

Video Article

VacuSIP, an Improved InEx Method for *In Situ* Measurement of Particulate and Dissolved Compounds Processed by Active Suspension Feeders

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Abstract

Benthic suspension feeders play essential roles in the functioning of marine ecosystems. By filtering large volumes of water, removing plankton and detritus, and excreting particulate and dissolved compounds, they serve as important agents for benthic-pelagic coupling. Accurately measuring the compounds removed and excreted by suspension feeders (such as sponges, ascidians, polychaetes, bivalves) is crucial for the study of their physiology, metabolism, and feeding ecology, and is fundamental to determine the ecological relevance of the nutrient fluxes mediated by these organisms. However, the assessment of the rate by which suspension feeders process particulate and dissolved compounds in nature is restricted by the limitations of the currently available methodologies. Our goal was to develop a simple, reliable, and non-intrusive method that would allow clean and controlled water sampling from a specific point, such as the excurrent aperture of benthic suspension feeders, *in situ*. Our method allows simultaneous sampling of inhaled and exhaled water of the studied organism by using minute tubes installed on a custom-built manipulator device and carefully positioned inside the exhalant orifice of the sampled organism. Piercing a septum on the collecting vessel with a syringe needle attached to the distal end of each tube allows the external pressure to slowly force the sampled water into the vessel through the sampling tube. The slow and controlled sampling rate allows integrating the inherent patchiness in the water while ensuring contamination free sampling. We provide recommendations for the most suitable filtering devices, collection vessel, and storing procedures for the analyses of different particulate and dissolved compounds. The VacuSIP system offers a reliable method for the quantification of undisturbed suspension feeder metabolism in natural conditions that is cheap and easy to learn and apply to assess the physiology and functional role of filter feeders in different ecosystems.

Video Link

The video component of this article can be found at <http://www.jove.com/video/54221/>

Introduction

Benthic suspension feeders play essential roles in the functioning of marine ecosystems¹. By filtering large volumes of water^{2,3}, they remove and excrete particulate (plankton and detritus) and dissolved compounds¹ (and references therein) and are an important agent of benthic-pelagic coupling^{4,5} and nutrient cycling^{6,7}. Accurately measuring the particulate and dissolved compounds removed and excreted by benthic suspension feeders (such as sponges, ascidians, polychaetes, and bivalves) is fundamental to understand their physiology, metabolism, and feeding ecology. Together with pumping rate measurements, it also enables a quantification of the nutrient fluxes mediated by these organisms and their ecological impact on water quality as well as on ecosystem scale processes.

Choosing the appropriate method of measuring removal and production rates of particulate and dissolved compounds by suspension filter feeders is crucial for obtaining reliable data concerning their feeding activity⁸. As pointed out by Riisgård and others, inappropriate methodologies bias results, distort experimental conditions, produce incorrect estimations of ingestion and excretion of certain substances, and can lead to erroneous quantification of the nutrient fluxes processed by these organisms.

The two most frequently employed methods to measure particulate and dissolved nutrient fluxes in filter feeders involve either incubation (indirect techniques) or simultaneous collection of ambient and exhaled water (direct techniques). Incubation techniques are based on measuring the rate of change in the concentration of particulate and dissolved nutrients in the incubated water, and estimating rates of production or removal compared to adequate controls⁸. However, enclosing an organism in an incubation chamber can alter its feeding and pumping behavior due to changes in the natural flow regime, due to a decline in oxygen and/or in food concentration, or due to accumulation of excretion compounds in the incubation water^{7,9} (and references therein). In addition to the effects of confinement and modified water supply, a major bias of incubation techniques stems from re-filtration effects (see for example¹⁰). Although some of these methodological problems have been overcome by using the right volume and shape of the incubation vessel¹¹ or with the introduction of a recirculating bell-jar system *in situ*¹², this

technique often underestimates removal and production rates. Quantifying the metabolism of dissolved compounds such as dissolved organic nitrogen (DON) and carbon (DOC) or inorganic nutrients, has proven to be especially prone to biases caused by incubation techniques¹³.

In the late 60s and early 70s, Henry Reiswig^{9,14,15} pioneered the application of direct techniques to quantify particle removal by giant Caribbean sponges, by separately sampling the water inhaled and exhaled by the organisms *in situ*. Due to the difficulty to apply Reiswig's technique on smaller suspension feeders and in more challenging underwater conditions, the bulk of research in this field was restricted to the laboratory (*in vitro*) employing mostly indirect incubation techniques¹⁶. Yahel and colleagues refitted Reiswig's direct *in situ* technique to work in smaller-scale conditions. Their method, termed InEx¹⁶, is based on simultaneous underwater sampling of the water inhaled (In) and exhaled (Ex) by undisturbed organisms. The different concentration of a substance (e.g., bacteria) between a pair of samples (InEx) provides a measure of the retention (or production) of that substance by the animal. The InEx technique employs open-ended tubes and relies on the excurrent jet produced by the pumping activity of the studied organism to passively replace the ambient water in the collecting tube. While Yahel and colleagues have successfully applied this technique in the study of over 15 different suspension feeders taxa (e.g.,¹⁷), the method is constrained by the high level of practice and experience required, by the minuscule size of some excurrent orifices, and by sea conditions.

To overcome these obstacles, we developed an alternative technique based on controlled suction of the sampled water through minute tubes (external diameter < 1.6 mm). Our goal was to create a simple, reliable, and inexpensive device that would allow clean and controlled *in situ* water sampling from a very specific point, such as the excurrent orifice of benthic suspension feeders. To be effective, the method has to be non-intrusive so as not to affect the ambient flow regime or modify the behavior of the studied organisms. The device presented here is termed VacuSIP. It is a simplification of the SIP system developed by Yahel *et al.* (2007)¹⁸ for ROV-based point sampling in the deep sea. The VacuSIP is considerably cheaper than the original SIP and it has been adapted for SCUBA-based work. The system was designed according to principles presented and tested by Wright and Stephens (1978)¹⁹ and Møhlenberg and Riisgård (1978)²⁰ for laboratory settings.

Although the VacuSIP system was designed for *in situ* studies of the metabolism of benthic suspension feeders, it can also be used for laboratory studies and wherever a controlled and clean, point-source water sample is required. The system is especially useful when integration over prolonged periods (min-hours) or *in situ* filtrations are required. The VacuSIP has been used successfully at the Yahel lab since 2011, and has also been employed in two recent studies of nutrient fluxes mediated by Caribbean and Mediterranean sponge species²¹ (Morganti *et al.* submitted).

The use of specific samplers, the prolonged sampling duration, and the field conditions, in which VacuSIP is applied, entail some deviations from standard oceanographic protocols for collecting, filtering, and storing samples for sensitive analytes. To reduce the risk of contamination by the VacuSIP system or the risk of modification of the sampled water by bacterial activity after collection, we tested various *in situ* filtration and storage procedures. Different filtering devices, collection vessels, and storing procedures were examined in order to achieve the most suitable technique for the analysis of dissolved inorganic (PO_4^{3-} , NO_x^- , NH_4^+ , SiO_4) and organic (DOC + DON) compounds, and ultra-plankton (<10 μm) and particulate organic (POC + PON) sampling. To further reduce the risk of contamination, especially under field conditions, the number of handling steps was reduced to the bare minimum. The visual format in which the method is presented is oriented to facilitate reproducibility and to reduce the time required to efficiently apply the technique.

System overview

To sample *in situ* pumped water from suspension feeders with exhalant orifices as small as 2 mm, the pumping activity of each specimen is first visualized by releasing filtered fluorescein dyed seawater next to the inhalant orifice(s) and observing its flow from the excurrent aperture¹⁶ (see also Figure 2B in¹⁸). The water inhaled and exhaled by the study specimen (incurrent and excurrent) are then simultaneously sampled with the use of a pair of minute tubes installed on custom-built manipulator or on two of the "arms" of an upside-down flexible portable tripod (**Figure 1** and **Supplementary Video 1**). The water inhaled by the study organism is collected by carefully positioning the proximal end of one tube inside or near the inhalant aperture of the study organism. An identical tube is then positioned inside the excurrent orifice. This operation requires good care to avoid contact or disturbance of the animal, e.g., by sediment resuspension. To begin the sampling, a diver pierces a septum in the collecting vessel with a syringe needle attached to the distal end of each tube, allowing the external water pressure to force the sampled water into the vessel through the sampling tube. The suction is initiated by the vacuum previously created in the vials and by the pressure difference between the external water and the evacuated sample container.

To ensure a clean collection of exhaled water and to avoid accidental suction of ambient water¹⁶, the water sampling rate needs to be kept at a significantly lower rate (<10%) than the excurrent flow rate. The suction rate is controlled by the length of the tube and its internal diameter (ID). The small internal diameter also ensures a negligible dead volume (< 200 μl per meter of tubing). Sampling over prolonged periods (minutes to hours) makes it possible to integrate the inherent patchiness of most substances of interest. To ensure that samples are adequately preserved in prolonged underwater sampling sessions as well as for transportation to the lab, an *in situ* filtration is recommended for sensitive analytes. The selection of sampling vessels, filtration assembly, and tubing are dictated by the study organisms and the specific research question. The protocol described below assumes that a full metabolic profile is of interest (for an overview see **Figure 2**). However, the modular nature of the protocol allows for easy modification to accommodate simpler or even very different sampling schemes. For a full metabolic profile, the sampling protocol should include the following steps: (1) Flow visualization; (2) Sampling ultra-plankton feeding (plankton < 10 μm); (3) Sampling inorganic nutrients uptake and excretion (using in-line filters); (4) Sampling dissolved organic uptake and excretion (using in-line filters); (5) Particulate feeding and excretion (using in-line filters); (6) Repeat step 2 (ultra-plankton feeding as quality check); (7) Flow visualization.

When logistically feasible, it is recommended that the metabolic profile measurements are combined with pumping rate (e.g., the dye front speed method, in¹⁶) as well as with respiration measurements. These measurements are best taken at the beginning and end of the sampling session. For respiration measurement, underwater optodes or micro-electrodes are preferable.

Protocol

1. Preparatory Steps and Cleaning Procedures

1. Cleaning solution

1. Wear protective gear, a lab coat, and gloves at all times. Carry out these preparatory steps in a clean space free of dust and smoke.
2. Prepare a 5-10% hydrochloric acid (HCl) solution with fresh, high quality, double distilled water.
3. Prepare a 5% highly soluble basic mix of anionic and non-ionic surfactant solution (See Materials List) with fresh, high quality, double distilled water.
4. Store all solutions in clean, acid washed containers.

2. Preparatory steps and cleaning procedures (in the lab)

NOTE: If phosphorus compounds are not of interest, the HCl wash can be replaced by high quality phosphoric acid (H₃PO₄) wash (8% H₃PO₄ final concentration).

1. Wear protective gear, a lab coat, and gloves at all times.
2. Wash the sampling apparatus (excluding the in-line stainless steel Swinney filter holder) with ample amount of high purity water. Leave the apparatus soaking in 5-10% HCl solution overnight. Rinse the apparatus again with ample amount of high quality double distilled water.
3. Wash the in-line stainless steel Swinney filter holders with ample amount of high purity water. Leave the filter holders soaking in 5% highly soluble basic mix of anionic and nonionic surfactants solution overnight. Rinse them again with ample high quality double distilled water.
4. Dry all the sampling apparatus, wrap them in aluminum foil and keep in a clean box until use.

3. Suction rate control

1. Control the sampling rate by adjusting the length and internal diameter of the intake tubing according to the planned work depth and water temperature. Use the following equation (derived from the Hagen-Poiseuille equation used for fully developed laminar pipe flow) as a guide: $F = \frac{\Delta P \cdot \pi \cdot r^4}{8KLV}$ where F = flow rate (cm³ min⁻¹), ΔP = differential pressure (bar), r = inlet tubing internal radius (cm), K = 2.417 × 10⁻⁹ (sec⁻²), L = tube length (cm), V = water viscosity (g cm⁻¹ sec⁻¹). See **Table 1** for more details.
2. Keep sampling rate below 1% of the pumping rate of the studied animal.

NOTE: Using evacuated containers, sometimes with unknown vacuum poses additional complications. Therefore, a field test is highly recommended. At 10 m depth and ~22 °C seawater (40 PSU), a 50 cm inlet tubing with an internal diameter of 254 μm delivers an average suction rate of ~26 μl sec⁻¹ (1.56 ml min⁻¹).

4. Sampling vessels

1. For small volume samples (3-20 ml, e.g., ultra-plankton for flow cytometry) use pre-vacuumed sterile plastic tubes.
NOTE: Pre-vacuumed sterile plastic tubes are routinely employed for standard blood tests in humans; make sure to use the sterile tubes with no additives. These vacuumed sterile plastic tubes are best sampled with the use of sterile, single-use tube holder with off-center Luer. While this is the safest and most efficient sampling apparatus, it has a slightly larger dead volume compared to a simple needle.
2. For larger water samples such as nutrients and dissolved organics, use 40 or 60 ml glass vials that meet the Environmental Protection Agency (EPA) criteria for volatile organic analyses. These vials include a polypropylene cap with a PTFE-faced silicone septum.
3. For even larger volumes, use penicillin bottles with rubber stoppers or vacuum flasks.
4. Use high-density polyethylene vials (HDPE vials) for silica samples.
5. To increase sample volume and to reduce the risk of the stopper dislodging during the ascent, evacuate (vacuum) items 1.4.2-1.4.4 before the dive with a vacuum pump. Vacuum manually by using a hand vacuum pump or even by sucking the air with a syringe. However, for best results, a good vacuum pump is recommended. Standard lyophilizers provides high vacuum.
NOTE: Pay special attention when using large vacuumed flasks to ensure that the faster initial suction rate would not contaminate the exhaled water samples.

5. Vessel cleaning procedures

1. For dissolved organics and NH₄⁺ analysis, use new pre-cleaned EPA vials.
2. Rinse the vials (glass and HDPE) for analysis of other nutrients as follows:
 1. Rinse the vials (glass and HDPE) and the polypropylene caps with high quality double distilled water. Install a new silicon septum.
 2. Soak the vials (glass and HDPE) in 10% of HCl for at least 3 days and rinse with ample high quality double distilled water.
 3. Combust the glass vials at 450 °C for 4 hr and allow to cool in the furnace. Install the cap, and wrap in aluminum foil until use.

6. Filters

1. Use binder-free glass fiber filters for filtration of all dissolved organic samples (e.g., DOC, DON) and for the collection of particulate organics (e.g., POC, PON). Pack each glass filter in a separate aluminum foil envelope. Combust at 400 °C for 2 hr to volatilize organic residues and store in a clean and dry vessel until use.
2. Use either a binder-free glass fiber filters as above, or 0.2 μm polycarbonate membranes for sampling inorganic nutrients (e.g., PO₄³⁻, NO_x⁻, NH₄⁺). Clean the latter once installed in the filter holder as explained below (1.7.3).
3. Use 0.2 μm polycarbonate membranes filters for silica sampling. Clean them once installed in the filter holder as explained below (1.7.3).

7. Preparation of filtration assembly

1. Filter the highly soluble basic mix of anionic and non-ionic surfactant solution and the high quality double distilled water through a 0.2 μm filter before using them to clean the filtration assembly.
2. Filtration assembly for nutrients and dissolved organics other than silica:
 1. Place a combusted binder-free glass fiber filters inside the cleaned in-line stainless steel Swinney filter holder.

2. Use an acid-cleaned syringe to run 100 ml of 5% highly soluble basic mix of anionic and non-ionic surfactant solution and then 100 ml of high quality double distilled water through the entire assembly.
3. Filtration assembly for SiO₄:
 1. Place the polycarbonate filter inside the cleaned polycarbonate filter holder (PC filter holder).
 2. Use an acid-cleaned syringe to run 30 ml of 5% HCl and 30 ml of high quality double distilled water through the entire assembly.

8. System assembly

1. Assemble the system for underwater work using PEEK (Polyether Ether Ketone) tubing with an external diameter (OD) of 1.6 mm and an internal diameter (ID) of 254 µm or 177 µm.
2. Use a sharp knife or PEEK cutter to cut the tubes to the required length.
3. At its distal end (sample container side), fit each tube with a male Luer connector attached to a syringe needle. Make sure you follow the manufacturer's instruction and align the flat side of blue flangeless ferrule with the end of the tube before tightening the green nut.
4. Attach the PEEK tubing to the tripod "arms" or custom-built manipulator by using an insulating tape.
5. Attach a disposable syringe needle to the male Luer connector. Keep the needle with its protective cap to prevent injuries.
6. Clearly label the sampling gear and color code all inhaled and exhaled components (e.g., green=In, red=Ex).
7. Similarly color code the sampling vessels with sets of paired sampling vessels sequentially numbered.

2. Working Underwater

1. Work site preparation

1. Preliminary survey and selection of specimens

NOTE: Due to the complex nature of the underwater sampling protocol, devoting the necessary time for preparation will ensure an efficient sampling dive.

1. Survey the work site and make necessary preparations ahead of time.
2. Select and mark suitable target organisms that can be accessed relatively easily. Since not all organisms may necessarily be active at the time of the sampling dive, prepare more workstations than you expect to sample.

2. Installation of base supports

1. When working on leveled substrate:
 1. Mount the original quick release clip of the flexible tripod on 1 kg diving weights and simply position it next to the target animal.
2. When working on vertical walls:
 1. Mount base support plates for the VacuSIP system, hooks for accessory gear, and hangers for the collecting vessel carrying tray during the work site preparation stage (2.1.1).
 2. When flexible portable tripods are used, use bolts or two-component epoxy resin to fix 10x10 cm PVC plates next to each target animal. Each plate needs to have a hole to attach the quick release clips of the flexible portable tripod.
 3. Once the resin has cured and the base plates are solidly attached to the wall, screw in the quick release clips, serving as a firm attachment point for the flexible portable tripod for the VacuSIP system.

2. Installing the VacuSIP

1. Check whether the specimen is pumping by releasing filtered fluorescein dye next to the inhalant orifice and confirm that the dye is emerging through the exhalant orifice as described in Yahel *et al.* (2005)¹⁶.
2. Install the VacuSIP device and place the inhalant (IN) sampling tube within the inhalant orifice or just next to it (within ~5 mm). Make sure that the inhalant tube is not in the proximity of another exhalant orifice.
3. Carefully direct the exhalant (EX) sampling tube toward the osculum/exhalant siphon and very gently insert it in, until it is positioned 1-5 mm inside the osculum/exhalant siphon (see **Figure 1** and **Supplementary Video 1**). Take great care not to make contact with or otherwise disturb the sampled organism.
4. Before and during the sampling double-check the location of both tubes.
5. After the sampling check whether the specimen is still pumping as described above (2.2.1).

NOTE: Because the movement of one arm of the tripod when manipulating the other might occur, make sure to firstly place the inhalant sampling tube and secondly the exhalant tube, which requires more precise manipulation. Following this order, even if the manipulation of the exhalant sampling tube might cause the movement of the inhalant tube, it will not affect the sampling.

3. Modular underwater sampling procedure

NOTE: Pending on the research question, each of the experimental steps below can be performed as a stand-alone experiment. The full metabolic profile sampling protocol described below is a lengthy process, requiring up to 8 hours per specimen (for an overview see **Figure 2** and **Table 2**). As diving conditions and regulations differ between sampling sites, regions, and institutions, diving plans are not included in this protocol. Nevertheless, devote extreme care and meticulous planning to the diving plan. Pay special care to avoid saturation and yoyo dive profiles. When possible, it is advisable to conduct these experiments at shallow depth (< 10 m). Closed circuit rebreathers can be very handy for such prolonged sampling schemes.

1. Before actual sampling begins, make sure no visible traces of fluorescein residue remain and that suspended sediments have been settled or have wafted away.
2. **Ultra-plankton, (no filter is installed in this step!)**

1. Use the needle to pierce the IN (inhaled) and EX (exhaled) vacuumed sterile plastic tubes septa. Verify that water is dripping in at the planned rate and collect 2-6 ml water samples.
2. On retrieval, keep samples in a cold box on ice. In the lab, preserve with 1% paraformaldehyde + 0.05% EM grade glutaraldehyde (final concentration), or 0.2% EM grade glutaraldehyde. Freeze cryovials in liquid nitrogen and store at -80 °C until analysis.

3. Silicate sampling and storing

1. Install the pre-cleaned in-line stainless PC filter holder containing a 0.2 µm polycarbonate membrane between the needle and the Luer male connector at the distal end of the tube.
2. Pierce the septum cap of the pre-cleaned high-density polyethylene vials (HDPE vials) to start sampling. Verify that both samplers are dripping and collect 15 ml of water in each vial.
3. Keep the samples refrigerated (4 °C) until analysis. If analysis cannot begin within two weeks, store at -20 °C. For analysis, make sure that the samples are thawed at 50 °C for at least 50 min to dissolve silica gels.
NOTE: The membrane can be preserved for microscopy or DNA analysis as needed.

4. Dissolved inorganics (PO_4^{3-} , NO_x^- , NH_4^+)

1. Replace the PC filter assembly with a pre-cleaned in-line stainless steel filter holder that contains a pre-combusted glass filter.
2. Before sampling, ensure that at least 20 ml of seawater samples were passed through the entire filtration system by using EPA, HDPE vials or other vacuum vessel to start the suction. Measure the volume of the collected water before they are discarded.
3. Pierce the septum cap of the appropriate EPA glass vials to start sampling, verify that both samplers are dripping, and collect 25-30 ml for nitrate and phosphate analysis.
4. Switch to new EPA glass vials for ammonia analysis, verify that both samplers are dripping, and collect 20 ml in each vial.
5. Keep the samples in a cold box on ice and store at -20 °C until the analysis.
NOTE: If only the filtrate is of interest, disposable syringe filters can be used for steps 2.3.3 and 2.3.4.

5. Dissolved organics (DOC+DON) sampling and storing

NOTE: Keep the samples as upright as possible throughout handling so that sample water does not come in contact with the silicon septa.

1. Continue using the stainless steel filter assembly and collect 20 ml of seawater samples into new EPA glass vials, as described above.
2. Upon retrieval, keep the samples in a cold box on ice. In the lab, use a pre-combusted glass Pasteur pipette to fix the samples with orthophosphoric acid (add 5-6 drops of 25% trace metal grade acid into a 20 ml sample, final concentration 0.04%) or hydrochloric acid (add 2 drops of trace metal grade concentrated acid into a 20 ml sample, final concentration 0.1%) and keep refrigerated.
3. Keep the samples refrigerated (4 °C) until analysis. If samples are not analyzed within a week of collection, store at -20 °C until the analysis.

6. Particulate organic matter (POC, PON, POP)

1. Continue using the stainless steel filter assembly and filter at least 500 ml of seawater into an evacuated 250 ml vacuum flask. Replace flasks if necessary.
2. On retrieval, use an air-filled syringe to evacuate all remaining seawater from the filter holder, wrap it in aluminum foil, and store in a cold box on ice. In the lab, remove the filters from the filter holders and store at -20 °C until the analysis.

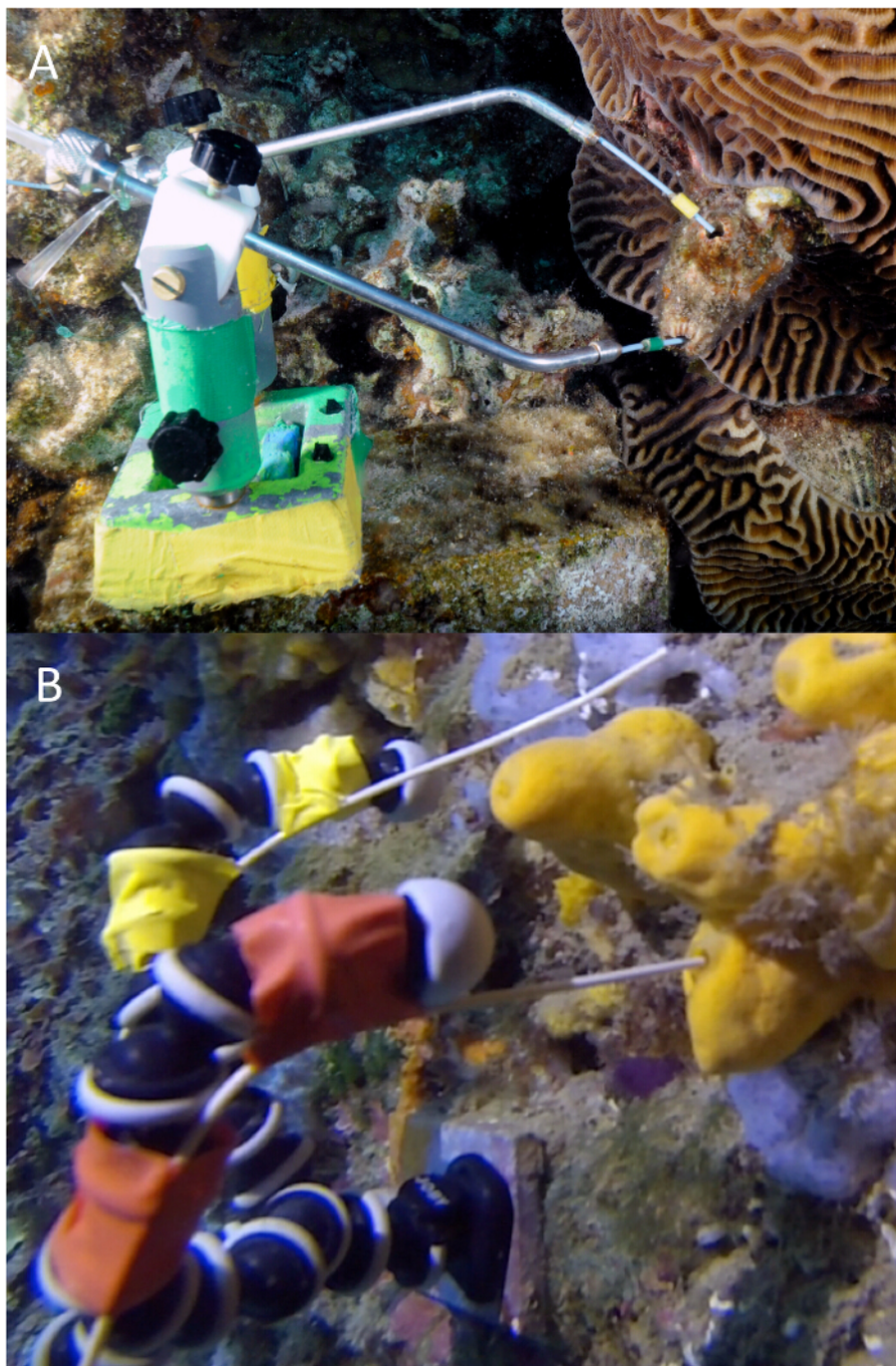


Figure 1. An example of correct installations of the VacuSIP: (A) sampling the ascidian *Polycarpa mytiligera* (Gulf of Aqaba, Red Sea) using a custom-built manipulator with the color code used green for inhaled and yellow for exhaled water samples (photo by Tom Shelizenger and Yuval Yacobi); (B) sampling the sponge *Agelas oroides* (NW Mediterranean Sea) with an osculum width of 3 mm, using the VacuSIP device. The color code used is yellow for inhaled and red for exhaled water samples. [Please click here to view a larger version of this figure.](#)

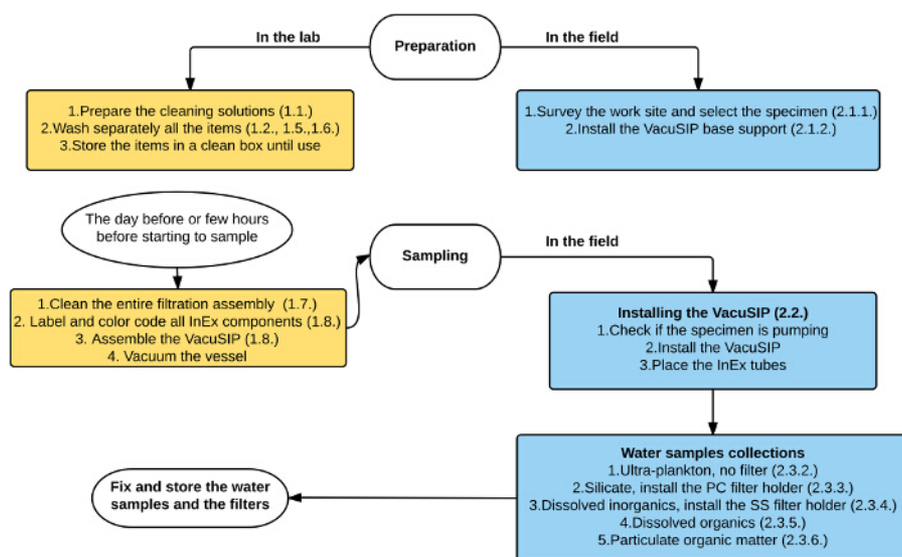


Figure 2. Overview of the VacuSIP technique described in the protocol section. The lab work is represented in yellow boxes, the fieldwork in blue boxes. [Please click here to view a larger version of this figure.](#)

Container	Flask Volume	Vacuum	Volume collected (mL)	Sampling rate (mL min ⁻¹)
Glass flask	250 ml	None	64.38 ± 33.06	1.13 ± 0.61
EPA glass vial	40 ml	1/2 volume	17.95 ± 2.81	0.59 ± 0.35
HDPE vial	40 ml	1/2 volume	7.73 ± 3.16	0.46 ± 0.27
Vacuumed sterile plastic tubes	9 ml	from manufacturer	8.78 ± 0.65	0.61 ± 0.32

Table 1. The overall average sampling rates (ml min⁻¹) obtained with different containers used for water collections and different vacuum levels: the flasks were not vacuumed (none); EPA glass vials and HDPE vials were vacuumed half of their volume (1/2 volume); sterile plastic tubes were already vacuumed by the manufacturer. Working at 5-8 m depth, water temperature of 18-22 °C, using PEEK tubes of 79 cm length and of 25 µm internal diameter.

Compound	Sampling vessel	Fixative	In-line filter assembly	Storage	Analytical method	Protocol
Ultra plankton	Pre-vacuum sterile plastic tube	Glut + Parafor	None	Liquid N (-80 °C)	Flow cytometry ²²	2.3.2
SiO ₄	HDPE vials		PC filter holder + PC membrane	-20 °C	Molivate colorimetry ²³	2.3.3
PO ₄ ³⁻ , NO ₂ ⁻ , NO ₃ ⁻	New/recycled EPA glass vials			-20 °C	Molivate colorimetry ²⁴ Colorimetry ²⁵ Cadmium reduction ²⁶	2.3.4
DOM	New EPA glass vials	H ₃ PO ₄ HCl	Stainless steel + 13 mm glass fibers filter	4 °C or -20 °C	High temperature combustion ^{27,28}	2.3.5
NH ₄ ⁺	New EPA glass vials			-20 °C	Fluorometric nanomolar technic ²⁹	2.3.4
POM	Vacuum flask			-20 °C	CHN analyzer	2.3.6

Table 2. Overview of the sampling vessel, fixative, in-line filter assembly, storage and analytical methods described in the protocol section. The analyzed compounds are: ultra-plankton abundance (plankton < 10 µm), silicate (SiO₄), phosphate (PO₄³⁻), nitrite + nitrate (NO₂⁻ + NO₃⁻), dissolved organic matter (DOM), ammonium (NH₄⁺) and particulate organic matter (POM). All the sampling vessels have silicon septum cap and are vacuumed before sampling. The fixatives are: paraformaldehyde + glutaraldehyde (Glut + Parafor), orthophosphoric acid (H₃PO₄) and hydrochloric acid (HCl). The in-line filter assemblies used are: polycarbonate filter holders and polycarbonate membrane 0.2 µm filters (PC filter holder + PC membrane) and stainless steel filter holders and binder-free glass fiber filters GFF.

Representative Results

Optimization of seawater collection methods

Selection of collector vials and cleaning procedure

VacuSIP-compatible collecting vessels should have a septum that allows sampling to be initiated by piercing with a syringe needle. They should withstand the elevated underwater pressure (2-3 bars at typical scuba working depths), and should hold a vacuum. Many (but not all brands) of vials approved by the EPA for the analysis of volatile organics meet these criteria. Pre-cleaned vials approved for DOC and DON analysis are also available. To test the suitability of these vials for the collection and analysis of nutrients and to optimize cleaning procedures, high quality double distilled water was collected in acid-cleaned polypropylene tubes (PP tubes), newly purchased, in acid-cleaned high-density polyethylene vials (HDPE vials), and in EPA glass vials, all equipped with a polytetrafluoroethylene (PTFE) septum cap. The HDPE vials and polypropylene tubes were cleaned as described in section 1.5.2 above, and the EPA glass vials were cleaned by the manufacturer.

The amount of NH_4^+ found in EPA glass vials was relatively minimal ($\leq 0.1 \mu\text{mol L}^{-1}$) and depends upon the high quality double distilled water standard quality. In contrast, NH_4^+ concentrations significantly increased (up to 3 and 7 fold, respectively) and exhibited a higher variability in acid-cleaned polypropylene tubes and in high-density polyethylene vials (ANOVA $F_{(5,53)}=7.183$, $p<0.001$, **Figure 3**). There was no effect of high quality double distilled water contact with the silicon septum on the ammonium analysis.

Comparison of new glass vials versus cleaned/recycled glass vials

To test whether EPA glass vials could be utilized for nutrient analysis more than once, the NO_x^- , PO_4^{3-} , and NH_4^+ concentrations in seawater samples collected in new EPA glass vials were compared to those collected in used EPA glass vials. The new EPA glass vials were pre-cleaned by the manufacturer, while the recycled glass vials were cleaned as described above (1.5.2). Recycled vials had significantly higher NH_4^+ concentration, up to 1.5 fold the level found in new glass vials (t test, $p<0.001$, $n=5$). No significant differences were found in NO_x^- and PO_4^{3-} content between the samples collected in recycled vials and the samples collected in new glass vials (**Figure 4**).

Silicate collection and storing procedures

To determine the best sampling vessel for the analysis of silicate, high quality double distilled water was collected in non-cleaned and in acid-cleaned polypropylene tubes (PP tubes), in acid-cleaned high-density polyethylene vials (HDPE vials), and in EPA glass vials. The expected silicate concentration was close to zero, so values that deviated from the expected concentration were considered contaminated. The silicate concentration significantly differed between the samples collected in the different vials (ANOVA, $F_{(3,19)}=210.047$, $p<0.001$), showing the lowest SiO_4 concentration in the acid-cleaned HDPE vials. Borosilicate glass vials contaminated the samples, with the final SiO_4 concentration increasing by up to $7 \mu\text{mol L}^{-1}$ (**Figure 5**).

Selection of filtration apparatus for dissolved organic matter (DOM) and nutrient analysis

To determine which filter apparatus produces the lowest blank in the analysis of dissolved organic (DOC and DON) and inorganic nutrients (NO_x^- , NH_4^+ , PO_4^{3-}), stainless steel filter holders were compared to polycarbonate in-line Swinney filter holders. With each filter holder type we tested both polycarbonate membrane and pre-combusted glass fiber filter. The combination of stainless steel filter holder and combusted glass fiber filter provided the lowest blanks, whereas the polycarbonate Swinney filter holder equipped with polycarbonate membrane clearly contaminated the samples by up to 9 fold. Increasing the wash volumes did not resolve this problem (**Figure 6**).

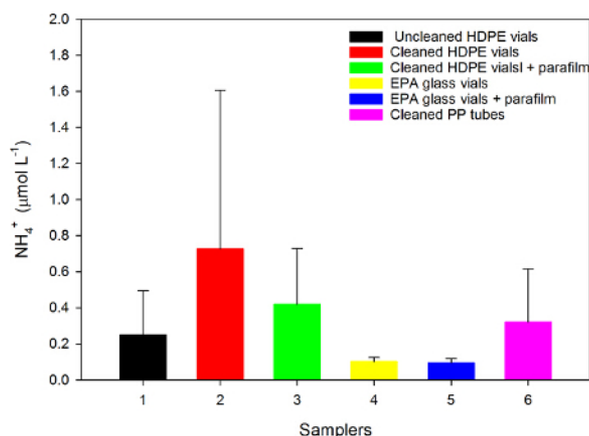


Figure 3. Ammonium concentration ($\mu\text{mol L}^{-1}$, average \pm SD) collected with different vials: (1) Uncleaned HDPE vial; (2) Cleaned HDPE vial; (3) Cleaned HDPE vial + Parafilm; (4) EPA glass vial; (5) EPA glass vial + Parafilm; (6) Cleaned PP tube. The Parafilm was placed to test whether the silicon septum may contaminate the water samples. For each treatment 9 samples of high quality double distilled water were analyzed. The samples were analyzed fresh. Significant differences were found between the four sampling vessels (ANOVA, $F_{(5,53)}=7.183$, $p<0.001$, power test=0.992). [Please click here to view a larger version of this figure.](#)

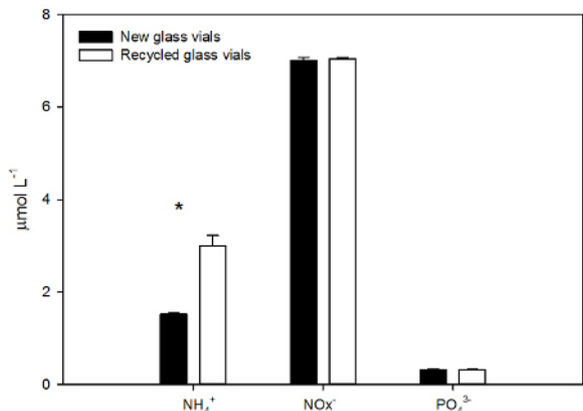


Figure 4. Ammonium (NH₄⁺), nitrite + nitrate (NO_x⁻), and phosphate (PO₄³⁻) concentrations (µmol L⁻¹, average ± SD) of seawater samples collected in new (dark) and recycled/cleaned (white) EPA glass vials. Seawater was collected at the Experimental Aquaria Zone of the Institute of Marine Science and was filtered with stainless steel filter holder and glass filter. The water samples were analyzed fresh. The asterisk (*) indicates that the difference is significant (*t* test, *p*<0.001, *n*=5, power test=1). [Please click here to view a larger version of this figure.](#)

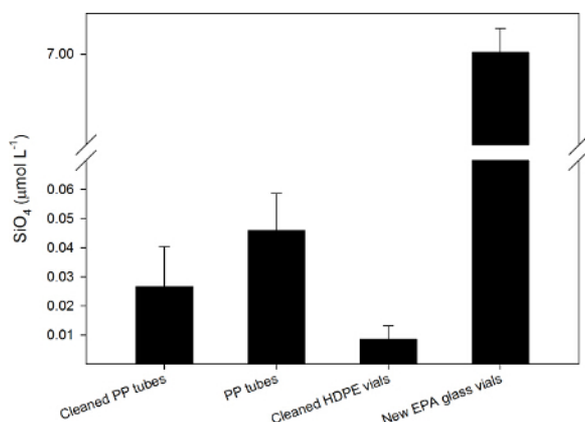


Figure 5. Silicate concentration (µmol L⁻¹, average ± SD) in high quality double distilled water collected in different vials: acid-cleaned PP tubes, PP tubes, acid-cleaned HDPE vials, new EPA glass vials. Significant differences were found between the four sampler materials (ANOVA, *F*_(3,19)=210.047, *p*<0.001, power test=1). [Please click here to view a larger version of this figure.](#)

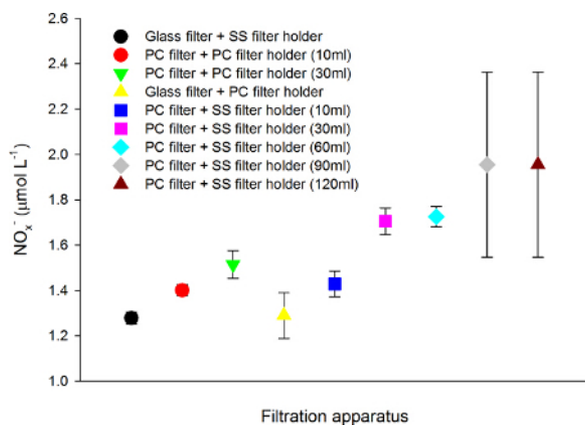


Figure 6. Examining the effect of different filtration assemblies and wash volumes on nitrite + nitrate (NO_x⁻ µmol L⁻¹). Samples for NO_x⁻ were obtained by filtering the seawater samples with stainless steel (SS filter holder) or polycarbonate in-line Swinney filter holders (PC filter holder) equipped with either a polycarbonate membrane (PC filter) or a pre-combusted glass fiber filters. For the PC filters, different volumes (10, 30, 60, 90 and 120 ml) of 5% HCl and high quality double distilled water were used for washing the filter assembly, the washing volume is given in parenthesis in the figure legend. Values are expressed as mean ± standard deviation (*n*=5). Seawater was collected at the Experimental Aquaria Zone of the Institute of Marine Science and the samples were analyzed fresh after the filtration. [Please click here to view a larger version of this figure.](#)

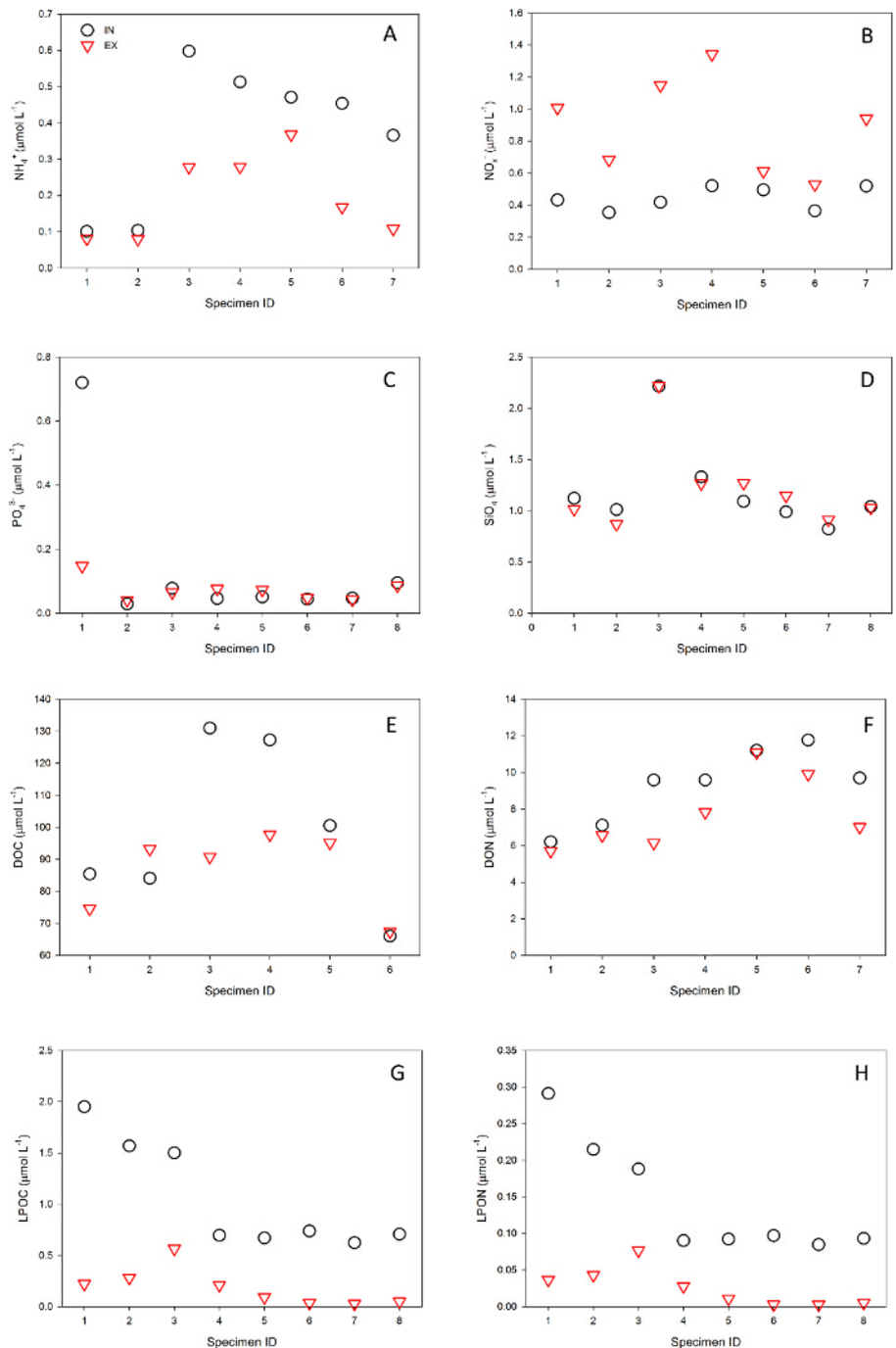


Figure 7. Example of experimental results: inhaled (IN, black circle) and exhaled (EX, red triangle) paired water sample concentrations ($\mu\text{mol L}^{-1}$) of different substances processed by the sponge *Chondrosia reniformis* in the Mediterranean Sea: **(A)** ammonium (NH_4^+); **(B)** nitrite + nitrate (NO_x); **(C)** phosphate (PO_4^{3-}); **(D)** Silicate (SiO_4); **(E)** dissolved organic carbon (DOC); **(F)** dissolved organic nitrogen (DON); **(G)** planktonic organic carbon (LPOC); **(H)** planktonic organic nitrogen (LPON). [Please click here to view a larger version of this figure.](#)

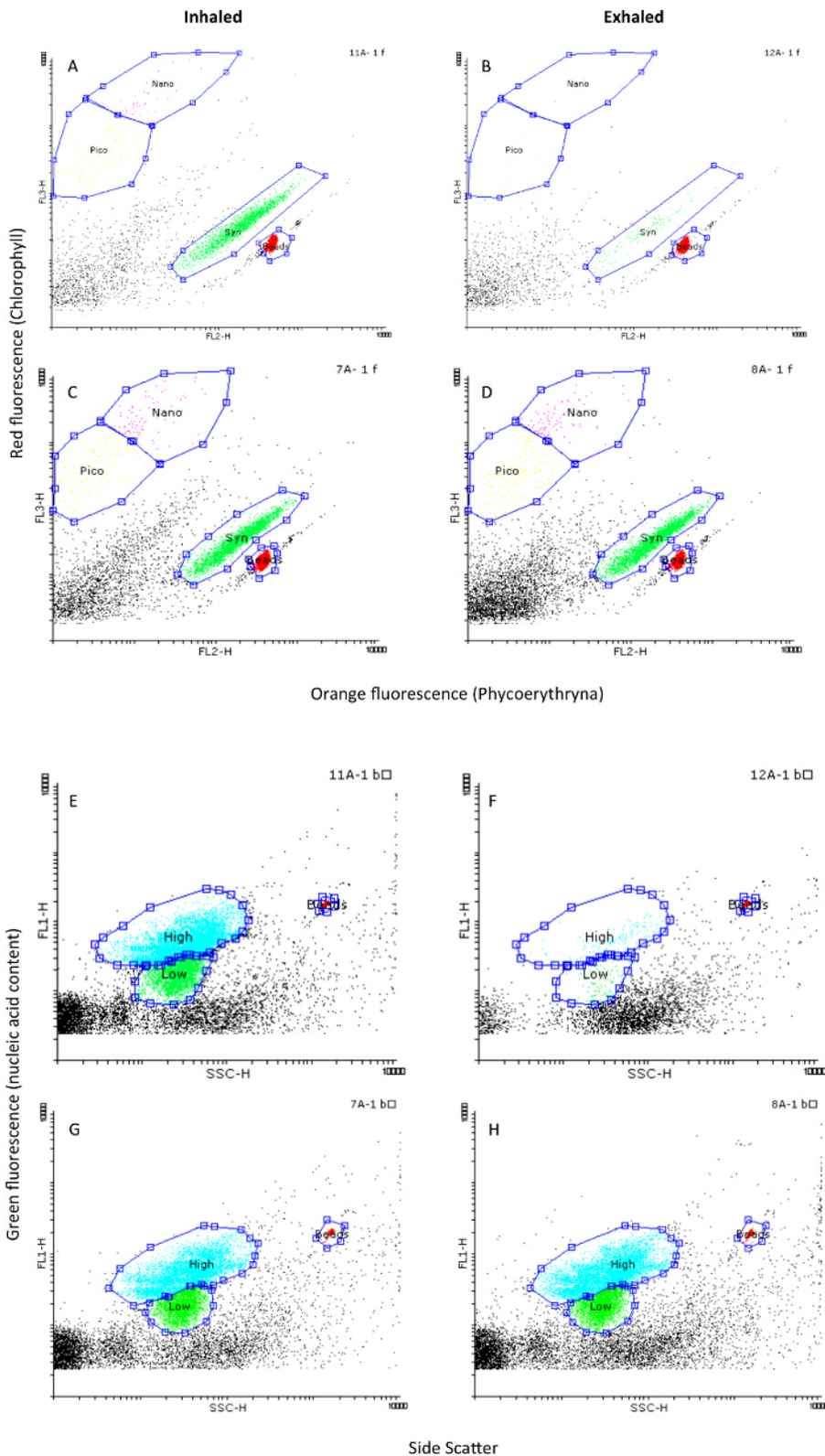


Figure 8. An example of a flow cytometry analysis of paired water samples drawn from the water inhaled (**A,C,E,G**) and exhaled (**B,D,F,H**) by the sponge *Chondrosia reniformis*: (A,B,C,D) phytoplankton populations; (E,F,G,H) heterotrophic bacteria. In A-B and E-F the sampling was clean and accurate (all planktonic groups were efficiently retained). C-D and G-H are examples of exhaled water contamination, showing low removal of all planktonic groups. Syn: *Synechococcus sp.*, pico: autotrophic picoeukaryotes, nano: autotrophic nanoeukaryotes, high: heterotrophic bacteria with high DNA content, low: heterotrophic bacteria with low DNA content. [Please click here to view a larger version of this figure.](#)

Supplementary Figure 1. Cell retention efficiency of different planktonic prey by the clam *Chama pacifica*: *Prochlorococcus* sp. (Pro), *Synechococcus* sp. (Syn), pico-eukaryotes (Pico Euk), nano-eukaryotes (Nano Euk). Error bars = 95% CI. [Please click here to download this file.](#)

Supplementary Video 1. Sampling the ascidian *Polycarpa mytiligera* using a custom-built manipulator with the color code used green for inhaled and yellow for exhaled water samples. Sampling tubes (PEEK, ID 54 μm , 75 cm long) are carefully placed in the exhalant and inhalant siphons of the ascidian. Water is then drawn into evacuated tubes at a rate of $\sim 1 \text{ ml min}^{-1}$. In this demonstration fluorescein dye is used to visualize the exhalant jet. Note that the exhalant sampling tube is placed well within the exhalant jet. [Please click here to download this file.](#)

Discussion

Preparatory steps

Collector vials for DOM and nutrient analysis

Since collector vessels may interact with dissolved micro-constituents and the sampler walls may be a substrate for bacteria growth³⁰⁻³⁴, different vials for DOM and nutrient collection were tested. Borosilicate is not recommended for silica quantification^{33,35}, since glass bottles can increase the initial concentration of silica by up to two fold if the samples are not quickly frozen³⁰. Our results demonstrate that using pre-cleaned EPA vials results in low concentration blanks for DOC, DON, and inorganic nutrients, most notably for ammonium.

DOC filtration and storage

Filtration is a required and, in many cases, is the first analytical step in marine chemistry and microbiology. While it is possible to filter the samples after collection in the lab, this procedure is not recommended for *in situ* work, where samples are collected underwater, often in remote locations, hours or days away from proper laboratory facilities. The use of in-line, *in situ* filtration minimizes sample handling and thus reduces the risk of contamination. *In situ* filtration also removes most of the bacteria and reduces the risk that the sample composition will be altered by bacterial metabolism during the prolonged sampling and transport time. The filtration assembly increases the dead volume of the sampling apparatus and may also be a source of contamination. A selection of the smallest possible filter holders (e.g., in-line Swinney filter holder 13 mm) and minute PEEK tubing (e.g., 254 μm ID) reduces the dead volume and the risk of contamination by ambient water.

If the proper filter is not used or if it is not washed carefully, artifacts and contamination of the water samples are likely to occur^{32,36-39}. Studies of DOC analysis showed that filters and filter holders made of organic compounds (polycarbonate and PFA-PTFE) may result in severe DOC contamination^{32,37}, especially when not thoroughly flushed with high quality double distilled water³⁸. The present protocol and results follow these guidelines and also indicate that polycarbonate filter holders should be avoided.

In situ work and interoperation

The VacuSIP system is a direct sampling technique that facilitates the study of the metabolism of undisturbed suspension feeders in their natural environment and the quantification of their ecological role in the system. For experienced and equipped divers, the application of the VacuSIP method is simple and requires only short training. InEx VacuSIP experiments are designed for a 'within'-design statistical analysis (*i.e.*, paired or repeated measure analysis), therefore controlling for most analytical artifacts including high blanks. The use of controlled suction ensures slow and adjustable sampling rates, thus preventing accidental contamination of exhaled water with ambient water. Where possible, the selection of work sites with low current and low turbidity is recommended and will ensure cleaner and more accurate results. The prolonged sampling time (minutes to hours) allows integration of the high patchiness that characterizes the benthic boundary layer. All these features ensure that when properly applied the VacuSIP method is highly robust, providing reliable and replicable results even when working with a small number of replicates. An example of typical results obtained from a Mediterranean sponge and Indo-Pacific clam species is shown in **Figure 7** and **Supplementary Figure 1**.

As with any technique, the VacuSIP is not free of potential pitfalls. The most common problem is contamination of the exhaled water sample with ambient water. Reasons for these artifacts include high suction rate, tube dislodgments, and animal behavior. Proper selection of the correct sampling rate is dependent on prior estimates of the excurrent flow rate. Such estimates can be obtained by using the dye front speed method¹⁶. Ideally, the suction rate should be kept below 1% of the pumping rate (e.g., 1 ml min^{-1} for a 6 L hour^{-1} pumping rate). To avoid contamination with ambient water, sampling rate should never be greater than 10% of the pumping rate.

To control for the sampling rate, the length and internal diameter of the intake tubing should be adjusted according to the planned work depth and water temperature. The Hagen-Poiseuille equation (see section 1.3.1 above) may be used as a guide. However, this equation should be considered as a first order approximation since ΔP and sampling rate decrease with sampling time and in-line filtration adds uncertainties. The use of evacuated containers, sometimes with unknown vacuum pressures, introduces further complications. An example of how sampling rates varies as a function of different evacuated containers with different vacuum, is shown in **Table 1**.

Reducing the sampling rate is easily achieved by adjusting the tube length and ID, with no technical limitations to this reduction (sampling rates of few microliters per hour are feasible). Nevertheless, experimenters should be aware of the slow sampling rate dictated by this limitation for animals with slow pumping rate and for small organisms or specimens. The immediate implication of slow sampling rate is the limited volume of water that can be collected during a single sampling session. This low volume will limit the number of analyses and replicates that can be run with these samples, and will thus also limit the information that can be obtained from these populations.

Tube dislodgement can be easily spotted and sampling can be aborted or restarted, provided that a diver is keeping constant watch. In contrast, cessation of pumping during sampling is not always easy to detect. This is true not only for sponges, but also for tunicates, bivalves, and polychaetes. In fact, contrary to common belief, events in which an ascidian or a bivalve stopped pumping were documented with no visible change in the siphon geometry (Yahel, unpublished data). Moreover, in some cases, tunicates can maintain active pumping with no mesh secretion (that is, no filtration is taking place).

Controlling the sampling rate is critical. In this respect the VacuSIP is better than other methods, especially when the study animals are relatively small or when they pump slowly. Syringes are particularly difficult to control². For instance, Perea-Blázquez and colleagues (2012a)⁴⁰ used a syringe to sample the water exhaled by several temperate sponge species and surprisingly did not find a general pattern of ingestion/excretion of particular nutrients (NO₂⁻, NO₃⁻, NH₄⁺, PO₄³⁻, SiO₄). The lack of a clear pattern is likely a result of contamination of the exhaled samples with ambient water due to syringe use. Contamination is evident from the extremely low retention efficiency of pico plankton reported by Perea-Blázquez and colleagues (2012b)⁴¹ for their sponges: 40 ± 14% of heterotrophic bacteria and 54 ± 18% of *Synechococcus sp.* For comparison, using the VacuSIP, Mueller *et al.* (2014)²¹ reported a removal efficiency of heterotrophic bacteria of 72 ± 15% in *Siphonodictyon sp.* and 87 ± 10% in *Cliona delitrix*.

To verify sample quality and ensure that no contamination of the ambient water occurs, we strongly recommend to first analyze the pico and nano plankton samples using flow cytometry. This fast, reliable, and cheap analysis will provide immediate information of sample quality. It is very common for some prey taxa such as *Synechococcus sp.* to be removed at close to 90% efficiency^{14,42} by sponges and ascidian. Significant deviations from this benchmark suggest that contamination might have occurred (Figure 8).

For reliable and clean sampling, make sure the experiment design satisfies seven simple rules: (1) perform a preliminary survey (including pumping rate estimates) and prepare the worksite well; (2) know the studied animals; (3) verify that the studied specimen has a well-defined excurrent aperture and accessible location; (4) verify that the studied specimen is pumping before and after each sample collection; (5) place the tube for the collection of the exhaled water slightly inside the excurrent aperture (Figure 1); (6) use sampling rate < 10% of the excurrent flow rate, 1% is highly recommended; (7) define a quality criterion and omit suspected InEx pairs.

Following these simple rules, the VacuSIP system offers a practical and reliable method of measuring how active suspension feeders process particulate and dissolved compounds in natural conditions, allowing accurate and comparable estimates that can be used to assess the functional role of filter feeders in different ecosystems around the world.

Disclosures

The authors have nothing to disclose.

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