}^{-3} \text{d}^{-1}$

Zooplankton community
fecal pellet production in
euphotic zone

$\text{mg-C or mg-N m}^{-2} \text{d}^{-1}$

* Each to be reported separately for each dominant species

SeaBASS submission fields and units:

$\text{/fields=Sample,sample_1id,date_start,time_start,date_end,time_end,elapsed_time,bin_diameter_lower,bin}$
$\text{diameter_upper,lightlevel,fecalpellet_production_carbon_dryweight,fecalpellet_production_carbon_carbon}$
$\text{onweight,namespace_manual,identification_manual,biotic_group}$

$\text{/units=none,none,yyyymmdd,hh:mm:ss,yyyymmdd,hh:mm:ss,seconds,um,um,\%,mg/mg/hr,mg/mg/hr,none}$

$\text{none,none}$

Key method references

Butler M, Dam HG (1994) Production rates and characteristics of fecal pellets of the copepod
Acartia tonsa under simulated phytoplankton bloom conditions: Implications for vertical
fluxes. Marine Ecology Progress Series 114, 81-91

fecal pellets to the carbon flux on Nordvestbanken, north Norwegian shelf in 1994.
Sarsia 84, 253-264.

Urban-Rich J (2001) Seston effects on faecal pellet carbon concentrations from a mixed
community of copepods in Balsfjord, Norway, and the Antarctic Polar Front. ICES
Journal of Marine Science 58, 700-710.

zooplankton faecal pellets on and off the Iberian shelf, north-west Spain. Progress in
Oceanography 51, 423-441.
Physical transport models  
Protocol document  
Andy Thompson, andrewt@caltech.edu  
Last updated: October 8, 2017

This document describes protocols for archiving any “physical” transport model output for which archiving is required for the EXPORTS project. This output is likely to be a combination of numerical forecast model output in the region corresponding to the EXPORTS field program as well as any idealized model output that is generated as part of the NASA-funded pre-EXPORTS data mining and process-study activities. Most model output will be used primarily to consider the advective (submesoscale) component of the EXPORTS wiring diagram (Pathway 4, Mixing of DOC and particles). However, some of the pre-EXPORTS process studies are also considering the relative importance of sinking and advection. The format of the model output is unlikely to follow a single protocol because of the diversity of the models that may be utilized. However, this document suggests some uniformity in terms of how the output fields should be saved.

Other contributing protocols: Forcing fields (wind stress, air sea fluxes); hydrography; dissolved oxygen, chlorophyll, optical backscatter. In reality, any of the data streams may be useful for future modeling efforts. Observations collected from the autonomous platforms will be of particular interest because they will provide information on states over a longer time period.

Uncertainties and quality control: There are no uncertainties or quality control issues with regard to the model output. However, documentation of the model runs will be essential. This may include information about spatial and temporal model resolutions, the type and location of forcing fields applied, documentation of specific parameterizations used for turbulent mixing and other subgrid-scale processes. In some cases where idealized process models are archived, the input files needed to run the model (for example using a standard model, such as the MITgcm) may be archived rather than all the output fields. It will be critical that meta-data files are submitted along with any archived model output or forcing/input files.

Data products originating from this method: Model output from simulations that were conducted outside of the EXPORTS project is likely to be archived elsewhere; clear documentation of where these files can be found will be sufficient. Archiving input files so that process-based models can be re-run will be critical because it is unlikely that we know a priori all the fields that will be useful for future analysis. Similarly, it is difficult to dictate a standard format for the output in terms of averaging duration (e.g. snapshots, daily/weekly/monthly averages); these differ for various parameters, such as eddy kinetic energy or various tracer fluxes. Standard output will be temperature, salinity, current velocities, tracer/particle concentrations; derived quantities will likely need to be calculated from other model output or re-running the simulation.
Observation-based vertical advection and mixing of organic carbon to depth by physical oceanographic processes (Pathway 5)

Andy Thompson & Melissa Omand; andrewt@caltech.edu, momand@uri.edu
Last updated: November 12, 2017

This document describes protocols for deriving the physical transport of particulate and dissolved organic carbon between the surface mixed layer and the seasonal thermocline. We nominally divide this pathway into three: 1) meso- and submesoscale subduction (the eddy-driven pump), 2) vertical mixing and seasonal restratification (the mixed-layer pump), and 3) diapycnal turbulent mixing.

1) The eddy-driven pump of organic carbon: Physical transport of organic carbon may occur either along sloping isopycnals that evolve rapidly in space and time or via a diapycnal flux across density surfaces. A net downward flux occurs when there is a correlation between the vertical velocity anomaly \( w' \) and the anomaly in organic carbon \( C' \), such that the mean vertical flux \( \langle w' C' \rangle \) is negative. We assume that the fluxes will not be resolved directly from observations due to insufficient spatial and temporal coverage. Therefore, initial fluxes will be derived from existing estimates/parameterizations of the vertical velocity in terms of observed properties. These techniques include the omega equation for the mesoscale velocity (Holton 2004, Vallis 2017) and mixed layer baroclinic instability (Fox-Kemper et al. 2008, Omand et al. 2015) and other more recent parameterizations (Bachman et al. 2017) for estimating \( w' \) at submesoscales. We recognize that our understanding of submesoscale dynamics is changing rapidly, so parameterizations may be superseded over the course of the EXPORTS project. Particulate organic carbon (POC) concentration, anomalies and gradients will be estimated from optical sensors on the autonomous platforms and ships (see POC proxies protocol). Dissolved organic carbon concentrations will be estimated from ship-based bottle measurements. We expect significant differences in the vertical and lateral stratification between the Atlantic and Pacific sites as well as in the temporal correlation between the seasonal cycle of POC and submesoscale velocities. In addition, we will leverage the on-going pre-EXPORTS modeling activity, led by Mahadevan with Thompson, Nicholson and Omand as co-PIs, to run numerical simulations based on the observed conditions, thereby validating or correcting the parameterized flux estimates. Our measurements of physical and ecological data on the same space and time scales, combined with models will address: SQ1d How do physical and ecological processes act together to export organic matter from the surface ocean? SQ2a (submesoscale component): How does transfer efficiency through the mesopelagic vary?

2) The mixed layer pump of organic carbon: Shoaling of the mixed layer caused by large-scale, 1-D surface heat fluxes can also cause organic carbon to be 'left behind' in the seasonal thermocline. This mechanism is different from eddy-driven subduction, which requires 3-D dynamics. It may be difficult to disentangle these two contributions because both may be enhanced at similar times, and lead to a shoaling of the mixed layer. For example during NAB08, eddy-driven subduction of POC was enhanced during spring-time restratification. Shortly after the onset of eddy restratification, the surface heat flux changed sign, continuing to stratify the upper ocean and likely enhancing the retention of subducted carbon below the mixed layer. Protocols to independently quantify the role of the mixed layer...
versus eddy-driven pump are not yet established, and will likely be part of the efforts of the autonomous team and others.

3) **Diapycnal mixing of organic carbon:** Turbulent mixing across isopycnals can also result in a vertical flux of organic carbon. If possible, the ADCPs on the Lagrangian float and glider will be used to quantify the turbulent vertical diffusivity ($\kappa_{di}$) as a function of depth and time. If this measurement is made, then the diapycnal turbulent flux can be parameterized from the local gradient in organic carbon (C) measured from the autonomous platforms and the ship according to

$$C Flux_{di} = -\kappa_{di} \frac{dC}{dz}.$$

**Other contributing protocols:** Upper ocean hydrography (temperature, salinity, derived quantities including density, mixed layer depth, potential vorticity); Surface forcing (wind stress, buoyancy fluxes); POC concentrations from optical sensors and any direct or derived calculations of turbulent dissipation.

**Uncertainties and quality control:** Concerns here are more likely to be with regard to spatial/temporal resolution and coverage as opposed to uncertainty in the actual measurements. It is important to note that parameterizations of submesoscale vertical velocities require a scale separation between the larger-scale hydrographic properties (e.g. lateral buoyancy gradients) and the scale of the submesoscale eddies. Depending on the scales that are resolved, the eddy-driven portion of the flux can be approached as a diffusive parameterization in a similar manner as the diapycnal mixing flux above

$$w'C' = -\kappa \frac{dC}{dz}.$$

Here the vertical diffusivity $\kappa$ encompasses the along-isopycnal and diapycnal mixing within some region.

**Data products originating from this method:** The goal is to generate a time series of the vertical POC flux at the center of the Lagrangian autonomous array for the duration of the deployment of this array. The temporal resolution of this time series can be re-visited, but we expect to be able to estimate the flux at least daily.

**Key method references:**
**Method:** Lipidomics of suspended and sinking particles

**Document author and contact information:** Ben Van Mooy (bvanmooy@whoi.edu)

**Brief description of protocol:** This protocol describes the basic steps for collecting particle samples at sea for subsequent lipidomic analysis in the laboratory ashore. For suspended particles, one-liter seawater samples are collected directly from Niskin bottles into round polycarbonate bottles after first rinsing the bottle 3x with 100 mL of sample seawater. Personnel conducting the sampling wear nitrile gloves since oils and detergents (e.g. fingerprints and hand soap residue) are the primary contamination risks. The samples are then filtered as quickly as possible; lipidomes are highly dynamic and can change drastically within an hour in response to changes in light and/or temperature. A custom vacuum filtration rig with glass filter supports/funnels accommodates up to 12 samples at once. Filters are 47 mm diameter 0.2 µm poresize Durapores. Vacuum in the rig is set to 200 mbar, and then the sample bottles are gently inverted into the funnels and held in place by the rig. These samples may then be ignored until the last 100 mL or so is filtered, which takes about 30 minutes. As soon as the sample is filtered (i.e. just as it “runs dry”), the filter is folded in half, and then placed in a foil envelope. The sample ID is written on the foil with a sharpie, and the sample is immediately immersed in liquid nitrogen, where it should reside for the duration of the cruise. A dry shipper as used as the liquid nitrogen vessel; at the dock prior to shipping the liquid nitrogen is simply dumped, leaving the samples in the shipper ready for transport. The protocol for sinking particles is essentially identical, except that the volume of the samples is dictated by the volume of the splits from the sediment traps.

**General sample requests:** Lipidomics is a powerful tool with the potentials for revealing both the sources of exported particles and the processes that control the magnitude of particle export flux. Key to realizing these potentials is integrating lipidomic data with other measures of microbes and microbial processes. Thus, we request that process-ship seawater samples for lipidomics of suspended particles be taken from the same depths/times as samples for the following other parameters (in rough order of preference): genomics, HPLC pigments, FCM, POC, nutrients. In addition, MOCNESS and MSC samples are also welcome (zooplankton and particles may simply be preserved on filters, as described above). Samples of sinking particles from sediment traps are also requested; surface moored PITs are OK. Finally, survey-ship samples of suspended particles from only the surface (i.e. mixed layer or satellite optical depth) are also highly sought (but we recognize this may be asking too much).

**Other contributing protocols:** genomics, HPLC pigments, FCM, POC, nutrients.

**Uncertainties and quality control concerns:** Usually, we take triplicate samples. Recognizing that this is not a possibility with EXPORTS, absolute concentrations of lipids will carry an assumed uncertainty (standard deviation) of 20%. Relative proportions of lipid (i.e. within sample %) will carry an assumed uncertainty of 10%. Perhaps triplicate samples could be obtained on a few occasions during the cruise if the opportunities present themselves? There are two major quality concerns: 1) contamination from soaps, detergents, oils, and greases, and 2) sample preservation (if the samples thaw, its over).

**Data products originating with this method:** Concentrations (mol L⁻¹) of approximately 1,000 lipid molecules, including: triacylglycerols, intact polar diacylglycerols, intact polar monoacylglycerols, chloropigments, carotenoids, diacylglycerols, monoacylglycerols, fatty acids, wax esters, and sterol esters.
Method: Measurements of $P_{11}$ (the volume scattering function, VSF), $P_{12}$ and $P_{22}$ elements of the scattering Mueller matrix at 517 nm from 0.1 - 155° (nominal) in water.

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Brief description of protocol: The small angle part of VSFs is measured using ring detectors, identical to the ones used in LISST-200X instrument. The range of angels covered by the rings is 0.088 to 14.8°. For large angles, the VSF and polarized components are measured using a rotating eyeball. The $P_{11}$, $P_{12}$ and $P_{22}$ are obtained for the particulates defined as the difference between the measurements of bulk and the 0.2 µm filtered seawater (serve as the background).

Deployment methodologies:
Discreet water samples will be used. At each station, the first measurement is to establish the background using 0.2 µm filtered water collected from the deepest depth. It takes about 10 – 15 min to finish this step. Twenty VSF measurements will be collected at each depth, and mean and standard deviation will be estimated.

Derived parameters:
Attenuation coefficient (m$^{-1}$, acceptance angle = 0.057°)
Volume scattering function at 515 nm [sr$^{-1}$ m$^{-1}$]. $P_{12}/P_{11}$ and $P_{22}/P_{11}$
Scattering coefficient by integrating VSF from 0.088 to 150° (m$^{-1}$)
Particle size distribution inferred from the VSF (m$^{-3}$ µm$^{-1}$)

SeaBASS fields and units:
/fields=VSF_###ang,PSD_DNSD_###umsize
/units=sr$^{-1}$ m$^{-1}$,Particles m$^{-3}$ µm$^{-1}$

Uncertainties and quality control concerns:
Instrument: The instrument performance will be validated/calibrated pre- and post- cruise in laboratory using standard beads. If necessary, mid-cruise validation will be conducted on the vessel. Measurement: Uncertainty in VSFs are computed from variability in 20 repeated VSF measurements at each depth.

Key method references
Measurement and instrument: Characterization of size distribution of nanoparticles with a Manta ViewSizer.

A brief description of the method: The ViewSizer tracks Brownian motion of nanoparticles and estimate the size of each particle, which is inversely proportional to the standard deviation of the particle’s random displacement, a theory first developed by Einstein. The methods for measuring PSD of particles generally include two main categories: (1) methods that measure a large number of particles simultaneously (ensemble methods), and (2) methods that measure particles individually (individual-particle methods). The nanoparticle tracking analysis deployed by ViewSizer belongs to the second method, whereas the MVSM method (summarized in another document) belongs to the first method. ViewSizer deploys three laser sources (447 nm, 532 nm and 655 nm) and record video of scattered light from wide-ranging sizes of individual particles simultaneously. Subsequent image analysis tracks the motion of each particle and estimates variation of the particles displacement under Brownian motion.

Data processing: The sample volume is 1.5 mL. A short video typically at a speed of 40 fps (frame per second) is recorded. Through test, we found 200 videos are needed to derive statistically valid estimates for oceanic samples. Between the consecutive video recording, the sample cuvette is automatically shaked. An internal software analyzes each of recorded videos and estimates the sizes of particles found within each video. The size estimated represents the dynamic diameter, of which a sphere would go through the same Brownian motion. The number of size bins and the width of each size bin can be decided by a user. We use 45 size bins, with the smallest bin from 1 nm to 8 nm and a center size of 4.5 nm and the largest bin from 1255 nm to 1300 nm and a center size of 1277.5 nm. The statistical analysis of particle size distribution can be repeated later, for instance, with a different bin setting.

Calibration: The instrument is validated using standard microbeads.

Uncertainties and quality control concerns: To eliminate potential interference from larger particles, which tend to settle instead of going Brownian motion, the samples should be filtered, say, using 0.7 µm filter. Based on our own lab test using standard beads of various sizes from 25 nm to 900 nm, we found consistent results for beads of sizes 100 nm to 700 nm. For smaller beads, the uncertainty is significant because, according to the manufacturer Manta, the sample volume is no longer 1 mL, instead, it should increase with decreasing size of small particles. We’re waiting for the company provide the volume correction for particles of sizes < 100 nm. But until then, only the size distributions between 100 and 700 nm should be used.

Data products originating from the method: videos of particles, particle counts within each video, and particle size distributions from 4.5 nm to 1277.5 nm (the size range from 100 – 700 nm should be used).

Key method references: