Sampling Protocols

Version 4.2

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1 Motivation

The collection of detailed sampling protocols is a crucial tool for the success of the Nansen Legacy, because they ensure:

- Methodological agreement between the involved researchers
- Continuity and comparable data throughout the 5 year sampling period
- An easily accessible overview over parameters sampled
- Easier cruise planning

Please forward any request/feedback/input/update for the sampling protocol to Miriam Marquardt (UiT, miriam.marquardt@uit.no).

2 Labelling of samples and data from cruises within the Nansen Legacy project

Method responsible: Pål Ellingsen (UNIS, data.nleg@unis.no), TBA

To secure a FAIR (Findable, Accessible, Interoperable, Reusable) data management within the project, a first step is to ensure that the collected samples are findable and that relevant metadata are logged along with the sample collection. The metadata need to be logged in a standardized manner and will be made accessible through a search interface as soon as possible after the cruises.

To help cruise and project participants keep track of their samples, a system where each sample is given a unique ID (a UUID; Universally Unique ID) has been developed. The UUID will follow each sample and is printed as a Data Matrix code on sticker labels which are placed on the sampling containers. The value of the Data Matrix (see Fig. 2-1) is read by hand-held scanners or cellphones and automatically written into an Excel sheet along with sample metadata. This allows for easy tracking of the samples using the Data Matrix.

Sampling within the project is to a large degree hierarchical, where we use a sampling gear (for instance a MIK net) which is subsampled to measure different parameters (community composition, sample biomass, single species collection etc). It is essential to be able to trace the individuals back to the sampling gear, station and cruise, and the UUID system used facilitates this through a hierarchical parent/child ID. The sampling gear is given a UUID which is used to trace all
subsamples taken from that one gear deployment. A subsample taken from the gear is given another UUID, but at the same time it is referred back to the parent UUID. Any sample taken from the subsample is given another UUID, and refers back to its parent (that is, the subsample UUID) as described in Fig. 2-1. Thus, we only need to keep track of two hierarchical levels of UUIDs for every sample but will be able to trace all samples back to sampling gear and the relevant metadata (station ID, cruise ID etc.) through the family tree.

**Fig. 2-1** The hierarchical system with sample/subsample IDs used to trace samples and data within the project.
2.1 STANDARDIZED METADATA

The metadata (and later on the data) needs to be logged according to accepted standards using a controlled vocabulary to allow for compatibility with the databases where our metadata and data will be made findable and accessible. We will use the Darwin Core standard wherever it is available for the biological/chemical/palaeontological data, and the NetCDF/CF standard for physical data. To build up an excel sheet where the standardized formats are used for the required metadata information, an excel template generator has been made available through the SIOS webpage:

https://sios-svalbard.org/cgi-bin/darwinsheet/index.cgi

During the 2018 cruises with “Kronprins Haakon” it will also be available on the ships internal network:

http://10.3.65.20/cgi-bin/darwinsheet/index.cgi

The required fields needed to log for every sample and subsample collected within the project is already selected in the excel template generator, and extra fields that may be relevant for different samples and where a controlled vocabulary already exists can be added (which is recommended to do). The required fields include a minimum of metadata information necessary, and most of this information can be taken directly from the screens/cruise logger onboard KPH. The excel sheet is set up to control the content of the cells, and thus it is not possible to write information in a wrong format (e.g. the date has to be written as YYYY-MM-DD). The generated excel sheet can be added additional fields which individual researchers want to log for their own samples. All the parameter definitions are explained in the excel template generator (when the mouse is held over the term), and there is also an explanation for the different parameters in the individual cells of the generated excel sheets.

2.2 THE UUID GENERATOR

In some cases, it is necessary to generate a UUID for a sample or a subsample which is not kept. This could for example be the sampling gear UUID in cases where all samples collected from the sampling gear represent unique subsamples and are labelled with their subsample ID. In such cases, UUIDs can be generated through the UUID generator:

KPH internal network:  http://10.3.65.20/cgi-bin/darwinsheet/uuid.cgi
SIOS portal:  https://sios-svalbard.org/cgi-bin/darwinsheet/uuid.cgi.

The UUID is copied into the excel sheet, but there is no labelled sample with the UUID.
Printing of sample labels
Sample labels can be printed in different sizes and label qualities depending on sample type. All labels must include the UUID which is printed as a Data Matrix using label printers. The label printers can be accessed here:  
http://10.3.65.20/cgi-bin/darwinsheet/print.cgi (via the KPH internal network). We recommend that sampling containers are also labelled according to individual researchers’ standard procedures, so that it is possible to identify them also without scanning the Data Matrix.

- **Small labels** – the smallest sized labels include only the Data Matrix on small stickers that can be added to e.g. Eppendorf lids. These labels must be put on a flat surface to allow the Data Matrix to be scanned. These are most relevant to use with pre-labelled small tubes, in particular in cases where many samples are collected from the same sampling gear/subsample.

- **Medium-size labels** – these labels include the possibility of adding four lines of written information to ease the identification of the samples without scanning the Data Matrix.

- **Large labels** – these labels are used for labelling of larger samples.

Reading of sample labels
Sample labels can be read with barcode scanners and mobile apps, as well as webcamera. At UNIS and UiT there is an opticon scanner available:

- You can change settings and see the reads of the scanner here: http://opticonfigure.opticon.com/
Mobile apps you can download:

And for the webcam:

Further reading:
The Nansen Legacy labelling protocol is ready for download on Yammer:
https://www.yammer.com/thenansenlegacy/ -
/uploaded_files/180071219?threadId=106708986347520
3 Transects, process stations, and mooring sites

The Nansen LEGACY transect and stations are defined below. It consists of 25 stations (of which 7 are defined as process study stations) covering the northern Barents Sea from south of the Polar Front to the Arctic Ocean (Fig. 3.1-1). Which, and how many of the stations that will be conducted on each Nansen LEGACY survey will vary among the surveys depending on ship time, sea ice conditions and focus of the survey.

Location of the process study stations are motivated by 1) they should cover both Arctic and Atlantic conditions for comparison, 2) some of them should have sediment conditions suitable for geological coring, and 3) some of them should have moorings to get data throughout the year (for seasonality).

Location of the main RF1 moorings (M1-M4) are motivated by the need of measuring the flow of Atlantic Water into the northern Barents Sea. These moorings will be deployed in 2018 and will be extensively equipped with instrumentation measuring the properties of the inflowing Atlantic Water as well as the rest of the water column. The process study stations to study the Arctic species must be placed at locations dominated by Arctic conditions. Moorings will be deployed at some of the locations in 2019 to obtain year-round time series at the Arctic process study sites.

3.1 BRIEF DESCRIPTION OF THE PROCESS STUDY STATIONS

P1 – located in Atlantic Water south of the Polar Front. P1 is suitable for comparing the Atlantic Water dominated, southern Barents Sea with the Arctic Water, dominated northern Barents Sea. P1 is likely to be free of sea ice during most of the winter. Being in a deeper trench, P1 is expected to have suitable sediments for geological coring.

P2 – located at the central/northern part of Storbanken in a region which usually have Arctic conditions in the intermediate part of the water column. P2 is likely to have varying sea ice coverage during winter, and a moderate surface layer of meltwater during summer.

P3 – located in the trench north of Storbanken. P3 will likely be characterized of Arctic conditions in the upper part water column and a deep northeastward flow of Atlantic Water (from the branch entering the southern Barents Sea) in the lower part of the water column. P3 is likely to have varying sea ice concentration during winter, and a surface layer of meltwater during summer. Being in a deeper trench, P3 is expected to have suitable sediments for geological coring.

P4 – located in the deeper trench south of Kvitøya. P4 will likely be characterized of Arctic conditions in the upper part water column and a deep southwestward
flow of Atlantic Water (from the branch going around Svalbard) in the lower part of the water column. It is expected that P4 will be covered with sea ice most of the winter, and by a surface layer of meltwater during summer. Being in a deeper trench, P4 is expected to have suitable sediments for geological coring.

**P5** – located on the shallow shelf northeast of Kvitøya. P5 is likely to have Arctic conditions below the seasonal varying surface layer. It is expected that P5 will be covered with sea ice most of the winter, and by a surface layer of meltwater during summer.

*Fig. 3.1-1* Map of the Nansen LEGACY process study stations (P1-7) and transect (yellow squares) and mooring locations (red dots, M1-4).
P6 – located at the shelf-break slope (at about 850m) towards the Arctic Ocean. P6 is situated in the core of the Atlantic Water flow in the shelf-break current. It is expected that P6 will have a varying sea ice cover during winter, and a varying surface layer of meltwater during summer. ATWAIN-moorings (not shown in map) can provide year-long measurements for P6. P6 might be suitable for geological coring.

P7 – Located in the deeper Arctic Ocean. P7 is suitable for comparing the shallow Barents Sea with the deeper Arctic Ocean. P7 will have extensive sea ice coverage during winter, and varying sea ice cover during summer. P7 might be suitable for geological coring.

Fig. 3.1-2 Temperature (color) and salinity (black solid lines) distribution in the Nansen LEGACY transect sampled in August 2012. Note different scales on the horizontal axes. The location of the process study stations (P1-7) is shown with dashed vertical lines.
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<th>Latitude (decimal)</th>
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Vardø-N, Hopendjupet
3.2 STANDARIZED SAMPLING DEPTHS DURING THE NANSEN LEGACY CRUISES

All sampling should take place at the assigned standardized sampling depths:

- 5, 10, 20 m
- DCM (Deep Chla max: only if present and significantly different from standard depths, i.e. ± 5m)
- 30, 40, 50, 60, 90, 120, 150/200, Bottom-10
- In addition to the following at the deep stations: 500, 1000, 2000, 3000, 4000 m

4 Bridge-based observations

4.1 ICE OBSERVATIONS

http://www.climate-cryosphere.org/media-gallery/881-icewatch-assist

Photos can be uploaded in the software directly.
Fig. 4.1-1 Introduction to commonly used ice observations by http://www.climate-cryosphere.org/media-gallery/881-icewatch-assist

5 Underway sensors/measurements

5.1 SURFACE WATER PARTIAL PRESSURE OF CO2 (PCO2)

**Measured parameter:** mole fraction of carbon dioxide (xCO₂), partial pressure of CO₂ (pCO₂, μatm) and fugacity of CO₂ (fCO₂ (μatm))

**Method responsible:** Agneta Fransson (NPI, agneta.fransson@npolar.no) and Melissa Chierici (IMR, Melissa.chierici@hi.no)

**Description of parameter:** Partial pressure of CO₂ is the dissolved CO₂ in seawater and atmosphere.
**Method description:**

Surface water xCO2 (or pCO2, fCO2) is continuously measured from the ships’ seawater intake at 4 meter using the instrumentation of General Oceanics® (showerhead equilibration and dispersed infrared detection). Discrete water samples are collected regularly to be analysed regarding DIC and AT (see section 8.2) for quality control of the instrument data. Measurements of atmospheric pCO2 are performed during a 24-hour cycle. A set of four standard gases of different CO2 concentrations (0, 350, 450, 600 µatm) are analysed in the 24-hour cycle and used for calibration of the measurements.

Seawater salinity and temperature are measured by a TSG Seabird sensor, and will be logged by the pCO2 instrument and used in the calculations of pCO2.

---

**5.2 SURFACE WATER DISSOLVED OXYGEN**

**Measured parameter:** dissolved oxygen (DO, O2 µmol/kg),

**Method responsible:** Agneta Fransson (NPI, agneta.fransson@npolar.no) and Melissa Chierici (IMR, melissa.chierici@hi.no)

**Description of parameter:** Dissolved oxygen is the oxygen gas dissolved in seawater.

**Method description:**

Dissolved oxygen is continuously measured from the ships’ seawater intake at 4 meter using an Optode from Aanderaa®. Oxygen concentration, saturation and water temperature are logged in the pCO2 data log.
6 Water column structure

6.1 CTD

General requirements:

As of 26 July 2018, the KPH CTD is equipped with double set of temperature and conductivity sensors, pressure sensor, oxygen and fluorescence sensors, and an altimeter. To be installed are a turbidity and a PAR sensor.

Annual calibration of all CTD sensors (in particular T, S, and O2) should be documented through calibration sheets available on request. Samples for calibration of the conductivity cell(s) should be taken regularly during a research cruise covering the observed salinity range. This might require several samples per CTD cast.

Setup of the CTD and postprocessing follows standard IMR routines. Raw data (ie. .hex-files and configuration files) should be available for own postprocessing e.g. in case conversion to other units or different depth bins is required, or LADCP postprocessing requires lat&lon information throughout the cast.

If LADCPs are mounted on the rosette, GPS position from the NMEA feed should be recorded throughout the cast: in Seasave, choose the setting to append lat&lon to every scan. As of 26 July, this setting was in place.

Metadata of the cast should be recorded in the CTD file and on the CTD sampling sheet.

Information to be logged:

For each cast, information to be noted down:

- date & time (UTC)
- position at start, bottom, and end
- bottom depth (in meter)
- operator
- ship
• sensor type incl. serial number for each component
• software used e.g. for SBE instruments incl version number
• meteorological conditions (barometric air pressure at sea level, wind speed & direction (degrees), air temperature (deg C), cloud cover (in 10th)

In case of extensive water sampling of CTD casts, a sampling log sheet should be used for overview of type of water sample taken and sample number (see Appendix for template).

Deployment routines:

• Start data acquisition just before the CTD leaves the deck (this will help with assessing surface pressure adjustment).
• Soak for 1 minute at 5 m (10 m if very cold and there is a risk of freezing during deployment or on deck), bring back up to surface (i.e. just submerged, rosette/CTD frame not breaking through the sea surface), leave there for 3 minutes.
• Lower CTD to max depth. Rest at max depth for at least 1 minute (preferably longer).
• Lowering speed: between 0.8 and 1 m/s. If LADCP is mounted, 0.7 m/s. Take care not to hit the bottom with the CTD but stop 5-10 m above ground, depending on conditions (steepness of topography, drift speed of the ship). Use the altimeter on the CTD to assess distance to bottom.
• Depending on capacity for measuring salinity samples onboard, water samples should be taken at various depths throughout the CTD cast. Suggested minimum number of samples is 3 – one at bottom/max depth, one in the middle of the water column in a region without strong salinity gradient, and one in the surface mixed layer. More samples will improve accuracy of the calibration. Depth should be chosen to cover the range of salinities and pressure throughout the cast/cruise.
• If water samples for other variables are taken at standard depths (e.g. following protocols from the Nansen Legacy project, salinity samples can be taken at some of those depths.
• If salinity samples are measured throughout the cruise, at least 5 samples should be taken from each cast.
• Close Niskin bottles on upcast: stop CTD at sampling depth (do NOT fire "on the fly"!), wait at least 1 minute, preferably 2 minutes, then fire Niskin bottle. Flushing time (i.e. wait before closing the bottle) should be longer in calm conditions and for big rosette systems or Niskin bottles with small openings.
• Fill in depths of the bottles on the log sheet and send the log sheet to the CTD sampling crew as soon as possible.
• Stop data acquisition when the CTD is back on deck.

CTD log sheet template

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6.2 NISKIN ROSETTE

If many different groups sample water from the rosette, use the CTD log sheet for overview of Niskin bottle depths, sampling order, and sample numbers.

**Taking water samples for salinity calibration:**

- Use appropriate sample bottle! With plastic insert.
- Rinse bottle and bottle cap (but not insert) three times. Dry off bottle neck (in- and outside) and cap with paper towel before closing the bottle. Pay special attention to the thread on the bottle and in the cap.
- Note down on the sampling log cast number/station number, Niskin bottle number, sample bottle number.
- IMR have a registration system for salinity samples, make sure the information is entered there.

**For the winch drivers on deck:**

- Check that wire length corresponds with CTD depth – give information about depths as necessary if there is a difference. As of 26 July, the wire length was increasingly too short, indicating a need for calibration on the measuring wheel.
- When bringing the CTD back on deck in CTD hangar, the CTD needs to be moved further in than the winch does automatically, otherwise there is not enough space around the rosette for sampling.

6.3 ADCP

**General requirements:** There are four ADCPs mounted: two 38 kHz and two 150 kHz RDI Ocean Surveyors. One of each are located on a drop keel, the other ones are in the hull (aka Arctic window aka flush mounted). There are two PCs to control them, one per frequency, dual boot to reach drop keel or flush mounted instrument.
You can run only one of the 150kHz and one of the 38kHz together, but not both 150kHz or both 38kHz ADCPs.

You can access the PCs from any of the work stations in the instrument room, use shift-shift-o to change between computers.

Something to remember: If you want to use the 38 kHz, turn off the 38 kHz frequency of the EK80. Likewise, if the EK80 takes priority, do not use the 38 kHz ADCP, there is significant interference both ways.

**Configuring the ADCPs:**

Configuration files for standard setup are stored in a folder on the desktop on each PC/boot.

There are 2x4 configuration files to cover:

- Narrowband versus Broadband
- Bottom track on versus off
- Synchronization through the K-Sync unit or standalone pinging.

Default setup should be:

- for ADCP only use: Narrowband, bottom track on;
- for concurrent ADCP/EK80 use: Narrowband, bottom track off.

In both cases:

- For 38 kHz ADCP: bin depth = 24 m, 16 m blanking distance, 65 bins
- For 150 kHz ADCP: bin depth = 8 m, 4 m blanking distance, 65 bins

Some of the settings in the configuration files might need adjusting:

1. Draft of the boat:

   Variance between minimum 7.2 m and maximum 8.5 m → check on bridge and adjust in the configuration files (command: ED).
Might need adjusting during a cruise if ballasting changes significantly, so check draft regularly.

If drop keel is extended, add the additional draft (max 3.4 m) → display in instrument room. There should be an event log when the drop keel is deployed and taken up again.

2. Set salinity to something sensible. It’s set to 35 in the configuration file – adjust there if needed (command: ES)

→ NOTE: Make a copy of the standard config files and do the changes in the copy!

The rest of the configuration and starting and stopping of the ADCPs are done in VmDas. An extensive guide for VmDas settings is available on the ADCP PCs. Contact the instrument technicians for help.

7 Water samples from Niskin bottles

The order of sampling form the Niskin bottles (see 6.2 Niskin Rosette) should be decided on as some samples need to be taken immediately. Here one should also agree on some general rules when sampling from Niskin bottles, e.g. use of Nitril gloves.

All water samples taken from the ships Niskin bottles should be described in an own sub-chapter.

7.1 SALINITY SAMPLES

For sampling of salinity samples, see chapter 6.2 Niskin Rosette.

Method responsible: Angelika Renner (IMR, angelika.renner@hi.no)

(description of Salinometer will come when instrument is installed on KPH)
7.2 SAMPLING OF DIC, AT AND PH

Sampling of DIC/pH from the Niskin-bottle rosette will take place first (if no other sensitive gases are sampled, such as dissolved oxygen). From the Niskin bottle, use a tubing to gently fill a 250 ml borosilicate bottle (glass bottle with blue plastic cap) to avoid air bubbles, overfill 1-2 volumes for rinsing and removing air bubbles.

7.2.1 Total dissolved inorganic carbon (DIC)

Measured parameter: Total dissolved inorganic carbon (DIC, µmol/kg)

Method responsible: Melissa Chierici (IMR, melissa.chierici@hi.no) and Agneta Fransson (NPI, agneta.fransson@npolar.no)

Description of parameter: Total dissolved inorganic carbon measures the sum of bicarbonate, carbonate and carbonic acid and dissolved CO₂ (the sum of carbonic acid and CO₂ is generally referred as H₂CO₃*) in seawater, melted sea ice and brine, snow. Other abbreviations that are used instead of DIC are TCO₂ and CT.

Method description:

→ Special laboratory requirements:
  - Dry-lab with as constant temperature as possible. 3 meters of bench place, with space on top of bench (preferably no cupboards) and room under bench for circulating water baths. Room needs a sink for seawater waste.
  - Nitrogen gas (20 L cylinder at 200 bar) to be secured near the instrument to allow control and short distance.
  - Refrigerator on short distance or in room, access to MQ water

→ Sampling depths:

→ Chemicals used: 10% phosphoric acid, cathode and anode solution (DMSO and thymol blue) potassium iodide salt (KI), nitrogen gas (high-quality)
DIC will be sampled and analysed directly onboard KPH and follow the state-of-the-art method described in Dickson et al. (2007). DIC was determined using gas extraction of acidified samples followed by coulometric titration and photometric detection using a Versatile Instrument for the Determination of Titration Alkalinity (VINDTA 3D, Marianda, Germany). Accuracy of the measurements is ensured by routine analyses of Certified Reference Materials (CRM, provided by A. G. Dickson, Scripps Institution of Oceanography, USA). Precision is derived from repeated measurements on a daily basis.

7.2.2 Total alkalinity (AT)

**Measured parameter:** Total Alkalinity (AT, µmol/kg)

**Method responsible:** Melissa Chierici (IMR, melissa.chierici@hi.no) and Agneta Fransson (NPI, agneta.fransson@npolar.no)

**Description of parameter:** Total alkalinity (AT) is a measure of the buffer capacity of a seawater towards acidic or basic input. Other commonly used acronyms is TA.

**Method description:**

→ **Special laboratory requirements:** Dry-lab with as constant temperature as possible. 2 meters of bench place, with space on top of bench (preferably no cupboards) and room under bench for circulating water baths. Room needs a sink for seawater waste.

→ **Sampling depths:**

→ **Chemical used:** Hydrochloric acid (0.1 N), electrolyte solution (saturated potassium chloride, KCl).

Sampling and analyses of AT will take place onboard and be determined by potentiometric titration with 0.05 N hydrochloric acid (HCl) using a Titrando (Metrohm, Swiss) system with a Dosino (Metrohm) to allow for precise volume addition (sample volume 40 ml). The equilibration point is evaluated using a Gran evaluation in addition to end-point detection in the Tiamo software (Metrohm). Accuracy of the measurements is ensured by routine analyses of
Certified Reference Materials (CRM, provided by A. G. Dickson, Scripps Institution of Oceanography, USA). Precision is derived from repeated measurements on a daily basis.

A selection of duplicate samples (20-40 samples) will be stored and analysed post-cruise at the IMR laboratory in Tromsø for quality and consistency check using the state-of-the-art instrumentation for potentiometric titration in a closed cell (100 ml) and 0.1 N HCl on a Versatile Instrument for the Determination of Titration Alkalinity (VINDTA 3S, Marianda, Germany).

7.2.3 pH

**Measured parameter:** pH (total scale, spectrophotometric)

**Method responsible:** Melissa Chierici (IMR, melissa.chierici@hi.no) and Agneta Fransson (NPI, agneta.fransson@npolar.no)

**Description of parameter:** pH is a measure of the acidity (hydrogen ion concentration/activity) of a seawater

**Method description:**

- **Special laboratory requirements:** Dry-lab with as constant temperature as possible. 2 meters of bench place, with space on top of bench (preferably no cupboards). Room needs a sink for seawater waste.

- **Sampling depths:**

- **Chemical used:** m-cresol purple (2 mM solution)

Sampling and analyses of pH will take place onboard and is determined spectrophotometrically (Diode array spectrophotometer, type for ex: HP8453, Cary (former Agilent) using a 2mM solution of the sulfonaphtalein dye m-cresol purple as an indicator (Clayton and Byrne, 1993). Perturbation of indicator pH was corrected using the formulation described in Chierici et al. (1999).
7.3 SAMPLING FOR $\Delta^{18}$O (DELTA-O-18)

**Measured parameter:** $\delta^{18}$O (or delta-O-18)

**Method responsible:** Mats Granskog (RF1, NPI, mats.granskog@npolar.no), Agneta Fransson (RF2, NPI, agneta.fransson@npolar.no)

**Description of parameter:** $\delta^{18}$O (or delta-O-18) is a measure of the ratio of stable isotopes oxygen-18 ($^{18}$O) and oxygen-16 ($^{16}$O). It is commonly used with measurement of salinity of seawater to infer the amount of different freshwater sources in a seawater sample (sea-ice melt water or river/glacial water).

**Methodological description:**

→ **Special requirements/gear:**

→ **Sampling depths** Standard depths (5, 10, 25, 50, 75, 100 and 200 m) and DCM. Collect from same Niskin and same depths as DIC/TA.

→ **Chemicals used:**

**Materials:**

- Plastic (HDPE) bottle (15ml) with plastic caps (VWR number 215-7503, Figure 1).
- Parafilm (50mm x 50mm squares)
- Labels

**Method:**

Bottle is completely filled with seawater and cap closed tightly. Afterwards the vial must be sealed with Parafilm. Large headspace (of air) will contaminate the sample during storage.

1) Fill the bottle with seawater from the Niskin bottle. As full as possible to leave as little air inside bottle as possible.
2) Close the cap without touching the inside. Turn bottle upside down, if there is a little bubble of air it is ok.
3) Dry the outside of the cap and the bottle. Tighten the cap.
4) Seal the bottle cap with Parafilm. The Parafilm must be stretched tightly around the cap. If it is loose it will fall off. Practice applying Parafilm to an empty vial until you can do it securely. It helps to warm the parafilm for 5 seconds between the palms of your hands to make it stretch better without breaking.
5) Label sample (preferable before sampling, to make sure label attaches well).

→ **Sample storage / Shipment address** Store the $\delta^{18}$O samples at +4°C in dark (in refrigerator onboard). Room temperature storage is fine if bottles are well sealed. DO NOT FREEZE!

**Shipment:**  
Norsk Polarinstittutt  
Framsenteret  
Hjalmar Johansens gate 14  
9007 Tromsø  
ATT: Mats Granskog/Agnete Fransson

![Fig. 7.3-1 Bottle and cap](image)

### 7.4 PARTICULATE ORGANIC CARBON (POC) AND NITROGEN (PON)

**Measured parameter:** Quantification of particulate organic carbon and nitrogen (in $\mu$g C or N L$^{-1}$).

**Method responsible:** Marit Reigstad (UiT, marit.reigstad@uit.no), Gunnar Bratbak (UiB, [gunnar.bratbak@uib.no](mailto:gunnar.bratbak@uib.no)), Murat Van Ardelan (NTNU, murat.v.ardelan@ntnu.no - protocol responsible)

**Description of parameter:**
Methodological description:

→ **Special requirements/gear:** Drying oven, pre-combusted GF/F filters

→ **Sampling depths:** Standard depths 5, 10, 25, 50, 75, 100 and DCM

→ **Chemicals used:** none

Method:

- Collect seawater from the Niskin bottles from standard depths down to 100 m. Rinse the plastic sampling bottle with sample water (~100 ml) before collecting the sample.
- Remember to gently mix the bottle (turn upside down for a couple of times) before filtering to ensure that no particles settle down (results in uneven concentration if subsampling the sampling bottle).
- Be sure to have placed the filter in the middle of the filter holder and that the funnel is thoroughly placed on top of it (if something is leaking the exact volume filtered or the filtration area on the filter is not known – these are needed for the calculations later).
- Filter ca. 50-2000 ml (depending on biomass – a light colour on the filter is enough) from each depth through pre-combusted 25 mm GF/F filters.
- Use low vacuum pressure (about -30 kPa). Always have the valve of the filtering funnel closed before turning the pump on or off.
- Cover the funnels with tin foil when filtering. If you expect to filter 1 L or 2 L, you can place the respective plastic bottle into the funnel and let it run. Ideally the 0.5 and 1 L mark should be indicated on the plastic sampling bottles.
- Rinse the funnel with filtered seawater (collect filtered sea water from previous filtrations) once the sample has been filtered. Do not let the filters dry out, close the valve.
- After filtration, each GF/F filter should be directly placed into Pall filter slides (see picture below) and dried at 60°C in a drying oven. Wrap filter slides from one station in tin foil and keep them in a labelled Ziploc bag.
- Rinse the filtration equipment with MilliQ before the next sample. Cover filtration funnel with tin foil when not in use (to avoid dust).
• For each sampling day or event, prepare a blank filter by filtering MilliQ water through a pre-combusted GF/F filter (similar volume than sea water for the samples) and treating it the same way than the samples. The blank filters get normal sample running number (but make an additional note “MilliQ” on the sample label/analysis slide) and are noted in the filtering protocol and log sheet. Make also a note on CTD log sheet to avoid confusion with numbering on following casts.

⇒ Samples storage/shipment address: POC/N samples should be wrapped in tin foil and stored at -20 C until further processing at UiT and UiB.

Fig. 7.4-1 Petri slide (middle) for storage of POC/N filters

7.5 TOTAL (TOC) AND ORGANIC CARBON

Measured parameter: Quantification of TOC - total organic carbon (mg C L⁻¹).

Method responsible: Murat Van Ardelan (NTNU, murat.v.ardelan@ntnu.no)

Description of parameter:

Methodological description:

⇒ Special requirements/gear: Drying oven, pre-combusted GF/F filters, acid-cleaned glass vials
Sampling depths: 3 depths (surface, midwater column and near bottom) corresponding to the depths for DOM/C

Chemicals used: HCl 30%, H3PO4 (35%)

Method:
All the 40ml vials used for the sample collection of TOC were cleaned as follows (gloves, goggles, and lab coat were used at all times, the cleaning was done under a fume hood when needed):

1. Remove paper, plastic, or marker labels from all vials
2. Immerse the vials in a HCL 30% bath for at least 8 hours
3. Rinse the vials three times with MiliQ water
4. Dry the vials in a drier until completely dry, usually around 2 hours at 80°C
5. Wrap the vials with aluminum paper in packages small enough to be placed inside a combustion oven
6. Combust the vials in an oven at 450°C for 8 hours

The vials plastic caps were cleaned separately with the following procedure (gloves, goggles, and lab coat were used at all times, the cleaning was done under a fume hood when needed):

1. Place the caps in a HCL 30% bath for about ten minutes
2. Transfer the caps into a methanol bath (do not rinse after the HCL bath with MiliQ water to avoid contamination)
3. Leave the caps in the methanol bath from 6 to 8 hours
4. Take the caps out of the methanol bath and place them in a semi-closed (to allow evaporation)
5. aluminium foil foiled container
6. Place the caps to dry in a drier at 60°C for 2 hours or until completely dry
7. Transfer the caps to a plastic bag and keep them sealed until needed
TOC samples directly transfer, without filtration, to 40ml pre-rinsed (3x with filtered water) glassvial (QEC 2112-40mlE) and filled up to the bottle neck. The vials are stored in a freezer at -20°C.

→ **Samples storage/shipment address:** the samples should be stored in a freezer at at -20°C.

### 7.6 DISSOLVED ORGANIC MATTER CHARACTERIZATION (DOM)

**Measured parameter:** Dissolved organic matter (DOM)

**Method responsible:** Murat Van Ardelan (NTNU, murat.v.ardelan@ntnu.no)

**Description of parameter:** Qualitative analysis of the major components and distribution of dissolved organic matter at discrete depths.

**Methodological description:**

→ **Special requirements/gear:** GO-FLO BOTTLES

→ **Sampling depths:** 3 depths (surface, midwater column and near bottom). To be collected from the GO-Flo bottle. For deeper samples (> 600 m) water is to be collected from the Rosette.

→ **Chemicals used:** Methanol (washing and HPCL grade), Hydrochloric Acid (32 %)

**Materials:**

- Standard Rosette sampler with Niskin bottles and GO-FLOW bottle
- Clean environment not required. Bench space with sink necessary for filtration and preconcentration
- Ca 1.5 m2 bench space is needed
- Deep freeze space (ca 50x50x100 cm)
- Refrigeration space (ca 50x50x50 cm)
- Funnels or Tubing: To collect from the GO-Flo/ Niskin Rosette
• Volumetric Flasks (2.5-10 L): To collect from the GO-Flo/ Niskin Rosette. 
  1 per depth +
• spares

NOTE: Use glass for all materials if possible!

Filtration system: (ideally 2 setups for replicates)

• Vacuum or water pump
• Glass funnels (500-1000 ml) (2):
• Clamps (2): To hold the flask the filter and the funnel
• Frits 47 mm (2): for filters
• Filtration Vacuum Flasks (2 L): to collect the filtrated seawater.
• Pre-concentration flask (2-4 L): for setting the C-18 columns
• GF/C Filters (47 mm): Pre-combusted filters at 450°C for 8 h in Aluminum packages

Reagents:

• Methanol (washing)
• Methanol (HPLC grade) (extraction)
• Hydrochloric Acid (32 %)

Pre-concentration system:

• Peristaltic pump: Peristaltic tubing black-black (diameter 0.90 mm or 0.86 mm)
• Glass pipettes (10 ml): To place inside the flasks and connect to the tubing and the C18
• Columns. Six to connect to conduct simultaneous filtration of 3 depths with duplicates.
• Silicon Tubing
• Chromatographic columns: C18 Sep-Pak plus
**Extraction system:**

- Extracting setup/rack: custom-made rack to place the Ch. Columns for extraction
- Scintillation vials (20ml): for extraction

**Material washing and cleaning:**

1. Immerse the vials in a HCl 30% bath for at least eight hours.
2. Rinse the vials three times with MilliQ water.
3. Dry the vials in a drier until completely dry, usually around two hours at 80°C.
4. Wrap the vials with aluminum paper in packages to be placed inside a combustion oven.
5. Combust the vials in an oven at 450°C for eight hours.

The plastic caps of the vials were cleaned separately with the following procedure:

1. Place the caps in a HCl 30% bath for about ten minutes.
2. Transfer the caps into a methanol bath (do not rinse after the HCl bath with MilliQ water).
3. Leave the caps in the methanol bath from six to eight hours.
4. Take the caps out of the methanol bath and place them in a semi closed (to allow evaporation) aluminum foil foiled container.
5. Place the caps to dry in a drier at 60°C for two hours or until completely dry.
6. Transfer the caps to a plastic bag and keep them sealed until needed.
Method:

- Collection of seawater from the Rosette is performed on the glass bottles, avoiding using extra tubing if possible (depending on glass volume).
- After collection of volume of seawater: 3 replicate of 50 ml TOC samples are collected in (clean pre-combusted) glass vials (do not fill to max, leave out enough headspace).
- Samples will be acidified with HCl (pH ~ 2) and preserved in deepfreeze (-20°C).
- Ca 2000- 3000 ml seawater is filtered through pre-combusted clean GF/C filters, which are stored in the deepfreeze for POC analysis. The filtrated seawater is acidified to pH ~2 using 3M or 32% HCl (reagent grade).
- In order to eliminate impurities, use the peristaltic pump to rinse with 5 ml methanol (HPLC grade) the chromatographic columns (C-18) and the glass pipettes. Start by the columns.
- Proceed to extraction at a rate of ~10 ml /min. Make sure to cover the mouth of the glass container with aluminum foil to prevent any kind of contamination to the DOC filtrate.
- DOC accumulated in the column should not be stored refrigerated but extracted immediately!
- After extraction flush the columns with 0.1 M HCL (10-12 ml) and MQ water.
- Extraction is carried by flushing each column with methanol in 10 ml (small syringes).
- The extract will be kept in clean pre-combusted glass vials in refrigeration (4°C)

Ancillary measurements:

- TOC: 3 replicates in 40 ml vials filled, and frozen
- POC: Volume filtered in the 47 mm filters will be used for this analysis
- Glass vials (40ml): Pre-combusted at 450°C for 6rs in Aluminum packages
7.6.1 DOC-characterization

**Measured parameter:** Dissolved organic carbon (DOC)

**Method responsible:** Murat Van Ardelan (NTNU, murat.v.ardelan@ntnu.no)

**Description of parameter:** Characterization of dissolved organic matters, its lability for bacterial decomposition, its recalcitrant/refractory character. Its elemental ratios (C/O, C/N and C/H)

**Methodological description:**

➔ **Special requirements/gear:** Standard Rosette sampler with Niskin bottles can be used this sampling. No need special clean environment but a bench space near to sink s necessary for filtration & pre-concentration on the chromatographic column activities

➔ **Sampling depths:** !!! *Subject to change depending on context, physical and biological in situ measurements* temporary depths; surface, 5, mid-depth and ca 2 m above sediment water interface

➔ **Chemicals used:** HCl or H3PO4 (35%)

**Material:**
- vacuum pumps
- peristaltic pumps
- filtration set ups
- pre-concentration set ups
- large glass jars/flasks
- Ca 1.5 m² bench space is needed
- deepfreeze Space (ca 50x50x100 cm)
- refrigerator Space (ca 50x50x50 cm)

**Method:**
- After collection of ca 2.5 Liter seawater 3 replicates of 50 ml TOC samples will be
- acidified with either HCl (or H3PO4, 35%, 250 µl to 50 ml) in glass vial preserved in refrigerator (see 7.5 TOC protocol for the details).
• Ca 2000 ml sw will be filtrate with GF-F filter and acidified to pH 2 (see Fig 1)-if there is high productivity in the sampling spots, use GFM prefiltration before GF-F filtration
• Acidify the filtrated seawater with UP HCL (12 M) bring pH to 2.0.
• Rinsed PPL and C-18 columns with 6ml of ultrapure methanol in order to eliminate impurities before placing them inside the sample.
• Acidified SW will run through columns (PPL, (Agilent, Bond Elut-PPL) and/or C-18 (Water)) with help of peristaltic pumps (see Fig.1) with a flow rate about 2-3 ml/s.
• Rinsed the column with 5ml of HCl 0.01M in order to eliminate the salt residue from the column.
• Extract the organic compounds from column by using twice with 5ml of ultrapure methanol (total 10 ml) with help of plastic syringe; the columns were put on a support
• while applying methanol for extraction and left the extraction methanol to drip into a 40 ml pre cleaned and combusted glass vials.
• The extracted samples in the vials should be kept in refrigerator and dark until the analysis with LC-MS / FT-ICR-MS in the NTNU

![DOM Characterization diagram](image)

**Fig. 7.6.1-1** Set up of solid phase extraction (SPE) with PPL and/or C-18 column
7.7 TRACE ELEMENTS SAMPLING VIA GO FLO BOTTLES

**Measured parameter:** Trace elements (in nmol or μmol L-1).

**Method responsible:** Murat Van Ardelan (NTNU, murat.v.ardelan@ntnu.no)

**Description of parameter:** Quantification of particulate, total acid leachable and dissolved metal/element concentration (in nmol L-1).

**Methodological description:**

→ **Special requirements/gear:**
  - GO–Flo bottles (Teflon coated)
  - Ste line or Kevlar winch line (polymer-non-metallic line)
  - Class 100-clean air laminar flow bench (place for this and additional lab space for clean sample treatment (this space will be decked by PE sheet, and it should be in the area where there is minimal human activity (minimum turbulence)
  - Lab space: minimum 4-5 m² lab space including bench and sink is needed (this space will be decked by PE sheet temporarily to create a “clean -lab / dust free environment

→ **Sampling depth(s):** Subject to change depending on context, physical and biological in situ measurements temporary depths, 10, 20, 40 60, 120, 300, 500 and ca 10 m above sediment-water interface

→ **Chemicals used:**

**NOTE:** 1-GO-FLO deployment with polymer line (either ste-line or kevlar) sampling is necessary

for clean trace metal sampling
Method:

- Steps after GO FLO cast completed
- GO FLO bottles will immediately be placed in the clean lab space into the GO FLO racks. Work will be performed under Class -100 laminar flow chamber
- Total acid leachable trace elements (TTE): replicates of 125 ml TTE sample (without filtration) will be collected and acidify to ca pH 1.7
- Dissolved trace elements (DTE): Ca 200 ml of seawater will be filtrated with Sartobran Sartorius double-layer (0.4-0.2 μm pore size) and acidified to ca pH 1.7
- TTE and DTE samples can be stored under room temperature in double PE bags, and send to NTNU for final SeaFAST preconcentration and HP-ICP-MS analysis

### 7.8 SAMPLING FOR CDOM

**Measured parameter:** Colored (or chromophoric) dissolved organic matter (CDOM)

**Method responsible:** Mats Granskog (RF1: NPI, mats.granskoog@npolar.no), Børge Hamre (RF1: UiB, borge.hamre@uib.no), Murat V. Ardelan (RF2: NTNU, murat.v.ardelan@ntnu.no)

**Description of parameter:** Colored (or chromophoric) dissolved organic matter (CDOM) for characterization of marine dissolved organic matter (DOM) using optical measurements (absorbance and fluorescence). DOM is here defined as that passing through a nominal 0.22 μm pore size membrane filter. Absorption is important for the optics of water masses (RF1) and fluorescence is used to characterize the material (RF2).

**Methodological description:**

→ **Special requirements/gear:**

→ **Sampling depths:** Standard depths (5, 10, 25, 50, 75, 100 and 200 m) and DCM. Whenever possible collect from same Niskin and same depths as DOC, Chl-a, and δ¹⁸O. Whenever optical properties (RF1) of
the water column are measured CDOM samples should be collected from a cast back to back with optical profiles.

➔ Chemicals used:

Materials:

• Amber EPA type glass vials with caps (20-40ml, burned and caps acid-washed)
• Filter capsule (Millipore Opticap XL4, Durapore 0.22 μm, KVGLA04HH3) and (clean) tubing Laboratory gloves
• Labels

Preparations:

Before a new filter cartridge is used the first time it has to be rinsed with 4 L of Milli-Q water or clean seawater (from great depth with little particles or DOM) with gravity to clean off any residuals on the membrane. Tubing used should also be acid soaked in weak acid and copiously rinsed before use.

Collecting the samples:

![Diagram of sample collection](image)

Fig. 7.8-1 Schematic of the collection of CDOM sample.

Collecting the CDOM sample with cartridge:

1) *Always use laboratory gloves when collecting the samples* to minimize the contamination risk, as any fats, oils etc. on your hands contaminates
the measurements. Do not to touch the insides of the vials and caps, or the tip of the nozzle on the Niskin bottle or the filter capsule inlet/outlet, as these parts come in contact with sample water. Avoid touching anything that is dirty, dirty gloves do not help! If you have dirty gloves they do not do their job. Switch to a new pair.

2) The same filter capsule can (normally) be used for hundreds of samples. **Always start with the deepest seawater sample** (as it normally is the “cleanest” one) and move towards the surface when collecting sample water from the rosette. It is always good to flush with extra seawater through the capsule before collecting the first sample at each station.

3) Attach the tubing to the Niskin nozzle (note that the capsule is correctly aligned, see Figure 2). Make sure the bottom valve is closed, and the top valve is open. Point the cartridge such that arrow points downwards to help water flow through the filter.

4) Open the Niskin nozzle and fill the capsule with seawater (i.e. until water runs out of the top valve).

5) Close the top valve and let about 200 ml seawater pass through by gravity.

6) Fill the CDOM amber vial to the shoulder. Leave some headspace in vial! Close cap firmly.

7) Close the Niskin nozzle. Remove tubing from Niskin. To empty the capsule before next sample open the bottom valve, turn it upside down (arrow pointing upwards), and let water drain out. Close the bottom valve.

8) Note down the sample ID in the CTD sheet after each sample has been collected to be sure of which Niskin bottle it was collected from!

9) Go to next Niskin bottle. Repeat steps 1 to 6 until you have collected all samples.

10) After station empty the filter cartridge of seawater (step 5), and rinse with 200-300 ml of Milli-Q.

11) Store filter capsule and the tubing in a plastic bag or container with Milli-Q between stations, to avoid it getting contaminated.

**Sample storage / Shipment address** Store the CDOM samples at +4°C in dark (in refrigerator onboard). DO NOT FREEZE! Plan is to measure samples
onboard, if that is not possible, the samples are to be shipped (in a well-insulated box) to Murat V. Ardelan, NTNU, Trondheim

![Filter capsule in protective bag](image)

**Fig. 7.8-2** Filter capsule in protective bag (note that the filter capsules might have somewhat different shapes depending on batch and year). *Note that an arrow shows the direction water should flow through the filter (check it before using).*

### 7.8.1 Measurement of CDOM absorbance

**Measured parameter:** Colored (or chromophoric) dissolved organic matter (CDOM).

**Method responsible:** Murat V. Ardelan (RF2: NTNU, murat.v.ardelan@ntnu.no)

**Description of parameter:** Colored (or chromophoric) dissolved organic matter (CDOM) measurement with Liquid Waveguide Capillary Cells (LWCC) for ultraviolet (UV) and visible (VIS) absorbance spectroscopy is done for ultra-sensitive measurements of absorbance. Hence, several factors may affect the data analysis and external interferences must be kept at minimal.
Methodological description:

→ **Special requirements/gear:** Dry lab with stable temperature.

→ **Sampling depths:** See 7.8 CDOM sampling protocol

→ **Chemicals used:** Methanol (HPLC grade), 1N HCl (HPLC grade), high grade NaCl solution

Materials:

- CDOM samples in vials
- 1N HCl (HPLC grade)
- Methanol (HPLC grade)
- Milli Q water or Ultra-Pure Water (UPW) *(For cleaning and washing purpose)*
- Reference UPW *(For analysis store in an amber coloured clean glass bottle)*
- High grade pure NaCl solution with salinity similar to water samples collected (* salinity = 35*)
- Ocean Optics Liquid Waveguide Capillary Cells (LWCC) absorbance spectrometer system with software uploaded PC
- Waterbath *(Room temperature)*
- Peristaltic pump

Method:

1) Fix the LWCC system stably on to workbench and try to minimize movements.

2) Switch on the light source of LWCC at least one hour before the start of the measurement for constant temperature and stable measurements.

3) A water bath maintained at constant temperature must be used to stabilize the temperature of reference UPW and NaCl solution before running through LWCC system.
4) A constant flow rate of 1 ml / minute is maintained during the absorbance measurement. Always remember to fill in the capillary tube completely with the solution before each measurement.

5) Connect the spectrometer to the PC with software and observe the readings online.

6) Use reference UPW to measure the baseline of absorbance and to keep track if there is any shift in the baseline absorbance during the course of sample analysis. Repeat replicates until stable results are obtained. This procedure need to be repeated as and when needed if the drift noted is high.

7) Next measure the NaCl solution at beginning and after samples from a station or during start and end measurements of a day. Repeat with replicates for stable values.

8) Run at least three times 10 ml each of UPW to clean the system after step 4.

9) The absorbance measurement is done from 300 nm to 750 nm. At least triplicates must be run for each sample. However, as the sensitivity is quiet high it is recommended to get at least 15 to 20 spectra for each of the replicate measured.

10) After each analysis wash the system three times with 10ml UPW. After each sample wash with HCl (10 ml) and methanol (10 ml) followed by at least five times 10 ml UPW rinsing or until the spectra is stable.

11) Repeat step 3 to obtain baseline value.

Calculation of CDOM absorption:

Absorbance (A) from spectra is converted into Napierian absorption coefficient \([a]\) in \(\text{m}^{-1}\), using:

\[
a = \frac{2.303 \cdot A}{L}
\]

where \(A\) is the absorbance at specific wavelength and \(L\) is the optical path length of the LWCC in meters. \(a\) is generally adopted as a proxy for assessing the CDOM content in a given water sample.
\[ a_{CDOM} = (a_{sample} - a_{reference UPW}) - (a_{NaCl} - a_{reference UPW}) \]

Where \( a_{CDOM}, a_{sample}, a_{reference UPW}, a_{NaCl} \) where are absorption coefficients of CDOM, sample, reference UPW and NaCl solution respectively.

**Fig. 7.8.1-1 LWCC system with connections attached**

**Ocean View Software** (https://oceanoptics.com/product/oceanview/): This software gives helps for the measurement of the absorbance, fluorescence. The software also has help contents that are self-explanatory for making required modifications necessary for measurement of samples.
**Fig. 7.8.1-2 Ocean View software**

**Fig. 7.8.1-3 Background absorbance measurements**

### 7.9 PARTICULATE ELEMENT CONCENTRATION USING XRF

**Measured parameter:** Concentration of total particulate O, P, Na, Mg, Si, S, Ca, Mn, Fe, Zn (µM) measured using X-Ray Fluorescence (XRF)

**Method responsible:** Jorun K. Egge (UiB, jorun.egge@uib.no), Tatiana Tsagaraki (UiB, tatiana.tsagaraki@uib.no)
Description of parameter:

Methodological description:

→ Special requirements/gear:

→ **Sampling depths:** Standard depths, all stations 10, 25, 50 75, 100 and 200m, DCM and deep sample (bottom -15m); **Sampling Volume:** Sampling Volume: For surface water 500ml should be sufficient. Below Chl max, increase volume to 1000ml if possible.

→ **Chemicals used:**

Before filtration:

1) Collect water sample from rosette into a plastic container. The container should be clean and rinsed with distilled water before use.
2) If samples will stand for a while before filtration ideally they should be stored in a fridge or cool dark place.
3) Gently agitate container before filtration to resuspend any sedimented material
4) Prepare petri slides (pictured below): We usually mark a piece of tape with the sample code (a number) and tape it onto the petri slide. Each sample should have a cruise code and unique number (e.g. CRE001, CRE002 etc).

5) Blanks (two types are necessary):
a. distilled water blank: Filter, dry and store 500ml of distilled water, in triplicate, following the protocol outlined in the next section.

NB: Distilled water blank only needs to be collected once per survey.

b. filter blank- Keep at least three unused filters from each box of filters used. Mark the box 1, 2 etc. and note on the info sheet which samples the blanks correspond to in the line filter box number (see example below).

6) When changing box make a note of info sheet of which sample number begins with a new box. We need to know which samples come from which box of filters for correcting the blank accordingly. Write down the filter batch number (see example of info sheet below):

<table>
<thead>
<tr>
<th>Sample no</th>
<th>Station</th>
<th>Depth (m)</th>
<th>Date</th>
<th>Volume (ml)</th>
<th>Filter box number</th>
<th>Filter type</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>cre097</td>
<td>1</td>
<td>1500</td>
<td>10.05.</td>
<td>4</td>
<td>Box 1</td>
<td>0.6 PC</td>
<td>f.ex. starting new filter</td>
</tr>
<tr>
<td>cre098</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>blank box 2</td>
</tr>
<tr>
<td>cre099</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>blank box 2</td>
</tr>
<tr>
<td>cre100</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>blank box 2</td>
</tr>
<tr>
<td>cre101</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Box 2 starts</td>
</tr>
<tr>
<td>cre102</td>
<td>2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
For filtration:

1) Use 47mm 0.6μm (or selected pore size) polycarbonate filters in triplicate
2) Always note filtered volume on info sheet. It is OK to adjust volumes as needed, as long as the information is noted.
3) Before filtration gently agitate sample to resuspend particles
4) Filter selected volume using glass filtration funnels and collectors NB if there is a lot of material on the filter consider decreasing volume
5) When filtering is almost complete rinse filter with 5ml distilled water using pipette
6) Let filters air dry on sheet of paper or bench liner sheets.
7) Filters should be “weighed down” on one edge using labelled filter holder (see picture on the right)
8) Plastic box or other cover should be used to cover filters while drying.
9) Once filters are dry (for PC a couple of hours is more than enough), place in labelled filter holder and box

Important tips:

1) Flat forceps should be used to avoid damaging filter
2) Filter must be as centered as possible on the funnel. If the material is towards the edge of the filter it may result in a blank area of the filter measured in the XRF
3) When rinsing filter, pour the distilled water down the sides of the funnel to avoid redistribution of the material. If you forget to rinse make a note of it. Rinsing removes salt crystals from the filter, which interfere with the measurement of salts (obviously). Other elements are measured even if you forget to rinse.
4) There should not be any water left on the filter, dried drops create a lot of noise in the XRF
5) Filter should not be left empty under vacuum for too long, cells will break
6) Depending on the temperature, drying of PC filters can take ca 15 mins to two hours.

→ **Sample storage/ shipment address:** Samples must be stored at room temperature, in their box. Under no circumstances should the samples be refrigerated or frozen. Ship to operators address at end of cruise. No special requirements for shipping.

      Ship to :

      Institutt for biovitenskap, Biologen
      Thormøhlens gate 53A, 2. etg
      5006 BERGEN

**7.10 INORGANIC NUTRIENTS**

**Measured parameter:** Inorganic nutrients

**Method responsible:** Melissa Chierici (IMR, melissa.chierici@hi.no)

**Description of parameter:**

**Methodological description:**

→ **Special requirements/gear:**

→ **Sampling depths:**

→ **Chemicals used:** Chloroform: Trichloromethane

**Materials:**

- Sampling bottles: Vials PE 25 ml.
- Boxes: Cardboard box for storage of 100 pc of samples
- Dispenser: automatic pipette 0,2-1,0mL
- (remember adapter to 500mL chloroform bottle)
• Chloroform: Trichloromethane for spectrophotometry (Uvasol) Merck no: 2447.0500.
• NB! Chloroform shall be treated with caution!

Method:

• Use labelled sample vials and add the number on the CTD-Rosette sample list.
• Sampling: Rinse the bottles and cork with water from the appropriate Niskin flask at least 3 times before filling up. Fill the bottle, but leave a little room left for preservation. Use only new and clean scintillation bottles.

Preservation after sampling:

1) Preserve the samples with 200 µL (0.2 mL) chloroform using an automatic pipette. Check that it is set on the correct volume. Test the pipette before preserving the samples to ensure that dosing takes place. Keep the equipment clean to avoid contamination!
2) Keep cool! Keep the preserved samples in the fridge in the dedicated cardboard boxes, label with cruise name/number, date.

SHORT VERSION:

1. Make sure the vials are numbered
2. Rinse vial and cap with water from the sampled Niskin 3 times before filling up leaving some headspace for preservation
3. Add 0,20mL (200 microliter) chloroform
4. Place the samples in fridge/cool storage in boxes

→ Sample storage/ shipment address: Samples have to be transported using cold freight and they should not be in room temperature or more for more than a few hours (1-3 hrs).
7.11 WATER COLUMN SAMPLING FOR ALGAL PIGMENT CONCENTRATIONS (CHLOROPHYLL A, PHAEOPIGMENTS)

**Measured parameter:** Algal pigment concentrations (Chlorophyll a and phaeopigments) from the water column

**Method responsible:** Rolf Gradinger (UiT, rolf.gradinger@uit.no), Anna Vader (UNIS, anna.vader@unis.no)

**Description of parameter:** Quantification of algal pigment concentration (chlorophyll a, phaeopigments; in µg L-1) using the fluorometric acidification method (see separate protocol for measurements).

**Methodological description:**

- **Special requirements/gear:** Avoid any direct sun light or strong light exposure of samples during any part of the sampling process because algal pigments are sensitive to light. For example, use plastic sampling bottles wrapped in tin foil.

- **Sampling depths:** Standard depths 5, 10, 25, 50, 75, 100 and DCM (deep chlorophyll fluorescence maximum; as detected by in situ fluorescence profile; only if present and significantly ±5m different from standard depths). One sample per depth as standard.

- **Chemicals used:** Methanol

**Method:**

1) Lower CTD mounted fluorescence profiler to below 100m (or bottom depth if shallower) to determine the presence/absence of a DCM.

2) Collect seawater (250 ml to 2l, depending on expected pigment concentration) from the Niskin bottles from standard depths and potentially DCM down to 100 m. Rinse the plastic sampling bottle with sample water three times (~100 ml in total, depending on water budget) before collecting the sample.

3) Store plastic bottles in dark and cold location until time for filtration; filtration should occur within one hour (the faster the better).
4) Remember to gently mix the bottle (turn upside down for a couple of times) before filtering to ensure that no particles settle down (results in uneven concentration if subsampling the sampling bottle).

5) Be sure to have placed the non-combusted Whatman GF/F filter (typically 25mm, depending on filtration rig) in the middle of the filter holder and that the funnel is thoroughly placed on top of it (if something is leaking the exact volume filtered or the filtration area on the filter is not known – these are needed for the calculations later).

6) Filter ca. 50-2000 ml (depending on biomass – a light colour on the filter is enough) from each depth through 25 mm GF/F filters. Too much volume can result in clogging of the filter, too little volume can cause poor readings on the fluorometer.

7) Use low vacuum pressure (about -30 kPa). Always have the valve of the filtering funnel closed before turning the pump on or off.

8) Cover the funnels with tin foil while filtering. If you expect to filter 1 L or 2 L, you can place the plastic sampling bottle directly onto the funnel and let it run until empty. Ideally the 0.5 and 1 L mark should be indicated on the plastic sampling bottles prior to the sampling.

9) Rinse the funnel with GF/F filtered seawater (collect filtered sea water from previous filtrations into squeeze bottle) once the sample has been filtered. Do not let the filters dry out, close the valve.

10) Use forceps to fold the filter once and then place it into the plastic tube for Chl a extraction (10 ml Polypropylene (PP) tubes or reusable »Dramsglass» tubes with labelled plastic lids).

11) Label the PP tube with a cryomarker with sample ID. NOTE: Sample ID rules for AeN TBD.

12) Cover the sample labelling on the extraction tubes with transparent scotch tape because the solvent (methanol) will remove all markers.

13) Rinse the filtration equipment with MilliQ before the next sample. Cover filtration funnel with tin foil when not in use (to avoid dust).

14) Prepare samples in PP tubes for extraction with organic solvent if samples can be processed the next day (see separate document for details, preferred way).

15) If samples can't be measured the next day, samples in PP tubes should be shock-frozen in liquid nitrogen in a Dewar container (e.g. 2L)
immediately after filtering and transferred to a -80°C freezer for storage once all samples from one station have been filtered. Wrap PP-tubes in tin foil prior to long term storage. Store all samples from one station within one Ziplock bag and label Ziplock bag with station ID, sample IDs and “Algal pigment samples.”

**Fig. 7.11-1** Example of a filtration set-up.

7.11.1 Measurements of algal pigment concentrations (chl a, phaeopigments) based on samples from water column and sea ice

**Measured parameter:** Algal pigment concentrations (chl a, phaeopigments) from the water column and sea ice

**Method responsible:** Rolf Gradinger (UiT, rolf.gradinger@uit.no), Anna Vader (UNIS, anna.vader@unis.no)

**Description of parameter:** Quantification of algal pigment concentrations (chl a, phaeopigments) (in µg L⁻¹) using the fluorometric acidification method.
Methodological description

→ Special requirements/gear:

- Approx. 1 m bench space for Turner Design AU10 fluorometer,
- Potentially fume hood for organic solvent addition to Polypropylene (PP) tubes, waste containers for organic solvent and HCl mixture after measurements.
- Waste container for gloves.
- Nitrile lab gloves recommended (to avoid potential LATEX allergic reactions).

→ Sampling depths:

→ Chemicals used: Methanol, HCl

Method:

1) **Work as dark as possible.** Fluorometer should be in the darkest possible space, potentially wearing a red head lamp.
2) Add 5 ml methanol to the PP tube containing the GF/F filter. Use a dispenser, put a lid on the tube and cover with tin foil or store in black box/bucket with or cardboard box.
3) Extract in the dark “over night“ (12 hours minimum, 24 hour maximum) in a refrigerator between 0 and 4 deg C. NB: Note the start and end time of extraction period.
4) Turn on fluorometer (model AU 10) at least 10 min before taking the first measurement. For greater details regarding Turner fluorometers, you can check the company website recommendations for chlorophyll measurements.
5) Check that the fluorescence reading of the fluorometer is at Zero, using pure methanol.
6) Vortex the tube for a few seconds and let the sample adjust to room temperature before fluorometer reading.
7) Transfer the sample to a clean borosilicate cuvette, and dry the cuvette on the outside using lense paper.

8) Place the cuvette in the cuvette holder of the fluorometer and wait until readings have stabilized. Press * button on fluorometer (see picture below), it will first show Delay, then Average and finally Done on the fluorometer display. Read the value on the fluorometer. This is the Rb value (Reading before acid addition).

9) Take the cuvette out of the cuvette holder and add 2 drops of 5% HCl using Pasteur pipette, cover the cuvette with parafilm and mix it gently 3 times. Wait ca 90 seconds, then take a second fluorescence reading with the fluorometer. This is the Ra value (Reading after acid addition).

10) Place all chemical waste in a labeled waste container bottle. Wash cuvette with clean methanol between each sample and let it dry (place upside down on tissue paper).

11) Remember: Before start, and in between (every 15 samples), use a methanol blank to check that cuvettes are clean and that the fluorometer blank reading is stable at zero for methanol.

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**Fig. 7.11.1-1** Display and buttons of the Turner Design AU10 Fluorometer, * button is indicated by the red arrow
Calculation of chlorophyll and phaeopigment concentrations:

For chlorophyll a - concentration C:

\[ C = F_s \frac{r}{(r-1)} (R_b - R_a) \frac{V_{ex}}{V_{sam}} \]

where:

- \( C \) = chlorophyll a concentration (\( \mu \)g/L),
- \( F_s \) = response factor for the sensitivity setting S (see section calibration),
- \( r \) = the before-to-after acidification ratio of a pure chlorophyll a solution (see section calibration),
- \( R_b \) = fluorescence of sample extract before acidification
- \( R_a \) = fluorescence of sample extract after acidification.
- \( V_{ex} \) = extraction volume (L, 0.005L suggested)
- \( V_{sam} \) = volume filtered sample (L)

For phaeopigment - concentration P:

\[ P = F_s \frac{r}{(r-1)} (rR_a - R_b) \frac{V_{ex}}{V_{sam}} \]

For abbreviations, see chlorophyll a calculation above.

7.12 CONCENTRATING PROTISTS <10 \( \mu \)M BY VIVAFLOW FILTRATION

Measured parameter: Concentration of protist cells <10\( \mu \)m by vivaflow filtration

Method responsible: Bente Edvardsen (UiO, bente.edvardsen@ibv.uio.no), Luka Supraha (UiO, luka.supraha@ibv.uio.no)

Definition of parameter: Concentration of protist cells <10\( \mu \)m for electron microscopy and culturing by tangential filtration.
Methodological description:

→ **Special requirements/gear:** Vivaflow Cartridge for plankton, Masterflex pump with adjustable speed

→ **Sampling depths:** DCM if detectable, Melt ponds (Ice stations)

→ **Chemicals used:** NaOH 1M, Ethanol

Material:

- Vivaflow Cartridge for plankton
- Masterflex pump with adjustable speed
- Bottle 3L for the sample
- Falcon tube 50ml
- Clamps with screw
- Containers for
- MiliQ water 1L
- NaOH 1M 500ml
- EtOH 10% 500ml
- FSW (0.2µm) 500ml

Method:

1. Prepare the Vivaflow setup by placing the filtrate and retentate tubes into a waste tank and the feed tube into MQ water container.
2. Rinse the cartridge with 250ml of MQ water.
3. Fill a 3L water bottle with water sample prefILTERED through ca 90 µm mesh sieve.
4. Replace MQ water with sample and rinse the cartridge with 250ml of sample.
5. Put the retentate tube in the sample bottle. Increase the speed.
6. Concentrate the sample until about 250ml remains. For 3L of sample it will take about 30min.
7. Transfer 250 ml of the sample to a smaller bottle and then to 50ml Falcon tube.
8. Concentrate at lowest speed until the volume is reduced to 20ml.
9. Clamp the filtrate tube to recirculate the sample. No change in volume should take place.
10. Remove the feed tube from the sample to collect all the material in the Falcon tube.
11. Store the sample in a cool room for later use.
12. Place the retentate tube and the filtrate tube in the waste tank and the feed tube in the FSW container. Rinse with 250ml of FSW.
13. Rinse with 250 ml of MQ water.
14. Rinse with 50ml of 1M NaOH, then place all three tubes in the 1M NaOH container and recirculate for 20min
15. Rinse with 250ml of MQ water.
16. Turn off the pump and clamp all three tubes. For storage longer than 1 day store in 10% EtOH.

→ Sample storage: Concentrated samples should be kept in a cool and illuminated room at ca. 4°C.

Literature:

- Daniel Vaulot Tangential flow filtration (TFF) concentration of phytoplankton. protocols.iodx.doi.org/10.17504/protocols.io.krmcv46

7.13 LIGHT MICROSCOPY PHYTOPLANKTON/PROTISTS

Measured parameter: Phytoplankton/protist abundance (in cells L⁻¹) using the Utermöhl method.

Method responsible: Rolf Gradinger (UiT, rolf.gradinger@uit.no), Philipp Assmy (NPI, philipp.assmy@npolar.no)

Description of parameter: Identification and quantification (abundance in cells L⁻¹) of phytoplankton/protists with an inverted light microscope using the Utermöhl method.

Methodological description:

→ Special requirements/gear: Fume hood for addition of fixatives. Wear personal protection. Use waste containers as needed.
→ **Sampling depths:** Standard depths at process stations 5, 10, 30, 60, 90 and DCM (see algal pigment sampling for specific comments regarding DCM)

→ **Chemicals used:** hexamethylenetetramine-buffered formaldehyde and acidic Lugol's solutions.

**Material:**
- 200 mL brown glass bottles, one per sample
- hexamethylenetetramine-buffered formaldehyde and acidic Lugol's solutions (see separate protocol for preparation of fixatives).

**Method:**

a) Label the bottles with sample ID.

b) Wear gloves. Add 2 mL acidic Lugol's to empty 200 ml brown glass bottles (final concentration 1%).

c) Fill ca 180 ml of seawater from each depth directly from the Niskin bottles into 200 ml measuring cylinder and transfer into the 200 ml brown glass bottles (see picture below).

d) Mix by gently turning the bottle up-side down three times.

e) Under the fume hood add 20 ml of 20% hexamethylenetetramine-buffered formaldehyde (final concentration 2%).

f) Mix by gently turning the bottle up-side down three times.
→ **Samples storage/shipment address:** Store samples in fridge or cold room during the cruise. Ship to operators address at end of cruise. Make sure that bottles are safely packed for transport (hard-casing boxes with bubble-foil filling).

![Image: 200 ml brown glass bottle]

**Fig. 7.13-1 200 ml brown glass bottle**

**Literature:**


### 7.14 LIGHT MICROSCOPY LARGE PROTISTS/RARE TAXA

**Measured parameter:** Protist (>20 µm) and mesozooplankton abundance (in cells L⁻¹) of larger protists and small mesozooplankton

**Method responsible:** Rolf Gradinger (UiT, rolf.gradinger@uit.no), Philipp Assmy (NPI, philipp.assmy@npolar.no), Bente Edvardsen (UiO, bente.edvardsen@ibv.uio.no), Luka Supraha (UiO, luka.supraha@ibv.uio.no)

**Description of parameter:** Identification and quantification (abundance in cells L⁻¹) of larger protists (>20 µm) and small mesozooplankton (e.g. copepod nauplii and *Oithona*) with an inverted light microscope using the Utermöhl method and the enriched water samples. Please note that naked/delicate protists are not quantitatively retained by this method. Those taxa are counted in directly fixed non-enriched water samples.
Methodological description:

→ **Special requirements/gear:** Requires potentially an extra CTD cast due to the large volume needs. Fume hood for addition of fixatives. Wear personal protection for fixation and sample handling. Use waste containers.

→ **Sampling depths:** Standard depths: 5, 10, 30, 60, 90 and DCM and 200m (if available, depending on bottom depth)

→ **Chemical used:** hexamethylenetetramine-buffered formaldehyde

→ **NOTE:** Suggested as sampling activity at all process stations only due to large water and time demand for sampling.

   a) Three Niskin bottles (10 L each) are closed per depth and their entire contents drained into a large carboy via long silicon tubing. Note the volume in the carboy.

   b) The contents of the carboys are then gently filtered over 20 µm mesh via a handnet or a filtration tower if available (see picture below).

   c) Samples are washed off the mesh with GFF filtered seawater using a squeeze bottle and concentrated to approx. 90 ml (measure exact volume with 100 ml cylinder)

   d) Sample is filled into 100 ml brown glass bottle and 10 mL 20% hexamethylenetetramine-buffered formaldehyde (final concentration of 1%) added.

   e) Wear personal protection when adding fixative!

   f) Label all bottles with sample ID.

   g) Store the samples dark and cold, do not freeze!

→ **Samples storage/shipment address:** Store samples in fridge or cool room during the cruise. Make sure that bottles are safely packed for transport (hard-casing boxes with bubble-foil filling).
**Fig. 7.14-1** 100 ml brown glass bottle (left hand side)

**Fig. 7.14-2** Example of filtration tower (AWI-made; contact Uwe John) to concentrate large protists and small mesozooplankton (right side)

**Literature:**


### 7.15 METABARCODING (DNA AND RNA) OF PROTISTS AND PROKARYOTES

**Measured parameter:** Metabarcoding (DNA and RNA) of protists and prokaryotes

**Method responsible:** Anna Vader (UNIS, anna.vader@unis.no), Bente Edvardsen (UiO, bente.edvardsen@ibv.uio.no), Lise Øvreås (UiB, lise.ovreas@uib.no)
Description of parameter: Cells collected on filter, to be used for DNA and RNA isolation. Will result in OTU table of organisms.

Methodological description:

→ Special requirements/gear:
  - Clean area for filtration/sample handling
  - Fume hood for adding Glutaraldehyde and Lugol’s
  - Minus 80°C freezer and liquid nitrogen tank

→ Sampling depths: 5m, DCM, “deep” (=15 m above-bottom), 200m (at deep stations)

→ Chemicals used: Glutaraldehyde EM grade 25% Merck and Lugol’s

Method:

IMPORTANT:

- Always use lab gloves (also when handling equipment, e.g. washing, storing).
- RNA degrades quickly due to hydrolysis and RNAses (which are everywhere!). Change gloves after touching surfaces, work quickly. Never leave RNA at room temperature.
- DNA is very stable. Cross-contamination of samples is thus an issue. Clean tweezers with ethanol before use. Change gloves between samples.
- Immediately after use, rinse all containers, tubes and filtration equipment with distilled water and leave to dry in clean area.
- Glutaraldehyde is very toxic and should be added in fume-hood. Also keep waste in fume hood.

→ NOTE: Filtration should be done in triplicate for each depth. In total 3x7L=21L of water should be filtered.
• Prior to sampling: Label and number Sterivex filters, prepare pump (insert tubing). Number one 10L bucket for each Sterivex filter. Label sample-containers.
• Collect 22-25L of water from Niskins into 10 or 20L clean carboys (labelled with depth).
• Before connecting Sterivex filters, rinse the tubes by pumping appr. 1L of sample water through them.
• Connect the Sterivex filter unit to the Luer-Lok fittings on the tube (with a twist, see photos). The female Luer-lok side is the inlet. The filter can be hung over a bucket; there is no need for a tube connected to the outlet of the filter.
• Set the pump speed (flow rate 6 mL per sec) and the pump in forward position. Filter appr. 7 liters of water through each Sterivex filter unit. Make sure the connections are tight and that there is no pressure build up (seen as considerable expansion of the tubing and leaks)!
• Disconnect the Sterivex filter unit from the tube and remove as much of the remaining water as possible using a 50 mL syringe containing air.
• Cap the filter in both ends with the inlet and outlet caps. Snap-freeze the filter in liquid N₂. Transfer to -80°C freezer for storage.
• Measure the volume of filtered water from buckets, using measuring cylinders.
• Repeat procedure with next Sterivex filter.
• For each filter, note down time of sampling (UTC), filtration volume, filtration time, size fraction, any deviation from protocol, observation (e.g. color on filter) in log-sheet.
• After filtering all samples, clean the tube by pumping through approx. 1 liter of sterile distilled water.

Extra procedure during bloom conditions (only surface and DCM samples):

➔ NOTE: this filtration is carried out to enrich for small cells that may be poorly represented due to clogging by larger cells. It should be done in addition to standard sampling.
• Pour water sample through pre-filtration funnel equipped with 20 µm net.
• Collect cells (from 20 µm flow-through) on labelled Sterivex filter using a peristaltic pump as above.
• Cut out the 20 µm mesh from the funnel using a sterile scalpel and a tweezer washed in 70% EtOH, and transfer to 50 mL of sterile filtered seawater in a 50 mL Falcon tube. Shake tube gently to wash off organisms from mesh. Transfer 14 mL of sample to a 15mL Falcon tube and add 560 µL 25% glutaraldehyde (1% final), and 14 mL of sample to another 15mL Falcon tube and add 150 µL Lugol's (1% final conc.). The remaining is kept alive (ca 4 °C in growth chamber or fridge) for microscopy and single cell PCR.

\textbf{Fig. 7.15-1} Filtering equipment overview, inlet and outlet
Material and labelling:

- **Amount of water needed:** 3 x 7000 mL (plus extra for rinsing, total 22-25L)
- **Sampling time:** appr. 1 hour per sample
- **Sample labelling:** Station, date, depth, volume
- **Parameters to be recorded:** Station, date, time of sampling (UTC), depth, size-fraction, volume, filtration time, protocol used (ID nr?), any deviations from protocol, filter colour, buffer added (volume and type), storage (i.e. LNG), operator, sample responsible

**Sampling equipment (on “deck”):**

- Niskin bottles (rosette)
- Silicon tubes to attach to the Niskin bottles to empty them, one per Niskin bottle, length ca 60 cm x 12
- Clean 10 or 20L containers (for collecting sample)
- Sterile lab gloves (all sizes)
- Plastic funnels lined with 20 µm mesh (KC Denmark, for pre-filtration), labelled with depth

**Filtration equipment (in lab):**

- Peristaltic pump (ideally one for each depth=3)
- Tubing with Luer-Lok fitting
- 0.2 µm Sterivex filters, SVGPL10RC
- Syringe (50 mL)
- Inlet and outlet caps for sterivex filters
- Zip-ties for tightening tube to Luer-Lok connection
- Sterile distilled water and sea water
- Sterile lab gloves (all sizes)
- Buckets (10 L) to collect flow-through
- Liquid Nitrogen (LNG) tank for snap-freezing
- Measuring cylinder (2 L)
- Labelling pens
- Sterile scalpels and tweezers
- 70% Ethanol in squeeze-bottle
• 25% Glutaraldehyde
• Lugol's solution (see protocol for microscopy fixatives)
• Micropipettes for Glutaraldehyde and Lugol's and pipette tips
• 50 mL and 15 mL Falcon tubes

→ Samples storage/shipment address: Store samples in minus 80°C freezer on ship (boxes labelled “metabarcoding/Anna Vader/UNIS”). Will be unloaded at UNIS by operator after each cruise (dry-shipper needed for transport).

7.16 METATRANSCRIPTOMICS (mRNA) OF PROTISTS AND PROKARYOTES

Measured parameter: Metatranscriptomics (mRNA) of protists and prokaryotes

Method responsible: Anna Vader (UNIS, anna.vader@unis.no)

Description of parameter: Cells collected on filter, to be used for mRNA isolation. Will result in catalogue of gene activities.

Methodological description:

→ Special requirements/gear:
  • Should be sampled at local noon (to allow comparison between stations).
  • Samples need to be processed immediately
  • Clean lab area for filtration and sample handling
  • Minus 80°C freezer and liquid nitrogen tank

→ Sampling depths: 5m, only process stations

→ Chemicals used:
Method:

**Important:**

- Always use lab gloves (also when handling equipment, i.e. washing, storing).
- RNA degrades quickly due to hydrolysis and RNAses (which are everywhere!). Change gloves after touching surfaces, work quickly. Never leave RNA at room temperature.
- The mRNA composition changes according to the environment of the cell. Process samples immediately after collection, and try to keep sample at conditions resembling in vivo environment (i.e. light, temperature)
- Immediately after use, rinse all containers, tubes and filtration equipment with distilled water and leave to dry in clean area.

**NOTE** that filtration should be done in triplicate, to process samples as quickly as possible.

- In total 3x10L=30L of water should be filtered.
- Prior to sampling: Label and number Sterivex filters, prepare pump (insert tubing). Number one 10L bucket for each Sterivex filter.
- Collect 35L of water from Niskins into 10 or 20L clean carboys.
- Before connecting Sterivex filters, rinse the tubes by pumping appr. 1L of sample water through each tube.
- Connect the Sterivex filter unit to the Luer-Lok fittings on the tubes (with a twist, see photos). The female Luer-lok side is the inlet. The filter can be hung over a bucket; there is no need for a tube connected to the outlet of the filter.
- Set the pump speed (flow rate 6 mL per sec) and the pump in forward position. Filter appr. **10 liters** of water through each Sterivex filter unit. Make sure the connections are tight and that there is no pressure build up (seen as considerable expansion of the tubing and leaks)!
- Disconnect the Sterivex filter unit from the tube and remove as much of the remaining water as possible using a 50 mL syringe containing air.
Cap the filter in both ends with the inlet and outlet caps. Snap-freeze the filter in liquid $\text{N}_2$. Transfer to -80°C freezer for storage.

Measure the volume of filtered water from buckets, using measuring cylinders.

For each filter, note down time of sampling (UTC), filtration volume, filtration time, size fraction, any deviation from protocol, observation (e.g. colour on filter) in log-sheet.

After filtering all samples, clean the tubes by pumping approx. 1 liter of sterile distilled water through each tube.

**Extra information:**

- **Amount of water needed:** 30L (plus extra for rinsing tubes, total 35L)
- **Sampling time:** appr. 30 mins
- **Sample labelling:** Station, date, depth, volume
- **Parameters to be recorded:** Station, date, time of sampling (UTC), depth, size-fraction, volume, filtration time, protocol used (ID nr?), any deviations from protocol, filter colour, buffer added (volume and type), storage (i.e. LNG), operator, sample responsible

**Sampling equipment (on “deck”):**
- Niskin bottles (rosette)
- Silicon tubes to attach to the Niskin bottles to empty them, one per Niskin bottle, length ca 60 cm x 12
- Sterile lab gloves (all sizes)
- Clean 10 or 20L containers (for collecting sample)

**Filtration equipment (in lab):**

- Peristaltic pump with 3-4 pump-heads
- Tubing with Luer-Lok fitting
- 0.2 µm Sterivex filters, SVGPL10RC
- Syringe (50 mL)
- Inlet and outlet caps for sterivex filters
- Sterile distilled water
- Sterile lab gloves (all sizes)
- Buckets (10 L) to collect flow-through
- Liquid Nitrogen (LNG) tank for snap-freezing
- Measuring cylinder (2 L)
- Labelling pens
- Zip-ties for tightening tube to Luer-Lok connection
- Squeeze bottle with EtOH for cleaning
- Measuring cylinder (2L)
- Labelling pens

➔ **Samples storage/shipment address:** Store samples in minus 80oC freezer on ship (boxes labelled “metatranscriptome/Anna Vader/UNIS”). Will be unloaded at UNIS by operator after each cruise (dry-shipper needed for transport).

**7.17 FILTRATION FOR DNA/RNA SAMPLES USING 0.22 µM DURAPORE FILTERS**

**Measured parameter:** Microorganisms are collected on a 0.22µm Durapore filter for molecular analyses

**Method responsible:** Oliver Müller (UiB, oliver.muller@uib.no)
Description of parameter/Aim: Water samples from experiments are filtered on 0.22µm Durapore filters to collected cells on the filter. The filter is later used for DNA and/or RNA isolation. The DNA/RNA can be used for different molecular analyses including community composition analysis which will have an OTU table of organisms as data output.

Methodological description:

- **Special requirements/Gear:** - Vacuum pump, 0.22µm Durapore filters, filtration bottles with 47mm funnel (including tubing and connections)
- **Sampling depths:** Only water from experiments is used
- **Chemicals used:** No chemicals

Method: Filtration of water on a 0.22µm Durapore filters

Important: Work sterile!!!

- Always use lab gloves (also when handling equipment, e.g. washing, storing).
- Clean tweezers with ethanol between different filters.

- Transfer the respective volume (depends on the time point of sampling T0: 1L; T3: 200ml; T6: 500ml) from the experimental container (1L PC bottle or 0.5L whirlpack bag) into sterile bottles (1L).
- Use vacuum filtration systems and 0.22µm Durapore filters to collect biomass on the filters.
- Retrieve filters, transfer them into marked sterile Eppendorf tubes, flash-freeze in LiqN2 and store in -80°C freezer.

- **Sample storage/shipping address:** Eppendorf tubes with Durapore filters inside are stored in cryoboxes in the -80°C freezer. The shipping address for all samples is:
7.18 FLOW CYTOMETRY SAMPLING

Measured parameter: Bacteria, virus and small protists abundance

Method responsible: Aud Larsen (Uni Research/Norce, aula@norceresearch.no) / Gunnar Bratbak (UIB, Gunnar.Bratbak.uib.no)

Description of parameter: Samples for enumeration of bacteria, virus and small protists (primarily <10µm) by flow cytometry.

Methodological description:

→ Special requirements/gear: Liquid nitrogen for flash freezing and -80 freezer for storing of samples. Fume hood for use of fixatives.

→ Sampling depths: Standard depths at process stations 5, 10, 25, 50 75, 100 and 200m, DCM and deep sample (bottom -15m)

→ Chemicals used: 25% glutaraldehyde (EM grade)

Method:

- When preparing for station: Label each cryovial (2mL) with running number using cryomarker and/or use pre-printed label – NB: If using label make sure the label sticks after snap-freezed in liquid nitrogen.
- Under the fume hood, add 38 µl of 25% glutaraldehyde (EM grade) to each cryovial.
- Add sample directly from Niskin bottle into 20 ml scintillation vial (same as used for nutrients).
- Under the fume hood, pipette 1.8 ml of sample into 2 ml cryovial (3 vials per depth).
• Fix the samples for 2 h in a fridge.
• Snap freeze in liquid nitrogen.
• Store in cryobox at -80°C.

→ **Samples storage/shipment address:** Store samples in -80C during the cruise. Ship samples on dry ice using :

  World Courier (Norway) AS  
  Industrivegen 20  
  2069 Jessheim (Oslo)

Tel: 47 6394 6200; Fax: 47 6394 6201; Email: ops@worldcourier.no

**OBS:** Make appointment for pick up well in advance of docking at the end of the cruise and coordinate with shipping of other frozen samples (e.g. Sterivex filters)

  **Ship to:**

  Institutt for biovitenskap, Biologen  
  Thormøhlens gate 53A, 2. etg  
  5006 BERGEN

7.19 BACTERIAL BIOMASS PRODUCTION

**Measured parameter:** Bacterial production of carbon biomass (gC per volume per unit time)

**Method responsible:** Gunnar Bratbak (UIB, Gunnar.Bratbak.uib.no)

**Description of parameter:** Bacterial production of carbon biomass (gC per volume per unit time) estimated from incorporation of tritiated leucine ([2,3,4-3H] leucine).
Methodological description:

→ **Special requirements/gear:** Isotope lab or area approved for use of 3H. Incubator if not *in situ* incubation. Operator must hold a radioisotope safety course certificate.

→ **Sampling depths:** Standard depths at process stations 5, 10, 25, 50 75, 100 and 200m, DCM and deep sample (bottom -15m)

→ **Chemicals used:** tritiated leucine, 100% TCA

Method:

- Use sterile 2 mL Eppendorf tubes labelled on top, 4 per sample (3 parallels + 1 blank)
- Dilute the isotope 1:10 in sterile saline (eye drops (physiological salt-water) ampules are convenient). Add xx µL isotope to each tube to give a final concentration of 50µM leucine.
- Start timer. For each sample (depth) add 1.5 mL water to 4 tubes and then 80µl 100% TCA to the blank. Close the tubes. Start each new sample with 30 sec intervals to keep track of the incubation time.
- Incubate in situ or at in situ temperature for xx hours.
- Add 80µl 100% TCA to all samples (except blanks) following the same sequence as used for starting the incubations.
- Label and secure tubes in the racks. Store the samples in the fridge and ship home for workup and scintillation counting as described in standard protocols.
- Clean the working area and perform wipe test.

→ **Samples storage/shipment address:** Store samples in fridge during the cruise. Ship to operators address at end of cruise following protocols for shipment of radioactive material.
Literature:


7.20 IRON CHLORIDE PRECIPITATION OF VIRUSES FROM SEAWATER

**Measured parameter:** Recover viruses from natural waters.

**Method responsible:** Ruth-Anne Sandaa (UiB, ruth.sanda@uib.no)

**Description of parameter:** Recover viruses from natural waters via iron chloride precipitation.

**Methodological description:**

- **Special requirements/gear:** -80 freezer for storing of samples.
- **Sampling depths:**
  
  - WATER: Chl a max (or 20m if not detectable) and deepest sample at process stations. At deep stations (> 1000 m) add one sample at 500 m. 2 parallel samples of 20L pr event.
  - SEA ICE: Bottom 10 cm of 2-4 ice cores.
- **Chemicals used:** FeCl₃, 0,1M EDTA-0,2M Ascorbate buffer (see recipe below)
  
  - 0.966 g FeCl₃ x 6H₂O in 50 mL falcon tubes (20 mL MQ gives 10g/L Fe Stock Solution
  - 10g NaOH (50ml MQ gives 5N NaOH)
• 1.51g Tris-base
• 3.72g Na2-EDTA dihydrate
• 4.07g MgCl₂
• 3.52g of ascorbic acid
• HCL 5N
• pH paper

Buffer preparation: 0.1M EDTA-0.2M Ascorbate buffer, pH 6.0-6.5 – 100ml

- Dissolve 1.51g Tris-base in 80ml Milli Q water.
- Dissolve 3.72g Na2-EDTA dihydrate into solution. (For info: pH will be ~10.0)
- Once EDTA is in solution, dissolve 4.07g MgCl₂. (For info: pH will drop to ~8.0)
- Add 3ml of 5N NaOH. (For info: This will drop the pH to ~4.5 and the solution will become cloudy which indicates that the EDTA is coming out of solution.)
- Dissolve the reductant (3.52g of ascorbic acid or 2.52g of oxalic acid). (For info: The pH will increase to ~8.3 and the solution will clear up.)
- Once the reductant is in solution, add the last 1ml of 5N NaOH.
- Check the pH using pH paper (the buffer should be at pH 6.0 - 6.5). The solution may need some minor adjusting with NaOH or HCl to achieve a pH of 6.0. pH 6.0 is ideal for good recovery of viruses.
- Check the volume and add MilliQ water for a total volume of 100ml.
- Store the buffer in the dark (bottle wrapped in foil) and visually inspect prior to use. It should be clear without precipitates. At this point, 10-15ml of buffer can be sacrificed for a final pH check using a pH meter (if available).

The buffer will start to change color after about 24 hours. It is okay to use if slightly discolored, but do not use after about 36 hours (eventually the buffer will turn almost orange!).
Method:

WATER SAMPLES

- Prepare pump and Tripod by mounting the tubing to the pump head (see Figure and Tripod assembly diagram, Fig. 7.20-1). Set occlusion adjustment to 3 and the pump in direction of the Tripod (see Figure). When filtering, start the pump at low speed and increase to pump speed to 10 (will give a flow rate of ca 400 mL per min) when air has been removed and the vet valve closed.

- Filter 20 L of sample through 0.45 µm, 142mm Durapore filters into clean 20 L containers. This is the water (filtrate) to be used for virus concentration, discard the filter.

- Prepare FeCl$_3$ stock solution (10g/L) by adding 20 mL MQ water to pre-weight 0.966 g FeCl$_3$ x 6H$_2$O in 50 mL falcon tubes.

- Add 1 mL of the FeCl$_3$ stock solution to each 20 L of sample. Shake vigorously for 1 min (roll the containers back and forth on the floor).

- Add additional 1 mL of the FeCl$_3$ stock solution to each 20 L of sample. Shake vigorously for 1 min (roll the containers back and forth on the floor).

- Let the FeCl$_3$ treated water sit for 1h at room temperature to precipitate.

- Filter the FeCl$_3$ treated water using 1.0 µm, 142 mm polycarbonate (PC) membrane on top of a 0.8 µm, 142 mm Supor Membrane filter (support). Change the PC filter 1-3 times during filtration if necessary (slow filtration rate, depending on biomass).

- Place all PC filters from the same 20L sample in one 50 mL Falcon tube, with the precipitate facing out. Discard the Supor filter.

- Prepare fresh 0,1M EDTA-0,2M Ascorbate buffer, pH 6.0-6.5 (See buffer recipe below)

- Add 20mL of 1x buffer to each of the Falcon tubes (precipitate from 20L seawater)

- Place tubes on a rotor o/n at 4°C.
• Transfer the liquid to two 15 mL marked Falcon tubes and store in cryobox at -80°C.

ICE CORE SAMPLES

• Melt the ice, filter through 0.45 µm Durapore filters, measure the volume filtered and add the appropriate amount of FeCl₃ stock solution (100µL/L) in c) and d), proceed as above.

→ **Samples storage/shipment address:** Store samples in -80 during the cruise and on dry ice during transport.

**NOTE:** Make appointment for pick up well in advance of docking at the end of the cruise and coordinate with shipping of other frozen samples (e.g. Sterivex filters)

Ship samples on dry ice using
  World Courier (Norway) AS
  Industrivegen 20
  2069 Jessheim (Oslo)
  Tel: 47 6394 6200; Fax: 47 6394 6201; Email: ops@worldcourier.no

Ship to:
  Institutt for biovitenskap, Biologen
  Thormøhlens gate 53A, 2. etg
  5006 BERGEN

**Literature:**

• John, S, Poulos, B and Schirmer C. 2015, Ion Chloride Precipitation of viruses from seawater, Protocols.io
**Fig. 7.20-1** Filtration set up for viruses using a Masterflex I/P pump with Easy Load head and a Millipore Tripod - Standing Stainless Steel Filter Holders (142 mm)
**Fig. 7.20-2** 142 mm stainless steel filter holder assembly diagram (© Merck Life Science).

7.21 VIRAL PRODUCTION AND DECAY (REV.1)

**Measured parameter:** Viral production and decay

**Method responsible:** Ruth-Anne Sandaa (UiB, ruth.sandaa@uib.no)
Description of parameter: Measure rate for viral production and degradation

Methodological description:

→ Special requirements/gear: -80°C freezer for storing of samples.

→ Sampling depths: Chl a max (or 20 m if not detectable) and deepest sample at process stations

→ Chemicals used: 0.5 M NaOH, Ethanol

Method:

Prepare two Vivaflow 200 cassettes, one 0.2 µm PES and one 100 000 MWCO PES, and flush the systems with 500 mL MQ water as described (see Vivaflow Assembly and operation, Fig. 21-1). After use wash cartridges according to producer manual and store 10% EtOH (4°C) (see Cleaning and storage of Vivaflow below)

Concentration of prokaryote and prokaryote free water (0.2 µm cartridge)

- Flush the Vivaflow 0.2 µm PES cartridge with 50 ml of a 500 mL sample
- Start concentrating the sample while collecting the prokaryote-free filtrate.
- At approx. 50 mL, turn off the pump and transfer the sample to a 50 mL Falcon tube before continuing concentration to approx. 20 mL.

Virus-free water (100 000 MWCO cartridge)

- Flush the Vivaflow 100 000 MWCO PES cartridge with 50 ml of a 200 mL aliquot of the prokaryote-free filtrate.
- Collect 150 mL of virus free filtrate

Virus production

- Dilute the prokaryote concentrate to 150 mL with virus-free filtrate
- Distribute ca 50 mL to each of three Falcon tubes
• Incubate at *in situ* temperature in the dark.
• Collect, fix and store 1.8 mL samples for viral and bacterial counts according to the FCM protocol at time zero and after approx. 3, 12, 24 and 48 h of incubation (make notes of exact incubation time).

**Viral decay**

1. Distribute 50 mL of the prokaryote-free water to each of three Falcon tubes
2. Incubate at *in situ* temperature in the dark.
3. Collect, fix and store 1.8 mL samples for viral and bacterial counts according to the FCM protocol at time zero and after approx. 3, 12, 24 and 48 h of incubation (make notes of exact incubation time).

**Vivaflow Assembly and operation**

1. Set up the system as illustrated below. Note the positioning of the flow restrictor on the return line.
2. Place 500 ml deionised water in a suitable reservoir and pump liquid through the system to purgee any air pockets. The recirculation rate should be 200–400 ml/min, and suitable flow should exit the filtrate line. If used, the pressure indicator should read approximately 2.5 bar.
3. Allow 400 ml to pass into the filtrate vessel. Check for any leakage at tubing connection points.
4. Drain the system and empty or replace the filtrate vessel
Cleaning and storage of Vivaflow

Cleaning

1. Flush the system with 200 ml of deionised water with the filtrate going to waste.
2. Place the feed, return and filtrate lines in a container with 250 ml 0.5 M NaOH (cleaning solution).
3. Recirculate at 50–100 ml/min for 30–40 minutes.
4. Drain the system and recirculate 250 ml of deionised water through the system for 5–10 minutes.
5. Drain and discard rinse solution and flush with a further 500 ml of deionised water with the filtrate going to waste. The system is now ready for further use.
To store Vivaflow after cleaning, fill module with deionised water and 10% ethanol. Seal inlet, outlet and filtrate ports and refrigerate at approximately 4°C.

For full description see Sartorius Lab Instruments Publication No.: SLU6097-e160609, Directions for Use Vivaflow 50 | 50R | 200

→ **Samples storage/shipment address:**

Store samples in -80°C during the cruise. Ship samples on dry ice using

World Courier (Norway) AS  
Industrivegen 20  
2069 Jessheim (Oslo)

Tel: 47 6394 6200; Fax: 47 6394 6201; Email: ops@worldcourier.no

**NOTE:** Make appointment for pick up well in advance of docking at the end of the cruise and coordinate with shipping of other frozen samples (e.g. Sterivex filters)

**Ship to:**

Institutt for biovitenskap, Biologen  
Thormøhlens gate 53A, 2. etg  
5006 BERGEN

**Literature:**

- Weinbauer, Rowe, Wilhelm, 2010, Determining rates of virus production in aquatic systems by the virus reduction approach., MAVE, chapter 1 pp 1-8
7.22 COCCOLITHOPHORES AND OTHER SMALL PLANKTON FOR SEM

**Measured parameter:** Qualitative and quantitative analysis of coccolithophores and other small plankton

**Method responsible:** Luka Supraha (UiO, luka.supraha@ibv.uio.no ), Bente Edvardsen (UiO, bente.edvardsen@ibv.uio.no)

**Description of parameter:** Qualitative and quantitative analysis of coccolithophores and other small plankton

**Methodological description:**

- **Special requirements/gear:** Vacuum filtration system for 25mm filters with a peristaltic pump.

- **Sampling depths:** Standard depths as for quantitative phytoplankton samples (including 5 m, DCM, 50, 100 m).

- **Chemicals used:** none

**Material:**

- Polycarbonate filters (25mm, pore size 0.8 µm)
- Cellulose nitrate membrane filters (25mm, pore size 3 µm)
- Vacuum filtration setup for 25 mm filters
- Tweezers, x2
- Gloves
- Plastic pipette, 2 mL
- Measuring cylinder (1L)
- Plastic bottle (1L)
- Millipore dishes (50 mm) for storing filters
- Waterproof marker
- Buffered freshwater (pH >7.4, buffered with ammonium) and filtered seawater (FSW)
Method:

- Use clean and dry equipment. Wear gloves.
- Mount the filtration unit. Add 2 mL buffered freshwater on top of the scint and turn on the vacuum pump shortly. Place the cellulose nitrate membrane on the scint, and then the polycarbonate filter on top of it. Filters should be well aligned and without wrinkles.
- Collect >1 liter of sample from the Niskin bottles in a plastic bottle. Mix the bottle gently (do not shake it) to ensure even distribution of material.
- Fill a cylinder with the desired volume, 250 mL -1 L, depending on cell density. The material should be visible, but not be as a thick layer on the filter.
- Pour the water into the filtration funnel. Turn on the pump. The flow should be steady and not too fast, and it is better to use weak vacuum. Write down the volume filtered per sample.
- When the filtration is done, remove the funnel and let the vacuum on, until no liquid is visible.
- Carefully transfer the upper polycarbonate filter to a labeled Millipore filter dish (sample side up) and discard the cellulose nitrate membrane. The filters should be dried in an incubator or oven set to ventilation program at 50°C for ca 2 h. The filter dishes with filters should be partially covered with a lid while in the incubator.
- Rinse all the equipment with freshwater and wipe off the tweezers between samples.

Samples storage/shipment address: Once dried, the filters should be stored in closed Millipore dishes. During transport and storage, it is important to keep the filters with the sample side up. The samples are brought to the University of Oslo, Department of Biosciences.

Literature:


7.23 MICROALGAL DIVERSITY BY CULTURING (SERIAL DILUTIONS)

**Measured parameter:** Microbial diversity

**Method responsible:** Bente Edvardsen (UiO, bente.edvardsen@ibv.uio.no), Luka Supraha (UiO, luka.supraha@ibv.uio.no)

**Description of parameter:** Isolation of microalgal cultures by serial dilutions

**Methodological description:**

→ **Special requirements/gear:** Cold room or cold plate, culture chamber or culture room

→ **Sampling depth(s):** Seawater samples from all sampling stations from 5 m and DCM depths, and diluted vertical net hauls.

→ **Chemicals used:** no other than algal medium (below).

**Material:**

- Multiwell cell culturing plates (96 wells, TPP), or 15 mL culturing glass tubes with screw lid. Each plate can be used for four samples.
- Suitable algal medium (e.g. IMR ½, L1, K or ES, 30 PSU) with salinity similar to the sea water sample. For isolation of species that thrive in oligotrophic waters the strength of the medium can, to start with, be reduced to 1/2-1/10 of the full medium. Dilute, if possible, the medium with sterile sea water from the actual locality.
- Automatic pipettes -1000 µL and -100 µL with tips
- Fresh seawater sample (or net haul sample for larger, abundant cells) or raw culture
Method:

- Distribute 270 µL algal medium to each well with the automatic pipette. Keep the plate at the same temperature as the water sample, e.g. by placing the plate in the fridge before use or on a cooling element. Mark a cross on the lid of the plate between column 6 and 7 and between row D and H, and the sample ID in each of the 4 squares.
- Add 30 µL of algal sample to the first well in a row (column A).
- Mix carefully the diluted sample in the first well in the row by pipetting up and down slowly with the 1 mL pipette. Transfer 30 µL from well A1 to A2. Mix, transfer 30 µL from well A2 to well A3 etc, all the way to well A6. Do the same with rows B-D with the same sample. Do the same with three other samples in the other squares of the plate.
- Mark the plate with sample ID and date.
- Put the plate in a clear plastic bag to avoid evaporation, and leave it in an incubator with white fluorescent light (ca 30-50 µmol photos m\(^{-2}\) s\(^{-1}\)) at suitable temperature (ca 4°C) for 2-4 weeks.
- Check the plates under an stereo microscope. From the wells with apparently uniform cultures a droplet is examined under the microscope at higher magnifications. If the culture is monoalgal and of interest, transfer it to a tube with algal medium.

→ **Samples storage/shipment address:** Multiwell plates containing serial dilutions should be kept in a culture room or an incubator with light/dark cycle at low temperature (4°C). During transportation put on a fitting lid and place in a cooling box with cooling elements and fill up with packing material to avoid tilting. Monoalgal cultures can be deposited and maintained at the Norwegian culture collection of algae (NORCCA) at UiO.

**Literature:**

7.24 DETERMINATION OF \textit{IN SITU} PRIMARY PRODUCTIVITY RATE \ USING 
THE 14C METHOD

\textbf{Measured parameter}: \textit{In situ} primary production rate

\textbf{Method responsible}: Rolf Gradinger (UiT, rolf.gradinger@uit.no), Philipp Assmy (NPI, philipp.assmy@npolar.no), Marti Amargant Arumi (UiT, marti.a.arumi@uit.no)

\textbf{Description of parameter}: Determination of the production rate of organic material by phytoplankton under \textit{in situ} conditions by using the $^{14}$C method.

\textbf{Methodological description}:

$\rightarrow$ \textbf{Special requirements/gear}: radioisotope container, permission to work with radioactive isotopes, in situ incubations through ice or deployed drifting buoys

$\rightarrow$ \textbf{Sampling depth(s)}: 5, 10, 20, 30, 40, 60, 90 m

$\rightarrow$ \textbf{Chemicals used}: 14C-sodium bicarbonate working solution (e.g. 20 µCi/ml), Hydrochloric acid (HCl 6M and 1M), NaOH (6N), Ethanolamine, Scintillation cocktail Ecotone

\textbf{Material}:

- Incubation bottles 250ml Polycarbonate
- Rope, buoy, carabiners
- GF/F filters
- Vacuum filtration manifold with waste bottles and vacuum pump and tubing
- Tweezers (2-3)
- Execator with fuming HCl (37%)
- Micropipettes and tips (One of each: 1000 µl, 200 µl and 20 µl)
- Cylinder (100 ml)
- Rinsing bottles (3x)
- Scintillation vials
• boxes to store scintillation vials for further processing back on land
• Trays for radioactive work
• Gloves (nitril size dependend on operator)
• Paper towels
• Radioactive waste containers (solid, liquid)
• Absorbing bench paper
• Light sensor on CTD
• Go Flow bottles or Niskin bottles on CTD
• 2 glass duran bottles 0.5L
• 14C-sodium bicarbonate working solution (e.g. 20 µCi/ml)
• Hydrochloric acid (HCl 6M and 1M)
• NaOH (6N)
• Ethanolamine

Assumption: DIC determined by other group, otherwise additional sampling needed.

➔ NOTE: Always wear safety goggles, a lab coat and gloves when working with 14C and any hazmat (e.g. scintillation cocktail). Collect all chemical/radioactive waste in clearly labeled waste containers. Keep a clear protocol on how 14C has been used. Do not mix radioactive waste with other chemical waste.

Method:
1) Preparation of incubation bottles flasks
   • Use the provided 250ml culture tissue flasks. Rinse flasks between measurements with 10%HCl, followed by distilled water. Dry before re-use.
2) Taking of water samples/ice sample:
   • Water samples are taken at the same depth as sediment traps are deployed (5, 10, 20, 30, 40, 60, 90 m). This should be matched by Chl and POC data from the same depths. The selection of depth should cover
the approximate light levels from >90 to ca 1% or less of surface PAR). PAR in the water should be determined during a CTD cast. If a Chl a maximum exists, one of the samples should be taken in the chl a max. If melt ponds exist on the ice, take on set of samples from the melt pond.

- For water sample form ocean or for melt pond sample: Take one light and one dark bottle from each sampling depth. Rinse bottles three times before filling. Keep the samples as shaded/dark as possible during treatments.
- For ice sample: take one complete ice core. Cut off and use only the bottom 1cm. Protect from direct sun light (black trash bag). Cut inside a dark tent. Crush the ice (likely it will be in small pieces anyway if skeletal layer present). Place the sample into a clean 500ml duran bottle. Add 400m filtered sea water at 0-2degC from the surface. Shake to distribute the ice evenly and/or wait until it is melted. Use the melted sample as a water sample. Important: Note the final volume of the melted water sample to calculate the volume of the added ice sample. Take a subsample for chl a and POC.
- Bring the samples in a dark container into the isotope lab for spiking.
- Note: make sure that for each sampling depth/melt pond/sea ice samples, also samples are processed for chl a, POC. Adjustment of sampling depths might be needed to match sampling by other groups. More important to cover the above given light profile range for the water column samples than precise depths.

3) Spiking with NaH\textsuperscript{14}CO\textsubscript{3} solution:

- Achieve a final concentration of 0.1µCi/ml inside your incubation bottle. In case of low biomass in high Arctic samples you can also use a final concentration of 0.4µCi/ml (e.g. Ferland et al. 2011, http://dx.doi.org/10.1016/j.jmarsys.2011.03.015).
- Incubation bottles are filled with a measuring cylinder. The flasks should be filled up to the neck, leaving an air bubble in the flask. One dark flask from each original sample should be incubated. All places on the incubator wheel must be taken by bottles. So empty spaces should be filled with bottles filled with DI water.
4) Determination of total labelled carbon added:
   • After mixing thoroughly add 250 µl (x2) from each spiked bottle into 20ml scintillation vials which contain 250µl of ethanolamine. Keep at room temperature until measurement with scintillation counter. Prior to counting: add 10 ml of scintillation cocktail–shake the sample.

5) Incubation
   • Incubate for 24 hours. Ideally incubation should go from dawn to dawn the next day to estimate net primary productivity. Attach the bottles with carabiners to the sediment trap rig for drifting incubations. For incubation in holes through the ice: use a bottom weight and a line with loops to attach the carabiners. Cover the hole in the ice with the white disk. Secure the line with two ice screws.

6) Filtration: particulate and dissolved PP
   • After incubation the flask contents are filtered immediately. Maintain darkness or very low light intensities while processing the samples. In case of high algal biomass or high sedimentation load it might be needed to filter a subsample. A defined portion should be taken and filtered.
   • Glass-fibre filters (GF/F, Ø 25 mm) should be used. To avoid any contamination of the filter edges, prewetted filters should be used. The suction pressure should not exceed 30 kPa during filtration. The filters should be rinsed once with a small volume (5 ml) of filtered seawater from the original sample (use filtrate of the chlorophyll-a measurement!). Note: This rinsing step is debated and some groups do not rinse the samples.

7) DOC production samples
   • If DOC is going to be measured use the following sub-sampling from the spiked sample:
   • Filter 3ml of the incubation from each incubation bottle through a GF/F or 0.2 or 0.4 µm filter into a scintillation vial. You can use a Syringe filter for this. Add 500 µl of 6N HCl. Leave in the fume hood for 24 hours to
release of $^{14}$CO$_2$. Neutralize the sample with addition of 500µl of 6N NaOH. Add 15ml of scintillation cocktail for immediate measurement in scintillation counter.

8) Post-Processing of particulate production filters:

- The filters are placed in the scintillation vials under the fume hood with a drop of fuming HCl (37% or 6N) for 24 h to eliminate all the non-incorporated $^{14}$C. Once they are dry 6 ml of Ecolmune Scintillation cocktail are added. Following addition of scintillation liquid, the samples should be kept in dark for at least 3 hours to reduce chemiluminescence.

9) Scintillation counting:

- Radioactivity of the samples is determined by a liquid scintillation counter of beta particles at the University of Tromsø.
- The total carbon uptake is calculated from the equation:

$$\frac{dP}{dt} (\mu g C L^{-1} hr^{-1}) = \frac{dpm (a) \times total^{12}CO_2 (c) \times 12 (d) \times 0.06 (e) \times k1 \times k2}{dpm (b)}$$

Where

(a) = Sample activity (minus back-ground), dpm

(b) = Total activity added to the sample (minus back-ground), dpm

(c) = Total concentration of $^{12}$CO2 in the sample water, µmol/L (or µM)

(d) = The atomic weight of carbon

(e) = A correction for the effect of 14C discrimination

$$k1 = \text{subsampling factor (e.g. sample 50 ml, subsample 10 ml: } k1 = \text{subsample factor } 50/10 = 5)$$
k2 = time factor (e.g. incubation time 125 minutes: k2= 60/125= 0.48)

The results will be given as $\mu g \cdot C \cdot L^{-1} \cdot h^{-1}$ per irradiance level.

10) Check of $^{14}$DOC in the stock solution:

- at beginning and end of the cruise fill 50µl of stock solution into scintillation vial.
- Add 0.5ml of 6N HCl and place under fume hood for 24 hours.
- Neutralize with 0.5ml of 6N NaOH.
- Add 15ml of Ecotone scintillation cocktail and keep vial in the dark until counting.

11) Calculation of total PP:

- Daily production rates from each depth are integrated to using trapezoidal integration. The rate nearest to the surface is assumed to be constant up to 0m depth. The difference of DIC uptake between the dark bottle and the dark bottle with nitrification inhibitor is the DIC uptake based on nitrification. The uptake of DIC in the bottle with nitrification inhibitor is assumed to be solely by anapleurotic heterotrophic processes.

7.25 DETERMINATION OF PRIMARY PRODUCTIVITY RATE IN RELATION TO LIGHT INTENSITY (P VS I CURVES) USING THE $^{14}$C METHOD

**Measured parameter:** Primary production rate

**Method responsible:** Rolf Gradinger (UiT, rolf.gradinger@uit.no), Philipp Assmy (NPI, philipp.assmy@npolar.no)

**Description of parameter:** Determination of the production rate of organic material by phytoplankton under defined light intensities using the 14C method.
Methodological description:

⇒ **Special requirements/gear:** radioisotope container, permission to work with radioactive isotopes, incubator onboard in cold room close to in situ temperature

⇒ **Sampling depth(s):** Subject to change depending on context, physical and biological in situ measurements temporary depths, 10, 25, 50 75, 100, DCM and ca 2 m above sediment-water interface

⇒ **Chemicals used:** 14C-sodium bicarbonate working solution (e.g. 20 µCi/ml), Hydrochloric acid (HCl 6M and 1M), NaOH 6N, Ethanolamine

Material:

- Hydrobios ICES P vs I Incubator
- Incubation bottles with light screens
- GF/F filters
- Vacuum filtration manifold with waste bottles and vacuum pump and tubing
- Tweezers (2-3)
- Execator with fuming HCl (37%)
- Micropipettes and tips (One of each: 1000 µl, 200 µl and 20 µl)
- Cylinder (100 ml)
- Rinsing bottles (3x)
- Scintillation vials
- boxes to store scintillation vials for further processing back on land
- Trays for radioactive work
- Gloves (nitril size depended on operator)
- Paper towels
- Radioactive waste containers (solid, liquid)
- Absorbing bench paper
- Light sensor
- Thermometer
- 2 glass Duran bottles 0.5L
- 14C-sodium bicarbonate working solution (e.g. 20 µCi/ml)
• Hydrochloric acid (HCl 6M and 1M)
• NaOH 6N
• Ethanolamine

→ Assumption: DIC determined by other group, otherwise additional sampling needed.

→ **NOTE**: Always wear safety goggles, a lab coat and gloves when working with $^{14}$C and any hazmat (e.g. scintillation cocktail). Collect all chemical/radioactive waste in clearly labeled waste containers. Keep a clear protocol on how $^{14}$C has been used. This needs to be reported after the cruise.

Method (based on ICES Primary Productivity Protocol):

1) Placement of the incubator:

• Place the incubator in a dark cold room (no additional light sources) close to in-situ temperature. Use 10 fluorescent tubes (will provide approximately a light intensity of 500µE m$^{-2}$ s$^{-1}$). Check with PAR sensor and note the correct 100% light value.

![HYDRO-BIOS ICES incubator](www.hydrobios.de)
2) Preparation of flasks

- Use the provided 50ml culture tissue flasks. Rinse flasks between measurements with 10%HCl, followed by distilled water. Dry before re-use.
- Do a minimum of six different light intensities for a P vs I curve from one water depth or one ice core sample.

3) Selection of sample

a. Water sample: take a water sample from either the chlorophyll a maximum (if present) or from the middle of the surface mixed layer (if no maximum present).

b. Ice sample: take one complete ice core. Cut off and use only the bottom 1cm. Crush the ice (likely it will be in small pieces anyway if skeletal layer present). Place the sample into a clean 500ml Duran bottle. Add 400ml filtered sea water at 0-2degC from the surface. Shake distribute the ice evenly and wait until it is melted. Use the melted sample as a water sample. Important: Note the final volume of the melted water sample to calculate the volume of the added ice sample. Take a sub sample for chl a and POC.

c. Melt pond sample: if melt ponds are available on the ice, it should be treated like a water sample from the ocean.

4) Spiking with NaH\(^{14}\)CO\(_3\) solution:

- Spike the entire volume of collected water with \(^{14}\)C in a Duran glass bottle (e.g. if you use six light intensities plus one dark bottle you will need 7*50=350ml spiked sample).
- Achieve a final concentration of 0.1µCi/ml inside your incubation bottle. In case of low biomass in Arctic samples you can also use a final concentration of 0.4µCi/ml (e.g. Ferland et al. 2011, http://dx.doi.org/10.1016/j.jmarsys.2011.03.015).
- Incubation bottles are filled with a measuring cylinder. The flasks should be filled up to the neck, leaving a small air bubble in the flask. One dark flask from each original sample should be incubated.
• All places on the incubator wheel must be taken by bottles. So empty spaces should be filled with bottles filled with DI water.

5) Determination of total labelled carbon added:

• After mixing thoroughly 100 μl (x2) of the spiked solution are placed in two small scintillation vials and 6 ml of scintillation cocktail is added immediately if available.
• In case direct counting is impossible the inorganic 14C should be mixed with ethanolamine by pipetting 0.25 ml of sample with added activity together with 0.25 ml of ethanolamine.
• 6ml of the scintillation cocktail can be added later and radioactivity determined in a scintillation counter.

6) Incubation

• The incubation time should be two hours, and the rotor should rotate ca 10 rotations/min. Note precise incubations times (by minutes). Check proper functioning of the incubator, so that it does not stop – can be tricky....

7) Filtration: particulate and dissolved PP

• After incubation the flask contents are filtered immediately. In case of high algal biomass or high sedimentation load it might be needed to filter a subsample. A defined portion should be taken and filtered.
• Glass-fibre filters (GF/F, Ø 25 mm) should be used, To avoid any contamination of the filter edges, prewetted filters should be used. The suction pressure should not exceed 30 kPa during filtration. The filters should be rinsed once with a small volume (5 ml) of filtered seawater from the original sample (use filtrate of the chlorophyll-a measurement!).

8) DOC production samples

• If DOC is going to be measured use the following sub-sampling from the spiked sample:
• Filter 3ml of the incubation from each incubation bottle through a GF/F or 0.2 or 0.4 µm filter into a scintillation vial. You can use a Syringe filter for this. Add 500 µl of 6N HCl. Leave in the fume hood for 24 hours to release of $^{14}$CO2. Neutralize the sample with addition of 500µl of 6N NaOH. Add 15ml of scintillation cocktail for immediate measurement in scintillation counter.

9) Post-Processing of particulate production filters:

• The filters are placed in the scintillation vials under the fume hood with a drop of fuming HCl (37% or 6N) for 24 h to eliminate all the non-incorporated $^{14}$C. Once they are dry 6 ml of Ecolume Scintillation cocktail are added. Following addition of scintillation liquid, the samples should be kept in dark for at least 3 hours to reduce chemiluminescence.

10) Scintillation counting:

• Radioactivity of the samples is determined by a liquid scintillation counter of beta particles at the University of Tromsø.

• The total carbon uptake is calculated from the equation:

$$\frac{dP}{dt} (\mu g C.L^{-1}.hr^{-1}) = \frac{dpm (a) \cdot total^{12}CO_2 (c) \cdot 12 (d) \cdot 1.05 (e) \cdot k1 \cdot k2}{dpm (b)}$$

Where

(a) = Sample activity (minus back-ground), dpm

(b) = Total activity added to the sample (minus back-ground), dpm

(c) = Total concentration of 12CO2 in the sample water, µmol/L (or µM).

(d) = The atomic weight of carbon

(e) = A correction for the effect of 14C discrimination.
k1 = subsampling factor (e.g. sample 50 ml, subsample 10 ml: k1=subsample factor 50/10= 5)

k2 = time factor (e.g. incubation time 125 minutes: k2= 60/125= 0.48)

- The results will be given as $\mu$g C·L$^{-1}·h^{-1}$ per irradiance level and as well as the photosynthesis at light saturation (Pmax), the maximum light utilization coefficient (a), and light saturation parameter $E_k$, from the P-E curve (see below for equation).

11) Check of $^{14}$DOC in the stock solution:

- at beginning and end of the cruise fill 50µl of stock solution into scintillation vial. Add 0.5ml of 6N HCl and place under fume hood for 24 hours. Neutralize with 0.5ml of 6N NaOH. Add 15ml of Ecotone scintillation cocktail and keep vial in the dark until counting.

12) Calculation of PP:

- The carbon uptake ratios (y axis) are plotted against the light intensity received by each bottle (x axis). Fit the curve based on the equation of Platt and Gallegos (1980).

$$P = P_{max} (1- e^{-\alpha I/P_{max}}) e^{(-\beta I/P_{max})}$$

- Where $P$ is the photosynthetic rate at irradiance $I$, $\alpha$ is the light-limited initial slope, $\beta$ is a parameter describing the reduction in photosynthetic rate at high irradiance, and $P_{max}$ is a parameter equivalent to the light-saturated rate of photosynthesis when there is no photoinhibition.
- Manual for Marine Monitoring in the COMBINE program of HELCOM
- Part C: Annex C-5 Phytoplankton primary production
- Bouman et al. 2018: Earth Syst Scie Data 10: 251-266
7.26 WATER SAMPLING FOR BENTHIC EXPERIMENTS (NISKIN ROSETTE)

**Measured parameter:** Water sampling for benthic experiments chapter 10.3

**Method responsible:** Elisabeth Alve (elisabeth.alve@geo.uio.no) and Silvia Hess (silvia.hess@geo.uio.no), Paul Renaud (APN, per@akvaplan.niva.no), Lise Øvreås (UiB, lise.overas@uib.no)

**Description of parameter:** Water sampling for benthic experiments chapter 10.3

**Methodological description:**

- **Special requirements/gear:** Niskin bottles on CTD
- **Sampling depth(s):** Bottles need to be fired as close to the seafloor as possible in order to obtain bottom water
- **Chemicals used:** none

**Method:**

- Approximately 100L of bottom water per station where respiration incubations will take place. Note: total volume need not be from a single deployment.
- Data needed: latitude / longitude, water depth.
- Bottles need to be fired as close to the seafloor as possible in order to obtain bottom water.
- Upon recovery to the ship, open bottles and use a funnel with a coffee filter to pour water into canisters.
- Keep canisters in the cold rooms in the dark where the respiration experiments will take place to get them to the temperatures that will be used for the experiments.
- The water will be used for filling up the cores used for the respiration experiments.
7.27 MICROBIAL COMMUNITIES EXPERIMENTAL DESIGN

7.27.1 Grazer-exclusion experiment

**Measured parameter:** Abundance, activity and diversity of microorganisms (<0.8µm; <3µm; <90µm)

**Method responsible:** Oliver Müller UiB (oliver.muller@uib.no); Lasse Mork Olsen (Lasse.Olsen@uib.no); Aud Larsen (aula@norceresearch.no); Gunnar Bratbak (Gunnar.Bratbak@uib.no)

**Description of parameter/Aim:** Surface water samples is gentle reverse filtrated to retain organisms of different size fractions (<0.8µm; <3µm; <90µm) and are incubated for six days. Samples for abundance and diversity analysis are taken at different frequencies throughout the incubation period.

**Methodological description:**

→ **Special requirements/Gear:**

- Reverse filtration tubes, nylon filter mesh (<3µm; <90µm) and 0.8µm (142mm) polycarbonate membrane filters
- Set up experiments (filtration and filling bottles) in a cold room (301-chilled lab)
- Incubation in Termaks-incubator with close to in-situ light (ca. 30-50 μmol photons m-2 s-1) and at temperatures close to in-situ at the different stations (from 0-6°C)

→ **Sampling depths:** Process stations P1, P4 and PICE at chlorophyll a maximum (DCM or 20m)

→ **Chemicals used:** For sub-sampling as described in respective separate protocols (7.18 Flow cytometry sampling; 7.19 Bacterial biomass production; 7.13 Light microscopy phytoplankton/protists)
Method:

**NOTE:** Since only trained personnel will conduct these experiments, the protocol provides only a shortened outline.

1. **Preparations**
   - Make sure that all the bottles, reverse-filtration equipment and carboys have been acid-washed (5% HCL) and rinsed in Milli-Q water (3 times) and are labeled accordingly (e.g. “P1, <90µm, A, 15.02.2020”).

2. **Sampling for the experiment**
   - Fill up each 20L in two carboys from the DCM at station P1, P4 and PICE and store the carboys cold until further processing when they cannot be processed at once.

3. **Setting up the experiment**
   - Take the carboys with the water you sampled first and gently mix it and transfer it into the large HDPE bucket. Then use the plexi-tubes for reverse filtration with the 90µm net mounted to it (always start with the largest size fraction). Transfer 12L of the filtrate from inside the tube into a new 20L container using silicone tubing and from there staggered fill 3+6 (including copepod addition) x 1L PC bottles (Fig. 7.27.1-1) controlled via an opening/closing system integrated to the tubing. The rest is stored cold until later to take samples for generating T0 data of different parameters as stated in Table 1.
Repeat reverse filtration process, now with the 3µm net and after that with the 0.8µm polycarbonate filter via gravity flow through a tripod filtration device. For these two size fractions only each 3 1L bottles have to be filled and thus only 5L have to be filtered and transferred into a smaller 5L container. Now you should have filled 9 (15) x 1L PC bottles (3 for each size fraction and 2x3 extra for the copepod treatment of the <90µm fraction) and have 3 times 2L left which is stored to be processed later.

- Transport all the bottles to the incubator where they will be incubated at close to in-situ conditions regarding temperature and light.

4. Taking sub-samples for different parameters at T0, throughout the incubation and at the end of the experiment (Important: different sampling frequencies for the different parameters)
Tab. 27.7.1-1 Overview of the parameters measured in course of the grazer exclusion experiment at different time points

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Protocol</th>
<th>Frequency</th>
<th>Volume</th>
<th>#Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>FCM</td>
<td>7.16 [Flow cytometry sampling]</td>
<td>T0, T2, T3, T4, T6</td>
<td>6ml</td>
<td>T0: 21</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2.: 102 (x5)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>T6: 102</td>
</tr>
<tr>
<td>Molecular sample</td>
<td>X.XX [Filtration for DNA samples using 0.22µm Durapore filters]</td>
<td>T0, T3, T6</td>
<td>T0: 1L</td>
<td>T0: 9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>T3: 200ml</td>
<td>T3: 34</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>T6: 500ml</td>
<td>T6: 34</td>
</tr>
<tr>
<td>Bacterial production</td>
<td>7.17 [Bacterial biomass production]</td>
<td>T0, T3, T6</td>
<td>T0: 5ml</td>
<td>T0: 18</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>T3: 5ml</td>
<td>T3: 9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>T6: 5ml</td>
<td>T6: 18</td>
</tr>
<tr>
<td>Flow-Cam/Microscope</td>
<td>7.12 [Light microscopy phytoplankton/protists]</td>
<td>T0 + T6</td>
<td>T0: 300ml</td>
<td>T0: 9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>T6: 100ml</td>
<td>T6: 10</td>
</tr>
<tr>
<td>Chl a</td>
<td>7.10 [Water sampling for chlorophyll a]</td>
<td>T0 + T6</td>
<td>T0: 300ml</td>
<td>T0: 9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>T6: 100ml</td>
<td>T6: 16</td>
</tr>
<tr>
<td>Macronutrients</td>
<td>7.9 [Inorganic nutrients]</td>
<td>T0 + T6</td>
<td>T0: 30ml</td>
<td>T0: 9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>T6: 30ml</td>
<td>T6: 16</td>
</tr>
<tr>
<td>TOC</td>
<td>7.4 [TOC]</td>
<td>T0 + T6</td>
<td>T0: 30ml</td>
<td>T0: 9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>T6: 30ml</td>
<td>T6: 16</td>
</tr>
<tr>
<td>POC</td>
<td>7.5 [POC]</td>
<td>T0 + T6</td>
<td>T0: 50ml</td>
<td>T0: 9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>T6: 50ml</td>
<td>T6: 16</td>
</tr>
</tbody>
</table>

→ Sample storage/shipping address: Samples that are taken during the experiment are stored according to the conditions stated in their respective protocols. The shipping address for all samples is:
Short-time drifting Sediment traps

**Measured parameter:** Sinking particle flux

**Method responsible:** Marit Reigstad (UiT, marit.reigstad@uit.no), Miriam Marquardt (UiT, miriam.marquardt@uit.no), Yasmin Bodur (UiT, yasmin.v.bodur@uit.no)

**Description of parameter:** Water collection via sediment trap deployment. Water will be used for analysis of several parameters: From each depth, Chla (size fractionated), POC/PON, stable isotope, phytoplankton and fecal pellet samples will be taken.

**Methodological description:**

- **Special requirements/gear:** Sediment trap type: KC-Denmark
- **Sampling depth(s):** 30, 40, 60, 90, 120 and 200m.
- **Chemicals used:** 4% GA-Lugol’s Solution, buffered formalin (2%), Ethanol

**Method:**

- Duration of deployment ~24h
- Depths of the sediment trap cylinders are 30, 40, 60, 90, 120 and 200m.
- 2-4 cylinders per depth
- Cylinders are filled with pre-filtered deep seawater (GF/F filtrate, about 0.7µm pore size) prior to deployment.
**Processing of core parameters:**

- From each depth, Chla (size fractionated), POC/PON, stable isotope, phytoplankton and fecal pellet samples will be taken. Water from the trap are pooled before subsampling. The pooled sample needs to be thoroughly but carefully inverted before taking subsamples as the particles can sink down very quickly.
- **Chla** 100-500ml (light coloration) in triplicate
- Total Chla is filtered onto GF/F filters, Chla > 10µm onto 10µm PC filters. Otherwise the protocol for Chla samples will be followed.
- **POC/PON** 100-500ml (light coloration) in triplicate onto pre-combusted GF/F filters, wrapped in pre-combuster aluminium foil, and stored at -20°C following the protocol for POC/PON samples.
- POC/PON and stable isotopes will be analyzed from the same filter simultaneously on an IRMS coupled to an elemental analyzer.
- **Phytoplankton** 100ml fixed with 4% GLA-Lugol's Solution (2ml) and stored at 4°C in brown glass bottles.
- **Fecal pellets** 100-200ml fixed with buffered formalin (2% final concentration) and stored at 4°C in Kautex bottles.

**For sampling of additional components:**

- This can be solved by deploying 4 cylinders at some depth to increase sample volume for extended sampling. Below extended sampling for 2018 is given.
- At 40, 90, and 120m 2 cylinders per depth will be deployed.
- At 30, 60 and 200m depth a full set of 4 cylinders will be deployed. At those depths, samples for the analysis of two types of extracellular polymeric substances (EPS) will be taken in addition, transparent exopolymeric substances (TEP) and Coomassie stainable particles (CSP).
- **TEP/CSP** concentration, 100-300ml in triplicate onto 0.4µm PC filters using a peristaltic pump. Filters are placed into filter holders and attached to the tubes of the pump. After the sample passed through. The filters are rinsed with air, MilliQ water, and air by attaching a syringe to the filter holder. Filters are stored in plastic cups and stored at -20°C.
The filtrate (50ml centrifuge tube) is collected to enumerate the dissolved concentration of TEP/CSP.

- **CARD-FISH** to identify bacteria living attached to exopolymeric particles. 100-300ml are fixed with buffered formaldehyde (ACS grade, 2% final concentration) for up to 24h at 4C in the dark. Samples are filtered onto 0.2µm PC filters (using filter holders), rinsed with 70% Ethanol using a syringe and placed surface up in a petri dish. Petri dishes are sealed with parafilm after all the liquid has evaporated. Samples are stored at -20C.
- In addition, samples for TEP/CSP will be taken from **suspended water** at the corresponding depths.

9 Net sampling

9.1 PHYTOPLANKTON NET HAUL SAMPLING

**Measured parameter:** Qualitative collection of protists with a phytoplankton net

**Method responsible:** Rolf Gradinger (UiT, rolf.gradinger@uit.no), Philipp Assmy (NPI, philipp.assmy@npolar.no), Bente Edvardsen (UiO, bente.edvardsen@ibv.uio.no), Luka Supraha (UiO, luka.supraha@ibv.uio.no)

**Description of parameter:** Qualitative collection of protists for: 1) documentation based on live material on board through live microscopy and video/imaging, 2) collection of concentrated material for scanning electron microscopy (SEM) analysis, 3) enriched fixed samples for sharing with taxonomic specialists, and 4) establishing of protist cultures.

**Methodological description:**

→ **Special requirements/gear:**

- Get permission from crew prior to deployment, especially if net is not towed by winch!
- Fume hood for addition of fixatives.
- MSDS sheets for formaldehyde and hexamethylenetetramine
- Waste container for gloves (preferred NITRILE)

**Sampling depth(s):** Vertical haul (upper 20 m) with a 10 µm hand-net (see below)

**Chemicals used:** 20% hexamethylenetetramine-buffered formaldehyde (final concentration 2%), Lugol's (final concentration 1%), glutaraldehyde (EM grade 25%, final concentration of 1%)

**Method:**

- Label three 100 mL brown glass bottles for fixed samples and one clear 200 mL bottle for live sample with ID.
- Connect the hand-net to appropriate rope and attach a weight (adjust weight based on type of net used; ask crew for e.g. shackles) below the cod end if needed. Use a bucket for the rope to avoid entanglement.
- Make sure that the valve of the cod end is closed when lowering the hand-net.
- Lower the net slowly to 30m depth (mark the rope), and retrieve with a slow and constant movement (<0.2 m/s) back to above the water surface.
- Wait until <300 ml are left in the cod end (less than entire cod end volume, dependent on hand net used), open valve and drain hand-net sample into 500 ml measuring cylinder. Fill up measuring cylinder containing the sample to 300 ml mark by flushing the mesh of the cod end with filtered sea water from squeeze bottle.
- The fixative should be added to the empty brown glass bottles prior to addition of the samples. Under fume hood, add to brown bottle 1) 10 ml of 20% hexamethylenetetramine-buffered formaldehyde (final concentration of 2%), brown bottle 2) 1 mL Lugol's (1% final concentration), and brown bottle 3) 1 mL Lugol's (1% final concentration).
• Wear gloves and work in a fume hood protection when adding formaldehyde and glutaraldehyde!
• Fill content of the measuring cylinder into three labeled and fixative containing 100 ml brown glass bottles (see picture below), ca 90 mL in each.
• Brown bottle 2: After ca 5 minutes add 4 mL glutaraldehyde (EM grade 25%, final concentration of 1%) under fume hood.
• Store fixed samples dark and cold (see below), do not freeze!
• For live samples for microscopy and cultures: Take separate net casts and fill sample into clear 200 mL bottle. Do not add fixatives and keep sample alive in cold room with white illumination (dark/light as needed).

→ Fixed samples storage/shipment address: Store glutaraldehyde fixed samples in fridge or cold room during the cruise (at 4°C). Make sure that bottles are safely packed for transport (hard-casing boxes with bubble-foil filling). Formaldehyde-fixed samples should be stored at room temperature and Lugol's fixed samples should be kept at temperature about 4°C.

**Fig. 9.1-1** HYDRO-BIOS phytoplankton hand net (www.hydrobios.de), left hand side, or IMR net with steel frame

**Fig. 9.1-2** 50 ml brown glass bottle (right hand side)
9.1.1 Microalgal diversity by culturing (capillary isolation)

Measured parameter: Capillary isolation of microalgal cells for culturing

Method responsible: Bente Edvardsen (UiO, bente.edvardsen@ibv.uio.no), Luka Supraha (UiO, luka.supraha@ibv.uio.no)

Methodological description:

→ Special requirements/gear: Inverted microscope, cold room or cold plate, culture chamber or culture room with white illumination.

→ Sampling depth(s): Vertical phytoplankton net hauls (0-30m depths) from all sampling stations

→ Chemicals used:

Material:

- Pasteur pipettes that are heated and pulled out into a very thin tube (capillary), preferably with a 30° angle about 5-10 mm from the point. The diameter of the point should be 3-5 x the diameter of the cells to be isolated.
- Silicon tubing fitting to the wide end of the pipette and with a tube tip as a mouthpiece in the other end.
- Sterile Petri dishes in plastic or glass slides that must be clean.
- 50 mL tube with distilled water and 50 mL tube with algal medium
- Multiwell plates (96 wells, TPP, tissue culture testplate) with fitting lid, or glass tubes with screw lid, in rack, filled with algal medium.
- Inverted microscope with 4/5x, 10x, 20x and 40x objectives.
- Temperature control as far as possible: keep samples, petri dishes and solutions cooled on cooling elements or in a cooling box if sample is from a cold environment. If available use a microscope with a lamp with low heat emission (led).
- Plankton concentrated by net haul or a raw culture. Collect and keep the sample in a non-toxic bottle and avoid rapid changes in temperature or light shock during transportation and storage.
- The algal medium IMR 1/2 can be used for diatoms in coastal waters. For dinoflagellates and other flagellates that can be difficult to grow it
may be worth to try other media such as ES- K- or L1-medium. Addition of soil extract can help. It is important to avoid salinity chock. For isolation of species that thrive in oligotrophic waters the strength of the medium can, to start with, be reduced to 1/2-1/10 of the ordinary level. Dilute, if possible, the medium with sterile sea water from the actual locality.

Method:

1. Add 300 µL suitable algal medium into each well of a 96 well plate. Add small droplets of medium in a row in a Petri dish (about 8) or on a glass slide (3). Add a droplet of the algal sample in the first droplet.
2. Fill some medium from a tube into the capillary by capillary-forces without sucking. By this it is easier to control the water movements in point 3.
3. Localize a cell under the microscope and pick up the cell (using 4 or 10x objective) with the capillary by gently sucking on the mouthpiece, together with as little medium and other cells as possible. Transfer the cell into the next sterile droplet of medium with a gentle blow.
4. Suck up distilled water into the capillary to kill other algae that stick to the pipette and blow it out again.
5. Repeat step 2-4 with the actual droplet, so that the cell is transferred to the next unused droplet and continue until the cell is seen alone in the next droplet, upon a quick look into the microscope.
6. Suck up the cell and transfer it to a well in the Multiwell plate with unused medium. Mark the used well and make a table with information about cell ID and origin.
7. When the desired cells are isolated place the plate in a clear plastic bag to avoid evaporation and place it for growth in a culture room or incubator with light and correct temperature.
8. Check the Multiwell plates in an inverted microscope (or a stereo microscope) after about 3-4 weeks depending on temperature and growth rate. From the wells with apparently uniform cultures a droplet is examined under the microscope at higher magnifications.
9. It is important to work fast to avoid heating and desiccation that will harm or kill the cell. It is better to take a chance and isolate many cells within a short time with the danger of including a “co-passenger” in some tubes, than to stress a few cells intensively to be safe that they are clean. You can also isolate several cells of the same species in one tube/well and let them start to grow, and from this culture repeat the isolation to obtain a clonal culture (originating from one single cell).

→ **Samples storage/shipment address**: Multiwell plates containing serial dilutions should be kept in a culture room or an incubator with light/dark cycle at low temperature (4°C). During transportation put on a fitting lid and place in a cooling box with cooling elements and fill up with packing material to avoid tilting. Transported to University of Oslo.

**Literature:**


### 9.1.2 Protist diversity by single cell PCR

**Measured parameter:** Isolation of protist cells for single cell PCR

**Method responsible:** Bente Edvardsen (UiO, bente.edvardsen@ibv.uio.no), Luka Supraha (UiO, luka.supraha@ibv.uio.no)

**Description of parameter:**

**Methodological description:**

→ **Special requirements/gear**: Inverted microscope, cold room or cold plate, freezer (-20°C or -80°C)

→ **Sampling depth(s)**: Vertical net hauls from all sampling stations

→ **Chemicals used**: none
Material:

- Inverted microscope with objectives x4 or 5, x10, x20 and x40 magnification and x10 in the oculars with camera for photographing picked cells.
- Sterile plastic Petri dishes (10 cm diameter)
- Automatic pipette with sterile filter tips, 0.5-10 µL
- Pasteur pipettes drawn out over a flame to a thin capillary with a ca 30 ° angle 1-2 cm from the tip. The diameter of the opening should be about 4-5 times the cell diameter.
- Silicon pipeline fitting to the wide end of the Pasteur pipette and a mouth piece (e.g. pipette tip) fitted in the other end.
- Sterile filtered sea water or algal medium in 50 mL tube in a rack
- Distilled water in a 50 mL tube (for cleaning capillary)
- PCR-water (Eppendorf) in 15 mL or Eppendorf tube
- 0.2 mL PCR-tubes in cooling rack
- Cooling blocks, x2 (to cool Petri dishes and racks with samples)
- Marking pen
- Diluted net haul sample (kept cool)

Method:

1. Pick cells by capillary isolation (see separate protocol).
2. For cells sticking to the bottom of a Petri dish, such as dinoflagellates: Transfer the cell into a droplet with sterile sea water (or medium) in a Petri dish. Remove most or all of the water (but not your dinoflagellate cell) in the first drop by a clean capillary (cleaned in sterile distilled water) and quickly add a new drop of sterile sea water on top of the cell (use a sterile pipette). Repeat this washing at least once with sterile sea water, or until no other cells are seen in the same droplet. Many dinoflagellates tend to stick to the bottom of the Petri dish and then this is the easiest way. If the cells do not stick to the bottom, e.g. planktonic diatoms or flagellated haptophytes or cryptophytes, transfer a single cell from one drop to another until you have only one cell in the droplet.
3. Prior to the last washing step the cells are examined and photographed in an inverted light microscope at 200-400 x magnification.
4. Remove all water with capillary and add 10 µl of PCR-water to the cleaned cell with an automated pipette, and the cell and water is then transferred to a 0.2 µL PCR tube and frozen to -20°C. Mark the tube with a code and collect information about the sample (sample ID, species etc) in a table. The cells can be stored up to several months prior to the PCR-reaction.

→ Samples storage/shipment address: PCR tubes with isolated single cells must be kept frozen at -20°C prior to the PCR reaction.

Literature:


9.2 ZOOPLANKTON SAMPLING

**Tab. 9.2-1** Overview of zooplankton sampling during Nansen Legacy cruises from August 2019 onwards.

<table>
<thead>
<tr>
<th>Gear</th>
<th>Depth</th>
<th>Haul</th>
<th>Station</th>
<th>Wire time (hrs)</th>
<th>No. depth</th>
<th>No. stations</th>
<th>No. samples</th>
<th>Taxa</th>
<th>Responsible</th>
</tr>
</thead>
<tbody>
<tr>
<td>Multinet 64 µm</td>
<td>bott-200-100-50-20-0m</td>
<td>vertical</td>
<td>all</td>
<td>1</td>
<td>5</td>
<td>7</td>
<td>35</td>
<td>Small mesozooplankton</td>
<td>RF3: Camilla Svensen</td>
</tr>
<tr>
<td>Multinet 180 µm</td>
<td>bott-200-100-50-20-0m</td>
<td>vertical</td>
<td>all</td>
<td>1</td>
<td>5</td>
<td>7</td>
<td>35</td>
<td>Mesozooplankton</td>
<td>RF3: Janne Søreide, Anette Wold</td>
</tr>
<tr>
<td>Multinet 64 µm</td>
<td>bott-200-100-50-20-0m</td>
<td>vertical</td>
<td>all</td>
<td>1</td>
<td>5</td>
<td>7</td>
<td>35</td>
<td>Gelatinous zooplankton</td>
<td>RF3: Janne Søreide, Anette Wold</td>
</tr>
<tr>
<td>Bongonet 180 µm</td>
<td>bottom-0m</td>
<td>vertical</td>
<td>all</td>
<td>1</td>
<td>1</td>
<td>7</td>
<td>7</td>
<td>Mesozooplankton</td>
<td>RF3: Janne Søreide, Kim Præbel</td>
</tr>
<tr>
<td>Bongonet 64 µm</td>
<td>bottom-0m</td>
<td>vertical</td>
<td>all</td>
<td>1</td>
<td>1</td>
<td>7</td>
<td>7</td>
<td>Small mesozooplankton</td>
<td>RF3: Camilla Svensen, Kim Præbel</td>
</tr>
<tr>
<td>WP2 90 µm</td>
<td>100-0m</td>
<td>vertical</td>
<td>P1, 4, 7</td>
<td>1</td>
<td>1</td>
<td>3</td>
<td>3</td>
<td>Mesozooplankton</td>
<td>RF3: Camilla Svensen</td>
</tr>
<tr>
<td>WP3 1000 µm</td>
<td>bottom-0m</td>
<td>vertical</td>
<td>P1, 4, 7</td>
<td>2</td>
<td>1</td>
<td>3</td>
<td>3</td>
<td>Mesozooplankton</td>
<td>RF2: Katrine Borgå, Doreen Kohlbauer</td>
</tr>
<tr>
<td>MIK 1.5 mm</td>
<td>bottom-0m v-haul</td>
<td></td>
<td>P1, 4, 7</td>
<td>2</td>
<td>1</td>
<td>3</td>
<td>3</td>
<td>Macrozooplankton</td>
<td>RF2: Katrine Borgå, Doreen Kohlbauer</td>
</tr>
<tr>
<td>MIK 1.5 mm</td>
<td>bottom-0m v-haul or vertical</td>
<td></td>
<td>x</td>
<td>1.5</td>
<td>1</td>
<td>7</td>
<td>7</td>
<td>Gelatinous zooplankton</td>
<td>RF3: Espen Bagbev</td>
</tr>
<tr>
<td>Macrozooplankton trawl 3 mm</td>
<td>bottom-0m v-haul</td>
<td></td>
<td>x</td>
<td>1.5</td>
<td>1</td>
<td>ad hoc</td>
<td>ad hoc</td>
<td>Macrozooplankton</td>
<td>RF3: Espen Bagbev</td>
</tr>
</tbody>
</table>

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9.2.1  Mesozooplankton

**Measured parameter:** Mesozooplankton abundance, biomass and species composition

**Method responsible:** Anette Wold, NPI (anette.wold@npolar.no), Janne Søreide, UNIS (janne.soreide@unis.no) and Camilla Svensen, UiT (camilla.svensen@uit.no)

**Description of parameter/Aim:** Provide information on mesozooplankton seasonal, annual and regional variations in abundance, biomass and species composition along a latitudinal environmental gradient in the Barents Sea from South to North.

9.2.1.1  Mesozooplankton abundance/taxonomy

**Measured parameter:** zooplankton abundance (ind/m3) and biomass (mg C/m3)

**Method responsible:** Anette Wold, NPI (anette.wold@npolar.no), Janne Søreide, UNIS (janne.soreide@unis.no)

**Description of parameter:** Provide information on mesozooplankton seasonal, annual and regional variations in abundance, biomass and species composition along a latitudinal environmental gradient in the Barents Sea from South to North.

**Methodological description:**

→ **Sampling depths:**
  - Bottom depth <600m (5 depths): bottom-200; 200-100; 100-50, 50-20 and 20-0 m
  - Bottom depth >600m (5 depths): bottom-600; 600-200; 200-50; 50-20; 20-0 m

→ **Chemicals used:** 37% formaldehyde free from acid, 96% non-denatured EtOH
Material and Method:

- **Stations**: All process stations
- **Gear**: Multinet midi 180 µm mesh size (opening 0.25m²), see Multinet manual section

**Sample treatment:**

- The nets should be rinsed and samples should be treated as gentle as possible throughout the process.
- Samples should be processed immediately after sampling.
- Gelatinous zooplankton should be removed prior to preservation, by use of plastic pipette or spoon with wholes, over a light table/ or white tray. See the section “Gelatinous zooplankton”
- Filter the rest of the content of the sample through a 180 µm sieve placed over a white tray in case something gets spilled.
- Move the samples from sieve into 125 ml bottle, fill the bottle to the neck with sea water
- Samples are preserved with Formaldehyde 37% (stabilised for histology free from acid).
- **Labels**: Use UUID labels including short human readable ID

**9.2.1.2 Small mesozooplankton abundance/taxonomy**

**Measured parameter**: zooplankton abundance (ind/m3) and biomass (mg C/m3)

**Method responsible**: Camilla Svensen, UiT (camilla.svensen@uit.no)

**Description of parameter**: Provide information on small mesozooplankton seasonal, annual and regional variations in abundance, biomass and species composition along a latitudinal environmental gradient in the Barents Sea from South to North.

**Methodological description:**

- **Sampling depths**: (same as Multinet 180 µm)
• Bottom depth <600m (5 depths): bottom-200; 200-100; 100-50, 50-20 & 20-0 m
• Bottom depth >600m (5 depths): bottom-600; 600-200; 200-50; 50-20; 20-0 m

→ **Chemicals used:** 37% formaldehyde free from acid, 96% non-denatured EtOH

**Material and Method:**

• **Stations:** All process stations
• **Gear:** Multinet midi 64 µm mesh size (opening 0.25m²) → See Multinet manual for details
• **Sample treatment:** Same as for mesozooplankton abundance
• **Labels:** Use UUID labels including human readable ID

### 9.2.1.3 Mesozooplankton genetics & biomass

**Measured parameter:** Molecular species diversity and relative abundance and biomass dry weight (mg/m3)

**Method responsible:** Metagenomics, Kim Præbel, UiT ([kim.præbel@uit.no](mailto:kim.præbel@uit.no)); biomass, Janne Søreide, UNIS ([janne.soreide@unis.no](mailto:janne.soreide@unis.no))

**Description of parameter:** Provide information seasonal, annual and regional variations in biomass and genetic composition of the mesozooplankton fraction along a latitudinal environmental gradient in the Barents Sea from South to North.

**Methodological description:**

→ **Sampling depth:** Bottom-0 m

→ **Chemicals used:** non-denatured 96% EtOH

**Material and Method:**

• **Stations:** Standard stations & process stations
• **Gear:** Bongo-net, two nets of 180 µm (60 cm dia./0.2827m² opening)
Sample treatment:

- One net for metabarcoding & one net for biomass.
- For the metabarcoding net, filter the content of the net through a 180 µm sieve placed over a white tray to remove as much water as possible. Then place the sieve on top of a generous amount of tissue paper to remove the capillary water trapped in the sieve and among individuals.
- Transfer the organisms to a canister containing approx. 10x volumes of ice cold non-denatured 96% EtOH to the amount of organisms.
- Place and store the canister at -20°C. After 24-48 hours, gently shake the canister, and change as much as possible of the EtOH with new ice cold EtOH.

- From the biomass sample, sort out larger specimen such as krill and amphipods and treat the rest as one sample. Larger specimen should be measured individuals. Gelatinous zooplankton should also be picked out prior to preservation in the same manner as for the abundance samples. If time permits take individual pictures of gelatinous zooplankton.
- The biomass samples should be dried at 60°C for 24 hours and weighted. For large organisms like medusae and ctenophores their volume fraction are determined by displacement volume on-board the vessels.
- If the biomass samples is not dried and weighted it should be frozen (-20°C)
- **Labels:** Use Pre-printed Mesozooplankton genetics & Mesozooplankton biomass UUID labels

### 9.2.1.4 Small mesozooplankton genetics & biomass

**Measured parameter:** Molecular species diversity and relative abundance and biomass dry weight (mg/m3)

**Method responsible:** Metagenomics, Kim Præbel, UiT (kim.præbel@uit.no); biomass, Camilla Svensen, UiT (camilla.svensen@uit.no).

**Description of parameter:** Provide information seasonal, annual and regional variations in biomass and genetic composition of the small mesozooplankton
fraction along a latitudinal environmental gradient in the Barents Sea from South to North.

Methodological description:

→ **Sampling depth**: Bottom-0 m

→ **Chemicals used**: non-denatured 96% EtOH

Material and Method:

- **Stations**: Standard stations & process stations
- **Gear**: Bongo-net, two nets of 64 µm (60 cm dia./0.2827m² opening)

**Sample treatment**: Same as for mesozooplankton genetics & biomass

### 9.2.1.5 Gelatinous zooplankton comments

**Measured parameter**: Gelatinous zooplankton volume (ml/m³), abundance(ins/m³) & genetics.

**Method responsible**: Anette Wold ([anette.wold@npolar.no](mailto:anette.wold@npolar.no)) & Sanna Majaneva ([sanna.majaneva@gmail.com](mailto:sanna.majaneva@gmail.com))

**Methodological description**:

Gelatinous zooplankton should be taken from MIK net and either Bongonet or Multinet cover all the size groups.

→ **Sampling depths**: (same as MIK net, Bongonet & Multinet)

→ **Chemicals used**: 96% non-denatured EtOH

**Sample treatment**

Gelatinous zooplankton should be removed prior to preservation

- keep the sample in a bit of water at all times in a white tray
- Be gentle when rinsing the samples
• Use wide mouthed pipettes or spoons to move the specimens from one container to another (do not pick them up with forceps or pour them from one container to another)

Features to consider when identifying ctenophores are:
• General size, shape and color
• Tentacles: length, location, tentacle bulbs, tentilla structures
• Comb row: lengths, length of individual cilia
• Radial canals
• See the «ID guide for pelagic Ctenophores in Norwegian Arctic” by Sanna Majaneva

**What to record**
• They should be identified to lowest taxonomic level possible, counted number of individuals and measure the volume for each taxa.
• Take a picture of each group of taxa including measuring tape or millimetre paper size reference.
• If time permits some individuals of each taxa group should be size measured and stored individually in Eppendorf, vials or falcon tubes in >96% non-denatured EtOH and stored at -20°C.
• Exchange as much of the EtOH as possible after 24-48 h

9.2.1 Macrozooplankton abundance, biomass & genetics

**Measured parameter:** Macrozooplankton biomass (wet-weight – g/m3 and/or g/m2) and abundance (ind/m3 and/or ind/m2), and metabarcoding

**Method responsible:** Espen Bagøien, IMR (espen.bagoien@hi.no)

**Description of parameter:** Provide information seasonal, annual and regional variations in abundance, biomass and genetic composition of the macrozooplankton fraction along a latitudinal environmental gradient in the Barents Sea from South to North.
Methodological description:

- MIK (Midwater Ring Trawl): standard gear for use at all stations (open water and in ice)
- Macroplankton trawl: use ad hoc – only in open water, if time and capacity permits. This is only a possible supplement to MIK – not a replacement.

→ Sampling depths: 700m (or near bottom) to surface

Mode of operation – 2 options:

- V-hauls: Possible in ice-free water: Oblique V-hauls down to near-bottom - or to 700 m when bottom-depth exceeds this depth – and then up to the surface again (see manual below for ship speed and deployment/heaving speeds).
- Vertical hauls: In Ice – or when V-hauls for other reasons can't be made: Use MIK net vertically from near bottom - or from 700 m when bottom-depth exceeds this depth - to surface (hauling speed 1.5 – 1.75 m/s). This mode of operation is rather simple.

In principle, V-hauls are preferable over vertical hauls as the first method will increase the volumes filtered and should provide better estimates for macroplankton, which often occur in relatively low concentrations. However, this may not be possible in areas with ice, when time is very limited, or for other practical reasons. In such cases, vertical hauls should be made.

→ Chemicals used: 4% formaldehyde free of acid

Material and Method:

Stations: Standard stations & process stations

Gear:
- MIK net (diameter 2m, opening 3.14m²). See MIK manual (chapter 9.3.2) for details regarding deployment
• Macrozooplankton trawl (ad hoc): opening area 38 m² mouth, 3 mm mesh size. See Macrozooplankton trawl manual (chapter 9.3.3) for details regarding deployment

Sample treatment:

• Weigh the total sample after removing excess water by use of sieve with mesh-size 0.5 mm (sieve-mesh must be <= 0.5 mm, as this is the smallest mesh-size of the MIK-net).
• Remove large jellyfish (cnidarians, ctenophores) and fish, as well as other particularly large organisms, from the total sample. Place these removed organisms aside in the refrigerator (ca. 4°C) until subsample for metabarcoding has been secured – see below).
• Weigh the remaining part of the sample (i.e. total sample without excess water, minus removed jellyfish and larger fish, etc.).
• Mix the remaining part of the sample gently, so that organisms of different species and sizes become “randomly distributed”.
• A certain fraction of the remaining (and now mixed) part of the sample is then removed “in a random way” and preserved for metabarcoding and thereby estimation of species-composition. A plankton-splitter can be used for the fractionating – or alternatively a fraction can be determined directly by weight-ratio if preferable. This fraction (depending on sample-size - smaller fractions for larger samples) can be for instance 1/4, 1/8, 1/16, etc, and must be noted in the journal. Note the weight (without excess water) of the subsample removed for metabarcoding. If possible, the subsample should weigh somewhere between ca. 10-20 g. Ensure that at least 3/4 of the sample that the metabarcoding-subsample is taken from is left for traditional, non-molecular analyses (described below). The subsample for metabarcoding is preserved in ca 10x excess ice-cold (best is -20 °C) 96% non-denatured ethanol (EtOH) in a 100, 250 or 500 mL canister. The sample can be distributed among several bottles if needed to ensure the 1:10 ratio of sample:EtOH, then label all bottles with the same UUID/sample name.
• Place and store the canister(s) at -20°C. After 24-48 hours, gently shake the canister(s), and change the EtOH with new ice-cold EtOH.

• The remaining part of the sample (i.e. total sample minus jellyfish and larger fish, etc., and after a fraction was removed for metabarcoding) is dedicated for estimation of species composition and abundance made the traditional, non-molecular way. This subsample is first weighed (without excess seawater), and thereafter preserved in seawater with final solution of 4% formaldehyde + borax) on 125, 250, 500- or 1000-mL sample-bottles. If necessary, use more than 1 bottle for large samples. To ensure a proper preservation of the organisms, their total volume should not exceed more than ca. ¼ of the bottle before adding sea-water and fixative.

• The organisms (jellyfish, larger fish, etc.) that were previously removed from the total sample and set aside in the refrigerator, are now sorted and identified to the lowest taxonomic level possible, length-measured and weighed (only one collective weight per species). For large catches of jellyfish, it may be too time-consuming to length-measure the individuals (can be skipped, then just weigh each species-group). To keep the sample quantitative, it is important to note all removals of organisms from the total sample – as well as the weight of each removed group - in the journal.

• Take pictures of the organisms sorted from the total catch for documentation – and organize the pictures in digital folders according to stations. Remember to add a scalebar in the pictures.

• Freeze the length-measured and/or weighed groups of removed large organisms at -20 °C, each species/taxon in a separate plastic bag. Jellyfish should be frozen flatly at -20 °C in plastic bags. For large catches of jellyfish, only freeze a representative subsample of each taxon.

• Note all the weights described in the protocol above in the journal so that we obtain quantitative results. This includes weights of total sample, removed large individuals (fish, jellies, etc.), remaining weight after removal of large individuals, subsample for metabarcoding, weight of sample for non-molecular taxonomic analyses, etc.

• PS. Remember to change ethanol on the samples for metabarcoding after 24-48 hours
• Registration of data: Keep detailed paper journal with all relevant information – including station and sampling information, flowmeter readings, sampling depths, sorting of samples with weights, splitting, etc. – i.e. all information that is needed to provide quantitative estimation of biomasses and abundances for the different taxa. Please also note when EtOH was changed in the bottle(s) for metabarcoding.

• All data from the MIK trawl must be registered in the IMR Plankton database (latest version of “RegPlankton”) – either during the cruise if the sampling is made by IMR employees – or elsewise after the cruise. In addition, keep the paper version in a journal.

• All samples and journals must be sent to IMR after the cruise.

• All samples and metadata are also to be registered and labelled according to the “Arven Legacy log system” – including UUID’s.

• NB! Requests for macroplankton for “other purposes” than here described will require additional and separate sampling – organisms must not be taken from the samples described above. This is to keep the samples quantitative.

**Guide for Macroplankton-trawl sampling - and processing of samples:**

• This comes in addition to the sampling with MIK, and is to be performed if - and only if - time and capacity permits, and only in open, ice-free water. To be used ad hoc.

• Sampling depths: 700 m (or near bottom) to surface

**Sample treatment:**

• Weigh the total sample after removing excess water by use of sieve with mesh-size <= 2 mm, as 3 mm is the mesh-size of the trawl).

• Remove large jellyfish (cnidarians, ctenophores) and fish, as well as other particularly large organisms, from the total sample. Set these removed organisms aside in the refrigerator (ca. 4°C) or on deck until subsampling of the remaining part has been performed (see below).

• Weigh the remaining part of the sample (i.e. total sample without excess water, and minus removed jellyfish and larger fish, etc.).

• Mix the remaining part of the sample gently, so that organisms of different species and sizes become “randomly distributed”.
• Collect and weigh 2 random subsamples (for instance 100-200 grams each) from the now mixed remaining sample described above (i.e. the Total sample minus large organisms that were removed (jellies, fish, etc.) and minus the optional sample for metabarcoding).
• These 2 subsamples are dedicated for estimation of species composition and abundance made the traditional, non-molecular way. The first subsample is first weighed (without excess seawater), and thereafter preserved in seawater with final solution of 4% formaldehyde + borax) on a 1 L sample-bottle. To ensure a proper preservation of the organisms, their total volume should not exceed more than ca. ¼ of the bottle before adding sea-water and fixative. The second subsample is just weighed and frozen directly in a plastic bag – preferably flatly.
• The organisms (jellyfish, larger fish, etc.) that were previously removed from the total sample and set aside in the refrigerator, are now sorted and identified to the lowest taxonomic level possible, counted and weighed (only one collective weight per species). It will probably be too time-consuming to length-measure these individuals. To keep the sample quantitative, it is important to note all removals of organisms from the total sample – as well as the weight of each removed group in the journal.
• If time permits, take pictures of the organisms sorted from the total catch for documentation – and organize the pictures in folders according to stations. Remember to add a scalebar in the pictures.
• Freeze the length-measured and/or weighed groups of removed large organisms at -20 °C, each species/taxon in a separate plastic bag. Jellyfish should be frozen flatly at -20 °C in plastic bags. For large catches of jellyfish, only freeze a representative subsample of each taxon.
• Registration of data: Keep detailed paper journal with all relevant information – including station and sampling information, sampling depths, sorting of samples with weights, splitting, etc. – i.e. all information that is needed to provide quantitative estimation of biomasses and abundances for the different taxa.
• All data from the Makroplankton trawl must be registered in the IMR Plankton database (latest version of “RegPlankton”) – either during the
cruise if the sampling is made by IMR employees – or elsewise after the cruise. In addition, keep the paper version in a journal.

- All samples and journals must be sent to IMR after the cruise.
- All samples and metadata are also to be registered and labelled according to the “Arven Legacy log system” including UUID's.

9.2.2 Zooplankton acoustics EK80

**Measured parameter:** Zooplankton acoustics

**Method responsible:** Randi Ingvaldsen (IMR, randi.ingvaldsen@hi.no)

**Description of parameter:**

**Methodological description:**

- Acoustic surveying on the Nansen Legacy surveys will be conducted using the six scientific Simrad EK80 echo sounders, all mounted on the drop keel, and simultaneously operated from a common computer. These are the 18kHz, 38kHz, 70kHz, 120kHz, 200kHz, 333 kHz split beam systems. When ice conditions are such that the keel transducers can be operated, they are the preferred systems used. When ice conditions are such that the keel must be retracted, and protected, the data collection will continue with similar systems mounted in Arctic tanks. As the echo sounders now must be operated through relatively thick protective windows, the noise level and measurement conditions will be worse. The decisions on when to switch systems will be taken by the captain or bridge personnel.

- EK80 should be run in CW modus during the first year of operation (until 2019) or until otherwise decided. The echo sounders should be calibrated at the start of the surveys, and when the survey will enter the above-mentioned conditions, both systems should be calibrated.

- Multi-frequency scrutinization and target strength analysis will be conducted daily for the 38kHz data with the Large Scale Survey System (LSSS) post-processing system (Korneliussen et al., 2006, 2016), which
also will be used for exporting files for subsequent analysis. The processing will involve manual removal of unwanted acoustic noise from e.g., trawl sensors during trawl operations. Simultaneous current measurements will be made with RDI 150 kHz ADCP, externally triggered by the echo sounder as a master. A fixed time delay in transmission should be implemented to prevent interference from the ADCP transmit pulse to the echo sounder data.

- Interpretations will be made per standard procedures where the total backscatter will be split into target categories like (see ICES, 2015; Korneliussen et al., 2016). Categories used for the Nansen Legacy will be: capelin, herring, polar cod, blue whiting, Norway pout, cod, haddock, redfish, saithe, O-group mixed, plankton mixed and other scatterers, in accordance with the protocol for acoustics during the ecosystem survey.

- The scrutinized acoustic backscattering data in the echo sounder output will be in the form of the Nautical area scattering coefficient (NASC) in standard units; (m² nmi⁻², MacLennan et al., 2002). They will be integrated to 10-m depth bins relative to the sea surface, starting from about 15 m depth, depending on the draft of the transducers. The scrutinized data will be stored in accordance with standard procedures at IMR: //kvalitet.hi.no/docs/pub/dok03896.

(Acoustic probing

- Detained inspections at short range of interesting layers in the Arctic may be made with acoustic / optic probing. A specially designed probe with full wideband capacity with carry 4 EK80 echo sounders with 5 selectable transducers, 38,70, 120, 200 and 333 kHz will be used in profiling mode (transducers in horizontal mode), and the multi-frequency echo sounder observing to 50 m to the side of the probe will be run at high PRF (3–4 Hz) while the probe is lowered from surface to the bottom (max 1500 m depth) at about 1 ms⁻¹. Full multi-frequency
echograms will be recorded during the profile, and still photo images from a stereo camera will be captured during retrieval. Scrutinizing and storing the probe data to a local database will be made according to procedures under development. The echo sounders will be calibrated at the start of the survey (or during the survey with stationary vessel for a few hours. The TS probe will be run on every primary level stations, where biological sampling is conducted to support the acoustic data collection. The probe may alternatively be used in vertical mode, for target strength measurements of specific organisms in pure concentrations. These numbers are needed for several of the Arctic categories for accurate density estimation from the vessels-based systems.

Literature


9.2.3 Mesozooplankton & macrozooplankton for ecotox, stable isotopes & fatty acid samples

**Measured parameter:** Organic pollutants, stable isotopes, fatty acids, Hg, Carbohydrates, Proteins and Energetics (see section 13)

**Method responsible:** Katrine Borgå (UiO, katrine.borga@ibv.uio.no)

**Description of parameter:** (see section 13)
Methodological description:

→ **Sampling depths:** Samples will be collected from two distinct depth layers containing Atlantic & Arctic water masses. The depth will be decided depending on CTD profile.

→ **Chemicals used:** No chemicals used. All samples will be frozen

Material and Method:

- **Stations:** Process stations only
- **Gear:** WP3 1000 µm & MIK net
- **Sample treatment:** Samples will be sorted to species or higher taxa, see protocol for RF 2 regarding the amount needed for the different sample types and pay attention to avoid contamination

### 9.2.4 Stable isotopes, Fatty acids & HBI

**Measured parameter:** Stable isotopes, fatty acids & HBIs of POM and main zooplankton species

**Method responsible:** Philipp Assmy (NPI, philipp.assmy@npolar.no), Doreen Kohlbach (NPI, tba) & Anette Wold (NPI, anette.wold@npolar.no)

**Description of parameter:**
Stable isotopes, fatty acids & HBIs of POM will be used to study coupling/de-coupling of sympagic and pelagic primary and secondary producers. In addition, fatty acids (together with C/N ratios) will be used as a measure of food quality for the planktonic grazer communities and will be linked to on board grazing experiment. At all the process stations, water for POM-samples will be collected in different water masses (Atlantic and Arctic water). In addition, POM will be sampled at the depth of Chlorophyll maximum, in order to link suspended fatty acid with grazing experiments.

Stable isotopes, fatty acids & HBIs of will be analysed in the dominating copepod species *Oithona similis, Calanus* spp. & *Thysanoessa* spp., Themisto spp. etc. At the ice-covered stations it may also be relevant to investigate FA in other dominating species, such as appendicularians
Methodological description:

→ Special requirements: The sampling will be done in collaboration with zooplankton ecotox sampling.

→ Sampling depths: Samples will be collected from two distinct depth layers containing Atlantic & Arctic water masses. The depth will be decided depending on CTD profile.

→ Chemicals used: 95 % Ethanol for rinsing (forceps, filtration equipment)

Material and Method:

Stations: Process stations only

Gear: Niskin bottle, Ice corer, WP3 1000 µm, MIK net &

→ Sample treatment sea ice & watersamples:

Equipment sea ice and seawater sampling:

• Niskin bottles
• Ice corer 14 cm (or 9 cm)
• Stainless steel handsaw, cutting board, ruler for cutting ice cores
• Melting cups for ice cores
• 20 L containers for seawater samples
• Whatman Nuclepore 0.22 µm membrane filters (for filtered seawater for ice core melting)
• 47 mm GF/F filters (pre-combusted: 6h, 500°C), for SI
• 2 mL acid-washed cryovials or 8 mL Wheaton glass vials (pre-combusted: 6h, 500°C) for filter storage

Procedure seawater samples:

• Sampling with Niskin bottles at different depths
• Collection the samples into 20L containers (rinse bottles 3 x with sample)
Procedure sea ice samples:

- Cut bottom 10 cm of the ice core with stainless steel handsaw
- Melt ice cores in container (cleaned with MilliQ) over a period of 24-36h
- Depending on biomass, ice core sections need to be pooled (particularly top and mid sections) in order to obtain sufficient sample material
- Melt ice cores without or with the addition of filtered seawater from the same station/day (Whatman Nuclepore 0.22 µm membrane filters; 500 mL per 5 cm bottom ice core) to mitigate cell stress; concentrations are corrected using a dilution factor

Procedure filtration of sea ice and seawater samples:

- Filtration on pre-combusted (6 h, 500°C) 47 mm GF/F filters
- Duplicate or triplicate filters per sample plus one blank per filtration day
- Filtration volume: for sea ice typically between 150 and 500 mL, for seawater typically between 2 and 7 L (filters need strong color) -> need approx. 50-60 ng of sample material
- Filter manipulation only with ethanol-rinsed forceps
- Wrap filters in into 2 mL cryovials or pre-combusted 8 mL Wheaton glass vials (6 h, 500°C)

→ Sample treatment zooplankton:

Equipment zooplankton sampling

- Forceps for handling filters, organisms (ethanol-rinsed)
- 8 mL or 25 mL Wheaton glass vials for zooplankton, benthic samples (pre-combusted: 6h, 500°C); vials should be pre-labelled with sampling ID and weight should be determined after combustion
Zooplankton, in-ice fauna:

- Collect samples using MIK net 1.5mm & WP3 1000 µm together with sampling for ecotox samples
- Sample two depth layer per station, one in the Atlantic layer and one in the Arctic layer
- Sort the samples to highest possible taxa under stereomicroscope using forceps or pipettes. Make sure to keep the samples cold during sorting by use of cold packs or ice.
- Transfer the samples into pre-combusted 8 mL Wheaton glass vials (animals) or 25 mL Wheaton glass vials (large animals, sediments)
- Sample size: small copepods: 15-30 individuals/sample & large amphipods: 1-2 individuals/sample
- Store the samples at -80°C until processing

Wear gloves at all time when manipulating sample material

---

9.3 MANUALS ZOOPLANKTON NETS

9.3.1 Multinet manual

**Specification:** Opening 50cm*50 cm = 0.25m², size of box: 80 cm*90 cm*95 cm, 5 net bags 250 cm long with mesh size of 180 µm & 64 µm, overall length 560 cm.

**Introduction:** The Multinet is deployed with all net bags closed and the water flowing freely through the frame. The instrument can be lowered with high speed to the greatest desired depth. It can either be operated by use of communication cable and deck unit and nets can be opened manually or it can be used off-line by pre-programming it as described below. All measuring data are stored inside the internal data memory during the operation and can be read out by a PC when the Multinet is back on board.
Deployment:

**Off-line preparation of Multinet:** Check that the small pin on the rotating cylinder is in the right position (if not see instructions below).

1. Turn on the Multinet
2. Connect Multinet to the Toughbook, open program “OceanLab3” from desktop.
3. Check the battery (3* Lithium 123A/ 3V status), should be >7.
4. Connect to Multinet using *connect symbol* (1\textsuperscript{st} from left in menu bar)

5. Open the *control mode* (6\textsuperscript{th} from left in menu bar)
6. Choose *Pressure programming* to program the releasing depth. Remember that the Unlock depth must be minimum 1 m below the depth of the 1\textsuperscript{st} net. The nets do not open if the Multinet has not been set below the unlock depth. Always go at least 5 m below to make sure it opens (if not the cast needs to be repeated).
7. Send to Multinet.

---

*Fig. 9.3.1-1* There is a small pin on the rotating cylinder which should point straight at the pin seen here.

*Fig. 9.3.1-2* Menu bar of the Ocean Lab software
8. Disconnect Multinet. **Remember to put dummy plug back on.**

→ If not all the net opened during the previous deployment, check that the small pin of the rotating cylinder that controls the opening of the nets, is in the right position, pointing straight up towards the pin at the net (see picture above). If not mark Net and reset the rotating cylinder either by using *reset counter* or *half step*.

9. Prepare the nets (strain the spring by using the bar to lift them into position).
• Net #1 samples the deepest layer, Net# 5 the surface layer.
• Make sure that cod ends are placed at the right net (check numbers)!
• Make sure that the nets are not twisted before the Multinet is lowered into the water.
• The weight of the lower part of the Multinet should be entirely supported by the ropes and not by the nets! (The nets have to hang loose, the ropes should be straight).
• More information can be found in the “Multiplankton Operational Manual from HydroBios”.

9.3.2 MIK

Specifications: Metal ring with diameter of 2 m in front, to which a 14 m long net is attached. Mesh-size of the net is 1.6 mm, except for the last 1.5 m of the rear part, where mesh-size is 500 µm.

For oblique hauls, a depressor is used, preferably a “saddle-shaped” Scripps-depressor, but an IKMT-depressor can also be used.
NB! A flowmeter is attached in the centre of the opening for estimation of filtered volume whenever sampling with MIK. This is particularly important when sampling by V-hauls, as a lack of flowmeter implies that no reliable estimation of volume filtered can be made.

For more information - see the protocols «ICES 2013» & «ICES 2016».

Oblique V-hauls:
• Deploy the MIK with 25 m per minute while keeping ship speed as low as possible (steering speed) until desired lower sampling depth is reached.
• Once at lower sampling depth, heave the MIK with 15 m per minute while holding a ship-speed of 3 knots throughout the haul.
• Flowmeter mounted in centre of the ring.
• Use a depressor – preferably saddle-shaped
• Hauling-wire 10 mm in diameter.
• Depth measured by Scanmar depth-sensor or trawl-probe (mounted on the ring).
• Deployed from shipside, or if required behind the ship.
• Use 10 m long «bridles» - which requires a large «block» in the crane.

Vertical hauls:
• 4 wires connected to the ring for increased stability
• Flowmeter mounted in centre of the ring.
• No depressor
• Attach weights to the rear end or cod-end of the MIK (20-30 kg)
• Depth measured by Scanmar depth-sensor or trawl-probe (mounted on ring).
• Deploy slowly (ca. 1 m/s) while keeping the ship “still”
• From desired lower sampling depth, heave with 1.5-1.75 m/s throughout the water-column

Literature:
9.3.3 Macroplankton trawl

**Specifications:** 38 m² mouth opening, mesh-size of 3x3 mm throughout the trawl – from opening to cod-end, total length of about 45 m.

**Background info:** The Macroplankton trawl (Melle et al., 2006; Wenneck et al., 2008, Krafft et al., 2010; Heino et al., 2011) has been used regularly by IMR since 2010 to obtain quantitative samples of macrozooplankton, particularly for krill, amphipods, shrimps, and mesopelagic fish. This trawl will also give improved quantitative estimates of various types of jellyfish, in particular for schyphozoans but also for ctenophores.

For safety reasons the trawl can only be used as deep as the trawl-floats are specified for!

**Deployment:** V-hauls are applied for quantitative estimation of macrozooplankton abundances. The aim is that the trawl moves through the surrounding water with a speed of ca. 2.5 knots (affected by both vessel-speed and release/hauling-speed of wire) – both on the way down and up. After the trawl has reached its maximum depth, ship speed should be approximately 2.0 knots while the trawl is hauled obliquely towards the surface - heave as slowly as possible. Make sure that the height of the trawl opening is ca. 5 m during trawling – large departures from this height are an indication that the trawl is not behaving as it should – and the sample may then not valid.
It is important to attach a Scanmar trawl-eye and speed sensor for monitoring trawl performance. Data from these sensors must be logged on the computer at the ships bridge for later determination of trawl profile and volume of water sampled. The trawl is to be used with standard pelagic trawl doors, sweep-length of 70 m, and 150 kg of weights (chain-weights - 75 kg on each side), and 8 m strapping between the wings. More details regarding rigging of the trawl can be found in the IMR's Plankton Manual (pages 13-15).

**Literature**


**9.3.4 WP2, WP3 & Bongo net manual**

**Specifications:**

- WP2 stainless steel ring of diameter 57 cm, mesh size 180 µm & 64 µm mesh size, length 2.6 m
- WP3 stainless steel ring of diameter 1 m, mesh size 1000 µm, length 2.6m

**Deployment:** Both WP2/ WP3 nets are deployed vertically. Both nets can be closed at specific depth by use of a messenger if needed.
How to fix the net: Connect the ring to the net (see picture guide below)

Connect the cup to the net
Insert the cup in the metal box

Connect the box to the net

**NOTE:** no twists (check before and during the net deployment)

Check presence of holes in the net (after each sampling) and eventually use tape to repair it

**NOTE:** Check the safety closure for the cup
10 Benthic sampling

This section contains information on planned seafloor sampling during joint AeN cruises. It has a focus on benthic organism sampling and processing as well as sediment sampling and processing where of joint biological-geological interest. We refer to sampling protocols provided by the geology team in several cases and for variables not of direct relevance to biologists in the project.

10.1 AEN TEAM MEMBERS INTERESTED IN BENTHIC INVERTEBRATE SAMPLES AND SEABED SEDIMENT SAMPLES

Sediment sampling is relevant for RF1 and RF3. Benthic invertebrate sampling is mostly relevant for RF3, with Foraminifera also of interest for RF1. Researchers interested in seafloor sampling (as of April 2018) are listed in Table 11.1-1. This list will obviously need updating as PhD students and Post-docs are hired into the project and other team members join.

Tab. 10.1-1 Researchers interested in seafloor sampling in the Nansen Legacy as of August 2018.

<table>
<thead>
<tr>
<th>RF / RA</th>
<th>Name</th>
<th>Inst.</th>
<th>Role</th>
<th>Topic</th>
<th>Mail address</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF1, 3</td>
<td>Elisabeth Alve</td>
<td>UiO</td>
<td>Sci</td>
<td>Foraminifera</td>
<td><a href="mailto:elisabeth.alve@geo.uio.no">elisabeth.alve@geo.uio.no</a></td>
</tr>
<tr>
<td>RF1</td>
<td>Murat Ardelan</td>
<td>NTNU</td>
<td>Sci</td>
<td>Trace metals</td>
<td><a href="mailto:murat.v.ardelan@ntnu.no">murat.v.ardelan@ntnu.no</a></td>
</tr>
<tr>
<td>RF3</td>
<td>Bodil Bluhm</td>
<td>UiT</td>
<td>Sci</td>
<td>Biodiversity &amp; community structure, food webs, population dynamics</td>
<td><a href="mailto:bodil.bluhm@uit.no">bodil.bluhm@uit.no</a></td>
</tr>
<tr>
<td>RF</td>
<td>Name</td>
<td>Institution</td>
<td>Position</td>
<td>Research Area</td>
<td>Email</td>
</tr>
<tr>
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<tr>
<td>RF3</td>
<td>Gunnar Bratbak</td>
<td>UiO</td>
<td>Sci</td>
<td>Sediment microbial communities</td>
<td><a href="mailto:gunnar.bratbak@uit.no">gunnar.bratbak@uit.no</a></td>
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<tr>
<td>RF2</td>
<td>Kathrine Borgå team?</td>
<td>UiO</td>
<td>Sci + PhD</td>
<td>Contaminants</td>
<td><a href="mailto:kathrine.borga@ibv.uio.no">kathrine.borga@ibv.uio.no</a></td>
</tr>
<tr>
<td>RF3</td>
<td>Raphaëlle Descoteaux</td>
<td>UiT</td>
<td>PhD (Arctic SIZE)</td>
<td>Meroplankton, benthic juveniles, barcoding, molecular diets</td>
<td>raphaë<a href="mailto:lle.descoteaux@uit.no">lle.descoteaux@uit.no</a></td>
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<tr>
<td>RF1</td>
<td>Matthias Forwick</td>
<td>UiT</td>
<td>Sci</td>
<td>Geology</td>
<td><a href="mailto:matthias.forwick@uit.no">matthias.forwick@uit.no</a></td>
</tr>
<tr>
<td>RF1</td>
<td>Katrine Husum</td>
<td>NPI</td>
<td>Sci</td>
<td>Geology, sediment properties (grain size, OC)</td>
<td><a href="mailto:katrine.husum@npolar.no">katrine.husum@npolar.no</a></td>
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<tr>
<td>RF3</td>
<td>Lis L. Jørgensen</td>
<td>IMR</td>
<td>Sci</td>
<td>Biodiversity &amp; community structure, traits, vulnerability, management</td>
<td><a href="mailto:lis.lindal.joergensen@hi.no">lis.lindal.joergensen@hi.no</a></td>
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<tr>
<td>RF1</td>
<td>Ulysses Ninnemann</td>
<td>UiB</td>
<td>Sci</td>
<td>Palaeoclimate, isotope geochemistry</td>
<td><a href="mailto:ulysses.ninnemann@uib.no">ulysses.ninnemann@uib.no</a></td>
</tr>
<tr>
<td>RF1</td>
<td>Tine Rasmussen</td>
<td>UiT</td>
<td>Sci</td>
<td>Geology, pore water profiles, pH</td>
<td><a href="mailto:tine.rasmussen@uit.no">tine.rasmussen@uit.no</a></td>
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<tr>
<td>RF3</td>
<td>Paul Renaud</td>
<td>APN</td>
<td>Sci</td>
<td>Benthic community respiration</td>
<td><a href="mailto:per@akvaplan.niva.no">per@akvaplan.niva.no</a></td>
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<tr>
<td>RF3</td>
<td>Luka Supraha</td>
<td>UiO</td>
<td>PhD</td>
<td>Phytoplankton spores</td>
<td><a href="mailto:luka.supraha@ibv.uio.no">luka.supraha@ibv.uio.no</a></td>
</tr>
<tr>
<td>RF3</td>
<td>Emmelie Åström</td>
<td>UiT</td>
<td>PD</td>
<td>Food web, stable isotopes</td>
<td><a href="mailto:emmelie.k.astrom@uit.no">emmelie.k.astrom@uit.no</a></td>
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<tr>
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<td>NN</td>
<td>IMR</td>
<td>PD</td>
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</table>
## 10.2 BENTHIC SAMPLING GEAR

Sampling gear targeting sediment and organismal samples in AeN is listed in Table 10.2-1. Benthic meiofauna will be sampled with a Gemini-corer (back up gear is a box core).

- **Macrobenthos** (terms used synonymously with infauna here) and microbes will be sampled with a box corer (back up gear is a van Veen grab),
- and **Megabenthos** (term used synonymously with epifauna here) will be sampled with beam and Campelen trawls.
- **Sediment** for shared biological and geological variables will be sampled with a box corer (again, back up gear is a van Veen grab).

Naturally, the sampling depth of all sampling gear matches the total depth of a given station. Appropriate knowledge of the bottom depth from the ship’s depth sounder is, therefore, a requirement. Knowledge on bottom topography and sediment type from multi-beam mapping is desirable.

⇒ **NOTE** that geologists prefer multi-corer samples for their measurements and will use that gear type during their cruises. In joint cruises where no geologists participate, select sediment properties will be sampled from box cores.

The gear will primarily be handled by the crew. Scientists, however, need to have the gear ready for deployment. Given that Kronprins

<table>
<thead>
<tr>
<th>RF3</th>
<th>NN</th>
<th>UNIS/UiT</th>
<th>PhD</th>
<th>Molecular diets plankton / meiofauna</th>
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<tbody>
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</tr>
</tbody>
</table>

| RF1, 3 | Marianne Risager Kjøller | UiO | PhD | Foraminifera | m.r.kjoller@geo.uio.no |
Haakon is a new vessel with a new crew, it is advisable to discuss the procedures of deployment with them. This section briefly summarizes key facts about each gear deployment. At the date of this draft, not all gear features have been clarified.

Tab. 10.2-1 Sampling gear for sediment and benthic organisms to be used during AeN (in alphabetic order). Method responsible and affiliation, potential users, sampling goals and cruise information also given.

<table>
<thead>
<tr>
<th>Gear type</th>
<th>Owner(s)/ responsible</th>
<th>Potential users</th>
<th>Sampling goals</th>
<th>Cruises</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beam trawl (10.2.1)</td>
<td>IMR/Jørgensen</td>
<td>RF3</td>
<td>Epifauna for population parameters / ECOPATH, fish, food web, barcoding, contaminants</td>
<td>All JC / seasonal cruises</td>
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<tr>
<td>Box corer (10.2.2)</td>
<td>Geology UiT/Forwick,  NPI/Husum</td>
<td>RF1, 3 (2?)</td>
<td>Sediment grain size, organic carbon content, pigments (chlorophyll, phaeopigments), macrofauna (juveniles / adults, for experiments, quantitative, for population parameters / ECOPATH), fauna, food web, pH (Geo cruises), oxygen (Geo cruises)</td>
<td>All JC / seasonal cruises</td>
</tr>
<tr>
<td>Campelen trawl (10.2.3)</td>
<td>IMR/Jørgensen</td>
<td>RF3</td>
<td>Epifauna (quantitative); for population parameters / ECOPATH, fish, food web, barcoding, contaminants</td>
<td>All JC / seasonal cruises</td>
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<td>Gemini core (10.2.4)</td>
<td>UiO/Alve</td>
<td>RF1/3</td>
<td>Foraminifera (quantitative, for experiments); all meiofauna for quantitative estimates, food web (isotopes, molecular diets)</td>
<td>Seasonal cruises</td>
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<tr>
<td>Van Veen grab (10.2.5)</td>
<td>BFE UiT/Bluhm, APN/Renaud</td>
<td>See box core</td>
<td>See box core</td>
<td>All JC / seasonal cruises</td>
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</table>
10.2.1 Galvanize steel beam trawl (Figure 10.2 – 1 A,B)

**Specifications:** Specs. in Jennings et al. (1999), Mesh size: 20 mm (10 mm knot to knot)

**Introduction:**
- 1 haul per station
- Effective opening: 2 m
- No sensors
- Trawl data needed: latitude / longitude at bottom and off bottom, water depth, time at bottom and off bottom, station number, ship speed (available on screens around ship)

**Deployment:**
- Deploy at scope ca. 2
- Bottom time 3 min
- Trawl at 1.5 kn
- Wire in / out at 1 m s\(^{-1}\)
- Empty on deck / sorting table

10.2.2 Box corer (Figure 10.2 – 1 E)

**Specifications:**
- Box size: 0.5 x 0.5 x 0.5 m
- Amount of weight on top: 4 x 25 kg or 4 x 50 kg

**Introduction/Requirements:**
- 3-4 cores per process station (where surface is undisturbed). This number will include ALL benthic sampling needs (respiration,
pigments/sediment parameters, community analysis, cores for experiments, etc.)
• Data needed: latitude / longitude, water depth

Deployment:
• To deploy, open spade
• Fix release mechanism to keep spade open
• Lower to seafloor at 1 m s\(^{-1}\)
• Lower speed to 0.5 m s\(^{-1}\) near seafloor to avoid loss of sediment surface layer and organisms
• Weight will push core into sediment
• Release will pop out upon the corer touching the seafloor
• Upon slow retrieval from sediment, spade will close
• Place on deck very carefully to reduce sediment surface disturbance
• Upon retrieval, unscrew box with sample, fix new box for next replicate
• For retrieved core, follow sediment sampling protocols

10.2.3 Campelen 1800 with Rockhopper ground-gear trawl (10.2 – 1 C,D)

Specifications: Mesh size: 80 mm in wings, 16-22 in the cod end;
Specs in Engås and Ona (1990)

Introduction:

• Trawl data needed: latitude / longitude at bottom and off bottom, water depth, time at bottom and off bottom, station number, ship speed, door opening
• scanmar sensors (available on screens around ship)

Deployment:
• 1 haul per main station
• Bottom time 15 min (should be equivalent to towing distance of ca. 0.75 nautical middles or 1.4 km)
• Trawl speed ~3 kn
• Target net opening ca. 17 m horizontal, 4 vertical
• Deploy at scope appropriate to depth and sediment (get from fish team)
• Wire in / out at 1 m s-1
• Empty into chute that leads to inside fish sorting table

10.2.4 Gemini corer (Figure 10.2 – 1 F)

Specifications: Corer contains 2 cores, 8 cm diameter each

Introduction:
• number of replicate hauls: 3 per station
• Deploy at 1 m s-1
• Data needed: latitude / longitude, water depth, core length

Deployment:
• Max. penetration depth about 60 cm
• Lower speed when the corer approaches the sea floor to about 0.5 – 0.8 m s-1 (depends on the substrate and may differ between stations)
• Weight may also be adjusted (0 to 3 pieces of lead which come with the corer)
• Place on deck very carefully, since undisturbed surface is essential

10.2.5 van Veen Grab (back up to Box corer) (Figure 10.2 – 1 E)

Specifications: Sample surface area: 0.1 m2

Introduction:
• In case box corer cannot be used, replicates per station: 4-5
• Data needed: latitude / longitude, water depth, fullness of grab
Deployment:
- Store and handle with safety pin in place, for deployment release safety pin
- open jaw and push pinch-pin with spring in place, keep pressure on pin (other types have a little metal hook instead)
- lift grab over board with winch
- lower at 1 m s-2, slower near bottom
- pinch-pin (or metal hook) will pop out when grab reaches the bottom and wire is no longer under tension
- pulling up cable gently will close jaws, swing grab over tub or sieve
- discard grabs that are less than half full, or where the sediment overflows through the windows
Fig. 10.2-1 Sediment / benthic sampling gears to be used during Arven etter Nansen cruises. A-B) Beam trawl, C-D) Campelen trawl (photos: B. Bluhm), E) box core (photo: R. Hjertenes), F) Gemini corer (photo S. Hess), G) van Veen Grab. (photo B. Bluhm).
10.3 SEDIMENT AND BENTHIC INVERTEBRATE SAMPLING

Sampling procedures for sediment and benthic invertebrates are outlined here. Not all sampling may be conducted on all cruises. **NOTE:** Additional sampling specific to a particular objective, requiring extensive training and/or only conducted on a single or few cruises is not listed here.

→ **Sample labelling:** A sample labelling system was developed for the entire project and was introduced on JC1-2 in August 2018. This system includes a labelling code and provides labels on chemical and water proof paper. See labelling protocol.

→ **Sample processing:** Lab processing notes are very brief at this point, full protocols are not provided here, but can be obtained from the responsible labs.

10.3.1 Sediment pigments (Box corer sampling)

**Measured parameter:** Sediment pigments (HPLC)

**Method Responsible:** Paul Renaud (APN, per@akvaplan.niva.no)

**Description of parameter/Aim:** concentration of pigments in the sediment. Measured in mg/m²

**Methodological description:**

→ **Special requirements/Gear:** Box corer

→ **Sampling depths:** Station bottom depths

→ **Chemicals used:** none, but samples frozen at -20°C
Method:

- Use Cut-off 60-ml plastic syringe
- Collect 3 replicates per station, but one per box core (ie, 3 samples from 3 different box cores per station)
- Insert syringe into the surface of the box core to a bit more than the sampling depth of 2 cm
- Pull out gently by turning such that the sediment does not get lost
- Place sediment “plug” in a plastic whirl-pak / zip lock bag
- Wrapped in aluminum foil
- Freeze at -20°C

→ Sampling storage: Akvaplan-niva, Tromsø

→ Shipping address: c/o Paul Renaud, Akvaplan-niva, Fram Centre, 9296 Tromsø (TRANSPORT: on plane, as checked in baggage (Arunima Sen))

10.3.2 Benthic chlorophyll a/phaeopigments (Box corer sampling)

Parameter measured/name: chlorophyll a, phaeopigments/ Sediment pigments (fluorometric)

Method Responsible: Paul Renaud (APN, per@akvaplan.niva.no)

Description of parameter/Aim: concentration of fluorometrically measured pigments in the sediment. Measured in mg/m².

Methodological description:

→ Special requirements/Gear: Box corer

→ Sampling depths: Station bottom depths

→ Chemicals used: none, but samples frozen at -20°C
Method:

- Use 4.7 cm inner diameter core
- Collect 3 replicates per station, but one per box core (ie, 3 samples from 3 different box cores per station)
- Insert core into the sediment in the retrieved box core.
- Use shovel or hands to dig in and remove the core, keeping bottom covered
- Extrude the core with specialized extruder in the following sections: 0-1 cm, 1-2 cm, 2-3 cm, 3-4 cm, 4-5 cm, 5-6 cm, 6-8 cm, 8-10 cm.
- Place each sediment section in a plastic whirl-pack/Ziploc bag.
- Wrap bags with aluminium foil
- Freeze at -20°C

→ Sample storage: Akvaplan-niva

→ Shipping address & Transport: c/o Paul Renaud, Akvaplan-niva, Fram Centre, 9296, Tromsø - on plane, as checked in baggage (Arunima Sen)

10.3.3 Sediment grain size and sediment carbon, nitrogen and stable isotopes (Box corer sampling)

Parameter measured/name:

- Sediment TOC: total organic carbon (TOC). Units: %
- Sediment TN: concentration of total nitrogen (TN). Unit: %
- Sediment grain size: fraction of the sediment corresponding to specific grain sizes. Measured as percentage of sizes (e.g., % < 63 µM).

Method Responsible: Elisabeth Alve (elisabeth.alve@geo.uio.no) and Silvia Hess (silvia.hess@geo.uio.no)

Description of parameter/Aim:

- Organic carbon and total nitrogen content of sediment and stable isotope ratios for both carbon and nitrogen. Units above
• grain size of sediment, or, fraction of the sediment corresponding to specific grain sizes. Units above.

**Methodological description:**

→ **Special requirements/Gear:** Box corer
→ **Sampling depths:** Station bottom depths
→ **Chemicals used:** none, but samples frozen at -20°C

**Method:**
- Use 5.5 cm inner diameter core
- Collect 3 replicates per station, but one per box core (ie, 3 samples from 3 different box cores per station)
- insert core into the sediment in the retrieved box core.
- Use shovel or hands to dig in and remove the core, keeping bottom covered
- Extrude the core with specialized extruder in the following sections: 0-1 cm, 1-2 cm, 2-3 cm, 3-4 cm, 4-5 cm, 5-6 cm.
- Place each sediment section in (60 ml Cerbo boxes).
- Freeze at -20°C

→ **Sample storage:** UiO

→ **Shipping address/Transport:** c/o Elisabeth Alve, Geologibygningen, University of Oslo, Sem Sællands vei 1, 0371 Oslo - Transport: on plane, as checked in baggage (Silvia Hess)

**10.3.4 Sediment microbes (Box corer sampling)**

**Parameter measured:** microbial community, parameter name: Sediment microbes

**Method responsible:** Lise Øvreås (UiB, Lise.Ovreas@uib.no)

**Description of Parameter/Aim:** Identification and quantification of sediment microbial community. Measured in abundance.
Methodological description:
- **Special requirements/Gear**: Box corer
- **Sampling depths**: Station bottom depths
- **Chemicals used**: none, but samples frozen at -80°C

**Method:**
- Use 4.7 cm inner diameter core
- Collect 3 replicates per station, but one per box core (ie, 3 samples from 3 different box cores per station)
- Insert core into the sediment in the retrieved box core.
- Use shovel or hands to dig in and remove the core, keeping bottom covered
- Extrude the core with specialized extruder in the following sections: 0-1 cm, 1-2 cm, 2-3 cm, 3-4 cm, 4-5 cm, 5-6 cm.
- When a section has been extruded, take a sample from the middle of the section with a 60ml syringe to avoid contamination.
- Place the sample in a plastic whirl pack or Ziploc bag.
- Freeze at -80°C

**Sample storage**: UiB

**Shipping address & Transport**: c/o Lise Øvreås, Centre for Geobiology, University of Bergen, Thormøhlensgate 53B, Bergen - needs to be shipped to shipping address with samples in dry ice.

10.3.5 Foraminiferal community and possibly meiofauna community (Box corer sampling)

**Parameter measured/name**: Benthic foraminiferal community. Measured in abundances/Sediment forams

**Method responsible**: Elisabeth Alve (elisabeth.alve@geo.uio.no) and Silvia Hess (silvia.hess@geo.uio.no)
**Description of Parameter/Aim:** identification and quantification of benthic foraminiferal community (remaining sample material is kept for metazoan meiofauna analysis)

**Methodological description:**
- **Special requirements/Gear:** Box corer
- **Sampling depths:** Station bottom depths
- **Chemicals used:** rose Bengal in 70% ethanol (2g/L)

**Method:**
- Use 5.5 cm inner diameter core liner
- Collect 3 replicates per station, but one per box core (i.e., 3 samples from 3 different box cores per station)
- Insert core into the sediment in the retrieved box core.
- Use shovel or hands to dig in and remove the core, keeping bottom covered
- Extrude the core with specialized extruder in the following sections: 0-1 cm, 1-2 cm, 2-3 cm, 3-4 cm, 4-5 cm, 5-6 cm.
- Place section in (180 ml Joni) container and add the same volume of rose Bengal stained 70% ethanol as sediment volume (1:1).
- Shake gently to homogenize so all sediment clumps are disintegrated.
- Do not freeze
- Note: in the lab, after sorting for forams, same samples can be used for determining meiofaunal community composition.
- Macrofauna in the >500 μm-fraction sent to Henning Reiss.

**Sample storage:** UiO

**Shipping address & Transport:** c/o Elisabeth Alve, Geologibygningen, University of Oslo, Sem Sælands vei 1, 0371 Oslo - needs to be shipped to shipping address. No special temperature needs (but do not freeze).
10.3.6 Benthic respiration experiments (Box corer sampling)

Parameter measured/name: Sediment community oxygen consumption

Method responsible: Paul Renaud (per@akvaplan.niva.no)

Description of Parameter/Aim: measuring benthic oxygen consumption. Measured as a rate (mmol O₂/m²)

Methodological description:

→ Special requirements/Gear: Box corer and Niskin water samples from bottom

→ Sampling depths: Station bottom depths

→ Chemicals used: sodium sulfite needed for calibration of oxygen system (and note that chemicals will be used for macrofaunal community post incubation)

Method:
- Use 11.7 cm inner diameter core.
- Collect 6 replicate cores per treatment per station (possibly leave out algal treatments at deep sites):
  - Treatment 1: respiration at ambient water temperature
  - Treatment 2: respiration at ambient water temperature + 4°C (+2°C in deep water)
  - Treatment 3: respiration at ambient water temperature + addition of 30 mg of labeled algae.
  - Treatment 4: respiration at ambient water temperature + 4°C + addition of 30 mg of labeled algae.
- Insert core liner into the sediment in the retrieved box core (ca 20 cm sediment).
- Insert all possible cores and sub-cores for other benthic sampling. Remove sub-cores first, to ensure the least amount of sediment is left when incubation cores have to be removed.
• Use shovel or hands to dig in and remove the core, keeping bottom covered. Do not shake. Attach core bottom. Make sure it is secure.
• Clean the core while on deck.
• Transfer core to water bath in the appropriate (dark) cold room (Tubs in the cold rooms should be prefilled with seawater in advance to maintain desired temperature).
• Use water from Niskin bottles to fill up the core to the top. This has to be done very slowly so as not to disturb the sediment surface. A paper circle (piece of water-resistant paper cut in a circle) can be used to help pour the water in slowly.
• Measure the height of the sediment in each core from at least three different locations (sides of the core).
• Add an air stone to each core (place as gently as possible so as not to disturb sediment surface) and allow to aerate for 1-2 h.
• Fix on the core top and attach to the electrical unit. Make sure that the stir bar is stirring (not too slowly, but also not too vigorously).
• Fill the last bit of water up via the small opening on the top of the core. Then plug it with the rubber stopper and try to avoid air bubbles.
• Check the temperature of the water in the water bath/tub.
• Take an oxygen reading and make sure that oxygen is at the max.
• Every few hours, take oxygen readings via the spot on the top of the cores using the PreSense Fibox 4 system according to manufacturer recommendations.
• Maintain cores IN THE DARK. Use headlamp for taking regular oxygen measurements.
• Check water temperatures when oxygen readings are taken.
• Continue taking measurements till oxygen comes down 15-30% from 100% saturation.
• After the termination of the experiment quantitative foraminifera sub-samples will be collected, and the rest of the sediment in the core is processed for macrofaunal analysis.

**Sample storage & Transport:** no long-term storage required, experiments will be terminated onboard. However, experimental cores will be saved for macrofaunal and foraminifera sampling. - **Transport:** same as above.
10.3.7 Quantitative macrofaunal assemblage (Incubation cores from Box corer sampling)

**Parameter measured/name:** Sediment macrofaunal: measuring macrofauna abundance

**Method Responsible:** Paul Renaud (per@akvaplan.niva.no) and Henning Reiss (henning.reiss@nord.no)

**Description of parameter/Aim:** identification and quantification of sediment macrofaunal community. Measured in abundance

**Methodological description:**
- **Special requirements/Gear:** Box corer and cores used for benthic respiration experiments
- **Sampling depths:** Station bottom depths
- **Chemicals used:** Formaldehyde 37% diluted to ~4% with seawater, mixed with buffer borax, Rose Bengal (add pre-weighed 4 g Rose Bengal vial to 5L of 37% formaldehyde. Dilute this mixture 9:1 with seawater and borax to obtain the working solution).

**Method:**
- Cores used for respiration experiments will be used.
- For treatments 1 and 2, sediment from cores will be gently sieved using a 500-μm sieve. For treatments 3 and 4, this will be done after sub-sampling for foraminifera.
- Transfer retained animals and detritus to a sample jar using forceps and squirt bottles
- Add buffered 4% formaldehyde-seawater-Rose Bengal solution to jars (2 parts formaldehyde solution to 1 part retained sediment)
- Never fill jar completely full with fauna/sediments – they will not preserve well
- Gently turn over sample a few times to ensure preservative fully penetrates the sample
• Seal lids with electrical tape (and possibly parafilm)
• Keep containers in chemical locker. DO NOT FREEZE.

→ **Sample storage**: Nord University

→ **Shipping address/Transport**: c/o Henning Reiss, Faculty of Biosciences and Aquaculture, Universitetsalléen 11, 8026, Bodø  -  **Transport**: needs to be shipped to shipping address. No special temperature needs (but do not freeze).

**10.3.8 Quantitative foraminiferal assemblage (Incubation cores from Box corer sampling)**

**Parameter measured/name**: Incubation experiment benthic foraminifera: Benthic foraminiferal abundance and carbon uptake

**Description of Parameter/Aim**: identification and quantification of benthic foraminiferal community, contribution of foraminiferal community to benthic respiration rates, incorporation of labeled algae.

**Method Responsible**: Elisabeth Alve (elisabeth.alve@geo.uio.no) and Silvia Hess (silvia.hess@geo.uio.no)

**Methodological description:**

→ **Special requirements/Gear**: Box corer and cores used for benthic respiration experiments

→ **Sampling depths**: Station bottom depths

→ **Chemicals used**: none, but samples frozen at -20 °C.

Method:

• Sub-cores from respiration experiments, Treatments 3 and 4 will be used.
• Use 5.5 cm inner diameter core OR subsample 0-1 cm and 1-2 cm sections (20 ml sediment in a 60 ml syringe) of the incubated cores (to be decided on the cruise).
• Insert core liner (or syringe) into the sediment.
• Place section in 60 ml Cerbo plastic container
• Freeze at -20 C.

→ Sample storage: UiO

→ Shipping address & Transport:
c/o Elisabeth Alve,
Geologibygningen
University of Oslo,
Sem Sælands vei 1
0371 Oslo

Transport: on plane, as checked in baggage (Silvia Hess)

10.3.9 Quantitative epibenthic megafauna (Beam trawl & Campelen trawl)

Parameter measured: Epibenthic megafauna abundance/biomass

Method responsible: Lis L. Jørgensen (IMR, lis.lindal.joergensen@hi.no)

Description of parameter/Aim: Quantitative investigation of epibenthic megafauna by use of Beam and Campelen trawl. Species identifications, counts and food web analysis.

Methodological description:

→ Special requirements/gear: Beam trawl & Campelen trawl
→ Sampling depths: station bottom depth
**Chemicals used:** Formaldehyde 37%, Formaldehyde solution ~4%, buffer borax or hexamethylenetetramine, Ethanol (95%, molecular grade), isotope information???

Megafauna assessment:

- Sorting, enumerating and weighing is done onboard with fresh samples
- Formalin vouchers will be transferred to 70% ethanol
- Taxonomic identifications from ethanol-preserved samples are completed in the home lab and with the help of experts
- Data recording of counts and weights by taxon and archiving follows IMR's standard protocol

Method:

1. Take photo of catch with sample label
2. record sediment type (rocks, shell hash, mud, etc.) if visible
3. If catch is too large to sort in its entirety, subsample for abundant taxa after mixing the catch, note subsampling factor by weighing total catch and subsample, then sort subsample (for rare and large taxa, sample 100% and note which taxa were sampled this way)
4. Sort the catch, separating all obviously different taxa, put each in a container (big or abundant specimens into in buckets/tubs, small or rare taxa in trays/petri dishes)
5. Keep specimens in seawater to keep them fresh
6. Identify taxa to lowest practical taxonomic level; give descriptive names to unidentified taxa and keep them consistent throughout the cruise
7. Count and weigh by taxon (note weight unit), including macroalgae
8. Do not include empty shells and tubes in counts / weights
9. Place 2-5 individuals in 4% formaldehyde seawater solution as voucher with label (at least all taxa where the identification is not certain, better all taxa once per cruise) – under hood, with nitrile gloves on, DO NOT FREEZE
10. Do not discard catch before sampling for all other objectives is done!
11. Record data on counts, weights (with unit), sub-sampling factor, station and date on data sheet, enter data into provided data file

→ Sample storage: Sample storage to be determined (museum voucher collection at the Tromsø Museum is being discussed)

10.3.10 Meiofauna (by using Gemini corer)

Parameter measured: Meiofauna abundance

Method Responsible: Foraminifera - Elisabeth Alve (UiO, elisabeth.alve@geo.uio.no), Metazoans - Bodil Bluhm (UiT, bodil.bluhm@uit.no) & Paul Renaud (APN, per@akvaplan.niva.no)

Description of parameter/Aim: Foraminifera are picked and the remaining sample is kept for metazoan meiofauna analysis.

Methodological description:

→ Special requirements/Gear: Gemini corer

→ Sampling depths: Station bottom depth

→ Chemicals used: rose Bengal, Ethanol
Method:

Sediment meiofauna community respiration experiments will be conducted during seasonal cruises. Since only trained personnel will conduct these experiments, **no protocol is provided here.**

- each core sectioned into 1 cm thick slices down to 6 cm, then 2 cm slices down to 10 cm
- each slice transferred to 180 ml Joni plastic containers add the same volume of rose Bengal stained 70% ethanol (2g/L) as sed volume (1:1) – DO NOT FREEZE
- mesh size of subsequent (post cruise) sieving: 63 um
- Post cruise: Forams are picked and the remaining sample is kept for metazoan meiofauna analysis
- if long enough (30-50 cm), core can be used for palaeo objective in RF1 (if not, need RF1 core)
- Foraminifera will be sorted from alcohol/rose Bengal samples
- Foraminifera will be identified morphologically in the home lab at UiO and enumerated
- Metazoan meiofauna samples will be archived at UiT for the time being until funding / an MS thesis is available

### 10.3.11 Food web studies (Beam trawl & Campelen trawl, Box corer and Gemini Trawl)

**Parameter measured:** Stable carbon and nitrogen samples, lipid analysis

**Method Responsible:** Lis L. Jørgensen (IMR, ), Bodil Bluhm (UiT, bodil.bluhm@uit.no), NN Postdoc (IMR/UiT), Emmelie Åström (UiT, emmelie.k.astrom@uit.no)

**Description of parameter/Aim:**
Methodological description:

→ Special requirements/Gear: Samples will be taken with various benthic gear: Beam and Campelen Trawl, Box Corer and Gemini corer

→ Sampling depths: Station bottom depth

→ Chemicals used: 1N HCl, chloroform, methanol, 13C

Method:

- Stable carbon and nitrogen samples will be run in replicates of n=4 per taxon and station, likely at UiO using an Isotope Ratio Mass-Spectrometer with V-PDB and atmospheric N2 as standards for carbon and nitrogen, respectively
- For δ13C, tissue samples containing carbonates will be treated with 1N HCl (drop by drop) until bubbling ceases, and dried again at 60 °C for at least 24h
- sediment samples will also be decalcified and undergo several washing steps
- Decide if Lipids will be removed or not. With 2:1 chloroform: methanol
- Samples for food web studies will be taken from Beam and Campelen Trawl, Box Corer and Gemini corer

For food web samples from trawl:

1. Once trawl is sorted and counted, stable isotope samples and contaminant samples will be taken and frozen at -20 °C (compound specific isotope samples at -80 °C)
2. pick out invertebrates and fishes from benthic sampling gear, at least 3 replicates per taxon and station of dominant / specific taxa (depending on objective)
3. If time is limited, put whole replicate specimens (clean, rinsed) in zip lock bags by taxon and label with station name and taxon
4. if time allows, dissect tissue samples (each roughly 1x1x1 cm) from 3 replicate organisms using muscle where possible (molluscs, crustaceans and such), otherwise body wall (annelids, cnidarians, ascidians, holothurians etc.), tube feet (asteroids), central disc (ophiuroid), organism pieces (sponges, bryozoans, hydroids etc.) or whole organisms (small taxa such as amphipods – take out gut where possible) – avoid guts whenever possible

5. Put tissue samples in microcentrifuge tubes / sampling vial, label on top and side with permanent marker

6. Record in sample log: Station number, date, isotope sample number (start at 1, consecutive numbers, species (or description), phylum, how many individuals, replicate number, tissue type

7. Prepare a voucher for all species collected for isotope analysis (at least 1, better several individuals in 4% formaldehyde-seawater solution with station label and number of isotope sample for cross-reference) (we do not need duplicate vouchers per species)

8. If time allows, dry tissues with open lids at 60 degrees C in drying oven for at least 24 hours. Close lids when done. Pack sample trays into ziplock bags and store in closed boxes so no additional carbon/nitrogen particles get in.

9. Work in clean area and with gloves during dissections, rinse dissection tools between species, avoid getting any additional carbon or nitrogen on the samples

10.3.12 Contaminants

Parameter measured: Contaminants

Method Responsible: Kathrine Borgå (UiO, kathrine.borga@ibv.uio.no)

Description of parameter/Aim:

Methodological description

→ Special requirements/Gear: Beam and Campelen Trawl
→ **Sampling depths:** Station bottom depth

→ **Chemicals used:**

**Method:**

- See Nansen Legacy workpackage RF2 protocols???

### 10.3.13 Barcoding samples

**Parameter measured:** Megafauna, Macrofauna and Meiofauna species identification by DNA barcoding

**Method Responsible:** Bodil Bluhm (UiT, bodil.bluhm@uit.no), PhD (tba)

**Description of parameter/Aim:**

**Methodological description:**

→ **Special requirements/Gear:** Beam and Campelen Trawl, Box and Gemini corer

→ **Sampling depths:** Station bottom depth

→ **Chemicals used:** 95% molecular ethanol

**Method:**

Megafauna (see 10.3.4), Macrofauna (see 10.3.3) and Meiofauna (10.3.5) assessment:
- **If time is limited:** For taxa from taxon list provided, place 4 specimens of same taxon in a ziplock bag and freeze at -20 °C with label.

- **If time allows:** Take picture of each of 4 specimens per species, place tissue sample (or whole individual if small enough) into separate sample vials/jars with label in ample amount of 95% molecular alcohol, shift alcohol after 24 h.
Fig. 10.3-1 Biological sample taking procedures. Top left) Box core sampling, top right) emptying van Veen grab into a tub for sieving, middle left) Sectioning Gemini core (and sampling container), bottom left) sorting trawl sample, bottom right) sieving macrofauna (from van Veen / box core). (Photos S. Hess, B. Bluhm).
**Tab. 10.3-1 Overview of different types of benthos data to be generated and parameter definitions.**

<table>
<thead>
<tr>
<th>Quantitative community samples</th>
<th>Process/rate measurements</th>
<th>Food web</th>
<th>Sediment properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Taxa lists from morphological taxonomic identifications (currently accepted name and taxonomic hierarchy for each taxon from World Register of Marine Species)</td>
<td>Sediment oxygen demand (mmol O$_2$ m$^{-2}$ d$^{-1}$)</td>
<td>Stable isotope values, i.e. ratios of carbon and nitrogen isotopes $\delta^{15}$N (per mil), $\delta^{13}$C (per mil)</td>
<td>Chlorophyll a (mg or $\mu$g m$^{-2}$)</td>
</tr>
<tr>
<td>Density/Abundance: number of individuals per unit area, by species or higher taxon (unit area may be m$^2$ for macrofauna, km$^2$ for megafauna)</td>
<td>Diet composition (benthic meiofauna) as genetic sequence data</td>
<td></td>
<td>Phaeopigments (mg or $\mu$g m$^{-2}$)</td>
</tr>
<tr>
<td>Biomass: weight per unit area, by species or higher taxon (unit weight may be grams wet weight, or grams / mg / $\mu$g carbon)</td>
<td></td>
<td></td>
<td>Total organic carbon (mg m$^{-2}$ and/or %)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sediment grain size → see geologist protocol (%fractions of silt, clay, sand; modal distribution)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Other sediment properties → see geologist protocol</td>
</tr>
</tbody>
</table>
10.4 SEDIMENT EXTRACTION FOR TRACE ELEMENTS

Parameter measured: Trace elements from sediments

Method Responsible: Murat Ardelan (NTNU, murat.v.ardelan@ntnu.no)

Description of parameter/Aim: Quantification of the trace element distribution and speciation in the sediment, through a four steps sequential extraction.

Methodological description:

→ Special requirements/gear: None. Samples can be collected from Box corer or Gravity corer

→ Sampling depths: Station bottom depth

→ Chemicals used:

Material:

- Plastic spatula
- Acid washed PE sampling tubes (15ml)

Method:

- Depending on the available instrument samples might collected for only surface sediment or different layers for a sediment depth profile.
- From each layer available, collect an amount of ~0.5 - 1 grams of sediment with the spatula
- Samples are collected in Acid washed PE tubes and stored frozen (-20 °C) until analysis.
- This is carried out in to obtain the following fractions:

Sequential Extraction:
1) Exchangeable, Acid Soluble
2) Easily Reducible
3) Oxidizable
4) Residual
10.5 DARK DIC UPTAKE IN SEDIMENTS

Parameter measured: Dark DIC uptake in sediments

Method Responsible: Tobias Vornahme (UiT, tobias.vornahme@uit.no), Rolf Gradinger (rolf.gradinger@uit.no)

Description of parameter/Aim: Estimation of dark DIC (Dissolved Inorganic Carbon) in sediments.

Methodological description:

→ Special requirements/gear: Isotope lab

→ Sampling depths: Station bottom depth

→ Chemicals used: 4% Formaldehyde, 1mCi/ml 14C-bicarbonate

Method:

• From the box cores 10ml surface sediment is diluted with 10ml filtered seawater and mixed into a slurry.
• The slurry is transferred into 2ml tubes for the uptake experiments and into 3 x 1.5ml tubes for later determination of the dry weight and normalization of the uptake rates.
• 5 tubes are filled with 1ml of the slurry using a cut off 1ml pipette tip.
• 2 of the tubes are killed immediately with 1ml 4% Formaldehyde in filtered seawater. All tubes are then enriched with 4ul 1mCi/ml 14C-bicarbonate.
• A second set of 5 tubes is treated in the same way with addition of 50ul saturated nitrification inhibitor solution.
• The tubes are incubated for 24h in the dark at in situ temperatures and killed with 1ml 4% Formaldehyde in filtered seawater.
• The samples will be processed as described by Molari et al. (2013).
• Conversion factors determined from biomass experiments in the central Arctic ocean (Molar et al. in prep) will be applied to calculate bacterial and archaeal biomass production.
• Nitrification will be determined as the difference between the treatments with and without nitrification inhibitor.

Literature:

11 Fish

Method Responsible: Elena Eriksen (IMR, elena.eriksen@hi.no → protocol responsible)

Fish sampling will be taken on process study stations, NLEG and additional station (chapter 3, Fig 3-1). The main variables measured will be fish distribution, species, biomass, length, weight and age (Stadstad procedure); and diet. Diet will be studied by traditional way (visual analyses and records of stomach content), isotopes (special procedure), chemical analyses (samples collected during the surveys and analysed on land at lab) and genetic analyses (samples collected during the surveys and processed on board, while father analysed will be taken on land at lab).

UPDATE: IMR will not conduct sampling for fish on the Nansen LEGACY cruises in 2019. Fish sampling will be conducted at the Joint Norwegian-Russian Ecosystem survey according to standard IMR/PINRO procedure, and these data will be used in the Nansen LEGACY. Additional fish stomach sampling for the Nansen LEGACY will be conducted at some stations during the Joint Norwegian-Russian Ecosystem survey. This sampling is described below.
**Tab. 11-1.** Overview over planned fish sampling activity during the Nansen Legacy cruises.

<table>
<thead>
<tr>
<th>Station name</th>
<th>Location</th>
<th>Bottom depth (m)</th>
<th>Activity at station</th>
<th>Standard activity at fish lab</th>
<th>Additional sampling</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test station/P1</td>
<td>76.0000° N, 31.2200° E</td>
<td>322</td>
<td>pelagic fish trawl (Harstad), demersal fish trawl (2 replicates; with and without fish lift)</td>
<td>Pelagisk og 1. bunntrål: Standard fish sampling, including stomach sampling of cod, polar cod and capelin.</td>
<td>Fish to diet_genetic: 10 ind of cod, polar cod and capelin for genetic.</td>
</tr>
<tr>
<td>NLEG03</td>
<td>77.0000° N, 34.0000° E</td>
<td>154</td>
<td>pelagic fish trawl (Harstad), demersal fish trawl</td>
<td>Standard fish sampling, including stomach sampling of cod, polar cod and capelin</td>
<td>If you didn't get samples at previous station, so take it here!!!</td>
</tr>
<tr>
<td>P2</td>
<td>77.5000° N, 34.0000° E</td>
<td>190</td>
<td>pelagic fish trawl (Harstad), demersal fish trawl (2 replicates; with and without fish lift)</td>
<td>Standard fish sampling, including stomach sampling of cod, polar cod and capelin</td>
<td>Fish to diet_genetic: take one sample (10 ind) of each fish species during the survey.</td>
</tr>
<tr>
<td>M3 (mooring location)</td>
<td>77.8700° N,</td>
<td>Acoustic surveying for mooring</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Location</td>
<td>Coordinates</td>
<td>Code</td>
<td>Sampled Fishes</td>
<td>Sampling Procedure</td>
<td>Genetic Analysis</td>
</tr>
<tr>
<td>--------------</td>
<td>-------------------</td>
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<td>---------------------------------------</td>
<td>------------------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>NLEG07 (P3 - but will not be sampled as P-station on this survey)</td>
<td>78.7500° N, 34.0000° E</td>
<td>301</td>
<td>pelagic fish trawl (Harstad), demersal fish trawl</td>
<td>Standard fish sampling, including stomach sampling of cod, polar cod and capelin</td>
<td>Fish to diet_genetic: Fish to genetic: take one sample (10 ind) of each fish species during the survey</td>
</tr>
<tr>
<td>NLEG09</td>
<td>79.2500° N, 34.0000° E</td>
<td>215</td>
<td>pelagic fish trawl (Harstad), demersal fish trawl</td>
<td>Standard fish sampling, including stomach sampling of cod, polar cod and capelin</td>
<td>Fish to diet_genetic: take one sample (10 ind) of each fish species during the survey</td>
</tr>
<tr>
<td>Mooring location M1</td>
<td>79.5930° N, 28.1000° E</td>
<td></td>
<td>Acoustic surveying for mooring deployment at later survey</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mooring location M2</td>
<td>79.6771° N, 31.9783° E</td>
<td></td>
<td>Acoustic surveying for mooring deployment at later survey</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P4</td>
<td>79.7500° N, 34.0000° E</td>
<td>332</td>
<td>pelagic fish trawl (Harstad), demersal fish trawl (2 replicates; with</td>
<td>Standard fish sampling, including stomach sampling of cod, polar cod and capelin</td>
<td>Fish to diet_genetic: take one sample (10 ind) of each fish species during the survey</td>
</tr>
<tr>
<td>Station</td>
<td>Latitude</td>
<td>Longitude</td>
<td>Sample Size</td>
<td>Methodology</td>
<td>Sample Description</td>
</tr>
<tr>
<td>---------</td>
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<td>-----------</td>
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<td>-------------</td>
<td>--------------------</td>
</tr>
<tr>
<td>P5</td>
<td>80.5000°N</td>
<td>34.0000°E</td>
<td>167</td>
<td>Pelagic fish trawl (Harstad), demersal fish trawl (2 replicates; with and without fish lift)</td>
<td>Standard fish sampling, including stomach sampling of cod, polar cod and capelin. Fish diet: freeze 10 ind of all captured small fish species. Fish to diet_genetic: 10 ind of cod, polar cod and capelin for genetic.</td>
</tr>
<tr>
<td>P6</td>
<td>81.5463°N</td>
<td>30.8548°E</td>
<td>865</td>
<td>Pelagic fish trawl (Harstad), demersal fish trawl (2 replicates; with and without fish lift)</td>
<td>Standard fish sampling, including stomach sampling of cod, polar cod and capelin. If you didn't get samples at previous station, so take it here!!! Fish diet: freeze 10 ind of all captured small fish species.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Short-term sea ice stations (5 hours each)</td>
<td>If possible, take a pelagic haul. If!!! So, standard fish sampling, including stomach sampling of cod, polar cod and capelin.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Sea ice stations (24 hours)/P 7 if sea ice conditions are suitable for it</td>
<td>If possible, take a pelagic haul. If!!! So, standard fish sampling, including stomach sampling of cod, polar cod and capelin.</td>
</tr>
</tbody>
</table>
11.1 FISH TRAWLING/SAMPLING OF POLAR COD, ATLANTIC COD AND CAPELIN BY DEMERSAL AND PELAGIC TRAWL

Parameter measured: See protocol “fish dissection” for more information

Method Responsible: Siv Hoff (UiO, s.n.k.hoff@ibv.uio.no), Sissel Jentoft (UiO, sissel.jentoft@ibv.uio.no) and Kim Præbel (UiT, kim.praebel@uit.no)

Description of parameter/Aim: Fish trawling and sampling for polar cod, atlantic cod and capelin by demersal and pelagic trawl

Methodological description:

→ Special requirements/gear:
  - Demersal trawl conducted with Campelen 1800# bottom trawl – provided by IMR
  - Pelagic trawl conducted with Harstad type mid water trawl – provided by IMR
  - Fish lift – provided by UiT
  - 2x 500 L seawater tanks on deck supplied with constant running seawater – Provided by UiT
  - Several 50 L buckets

→ Sampling depths:
  - Demersal trawl (with the fish lift) at sea floor (approx. 200 m depth) in sound scattering layer on shelf break
  - Pelagic trawl in sound scattering layer in the mesopelagic
  - Demersal and pelagic trawl should be conducted on all possible stations (at least one station south of polar front, two at the central shelf and one at the northern shelf)

→ Chemicals used: See protocol “11.2 Fish dissection” for more information
Method:

**Demersal trawl and pelagic trawl:**

- Demersal trawl for 5-15 min (trawl time depends on intensity of backscatter in sound scattering layer, checked on echosounder) at a towing speed of 2.2 – 3.4 knots
- Demersal trawl should be brought up slowly to avoid barotrauma in fish
- Pelagic trawl towed for 45 minutes at a towing speed of 2.2 – 3.4 knots (depending on intensity of backscatter in sound scattering layer, checked on echosounder)
- Empty trawl on deck in prefilled black buckets
- Quickly transfer desired fish (polar cod, Atlantic cod, capelin and other relevant species if we see that we can get nice sample size(s) of these) to seawater tanks (using small nets) for further processing and sampling
- Assess possibility of taking a second trawl to have enough fish samples for all participants
- Fish are kept in tank covered with a lid on deck and sampled successively during the next 24 hours after the trawl

→ Sample storage/ Shipment address

Fish will be sampled, see protocol “11.2 Fish dissection”

11.2 FISH STOMACH SAMPLING

**Measured parameter:** Stomach content analysis of fish

**Method responsible:** Institute for Marine Research (IRM, Tromsø/Bergen)

**Parameter description:**

Fish stomach sampling is a key element in study trophic interaction in the Barents Sea. In 2018, 203 stomachs were collected from 13 fish species sampled at the Nansen LEGACY cruise C1-2. I 2019, we plan fish stomach
sampling during the Joint Norwegian-Russian Ecosystem survey with “J. Hjort” in September. Focus is on fish species with limited knowledge of feeding ecology. Diet description based on stomach analyses will be supplemented with environmental data to give better understanding to species habitat and feeding preferences.

Stomach sampling will be conducted on species according to the prioritized list given below, and on stations during the standard monitoring survey. After standard processing of fish samples (standard procedure at IMR/PINRO, https://hinnsiden.no/tema/forskning/PublishingImages/Sider/SPD-gruppen/H%C3%A5ndbok_5.0_februar%202019.pdf), fish stomach will be studied by visual analyses and records of stomach content. The processing will be conducted on board if expertise is available. If not, frozen samples will be analysed at the laboratories at IMR.

Methodological description

- Special requirements/gear:
- Sampling depths:
- Chemicals used:

Sampling priority list:

Amblyraja hyperborean, Arctic skate
Bathyraja spinicauda, Spinetail ray
Rajella fyllae, Round ray
Lampanyctus macdonaldi, Rakery beaconlamp
Macrourus berglax, Rough-head grenadier
Enchelyopus cimbrius, Fourbeard rockling
Gaidropsarus argentatus, Arctic rockling
Phycis biennoides, Greater forkbeard
Gasterosteus aculeatus, Three-spined stickleback
Lophius piscatorius, Anglerfish
Sebastes norwegicus, Golden redfish
Sebastes viviparus, Norway redfish
Eutrigla gurnardus, Grey gurnard
Gymnocanthus tricuspis, Arctic staghorn sculpin
Myoxocephalus scorpius, Shorthorn sculpin
Aspidophoroides olrikii, Northern alligatorfish
Eumicrotremus derjugini, Leatherfin lumpsucker
Liparis bathyarcticus, Arctic snailfish
Lycodes eudipleurostictus, Doubleline eelpout
Lycodes polaris, Canadian eelpout
Lycodes seminudus, Longear eelpout
Anisarchus medius, Stout eelblenny
Lumpenus fabricii, Slender eelblenny
Ammodites marinus, Lesser sandeel
Glyptocephalus cynoglossus, Witch flounder
Hippoglossus hippoglossus, Atlantic halibut
Lycodes reticulatus, Arctic eelpout
Lycodes rossi, Threespot eelpout

11.3 FISH DISSECTION/BIOLOGICAL SAMPLING FOR POLAR COD, ATLANTIC COD AND CAPELIN – GENOMIC STUDIES

Parameter measured: Biological sampling for polar cod, Atlantic cod and capelin for genomic studies

Method Responsible: Siv Hoff (UiO, s.n.k.hoff@ibv.uio.no), Sissel Jentoft (UiO, sissel.jentoft@ibv.uio.no) and Kim Præbel (UiT, kim.praebel@uit.no)
Description of parameter/Aim:

- Fork length: the length of a fish measured from the most anterior part of the head to the deepest point of the notch in the tail fin.
- Total length: the length of a fish measured from the most anterior part of the head to the tip/end of the tail fin.
- Total weight: wet weight of fish in gram before dissection starts
- Presence of ectoparasites: visual check of the presence of ectoparasites on the fins or gills of the fish, record the number of ectoparasites
- Sex: is the fish male or female or immature (no gonads present in the fish)
- Gonad weight: weight of the gonad
- Liver weight: weight of the liver
- Somatic weight: weight of the fish when all inner organs are removed from the fish
- Maturation stage: On the basis of shape, size, color of the gonads and other morphological features, at least six maturity stages can be recognized
- Phenotype measurements: All parameters described above, i.e. fork length, total weight and length, presence of ectoparasites, sex, liver weight, gonad weight, maturation stage and somatic weight.

Methodological description:

→ Special requirements/gear:
  - Dissection should start right after the trawl comes up to minimize stress/degradation of DNA/RNA
  - We aim for dissection of 40 of each species (i.e. polar cod, capelin and Atlantic cod) in order to allow a balanced sex ratio at each station (P1, P2, P3, P4, and P5)

→ Sampling depths: see protocol “11.1. fish trawling”

→ Chemicals used: Ethanol (96%), RNAlater
Method:

- Record fork and total length, total weight, presence of ectoparasites
- Remove otoliths, place them in envelopes (for ageing later on)
- Open the fish -> record sex, maturation stage, sample tissue samples for DNA/RNA (see details below) and at then lastly record gonad weight.
- If sampling from same individuals as the ecotox-team we will also record:
  - liver weight and somatic weight in addition.

**Sampling priority list (for each station):**

<table>
<thead>
<tr>
<th>Priority after catch</th>
<th>Application</th>
<th>N</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>RNAseq – full set (Sissel)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>RNAseq – reduced set (Sissel)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>DNA – Genomics (Sissel)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Phenotype (Sissel)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>+</td>
<td>+</td>
<td>40</td>
</tr>
</tbody>
</table>

**RNAseq reduced set**

- Fish number 1 – 40: RNAseq – samples
- important to sample as soon as possible after death
- place in 5ml-15 ml cryovials with RNALater:
- Spleen*

*NB NB: For capelin we will instead of spleen sample liver as standard RNA sample.

**DNAseq sampling set**

- Fish number 1 – 40: Muscle/skin + spleen*
- place in 5ml-15 ml cryovials with 96% ethanol

*For polar cod we will sample gills instead as an additional DNAseq sample due to small size of the spleen.
Sample storage/ Shipment address:

- **RNAseq samples RNAlater (fish number 1-40)**
  stored at +4°C during the cruise and placed at -80°C for long term storage after shipment to UiO
- **DNA/genomic samples 96% ethanol (muscle & spleen of fish number 1-40)**
  stored at +4°C during the cruise and placed at -80°C for long term storage after shipment to UiO
- **Age determination on otoliths at room temperature**
- **Shipment of all samples to UiO**

11.4 PRECISION-CUT LIVER SLICE (PCLS) PREPARATION AND CULTURE FOR FISH

**Measured parameter:** The target species are Polar cod, Atlantic cod, capelin and other fish available during trawling.

**Method responsible:** Fekadu Yadetie (UiB, fekadu.yadetie@uib.no)

**Parameter description:** This is an ex vivo tissue culture method modified from Eide et al., 2013. The method describes preparation of liver slices, culturing and exposure to chemicals onboard for later analysis in the laboratory. The target species are Polar cod, Atlantic cod, capelin and other fish available during trawling. After the trawling, fish will be kept in running seawater for few hours while processing. Depending on availability, 10-20 fish will be sampled per station.

**Methodical description:**

→ **Special requirements/gear:**

- Clean space for vibratome/sample handling
- Fume hood for adding chemicals
- Access to cold room
- Access to liquid nitrogen and -80 °C freezer
Sampling depths:

Chemicals used: PCLS buffer, L-15 medium (L1518 Sigma), Charcoal-stripped fetal bovine serum, Antibiotic antimycotic solution (100x), Test chemicals for exposure studies, Ethanol, DMSO, PBS buffer, Heparin, Liquid Nitrogen, RNA later

Material:

- Bucket and water tanks with flow-through seawater supply
- Vibratome
- Ice
- Culture plates (12 and 24 well)
- Pipettes and tips 0.1-1000ul
- Superglue
- Petri dishes
- Syringes and needles
- Dissection set (knives, scalpel, scissors, forceps, tweezers, blades...)
- Aluminum foil
- Microcentrifuge tubes and racks
- Cryotubes and boxes
- Falcon tubes (15ml and 50ml) and racks
- Liquid nitrogen
- Foam dewars for LN2
- Weight scale
- Incubator
- Chemical hood
- Protection equipment and consumables (gloves, lab-coats, Faceshield, face masks, Safety Glasses)
- Microcentrifuge (refrigerated)
- Refrigerator/cold room and -20 and -80 freezers
- Distilled water

NOTE: Keep sterile working conditions. Use 70% ethanol to disinfect surfaces and equipment. Change gloves as often as necessary. Work on ice. The fish tissue should not be kept at room temperature.
Method:

PREPARATIONS:

- Put PCLS buffer and on ice or in the fridge.
- Label 15 and 50 ml falcon tubes.
- Thaw FBS and anti-biotic-antimycotic solutions and prepare the complete culture medium: (44.5 ml L-15 medium + 5 ml FBS + 0.5 ml anti-biotic-antimycotic 100x solution). Keep on ice.
- Fill a big petri dish with the PCLS buffer.
- Label and fill small petri dishes with the complete culture medium for liver blocks and keep on ice.
- Prepare small petri dishes with the complete culture medium for collection of slices.
- Prepare labeled cryotubes, microcentrifuge tubes and aluminum foil for collecting blood and tissue (liver, bile).
- Label 15 ml tubes for histology of gonads and liver if necessary.
- Prepare dissection set (sterilized): Scissors, razor blade, knives, tweezers, etc), different size needles and syringes (5 ml, 10 ml and 50 ml), heparin, kim-wipe, tissue paper, aluminum foil, bench paper. Washing ethanol (70%), superglue.
- Spray surfaces and the vibratome with 70% ethanol.

PREPARATION AND SLICING FOR LIVER TISSUE:

1. Kill the fish by blowing in the head and measure the length and weight. After the fish is dead keep it cold (on ice).
2. Take blood samples from caudal vein, in heparinized syringe and keep it on ice in microcentrifuge tube. Centrifuge the blood at 2000g for 10 min and freeze plasma samples at -80°C). Take a picture of the fish.
3. Keep the fish on ice and open the peritoneal cavity taking care not to puncture the intestine. Dissect out the liver and put it in a big petri dish with ice-cold PCLS buffer. Take the bile sample using syringe and needle. Record the liver weight and keep it on ice. The liver should not come in contact with non-sterile objects. Use clean (sterile) gloves and sterilized equipment to handle the liver. Change to new ice-cold PCLS buffer (to rinse the liver), keep on ice. Take a picture of the liver.
4. Dissect the gonad and record the sex and weight. Note the developmental stage (juvenile, mature, etc). Take a picture of the gonad.
5. Carefully cut out blocks of liver tissue (approximately 3 cm long and 2 cm wide; the height may vary 1-2 cm) and keep in ice-cold **complete culture medium**.
6. Using superglue to fix the blocks of liver tissue on the specimen plate of Leica vibrating blade microtome VT1200 (Leica, Wetzlar, Germany). Keep on ice until the superglue is dry and the liver tissue is stably fixed on the sample holder.
7. Assemble the specimen plate (with the tissue) into the buffer tray and the ice tray, fill the ice tray with ice and mount it on the microtome.
8. Mount and position the blade and cut 250 μm thick slices/strips at a speed of 0.9 mm/s and amplitude 3 mm. The tissue should be kept submerged in ice-cold PCLS buffer during slicing.
9. Collect the sliced 250 μm strips in a petri dish with ice-cold complete culture medium.
10. Using razor blade, cut the strips into smaller (approximately 4x4 mm) slices and pre-incubate at appropriate temperature for the species (10°C for cod) for 2 hours in the **complete culture medium** before exposure. About 4-10 slices can be cultured in 12-well plate with 2 ml **complete culture medium** per well. Divide equal number of slices into each well.
11. In the fume hood, prepare the exposure chemical in DMSO at 1000x final concentration of each dose-group (e.g. 1mM, 10mM and 100mM for final concentration of 1μM, 10μM and 100μM, respectively). Then prepare **complete culture medium** with 2x the final concentration (2uL/ml) of each dose-group.
12. After pre-incubation for 2 hours, replace half of the volume of the medium in each well (1ml) with the **complete culture medium** containing 2x the final concentration of the exposure compound and incubate at appropriate temperature (10°C for cod) with shaking at 50 rpm, for the required length of time (24 hours to 1 week). All wells including vehicle control should contain equal volume of DMSO (not exceeding 0.5% (v/v)).
13. At the end of the culture, carefully pick the slices with fine tweezers, rinse briefly in cold PBS or PCLS buffer and put in pre-weighed (and tared) cold cryotube and record the weight. Snap-freeze in liquid N2 and store at -80 °C. If liquid N2 is not available, collect the slices in a cryotube.
tube with RNAlater solution for RNA extraction. Incubate the samples in RNAlater solution overnight at 4°C and then transfer to -80°C for storage.

14. Store the media (in cryotubes) at -80 °C for LDH assay and other purposes as necessary (e.g. vitellogenin assay).

12 Marine mammals

Marine mammals related research, while part of the Nansen Legacy, is not a part of the RV Kronprins Håkon cruises. For further information on ongoing research contact:

**Method responsible:** Marin Biuw (IMR, martin.biuw@hi.no), Kit Kovaks, (NPI, kit.kovacs@npolar.no), Christian Lydersen (NPI, Christian.lydersen@npolar.no), Tore Haug (IMR, tore.haug@hi.no)

13 Ecotoxicological samples

**Measured parameter:** Several parameters are sampled for ecotoxicological examination

**Method responsible:** Katrine Borgå (UiO, katrine.borga@ibv.uio.no), Julia Giebichenstein (julia.giebichenstein@ibv.uio.no), Robynne Nowicki (robynneno@unis.no)

**Parameter description:** Collection of meso- and (low priority for II) - macrozooplankton, benthos and fish for stable isotopes, fatty acids, carbohydrates, protein and energy content during the Nansen Legacy biology seasonal process cruises. The taxa sampled will be the same for Aim I and Aim II, and data will be shared when overlapping parameters are analysed.
Aim I: Investigate seasonal food web biomagnification of contaminants in the present climate conditions compared to 2 decades earlier. Samples will be taken of water, zooplankton, fish (and benthos) for pollutants, stable isotopes and Hg. For future cruises: birds and mammals if possible to include logistically.

Aim II: Investigate seasonal and spatial patterns of energetic level and contaminants (organics and Hg) of main species at different trophic level in benthic and pelagic food chains.

IMPORTANT:

- NOTE: very important to avoid cross-contamination during handling.
- Samples stored in -20°C freezer.
- We will also sample opportunistically for microplastics screening, unless this is already part of the HI/NP sampling plans.
- Focus will be on the process stations which cover different areas.

13.1 WATER SAMPLING

13.1.1 POPs – Persistent Organic Pollutants

Parameters measured: POPs – Persistent Organic Pollutants

Method responsible: Julia Giebichenstein (julia.giebichenstein@ibv.uio.no), Katrine Borgå (katrine.borga@ibv.uio.no)

Parameters description: This water sampler is built at the workshop at IBV based on the design of a similar device used at NIVA. The water sampler has a pump that actively pumps water through the system and measures the water flow with a flow meter connected inside the pump. The system is suitable for use with a glass fiber filter (293mm) and polyurethane filters (PUFs). Filter and PUFs may be extracted and analyzed for organic contaminants. It is made for use at depths up to 200 m in both fresh and saltwater.
Methodical description:

→ **Special requirements/gear:** IBVs water pump, deploy water sampler for 3-4 hours to filter water (1-4 L/min). The water sampler should be placed upstream of the ship, attached to a buoy or to the ship with wire.

→ **Sampling depths:** Process stations only, via IBV water pump

→ **Chemicals used:**

Method:

Make sure to not wear any water repellent- or new/unwashed clothing, treated/water resistant boots, cosmetic skin products (sunscreen, moisturizer etc.). Do not handle any prepackaged food or products mentioned before when sampling. Wash hands well before handling sampling equipment.

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*Fig. 13.1.1-1* Location of glass fiber filter holder (1), and PUF holder (2)

*Fig. 13.1.1-2* Location of connection plug to USB cable.
Filter 6x 500L (3000L per area), make sure to have field blanks (normal sampling procedure without actual filtering), and get a blank from the water hose used onboard.

Label two Ziploc bags per water sample with Date, Station ID, Coordinates, leave a blank sheet inside the bag for notes to take during the sampling (e.g. temperature, if anything didn’t work out, etc.). Precut a large piece of Aluminum foil and place it in the bag (together with the second Ziploc bag).

1. Wearing gloves, install glass fiber filter (1) and polyurethane filters (PUFs) (2), (Figure 13.1.1-1).
2. Plug in the DELL Pad using the USB cable (3), (Figure 13.1.1-2). Turn on Pad (does not require any password) and check if the screen shows “BATTERY OK”.
3. Programming the pump is done using the software named AquaFilter
4. Remove the silicone tubes and make sure the system is soaked in water before start and to avoid air disrupting the flow meter.
5. The USB plug may be inserted at any time if you want the process to stop.
6. Ensure that enough water is constantly flowing through the system: the pump will stop if the water flow is not good enough (too light flow, or particles clogging the filter).
7. The pump will stop after the desired volume is reached (or if something has gone wrong).
8. Plug in USB and open the program on the Pad. The screen will show “MOTTAR DATA”. A .txt file with information on the last run will be saved under “Aquafilter” on the C-disk.
9. The water sampler should be rinsed with clean, fresh water after use – especially if used in salt water or dirty water.

**Sample storage**

Place the filters in Aluminum foil in two ziplock bags, store cool for transfer processes and freeze at -20°C for long-term storage.
13.1.2 Total Mercury (TotHg)

**Measured parameter:** Total Mercury (TotHg) – low priority

**Method responsible:** Julia Giebichenstein (julia.giebichenstein@ibv.uio.no), Katrine Borgå (katrine.borga@ibv.uio.no)

**Parameters description:**

**Methodical description:**

→ **Special requirements/gear:** Water from Niskin rosette

→ **Sampling depths:** Process stations only

→ **Chemicals used:**

**Method:**

Collect sample water (250 mL) from CTD mounted water collector (or similar).

1. Wearing gloves, put the 250 mL FLPE bottle inside an open ziplock plastic bag and unscrew the lid.
2. Hold the bottle underneath the water collector and fill it to the rim. There should be no air left in the bottle.
3. Screw lid back on, wrap it in the ziplock bag, close the zip, then wrap a second ziplock plastic bag around the bottle and close.
4. Refrigerate at 4°C.
5. Label the bottle with sample ID, station ID, date.

*Figure 13.1.2-1 250 mL FLPE bottle*
13.1.3 PFAS – perfluorinated alkylated substances

**Measured parameter:** PFAS – perfluorinated alkylated substances

**Method responsible:** Julia Giebichenstein (julia.giebichenstein@ibv.uio.no), Katrine Borgå (katrine.borga@ibv.uio.no)

**Parameters description:**

**Methodical description:**

- **Special requirements/gear:** Water from Niskin rosette
- **Sampling depths:** Process stations only
- **Chemicals used:**

Method:

Make sure to not wear any water repellent- or new/unwashed clothing, treated/water resistant boots, cosmetic skin products (sunscreen, moisturizer etc.). Do not handle any prepackaged food or products mentioned before when sampling. Wash hands well before handling sampling equipment.

Collect sample water (1L) from CTD mounted water collector (or similar).

1. Wearing powderless nitrile gloves, hold the 1L PP bottle underneath the water collector and fill it to the rim. There should be no air left in the bottle.
2. Label the bottle with sample ID, station ID/position, Date using an ink pen (avoid sharpies and other black markers)
3. Screw the lid back on and refrigerate at 4°C
13.2 ZOOPLANKTON SAMPLING

**Measured parameter:** Collection of meso- and microzooplankton to analyse for persistent organic pollutants (POPs), mercury (Hg), stable isotopes, fatty acids, carbohydrates, protein and energy content

**Method responsible:** Julia Giebichenstein (julia.giebichenstein@ibv.uio.no), Robynne Nowicki (robynneno@unis.no), Katrine Borgå (katrine.borga@ibv.uio.no)

**Parameters description:** Collection of meso- and macrozooplankton to analyse for persistent organic pollutants (POPs), mercury (Hg), stable isotopes, fatty acids, carbohydrates, protein and energy content during the Nansen Legacy biology seasonal process cruises. Additional samples are taken for Chlorinated paraffins (CPs) & perfluorinated alkylated substances (PFAS). The taxa sampled will be the same for Nansen Legacy workpackages 2.2.1 & 2.2.5, and data will be shared when overlapping parameters are analysed.

**Methodical description:**

→ **Special requirements/gear:**

- Macrozooplankton trawl (open water): opening area 38 m² mouth, 3 mm mesh size.
- Macrozooplankton ring net (in ice): opening area 2.01 m², 7 m long net bag with 1.55 mm or 4 mm mesh size.
- WP2 180 µm (opening 0.25m²) (NP/UiT) will be used for sampling mesozooplankton (only if time permits)
- WP3 1000 µm (opening 1m² (NP) will be used for sample larger mesozooplankton (only if time permits))
- Specification regarding the deployment of the different nets can be found in “Zooplankton sampling protocol AeN RF3”
- We need information about mesozooplankton and macrozooplankton community composition, obtained in RF3.

→ **Sampling depths:** Process stations only, 3-5 replicates per area. Zooplankton samples should be sampled from the entire water column.
Two samples per station, one in the Atlantic layer and one in the Arctic layer.

**Chemicals used:** Ethanol (for cleaning)

**Method:**

Sorting of zooplankton samples should be done as soon as possible after sampling. Samples should be diluted with *in situ* sea water and stored in 50L buckets in a cold room, to be processed as soon as possible. Keep the samples cold while sorting by use of cold packs or ice. Zooplankton should be sorted into species level if possible and larger macrozooplankton should be grouped into length groups (see table below). Samples can be stored in cryo vials, aluminum foil or zip-bags depending on the size of the sample. All samples should be stored at -20°C except for fatty acid samples that should be stored at -80°C (the other samples could also be stored at -80°C).

**Organic pollutants (>2 g):**

Only for Nansen Legacy work packages: RF 2.2.1
**Tab. 13.2-1** Target taxa for the different sample types. Minimum number of individuals per sample is indicated for zooplankton taxa.

<table>
<thead>
<tr>
<th>Target taxa</th>
<th>Organic pollutants</th>
<th>Stable isotopes</th>
<th>Hg</th>
<th>Fatty acid</th>
<th>Energetics</th>
<th>CPs / PFAS</th>
<th>Protein</th>
<th>Carbo-hydrates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sampled for 2.2.1</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Sampled for 2.2.5</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Sample size</td>
<td>5-10 g</td>
<td>0.5 g</td>
<td>0.5 g</td>
<td>0.5 g</td>
<td>1 g</td>
<td>xx g</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of replicates</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Priority</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Calanus spp.</strong></td>
<td></td>
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<td></td>
<td>20-30 ind</td>
<td>20-30 ind</td>
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<td>20-30</td>
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<tr>
<td>Other dominant species</td>
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</tr>
<tr>
<td><strong>Thysanoessa spp. 0-10 mm</strong></td>
<td>5 ind</td>
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<tr>
<td><strong>Thysanoessa spp. 10-20 mm</strong></td>
<td>5 ind</td>
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<tr>
<td><strong>Thysanoessa spp. 20-30 mm</strong></td>
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<tr>
<td><strong>Thysanoessa spp. &gt;30 mm</strong></td>
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<tr>
<td><strong>Themisto libellula 0-10 mm</strong></td>
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<tr>
<td><strong>Themisto libellula 10-20 mm</strong></td>
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<tr>
<td><strong>Themisto libellula 20-30 mm</strong></td>
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<td><strong>Themisto libellula &gt;30 mm</strong></td>
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<tr>
<td>Species</td>
<td>0-10 mm</td>
<td>10-20 mm</td>
<td>20-30 mm</td>
<td>30-40 mm</td>
<td>5 ind</td>
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<tr>
<td>Themisto abyssorum</td>
<td>5 ind</td>
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<td>5 ind</td>
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<tr>
<td>Themisto abyssorum 10-20 mm</td>
<td>5 ind</td>
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<tr>
<td>Themisto abyssorum 20-30 mm</td>
<td>5 ind</td>
<td>5 ind</td>
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<td>5 ind</td>
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<tr>
<td>Themisto abyssorum 30-40 mm</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
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<tr>
<td>Sagitta spp.</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mertensia ovum</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Beröe cucumis</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td></td>
<td></td>
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<tr>
<td>Other dominant species</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td></td>
<td></td>
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<tr>
<td>Clione limacina</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Limacina helicina</td>
<td>5 ind</td>
<td>5 ind</td>
<td>5 ind</td>
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<td></td>
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</tr>
</tbody>
</table>

**Stable isotopes (0.5 g) and Hg (0.5 g):**

Mesozooplankton and small macrozooplankton samples should be stored in small cryo vials. Larger macrozooplankton, benthos and fish liver and muscle samples should be stored in aluminum foil into plastic bags (dorsal muscle fish and larger zooplankton). Store frozen at -20°C.

**Fatty acids (1 g):**

We will assist others and are interested in these data, but will not have capacity for a full scale sampling and analyses on our PM and budget. Samples should be sorted as soon as possible and cryo vials should be frozen immediately in
at -80°C. Larger macrozooplankton, fish samples and benthic samples should be stored in cryo zip bags.

**Carbohydrates (1 g):**

Similar procedure for sorting and storing the samples as for stable isotopes.

**Protein (1 g):**

Similar procedure for sorting and storing the samples as for stable isotopes.

**Energetics (1 g):**

Analyses will be done using a Micro-bomb calorimeter. Alternative analyses will also be considered in order to compare methods.

### 13.3 FISH SAMPLING – INDIVIDUAL AND TISSUE SAMPLING

**Measured parameter:** Collection of fishes for analyses of persistent organic pollutants (POPs), mercury (Hg), stable isotopes, fatty acids, carbohydrates, protein and energy content

**Method responsible:** Julia Giebichenstein (julia.giebichenstein@ibv.uio.no), Robynne Nowicki (robynneno@unis.no), Katrine Borgå (katrine.borga@ibv.uio.no)

**Parameters description:** Collection of fishes for analyses of persistent organic pollutants (POPs), mercury (Hg), stable isotopes, fatty acids, carbohydrates, protein and energy content during the Nansen Legacy biology seasonal process cruises. Additional samples are taken for Chlorinated paraffins (CPs)
& perfluorinated alkylated substances (PFAS). The taxa sampled will be the same for Nansen Legacy workpackages 2.2.1 & 2.2.5, and data will be shared when overlapping parameters are analysed. We will also sample opportunistically for microplastics screening, unless this is already part of the HI/NP sampling plans.

**Methodical description:**

→ **Special requirements/gear:**

- Beam and Campelen trawl
- Dissection should start right after the trawl comes up to minimize stress, avoid cross contamination by working as cleanly as possible
- We need information about fish community composition as background information, obtained in Nansen Legacy work packages: RF3.

→ **Sampling depths:** Process stations only, 3-5 replicates per area (sampling depth: see protocol HI s protocol “fish trawling”)

→ **Chemicals used:** Ethanol (for cleaning)

Method:

**NOTE: very important to avoid cross-contamination during handling!!!**

Fishes should be sampled with a pelagic and beam trawl. The target species are polar cod (*Boreogadus saida*), capelin (*Mallotus villosus*) and Atlantic cod (*Gadus morhua*). If other fish species (that can form seabird prey) are abundant, then these should be considered as well. Samples should be taken of muscle and liver of fishes (maximum size <25). Biometrics, such as total length, whole fish weight, liver weight, age (otoliths), sex, maturation, reproductive stage and gross pathology (parasites, etc.) should be recorded. GIT should be sampled for diet and microplastic analyses.
Carefully write location and type of tissue (M - muscle, L - liver) on the bags and vials. Fill in data at the end of each day in an excel file.

Always wear gloves and clean the dissection equipment between each sample with Ethanol.

- Record total length, total weight, presence of ectoparasites
- Carefully dissect the fish and remove internal organs
- Freeze GIT
- Weigh liver, take samples for fatty acids
- Weigh the gonads and note the sex
- Weigh the empty fish (somatic weight)
- Take muscle (or whole fish) samples for POPs, stable isotopes, mercury
- Remove brain and otoliths

**Tab. 13.3-1 Ecotox fish sampling overview with sampling priority**

<table>
<thead>
<tr>
<th>Target taxa</th>
<th>POPs</th>
<th>Stable Isotopes</th>
<th>Total Hg</th>
<th>Fatty acids</th>
<th>PFAS /CP</th>
<th>Energy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polar cod (Boreogadus saida)</td>
<td>muscle / whole fish (w/o GIT), brain, liver</td>
<td>muscle</td>
<td>muscle</td>
<td>liver</td>
<td>whole fish</td>
<td>whole fish</td>
</tr>
<tr>
<td>Capelin (Mallotus villosus)</td>
<td>muscle / whole fish (w/o GIT), brain, liver</td>
<td>muscle</td>
<td>muscle</td>
<td>liver</td>
<td>whole fish</td>
<td>whole fish</td>
</tr>
<tr>
<td>Atlantic cod (Gadus morhua)</td>
<td>muscle / whole fish (w/o GIT),</td>
<td>muscle</td>
<td>muscle</td>
<td>liver</td>
<td>whole fish</td>
<td>whole fish</td>
</tr>
<tr>
<td>Opportunistic species</td>
<td>brain, liver</td>
<td>muscle / whole fish (w/o GIT), brain, liver</td>
<td>muscle</td>
<td>muscle</td>
<td>liver</td>
<td>whole fish</td>
</tr>
<tr>
<td>-----------------------</td>
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<td>---------------------------------------------</td>
<td>--------</td>
<td>--------</td>
<td>-------</td>
<td>------------</td>
</tr>
<tr>
<td>Sample size</td>
<td>&gt;5 g</td>
<td>0.5 g</td>
<td>1 g</td>
<td>1 g</td>
<td>&gt;5 g</td>
<td>&gt;5 g</td>
</tr>
<tr>
<td>Number of replicates per area</td>
<td>10</td>
<td></td>
<td></td>
<td>10</td>
<td></td>
<td>10</td>
</tr>
<tr>
<td>Priority</td>
<td>1</td>
<td></td>
<td></td>
<td>2</td>
<td>1</td>
<td></td>
</tr>
</tbody>
</table>

**POPs:**
- 10 replicates / species
- Muscle tissue (>2g) or whole fish
- Place in precut Aluminum sheet in Ziploc bag
- Freeze at -20°C

**Mercury:**
- 10 replicates / species
- Muscle tissue (1g)
- Place in Cryotube / Ziploc bag
- Freeze at -20°C

**Stable isotopes:**
- 10 replicates / species
- Muscle tissue (1g)
- Place in Cryotube / Ziploc bag
- Freeze at -20°C

**Fatty acids:**
- 10 replicates / species
- Liver tissue
- Place in precut Aluminum sheet in Ziploc bag
- Freeze at -80°C

**CPs / PFAS:**
- 10 replicates / species
- Wrap the whole fish individually in Aluminum foil
- Freeze at -20°C

**Energetics:**
- 10 replicates / species
- Wrap the whole fish individually in Aluminum foil
- Freeze at -20°C

→ **Sample storage & shipment address**
- POP, stable isotope and mercury samples: Stored at -20°C during the cruise and at -20°C after shipment to UiO
- Fatty acids: Stored at -80°C during the cruise and at -80°C after shipment to UiO
- Energetics: Stored at -20°C during the cruise and at -20°C after shipment to UiO
• CPs / PFAS: Stored at -20°C during the cruise and at -20°C after shipment to UiO

14 Sea ice work

14.1 SAFETY BRIEF
Prior to any ice station, a safety briefing is suggested to occur with the ship leadership, chief scientist and two to three scientists leading the ice work. The briefing should:

• Discuss the activities at the station, the needed time and safety precautions.
• Include a risk evaluation (e.g. wind, temperature, ice conditions, visibility (e.g. fog), swell, bears).
• Outline, when and how many people will be on the ice, and how long breaks will be with instrumentation on the ice and no people on the ice.
• Clarify communication on the ice/with the bridge (VHF channel, clear communication who is on/off the ice).

For safety reasons signals from the ship should be agreed upon (e.g. three times the ship's horn blowing) which will lead to immediate evacuation of the ice floe by scientists.

14.2 SELECTION OF THE ICE FLOE
The selection of the ice coring site should start ca 1 hour prior to the arrival of the ship at station with a 10-20-minute ice observation by a deciding person (e.g. chief scientist with one or two leaders of the ice teams). The task is to establish what kind of ice is typical for the area, what kind of different ice types exist. The team will pick an ice type where:

a) the ship can safely be moored, and

b) a typical ice type can be sampled.
While parking the ship alongside the floe or within the floe, ice thickness can typically be evaluated. Before any person enters the ice, it should be clarified on the bridge, where sampling will occur and what an acceptable distance from the ship is.

The first people going onto the ice might wear dry suits to check ice thickness with a thickness drill at sampling sites and transect.

Some general comments regarding the ice station selection and length:

- Try to avoid ship shadow in sampling area. Keep a distance of at least one ship length from ship for sampling if possible.
- Relatively homogenous level ice should be the preference for the short ice stations and the long ice stations.
- Once the main coring site for physical/biological sampling is decided upon, a clear plan should be made, where people can walk not to interfere with measurements.
- At a short ice station (5 hours):
  - Only one type of sea ice will be sampled.
  - No in situ BP/PP and sediment trap deployments will be conducted.
  - Additional sampling of melt ponds can be done.
- For the long ice stations (24 hours):
  - Ideally a second and third typical ice type (e.g. refrozen lead, thicker snow cover or other features such as ridges or multi-year ice) or regional patchiness in same ice type could be sampled by coring and covered in the ice thickness profiles.
  - Also in situ deployment of PP/BP production measurements and sediment traps occur as well as extended ice thickness profiles.
  - Several types of melt ponds could be sampled.
14.3 ICE WORKING TEAMS

On each AeN cruise, a core group needs to be assigned, who are responsible for sampling, note keeping, and then those who are responsible for processing and subsampling once samples are melted (and take care that melting is done properly).

Two ice coring/sampling teams are suggested for each ice station. Each team should consist of a minimum of four ideally five people (three/four working scientists plus one polar bear guard).

NOTE: Avoid making the team too big (for safety and for logistical reasons).

A) The following teams will work on a short ice station (5 hours):

Team 1:
- coring cores for physics, chemistry, nutrients, stable isotopes,
- ice thickness transects

Team 2:
- coring for biological parameters
- melt pond sampling
- under-ice water sampling

B) The following teams will work on long ice stations (24 hours):

Team 1:
- coring cores for physics, chemistry, nutrients, stable isotopes,
- ice thickness transects
- snow sampling

Team 2:
- making hole for PP and sediment traps
- coring for biological parameters
• deployment of sediment traps and PP/BP as soon as ready
• continue coring for biological parameters
• sampling melt ponds
• sampling under-ice water
• recovery of in situ deployments of sediment traps prior to any sediment/benthic sampling
• recovery of PP after ca 24 hours or end of station. BP could be recovered after 1-2 hours.

14.4 DEPLOYMENT OF IN SITU INCUBATIONS/SEDIMENT TRAPS
These deployments should be done as fast as possible during the long stations to allow long incubation times.

Making Ice hole:
• Use ice auger and ice saws to cut a hole into the sea ice (size: 75x75cm).
• If ice is thicker than 1.5m likely no hole big enough for sediment traps can be made.
• Than deploy only PP through ice auger hole as soon as the incubations are ready (spiked water samples from CTD and from above mentioned).
• For sediment traps you can use an open lead close to the ship (not for PP/BP) (risk: leads can close).

14.5 SAMPLING OF ICE CORES
The minimum team for ice coring consists of 3 (ideally 4) people taking and processing cores and samples, one person on polar bear watch and one person taking notes.

→ Responsibilities should be clearly identified.
→ Proper labelling of all taken samples is essential.

For handling the corer:
• use sturdy field gloves that are water proof.
• Avoid touching the cores with bare hands.
• Be careful when handling the corer to not cut any person.
• Be very careful during the coring procedure as you deal with rotating gear that can break fingers...
• All cores should only be handled with lab gloves to avoid contamination.
• Also avoiding pieces from the cutting boards and clothing is instrumental (thin woolen gloves can be worn below lap gloves to protect from the cold).
• **For genomic work, rinse the corer with ethanol prior to sampling.** Also make sure that the cutting board is clean.
• All biological process samples (PP, BP, genomics) should NOT be exposed to bright sunlight. Use dark foil or ice tent to protect from high light intensities.

→ **Clean hand & dirty hands approach:** DIRTY HANDS only use the corer and do not touch the core at all; the CLEAN HANDS get the core from the barrel and cut it.

### 14.5.1 Snow sampling

Should be done mainly for physical and nutrients properties, potentially also for biology.

**BEFORE ICE CORING:**

• Snow samples are collected at each sea ice collection area.
• Depending on the thickness of the snow, 1-2 snow samples are collected, e.g. when snow cover is 10 cm thick, one sample 0-5 cm (ice surface to 5 cm) and one sample 5-10 cm (5 cm to snow surface) are collected.
• If there is a slushy layer, this layer is collected as well.
• Temperature is measured in each snow/slushy layer.
• The snow is collected using a plastic spoon and put into gas-tight bags, then air is removed with the air pump, then the snow sample is melted in cool and dark place.
• Same procedure is used as for the sea ice samples.
• When melted, samples for DIC/AT, $\delta^{18}O$, nutrients are taken according to the Sampling protocol.

14.5.2 Cutting of the ice core and core parameters to be sampled

Ice cores should generally be cut into 10 to 20 cm long sections starting from the top if the core is longer than one corer length (so you process the first part of the total core from the top). HOWEVER, the lowermost part of the core should be cut starting from the bottom with the following bottom segments (bottom 10cm):

- 0-1cm, 1-3cm, 3-10, 10-20cm
- followed by 10 to 20cm long sections.

**Tab. 14.5.2-1** Table showing the core names, number of cores that has to be sampled and which parameters will be measured from it. The cores should be taken in the following sequence:

<table>
<thead>
<tr>
<th>No.</th>
<th>Core name</th>
<th>Amount</th>
<th>Parameter measured</th>
<th>Cutting scheme</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Physics core</td>
<td>2</td>
<td>1: temperature, salinity; 2: stratigraphy, density</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Nutrient chemistry core</td>
<td>2</td>
<td>1: One core for gases/CO2 (DIC/pH/AT), $\delta^{18}O$, nutrients and other tracers; 2: Backup (frozen whole)</td>
<td>cut into 10 cm sections, start from top</td>
</tr>
<tr>
<td>3</td>
<td>Chemistry core</td>
<td>2</td>
<td>1: tracers and gas measurements; 2: Backup (frozen whole)</td>
<td>cut into 10 cm sections</td>
</tr>
<tr>
<td>4</td>
<td>Backup core</td>
<td>1</td>
<td>Frozen whole</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Primary/bacteria productivity cores</td>
<td>3</td>
<td>Primary and bacteria productivity</td>
<td>cut off the bottom 1cm</td>
</tr>
<tr>
<td>7</td>
<td>Primary production P vs I curve core</td>
<td>1</td>
<td>Primary production</td>
<td>cut off the bottom 1cm</td>
</tr>
<tr>
<td>8</td>
<td>Biology cores</td>
<td>3</td>
<td>Chl a, POC/N, particle absorption, DNA/RNA?</td>
<td>Pooled! cut into 0-1, 1-3, 3-10, 10-20 and 20 cm</td>
</tr>
<tr>
<td></td>
<td>Sections</td>
<td>Meiofauna and Protist Cores</td>
<td>Abundance</td>
<td>Only the Bottom 30 cm! Cut into 0-1, 1-3, 3-10, 10-20 and 20 cm Sections Thereafter Starting From Bottom</td>
</tr>
<tr>
<td>-------</td>
<td>----------</td>
<td>-------------------------------</td>
<td>-----------</td>
<td>--------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>9</td>
<td>Meiofauna and Protist Cores</td>
<td>3</td>
<td>Meiofauna and Protist Abundance</td>
<td>Only the bottom 30 cm! cut into 0-1, 1-3, 3-10, 10-20 and 20 cm sections thereafter starting from bottom</td>
</tr>
<tr>
<td>10</td>
<td>Stable Isotope Core</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>Genomic Cores</td>
<td>3</td>
<td></td>
<td>Pooled!</td>
</tr>
<tr>
<td></td>
<td>Total Number of Cores</td>
<td>21</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Physics Core (2):**

- One core for temperature and salinity
- One core for stratigraphy and density

**Nutrient Chemistry Core (2):**

→ Cut into 10 cm sections, start from top:

1) One core for gases/CO2 (DIC/pH/AT), δ18O, nutrients and other tracers (9 or 14-cm corer). Melt each ice section in special gas-tight bags (see Fig. 14.5.2-1).
2) Plus one chemistry backup core (frozen whole)

→ For further processing see protocol 14.5.3
Chemistry core (2):

→ cut into 10 cm sections:

1) One core for tracers and gas measurements (14-cm corer). Merge lot of the tracers/biogeochemistry into one sample volume. Melting in special gas-tight bags (see Fig. 14.5.2-1).
2) One chemistry backup core (frozen whole)

→ for further processing see protocol 14.5.3

Backup core (1):

- Collect and keep frozen

Primary / bacterial productivity cores (3):

- Three cores, and cut off the bottom 1 cm (maximum biomass layer), place into clean zip-lock bag, add 500ml of 0.2μm filtered sea water and mix thoroughly.
• Than transfer into incubation bottles for PP and BP.
• Do all work in ice tent.
• Immediately prepare the incubations by adding the tracers in the isotope lab or if possible directly in the field.
• Transport always in a dark cooler.

**Primary production P vs I curve core (1):**

• Take one ice core, cut off bottom 0-1cm.
• Take back to ship and add 450ml of GF/F filtered sea water in Duran bottle.
• Use for P vs I curve.

**Biology cores (3):**

Three 9-cm cores pooled into one sample for biology (bulk variables: Chl a, POC/N, particle absorption, DNA/RNA?)

Biology cores BULK (cut into 0-1, 1-3, 3-10, 10-20 and 20 cm sections thereafter starting from bottom):

• Ice core sections are transferred into melting buckets (1 – 6 L depending on section length)
• Ice core sections are chopped up inside buckets to speed up melting process (make sure chopping device is clean)
• Ice cores are melted directly (without addition of FSW) at room temperature if constantly monitored or in a cold room (4°C). Needs to be done in the dark
• Swirl bucket contents regularly to ensure homogenous, low temperature
• As soon as ice cores have melted proceed with sample processing
• First measure total volume of sample, take salinity reading with salinity probe and make sure melting buckets are well mixed before subdividing into various parameters
• Measure exact volume with measuring cylinder for each parameter
• Proceed as outlined in sampling protocols for each variable

After cutting all biological samples need to be stored dark and in a cooler for transport back to the ship.

Meiofauna and protist cores (3):

→ **Chemicals used**: 37% formaldehyde (buffered), Lugol
→ cut into 0-1, 1-3, 3-10, 10-20 and 20 cm sections thereafter starting from bottom (process only the bottom 30cm)

• Three extra cores are taken for meiofauna.
• Place into zip-loc bags and add 100ml of GF/F filtered sea water for each cm of ice core length to avoid osmotic stress, which leads to organism loss.
• After complete melt, take a 100ml subsample for protist count (fix with Lugol) and concentrate the remaining volume for meiofauna over a 20μm sieve (fix with buffered Formaldehyde).
• Core sections are melted at room temperature in filtered seawater in melt buckets or in ziplock bags (double-bagged).
• Add 100 ml per 1 cm of section, meaning 1000ml for the 0-10 cm section.
• Filtered seawater and cool temperatures will ensure that organisms will not burst. Concentrate the organisms on the 20 μm sieve (small metal sieve) and rinsed into a vial or jar (~20 ml or larger) using a spray bottle with filtered seawater.
• Preserve the sample with 37% formaldehyde diluted 10x with filtered seawater to 4%. Do not freeze!

Stable isotope cores (1):

• One core.
**Genomics cores (3):**

- Three cores pooled.
- Use ethanol washed corer and cutting board or very clean sampling area.

**14.5.3 Ice samples for iron, other trace elements and nutrients**

**Measured parameter:** Iron, other trace elements and nutrients

**Method responsible:** Murat Van Ardelan (NTNU, murat.v.ardelan@ntnu.no)

**Parameter definition:** Quantification of total iron and other elements, (in nmol or µmol L⁻¹) and macronutrients (nitrate/phosphate/silicate in µmol L⁻¹). Measurements are taken from Chemistry and Nutrient cores mentioned in 14.5.2 as well as floating clumps of ice collected.

**Methodical description:**

→ **Special requirements/gear:**

- Plastic gloves
- Wood or plastic equipment (some wood broom handles and nylon nets are very useful).
- (Plastic coated metal poles are ok)
- !!!! avoid rusty metal equipment
- plastic bags and containers
- clean plastic syringes
- 125 mL nalgene LDPE bottles
- 125 ml plastic bottles for macronutrient samples
- syringe filter (0.2 µm) for macronutrient sample
- SPACE for Class-100 laminar flow clean-air chamber (2x2x1.5 m)
- Methodological description

→ **Chemicals used:**
Collection of ice:

- Ice samples should be collected from floating samples of ice by ‘fishing’ with wood or plastic equipment (some wood broom handles and nylon nets are very useful). Plastic coated metal poles are ok, but not rusty metal equipment.
- Plastic gloves should be worn when handling ice or collecting samples.
- It is important to randomize sample collection eg collecting every piece of ice in a suitable size range (approximately 1-4 kg), or collecting every \( x^{th} \) sample, to avoid bias towards ice that is not an ‘average’ color.
- After collecting an ice sample keep it in a plastic bag or container until ready to process the sample. Large samples can be smashed to collect a smaller piece for melting.
- Record the location where ice was collected and give every sample a unique label.

Melting the ice:

- Place each ice sample in a clean, airtight plastic bag.
- Allow the bagged ice to melt. After 5-10% of the ice has melted, swill the melt water around the bag and discard. Then re-seal the bag and continue the melting. We do this to remove films that may be present on the ice surface.
- Repeat this melt-swill-discard process 5 times. If the bag tears, move the ice to a new bag.
- After throwing away the meltwater 5 times, allow the water to accumulate until about 200 mL is in the bag.

Sampling for iron and nutrients:

- Using either a clean syringe (provided) or by gentle tipping, fill a pre-cleaned 125 mL nalgene LDPE bottle (provided) for analysis of iron.
• Ideally, up to the 125 mL mark, but anything from 25-125 mL is ok. Seal the LDPE bottles quickly and then store in clean plastic bags in a cool, dark place. Label the bottles with the sample name.
• The remaining water can be filtered (0.2 µm) and preserved for nutrient analysis. If keeping nutrient samples for later analysis, preserve by freezing and/or addition of Hg. Label the bottles with the sample name.

14.6 MELT POND SAMPLES
If melt ponds exist, one melt pond should be selected for a full set of samples similar to ice cores, see Table 14.5-1).

14.7 ON-ICE CTD
Use hand-held on-ice CTD (e.g. UiT CTD) for one deployment from the ice through auger hole down to 90m water depth. If equipped with light sensor, than make sure, that you are not in the ship shadow.

14.8 ON-ICE WATER SAMPLER
If a small hand-held water sampler is available, one water sample depth should be selected (ca 0.5m under ice bottom) for full set of variables/parameters (similar to ice cores, see Table 14.5.2-1). Some hand-held water samplers can be deployed through a 14cm ice corer hole, otherwise use larger ice auger.
15 Geological sampling

**Method responsible:** Tine Rasmussen (UiT, tine.rasmussen@uit.no), Matthias Forwick (UiT, matthias.forwick@uit.no), Elisabeth Alve (UiO, Elisabeth.alve@geo.uio.no), Katrine Husum (NPI, katrine.husum@npolar.no), Ulysses Ninnemann (UiB, ulysses.ninnemann@uib.no)

**Tab. 15-1** Overview of geological sampling effort and institutions responsible. TR: Tine Rasmussen. MF: Matthias Forwick. BF: Benthic foraminfera. PF: Planktic foraminifera. BF $d^{18}O$, $d^{13}C$: stable isotope analysis ($d^{18}O$, $d^{13}C$) measured on benthic foraminifera. PF $d^{18}O$, $d^{13}C$: stable isotope analysis ($d^{18}O$, $d^{13}C$) measured on planktic foraminifera.

<table>
<thead>
<tr>
<th>Speciality &amp; analysis</th>
<th>UIT (TR)</th>
<th>UiO</th>
<th>UIB</th>
<th>NPI</th>
<th>UIT (MF)</th>
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<tbody>
<tr>
<td>AMS 14C</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
<td>X</td>
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<tr>
<td>210Pb</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
<td>X</td>
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<tr>
<td>BF 0 - 2 ka</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>PF 0 - 2 ka</td>
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<td>BF &gt; 2ka</td>
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<td>X</td>
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<td></td>
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<td>BF $d^{18}O$, $d^{13}C$ 0 - 2 ka</td>
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<tr>
<td>BF $d^{18}O$, $d^{13}C$ &gt; 2ka</td>
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<td>HBIs (eg IP25)</td>
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<tr>
<td>Diatoms</td>
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<td>Trace elements/Ca (ikke Mg/Ca)</td>
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<td>Physical properties</td>
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<td>XRF core scanning</td>
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</tr>
</tbody>
</table>
15.1 SEDIMENT SAMPLING

15.1.1 Short sediment cores (2 ka) – Multicorer

The multi corer automatically gives four multicorer tubes:

- At least two casts will be carried out with multicorer in order to obtain sediment cores for the analyses and proxies (Table 15.1.1-1).
- Three casts will be carried out at stations where it is also planned to analyze porewater and sediment geochemistry (ChAOS).
- Samples from sediment cores that are sampled onboard will be kept frozen. Other cores will be stored cold (0-5°C).
- Further processing and analysis will be carried out onshore at NPI, UiB, UiO, UiT and UniRES (see table below).

<table>
<thead>
<tr>
<th>Proxy – parameters</th>
<th>Number of multicores</th>
<th>Partner</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paleo benthic foraminifera, planktic foraminifera</td>
<td>1</td>
<td>UiO, UiT</td>
</tr>
<tr>
<td>Living benthic foraminifera</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Stable isotopes</td>
<td>1</td>
<td>UiB</td>
</tr>
<tr>
<td>IP25, diatoms, aDNA, dinocysts</td>
<td>1</td>
<td>NPI, UniRES</td>
</tr>
<tr>
<td>Sedimentology</td>
<td>1</td>
<td>UiT</td>
</tr>
<tr>
<td>Reference</td>
<td>1</td>
<td>UiO/UiT/UiB/NPI</td>
</tr>
<tr>
<td>Porewater, sediment geochemistry</td>
<td>2</td>
<td>ChAOS, UiT</td>
</tr>
</tbody>
</table>
**Multicorer - Sampling protocol:**

- Remove water from multicorer tube.
- Prior to sub-sampling take a photograph of surface and one of the sediment core.
- Determine texture (clay, silt/sand/rocks) and colour using Munsell colour chart. Make notes of any disturbances, presence of benthic biota.
- Sub-sample in 0.5 cm steps down core using metal plates.
- If the sediments are very fluid spoon may also be applied.

15.1.1.1 *Living (stained) and fossil benthic foraminifera*

**Measured parameter**: Living and fossil benthic foraminifera

**Method Responsible**: Elisabeth Alve (UiO, Elisabeth.alve@geo.uio.no)

**Description of parameter/Aim**: 

**Methodological description**:

- **Special requirements/gear**: Multi corer (105 mm diameter)
- **Sampling depths**: Station bottom depth
- **Chemicals used**: Ethanol (96 %), Rose Bengal.

**Method**:

- Multi corer (105 mm diameter), which allows for the recovery of intact sediment-water interface.
- Stand with a piston.
- Shorter section of the core tube graduated with 1 cm mark.
- Slicing plates.
- Siphon.
- Plastic pipettes.
- 180 ml Joni plastic containers.
- Zip log bags.
• Ethanol (96 %).
• Rose Bengal.
• Notebook.
• Camera.
• Permanent marker pen.

Short multicores (~ 2 kyrs):

• Note down weather condition (if working outside) and sample process time.
• Place the core on a stand with a piston that fits the plastic core tube (Figure 15.1.1.1-1).
• Remove excess water above the sediment surface by using a siphon before pushing the core liner down on the piston in order to avoid sediment in suspension. Remove residual surface water using plastic pipettes.
• Prior to sub-sampling, take a photograph of the surface with label.
• Note down colour, texture, biota and possible disturbances, e.g. bioturbation, during slicing.
• If the sediment surface is irregular, define the sub-sample as the midpoint between the highest and the lowest point (measure the distance between these) (Schönfeld et al. 2012).
• Slice the sediment core using a slicing plate and a graduated, shorter section of the core tube (Figure 15.1.1.1-1).
• It is important to move the plate and sample horizontally when slicing to avoid dragging up sediment from the lower layer.
Sampling for palaeoceanographic and fossil benthic foraminifera analyses:

- Sub-sample one sediment core in 1 cm intervals down to 40 cm.
- Due to high water content in the sediment surface, transfer samples down to 5 cm to 180 ml Joni plastic containers, and in zip log bags from 5-40 cm. Store the zip log bags in one large zip log bag to separate samples from the different stations.
- Store the sub-samples frozen (-20 °C).
- It is important to avoid adding too much pressure on the lid when closing the plastic containers, as the containers might break. Instead, push the lid down in one place and drag your finger around until you hear a click.

Fig. 15.1.1.1-1 Schematic representation of a stand with a piston for sediment slicing (Licensed Material: Murray 2006, Cambridge University Press).
Sampling for living (stained) benthic foraminifera:

- Sub-sample three sediment cores in 1 cm intervals;
- One core down to 10 cm. pH may be measured in the sub-samples.
- Two cores down to 4 cm.
- Transfer the sub-samples to 180 ml Joni plastic containers and add at least an equal volume of rose Bengal (2 g/L) stained 80 % ethanol.
- Mix the samples thoroughly (but gently) until homogenized - examine the bottom of the container to see if any sediment is still clumping together (Figure 15.1.1.1-2).
- It is important to avoid adding too much pressure on the lid when closing the plastic containers, as the containers might break. Instead, push the lid down in one place and drag your finger around until you hear a click.

![Well-mixed sample (left) and not sufficiently mixed sample (right).](image)

**Fig. 15.1.1.1-1** Well-mixed sample (left) and not sufficiently mixed sample (right).

Literature

15.1.1.2 Stable isotopes

- One multicorer tube will be sampled from top of the core to the bottom for every 0.5 cm.
- Samples will be kept in plastic zip log bags and frozen (< 0° C).

15.1.1.3 Diatoms

- One multicorer tube will be sampled from top of the core to the bottom for every 0.5 cm.
- Samples will be kept in plastic zip log bags and frozen (< 0° C).

15.1.1.4 IP25 and Dinocyst sampling protocol

**Measured parameter:** Sea ice biomarkers (IP25) and dinocysts

**Method responsible:** Katrine Husum (NPI, katrine.husum@npolar.no), Stijn de Schepper (UNIResearch, stde@norceresearch)

**Description of parameter/Aim:** Sea ice reconstruction

**Methodical description:**

→ Special requirements/gear:

→ Sampling depth:

→ Chemicals used:

**Material:**

- Spatulas
- Plastic bags
- Marker
Method:

1) Amount: ca. 1/3 of opened core half.

2) Which interval:
   - Multicore and Gravity core: 1 cm slice samples in 1 cm intervals, starting at the top.
   - IP25 and Dinocyst samples were taken from multicores at stations NPAL04, NPAL05, NPAL07, NPAL08, NPAL12, NPAL14, NPAL15, NPAL17 and from gravity cores at stations NPAL04, NPAL08 and NPAL15.
   - No Calypso core was sampled on the cruise.
   - Diatoms were not sampled on the cruise. This is planned to be done in the same manner as IP25 and Dinocysts at a later stage (on archive half of the same cores).

3) Pre-sampling
   - Cores on deck are brought to the lab.
   - Cores are split. One half (work) is taken away for aDNA sampling. Another half is logged (archive). After aDNA sampling, the remaining material is sampled for IP25 and Dinocysts (multicores only).

→ Sample storage:
   - Samples are stored in a cooling room at approx. 4°C.
   - Transport:
     Bergen (NORCE): Dinocyst
     Tromsø (NPI): IP25
     Samples should be transported to Bergen/Tromsø in a cooling container.
15.1.1.5 Sedimentology

- One multicorer tube will be sampled from top of the core to the bottom for every 0.5 cm.
- Samples will be kept in plastic zip log bags.

15.1.1.6 14C-Dating sampling protocol

**Measured parameter:** 14C radiocarbon dating (of calcareous microfossils)

**Method Responsible:** Katrine Husum (NPI, katrine.husum@npolar.no), Ulysses Ninnemann (UiB, ulysses.ninnemann@uib.no), Elisabeth Alve (UiO, Elisabeth.alve@geo.uio.no), Matthias Forwick (UiT, matthias.forwick@uit.no), Tine Rasmussen (UiT, tine.rasmussen@uit.no)

**Description of parameter/Aim:** 14C radiocarbon dating (of calcareous microfossils). Dating the sediment

**Methodological description:**

- Special requirements/gear:
- Sampling depths:
- Chemicals used:

**Material**

- Spatulas
- Plastic bags
- Marker

**Method:**

1) Amount: ca. 1/3 of opened core half.
2) Which interval:
- Multicore: 2 samples, 1 cm slices ca. 5 cm from the bottom and 5 cm from the top (or where it seems reasonable). Samples were taken from the archive half of multicores sampled for IP25 and Dinocysts.
- Calypso/ gravity core: 3 samples per section. 1 cm slices ca. 5 cm from the bottom, in the middle and 5 cm from the top (or where it seems reasonable). Samples were taken from the work half after aDNA-sampling, or pore water sampling (ChAOS)
- Dating samples were taken from multicores at stations NPAL04, NPAL05, NPAL07, NPAL08, NPAL12, NPAL14, NPAL15, NPAL17 and from gravity cores at stations NPAL04, NPAL08, NPAL14 and NPAL15.

3) Pre-sampling:
- Cores on deck are brought to the lab.
- Cores are split. One half (work) is taken away for aDNA sampling or pore water sampling. Another half is logged (archive). After, the work sections are sampled for dating (GC only). For multicores, dating samples are taken from the archive half.

→ Sample storage/Transport:
- Samples are stored in a freezer at approx. -20 - -30 oC.
- Transport: Samples should be transported to Tromsø on dry ice or in a freezing container to ensure that the samples.

15.1.1.7 Sediment geochemistry (ChAOS project)
- Sediment samples will be taken from one multicorer tube.
- Sediments will be sliced for every 0.5 cm resolution down to 20 cm, and in 2 cm resolution below 20 cm sediment depth.
- Samples will be kept in standard zip lock bags.
• It is critical that these samples are frozen as soon as possible after sampling (ideally at -80°C, otherwise at -20°C) and are directly transferred into a freezer back at shore.
• Data to be generated onshore at University of Leeds, UK: TOC, TN, TS, C/N ratios, Fe speciation.

15.1.1.8 Ancient DNA (aDNA) sampling protocol

**Measured parameter:** DNA (ancient DNA from bulk sediment samples)

**Method Responsible:** Stijn de Schepper (UNIResearch, stde@norceresearch), Katrine Husum (NPI, katrine.husum@npolar.no)

**Description of parameter/Aim:** DNA (ancient DNA from bulk sediment samples). Purpose: Sea ice reconstruction.

**Methodological description:**

→ **Special requirements/gear:**

→ **Sampling depths:**

→ **Chemicals used:**

**Material:**

- single-use sterile 12ml polypropylene syringes (n= depends on sampling strategy)
- knife
- scissors
- disposable gloves (good to have both large and medium)
- single use large plastic spoons
- single use small plastic spoons
- -OR- metal spatula, wiped clean and flame-sterilised between samples
- sampling bags
- permanent marker for labelling
• sticky labels for sampling bags
• aluminum foil to work on a clean working desk and/or ethanol for cleaning workspace
• pre-cut “skumplast/oasis” (rubber foam) to fill in holes in sediment core
• Bunsen burner or lighter for flame-sterilizing knife

Method:

• Amount: At least 10cc per, more if available and if sediment is “soupy”.
• We need minimum 3 to 5 gram dry sediment for our DNA analyses.

1) Which interval:
   • Multicore: 2 samples at 0-1.5 cm, one to be stored in the freezer (-20- -30oC) and one to be stored in the fridge (ca. 4 oC).
   • All samples are taken with syringes with 1.5 cm diameter.
   • Multicores from sites where gravity cores also are taken are sampled downcore at same intervals as the gravity core to ensure overlap between the cores.
   • Calypso/gravity core: Sampled downcore at 1.5 cm slices in intervals reflecting ca. every 200-250 years.
   • Multicores were sampled at the top at every station. At station NPAL04, NPAL08, NPAL15 and NPAL17 one multicore half was sampled downcore.
   • Gravity cores from station NPAL04, NPAL08 and NPAL15 were sampled downcore.
   • One piston core (KH18-10-15-PC03) was sampled. Samples were taken at the top and bottom of each section (except for section 1, where we only took bottom samples). The length of the material was measured and represents the approx. sampling interval.

2) Pre-sampling:
   • Cores on deck are brought to the lab.
   • Cores are split using clean tools.
   • Split halves are cleaned by scraping off top sediment perpendicular to the main core axis, to avoid contamination from opening the core.
• If needed for cleaning the core/core liner: use fresh water or sterile, filtered (<0.2 µm) sea water.

3) Preparation:
On as clean workspace as possible (use aluminum foil as cover):

• Wear *clean* disposable gloves
• Label sample bags
• Label syringes with permanent marker on the tube
• Use unique labels that include cruise, station, site, core type and depth (cm):
  • e.g. “SIMEP–XX–STATION–CORE–xx” or similar
• Keep a log of all sediment samples taken
• Remove top from syringe using a sterilized knife
• Pull up (not out!) inner part of syringe and put in a plastic bag.

4) Core sampling:
• Sampling control. Place an Eppendorf tube open, on the bench, close to the core and your sampling spot. Give the sampling control an ID and make sure to track which sample(s) while the tube was open.
• Press syringe into sediment. Avoid to reach the core liner – i.e. leave some sediment – to prevent contamination from the core liner
• Pull syringe out of sediment
• When syringes do not work optimally (soupy sediments) use disposable spoons or metal spatula, wiped clean and flame-sterilised between samples
• Fill hole in sediment core with “skumplast”
• Close the sampling control (i.e. keep open during process of 1-3 samples (or what seems reasonable), and freeze the sampling controls in a tube/box (the same way you treat the syringes).

→ Sample storage/Transport:
• Put complete syringe with sediment into labeled sampling bag. Put syringe with sample into the -20°C freezer asap
• Transport: Samples should be transported to Bergen on dry ice. At least 5-6 kg dry ice will be necessary to keep samples frozen during transport.

15.1.2 Long sediment cores (14ka) – gravity/calypso corer

The long cores are cut for every m (GC) and every 1.5 m (PC) and carefully labelled. Those sediment cores that are sampled for pore water and/or for aDNA are opened and described onboard. Further sampling onboard is problematic due to carbonate dissolution which most probably will occur when the sediments are oxygenated. All cores will be logged with regard to magnetic susceptibility, color and photographed.

15.1.2.1 Laboratory analysis (opening cores, MSCL, lithostratigraphy)

Opening cores

Cores that will be opened will be extensively sampled onboard due to the potential loss of carbonate fraction after opening. All samples will be frozen immediately. Depending upon time and capacity we will try to process all cores completely, but in the case of backlog some (e.g. longer PC) cores may be stored for post-cruise processing.

Multi Sensor Core Logger (MSCL)

The magnetic susceptibility of the sediments will be measured on the surfaces of opened cores using a point sensor attached to a GEOTEK Multi Sensor Core Logger. Simultaneously, the colour properties will be measured with a spectrophotometer. Measurements will be made for every cm onboard and with a higher resolution onshore.
Lithological description
Lithostratigraphy will be established from visual descriptions of the surfaces of opened cores. Parameters of particular interest include lithological variations, colour changes, structures, signs of biological activity (bioturbation, occurrence of whole shells and shell fragments, as well as microorganisms).

15.2 GEOLOGICAL WATER SAMPLING

15.2.1 Geology proxy development of stable isotope analysis

Measured parameter: H, O, C, ³¹³C of DIC

Method responsible: Ulysses Ninnemann (UiB, ulysses.ninnemann@uib.no), Pål Tore Mørkved (UiB, pal.morkved@uib.no)

Description of parameter/Aim:

Methodological description:

→ Special requirements/gear:

→ Sampling depths: Upper 150m: 150, 100, 50, 25, 10, 5, 0m (adjust depending on local mixed layer and near surface stratification features). Maximum 12 bottles

→ Chemicals used: Mercury(II)chloride

CTD water sampling:

- 12 x 8L Niskin bottles (bottle #1 is from the deepest depth)
- Sensors: Chl & CDOM fluorescence, O2, transmissometer 660 nm, PAR, sPAR
- One cast whole water column for biology & chemistry
Sampling from ship CTD:

- Stable isotopes 60 ml serum vials with septa and crimpcaps

Sampling depths stable isotopes (H, O, C):

- Upper 150m: 150, 100, 50, 25, 10, 5, 0m (adjust depending on local mixed layer and near surface stratification features). Maximum 12 bottles
- Lower resolution sampling to (and including as close as possible and within 10m of) seafloor to define any major water masses.
- Stable Isotopes (\(\delta^{18}O - \delta \text{_D}\))

Collecting the sample:

- Sample evaporation and/or moisture condensation in the sample bottle must be avoided at all costs!
- Sampling from the Niskin bottles are done into 60 ml serum vials with butyl rubber septa and crimpcaps.

1. Rinse the vial and septum with sample three (3) times. This removes any water than may have condensed inside the bottle/cap.
2. Fill the vial and cap with seawater from the Niskin bottle (it is practical to use a silicon tube (i.d. 7mm), similar to the DIC sampling.
3. Apply the septum to the vial without touching the inside. Do not crimp the crimp cap. Press the septum sideways to release overpressure so the cap does not pop off.
4. Turn the bottle upside down and check for a small air bubble. If the bubble is too small loosen the septum and tighten it again.
5. When all the \(\delta^{18}O\) samples have been collected from one CTD, dry the vials, crimp the caps (and seal with Parafilm following the instructions below / on the next page.)
6. Store the \(\delta^{18}O\) samples at room temperature or in a fridge.
Alternative for onboard measurements with a Picarro:

4b) Bring the bottles to the lab. For each sample, transfer 1.7 ml to 2 ml labelled GC vials with screwcaps and PTFA lined butyl rubber septa using a 1 ml pipette. GC vials are stored upside down in boxes with lids in the fridge.

5b) Serum vials and septa can then be emptied, washed with DI water and dried for reuse or capped and crimped for refrigerated storing of sample for backup or further onshore measurements.

Sampling for δ13C of DIC in seawater:

SAMPLING DEPTHS (ocean acidification studies) are as above.

Equipment: Glass bottles 60 ml with butyl rubber septa and crimp caps, silicon tube (i.d. 7mm at KH), Mercury(II)chloride (in saturated solution in DI water, >7.4 g/100 mL, 20 °C), gloves.

1. Use gloves and goggles when handling the saturated Mercury(II)chloride and the fixed samples.
2. The bottles should be numbered before sampling, please use this number as reference.
3. Rinse the vial and septum with sample three (3) times. This removes any water than may have condensed inside the bottle/cap. Use the silicone tube to transfer water from the Niskin bottles to the sampling bottles.
4. The tube should be placed in the bottom of the sampling bottle when filling. Make sure there are no bubbles inside the tube when filling. This might require reducing the flow from the Niskin bottle.
5. Overfill the sample bottle with one bottle volume. Make sure no bubbles are trapped inside the sample bottle. Put the lid on so excess water runs out (squeeze sideways and down). When you have filled all sample bottles transfer them back to the laboratory.
6. Dry all the bottles with paper towel. Paper towels go in a black or clear waste bag (normal waste).
7. Using gloves and goggles and working in the fume hood add 5 drops of saturated Mercury(II)chloride from the small drop-bottle or using syringe w/needle to each bottle. Put the septum back on and secure it with a Crimp cap. Make a permanent, separate working area for Hg work (e.g. a fume hood) to avoid spreading Hg spill to the rest of the lab. Use bench paper and change gloves often to avoid spreading the mercury.

8. Dry of the bottles with paper towel in case of spillage. This paper goes in the zip bag labelled hazardous waste.

9. Put the dry bottles in the cooling room (NO FREEZING!). In case any spill dry with paper towel and put in the yellow plastic bag. All used gloves go in the hazardous waste.

For onboard analyses on a Thermo Delta Ray instrument:

5b) One extra bottle is sampled per sample. The vial is filled as in 5).

6b) as 6).

7b) Transfer 1 ml sample, using a 1 ml syringe, to each of 3 exetainers (Labco,UK) pre-prepared with 5 drops of 99-100% phosphoric acid each. Exetainers w/acid must be flushed with synthetic air before adding the sample (usually done onshore). No Hg is needed.

8b) stir (rotate glass) gently with vial remaining near vertical to mix water and acid and measure after the water reaches constant temperature (room temperature).
16 Methods guest researchers and previous cruises

As many researchers only join 1-2 cruises and then their sampling will not be continued, we opened this chapter for protocols of guest researchers.

16.1 TOC, TN, TS, C/N RATIOS, FE SPECIATION

**Measured parameters:** TOC, TN, TS, C/N ratios, Fe speciation

**Method responsible:** Christian Maerz (University of Leeds, c.maerz@leeds.ac.uk), Matthias Forwick (UiT, matthias.forwick@uit.no), Katrine Husum (Katrine.husum@npolar.no), Ulysses Ninnemann (UiB, ulyssses.ninnemann@uib.no)

**Description of parameter/Aim:**

**Methodological description:**

- **Special requirements/gear:**
  - **Sampling depths:** Multicore (pore water) sampling: BW, 0.5, 1.5, 2.5, 4.5, 6.5, 8.5, 10.5, 12.5, 14.5, 16.5, 18.5, 20.5, 25.5, 30.5 etc.
  - **Chemicals used:** HCl

16.1.1 CTD sampling for nutrients and ICP analysis

**Material:**

- 12 MQW rinsed (5x) 500 mL or 1 L bottles
- 12 MQW rinsed 50 mL syringes
- 12 centrifuge tubes
- 12 60 mL Nalgene
- 12 syringe filters
- HCl (conc.)
Method:

1. Label 12 acid washed and MQW rinsed (5x) Nalgene bottles (0.5 to 1 L) for sampling from the CTD Niskin bottles.
2. When sampling from the Niskin, rinse 3 times while sampling before filling the bottle.
3. In the lab, take a dry, MQW rinsed 50 mL syringe and using the CTD water, rinse the outside of the syringe.
4. Next, take up 10 mL of CTD water in the syringe. Hold the syringe vertically and pull down on the plunger to rinse the inside of the syringe 3 times.
5. Fill the syringe with CTD water and add the syringe filter to the syringe tip. Push out 10 mL of CTD water to rinse the syringe filter.
6. Rinse the Nalgene bottle with water from syringe 3x and then fill the bottle to just below the shoulder. Close the bottle and parafilm the cap. These samples will be flash frozen at -80°C for several hours and then stored in -20°C for the remainder of the cruise.
7. Rinse a centrifuge tube with water from the syringe 3x and then fill the tube. Ten uL of HCl will be added to each tube and they should be stored at 4°C.

16.1.2 Multicore sampling

1) Pore waters:

- **Sample depths**: BW, 0.5, 1.5, 2.5, 4.5, 6.5, 8.5, 10.5, 12.5, 14.5, 16.5, 18.5, 20.5, 25.5, 30.5 etc.

- **Chemicals used**: HCl
Material:

- Supplies needed
- Rhizons
- Rhizon spacers
- Syringes labelled with appropriate core depths
- Centrifuge tubes (15 mL)
- Nalgene bottles (1-60 mL and 15 mL for all sediment depths)
- Isotope vials
- HCl (conc.)

Method:

1. Core tubes should be drilled (0.4 cm) every 1 cm. The holes should be covered with clear office tape to avoid leaking.
2. Once cores are brought on deck, they should be cleaned and measured. The sediment core should then be transferred to the sink in the wet lab and secured with a bungee cord.
3. Using a pipette tip, a hole above the sediment water interface should be punctured and the rhizon inserted and attached with a syringe and a spacer. Once the syringe is filled, transfer water into the vials and tubes (and repeat) until a 60 mL Nalgene is filled for nutrients, a 15 mL centrifuge tube is filled for ICP work, a 15 mL centrifuge tube is filled for IC work, and 1 mL is transferred to the vial for isotopes. After these are full, remove the syringe and drain the remaining bottom water.
4. While bottom water is being removed, holes should be punctured and rhizons inserted starting from the bottom sample to 4.5 cm. Syringes can then be added. The 0.5 cm horizon will be set at the first sample below the sediment water interface. Once all overlying water has been removed, the holes at 2.5, 1.5 and 0.5 cm depth were opened, and rhizons were inserted very quickly to avoid the loss of pore water from the very water-rich uppermost sediment horizons. Rhizons will be left in the core tubes for up to ~2 hours, depending
on the efficiency of pore water extraction (very fast in the top layers, much slower in deeper, clay-rich layers).

5. Once all rhizons have finished extracting water, the water should be divided as follows:

<table>
<thead>
<tr>
<th>Sample type</th>
<th>Min volume</th>
<th>Max volume</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICP</td>
<td>3 mL</td>
<td>15 mL</td>
<td>10 μL HCl in centrifuge tube, stored at 4°C</td>
</tr>
<tr>
<td>Nutrients</td>
<td>11 mL (6 for dilution)</td>
<td>12-13 mL</td>
<td>Nalgene, capped tightly and parafilmed, flash frozen at -80°C standing up before storage at -20°C</td>
</tr>
<tr>
<td>Isotopes</td>
<td>0.5 mL</td>
<td>0.8-1 mL NO MORE</td>
<td>Transferred by syringe to vial for dry lab for analysis</td>
</tr>
</tbody>
</table>

**IMPORTANT:**

If less than 3 mL —> all sample to ICP

If 3-8 mL —> 1 mL to isotopes (air); remainder to ICP

If 8-15 mL —> 3 mL to ICP; remainder to Nut

If 16 mL —> 3 mL to ICP, 12 mL to Nut; 1 mL to isotopes (air)

If >16 mL —> 12 mL to Nut; 1 mL to isotopes (air); 1 mL to isotopes (He); remainder to ICP

2) **Sediments:**

Material:

- Whirl pak bags
- Sampling tools (plastic plates and rings)
- pH meter (calibrated)
- pH probe
- DI bottle and Kim Wipes
Method:

1. Sediment samples for both multi- and gravity cores should be sampled and frozen soon after cores are opened to avoid oxidation of sulfides and Fe in pore waters.

2. The core should be transferred to a core extruder on deck. Remove most overlying water using a silicon tube. The pH meter should be calibrated prior to sampling. Clean pH probe with MQW water. Be sure to remove any salt from probe. Measure pH within the overlying water. The remaining water should be removed with a disposable pipette.

3. Prior to extruding core, measure pH on the surface 3 times and record by hand in notebook. Clean pH probe thoroughly with MQW water.

4. Extrude core 1 cm by turning the crank two full turns, slice and transfer to labelled bag. Sample entire core at 1 cm intervals. Samples should be stored at -20°C.

16.1.3 Gravity core sampling

1) Pore waters:

Material:

- Drill and 0.4 cm drill bits
- Rhizons
- Rhizon spacers
- Syringes labelled with appropriate core depths
- Centrifuge tubes (15 mL)
- Nalgene bottles (1-60 mL and 15 mL for all sediment depths)
- Isotope vials
- HCl (conc.)
Method:

1. Gravity core sections should be transferred to wet lab counters and secured using bungee cords.
2. Holes should be drilled every 15 to 30 cm (based on section length). A rhizon should be inserted into the hole and a syringe should be attached and a spacer added. Rhizons should be left for 2-3 hours.
3. Pore waters should be divided and treated in the same way as the multicore pore waters above.

2) Sediments:

Material:
- Sample bags
- Sampling tools (spatulas)

Method:

1. Sediment samples for both multi- and gravity cores should be sampled and frozen soon after cores are opened to avoid oxidation of sulfides and Fe in pore waters.
2. Split core should be sampled at 1 cm intervals (a third to a half of the working half).
3. Samples should be immediately frozen at -20°C.
The Nansen Legacy in numbers

6 years
The Nansen Legacy is a six-year project, running from 2018 to 2023.

1,400,000 km² of sea
The Nansen Legacy investigates the physical and biological environment of the northern Barents Sea and adjacent Arctic Ocean.

>10 fields
The Nansen Legacy includes scientists from the fields of biology, chemistry, climate research, ecosystem modelling, ecotoxicology, geology, ice physics, meteorology, observational technology, and physical oceanography.

>350 days at sea
The Nansen Legacy will conduct 15 scientific cruises and spend more than 350 days in the northern Barents Sea and adjacent Arctic Ocean between 2018 and 2022. Most of these cruises are conducted on the new Norwegian research icebreaker RV Kronprins Haakon.

250 people
There are about 210 researchers working with the Nansen Legacy, of which 50 are early career scientists. In addition, 40 persons are involved as technicians, project coordinators, communication advisers and board members.

10 institutions
The Nansen Legacy unites the complimentary scientific expertise of ten Norwegian institutions dedicated to Arctic research.

50/50 financing
The Nansen Legacy has a total budget of 740 million NOK. Half the budget comes from the consortiums’ own funding, while the other half is provided by the Research Council of Norway and the Ministry of Education and Research.