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Dissolved organic carbon (DOC) is essential to balance the metabolic demands of four dominant North-Atlantic deep-sea sponges

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Abstract

Sponges are ubiquitous components of various deep-sea habitats, including cold water coral reefs, and form deep-sea sponge grounds. Although the deep sea is generally considered to be a food-limited environment, these ecosystems are known to be hotspots of biodiversity and carbon cycling. To assess the role of sponges in the carbon cycling of deep-sea ecosystems, we studied the carbon budgets of six dominant deep-sea sponges of different phylogenetic origin, with various growth forms and hosting distinct associated microbial communities, in an ex situ aquarium setup. Additionally, we determined biomass metrics—planar surface area, volume, wet weight, dry weight (DW), ash-free dry weight, and organic carbon (C) content—and conversion factors for all species. Oxygen (O₂) removal rates averaged $3.3 \pm 2.8 \ \mu \text{mol O}_2$ g DW_{sponge} h⁻¹ (mean \pm SD), live particulate (bacterio- and phytoplankton) organic carbon removal rates averaged $0.30 \pm 0.39 \ \mu \text{mol C}$ g DW_{sponge} h⁻¹ and dissolved organic carbon (DOC) removal rates averaged $18.70 \pm 25.02 \ \mu \text{mol C}$ g DW_{sponge} h⁻¹. Carbon mass balances were calculated for four species and revealed that the sponges acquired 1.3–6.6 times the amount of carbon needed to sustain their minimal respiratory demands. These results indicate that irrespective of taxonomic class, growth form, and abundance of microbial symbionts, DOC is responsible for over 90% of the total net organic carbon removal of deep-sea sponges and allows them to sustain themselves in otherwise food-limited environments on the ocean floor.

The oceanic seafloor constitutes by far the largest, and least studied, part of the Earth's surface area. It covers an area of 361 million km² of which over 90% is found at water depths greater than 150 m (Costello et al. 2010; Ramirez-Llodra et al. 2010). On the northern Atlantic continental shelf, the seafloor is abundantly inhabited by sponges that form large mono-specific sponge grounds, create sponge reefs by depositing thick spicule mats (i.e., layers of skeletal needles derived

fauna (Klitgaard 1995; Beazley et al. 2013; Hawkes et al. 2019).

Moreover, the first estimations on respiration and organic carbon uptake of deep-sea sponges (e.g., Pile and Young 2006; Yahel et al. 2007; Kahn et al. 2015) suggest that they play a crucial role in benthic-pelagic coupling. However, due to technical restrictions inherent to deep-sea work (e.g., costly shipbased expeditions, sampling under extreme conditions), data on the ecology and physiology of deep-sea sponges is still scarce. The few available studies on deep-sea sponge physiology consist of a mix of in situ and ex situ studies using differ-

from dead and damaged sponges), and are major components

of deep-sea coral reefs (Thomson 1873; Buhl-Mortensen

et al. 2010; Beazley et al. 2015). Deep-sea sponges fulfill

important ecological roles in these habitats, as they provide substrate and habitat complexity to both mobile and sessile

ent direct (collecting inhaled and exhaled water samples [Pile

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and Young 2006; Yahel et al. 2007; Leys et al. 2018]) and indirect (using flume experiments (Witte et al. 1997), or incubation chambers (Kutti et al. 2013, 2015; Rix et al. 2016) methodologies. Still, data on the metabolic rates of deep-sea sponges is often incomplete, and does not reflect the diversity and wide array of morphological traits found in deep-sea sponges.

Deep-sea sponges mainly belong to two taxonomic classes: demosponges (Demospongiae) and glass sponges (Hexactinellidae) (Lancaster 2014). Demosponges come in a wide variety of shapes and sizes-ranging from mm-thin encrusting sheets to m-wide barrels—, occur in freshwater and marine ecosystems, and their skeleton can consist of siliceous, calcium carbonate, or collagenous components (e.g., Müller et al. 2006; Ehrlich et al. 2010; Bart et al. 2019). Hexactinellids are exclusively marine, tubular, cup-, or vase-shaped, predominantly inhabit deep-sea habitats, and their skeleton consists of silica spicules (e.g., Schulze 1887; Mackie and Singla 1983; Leys et al. 2007). Sponges can be further classified as having either low or high abundances of associated microbes (LMA or HMA, respectively [Hentschel et al. 2003; Weisz et al. 2008]). LMA sponges contain microbes with abundances and sizes comparable to ambient seawater ($\sim 0.5-1 \times 10^6 \text{ cells mL}^{-1}$), while HMA sponges can harbor up to four orders of magnitude more and generally much larger microbes (Vacelet and Donadev 1977; Reiswig 1981; Hentschel et al. 2003). These symbionts are involved in various metabolic processes, including carbon (C) and nitrogen (N) metabolism (reviewed by Pita et al. 2018).

Sponges, including deep-sea species, are well-established filter feeders, efficiently capturing and processing nano- and picoplankton (reviewed by Maldonado et al. 2012). Additionally, it has been shown that many shallow-water sponges primarily rely on dissolved organic matter (DOM) as a food source (reviewed by de Goeij et al. 2017). DOM, often measured in the form of dissolved organic carbon (DOC), is the largest potential food source in the oceans (Hansell et al. 2009). However, DOC uptake has only been confirmed for one deep-sea sponge species (Hymedesmia coriacea [Rix et al. 2016]), but only qualitatively, using laboratory-made stable-isotope-enriched DOM. Consequently, direct evidence of ambient DOM uptake by deep-sea sponges is still not available at present. For some species DOM uptake has been suggested (Leys et al. 2018), for others it was not found (Yahel et al. 2007; Kahn et al. 2015). However, these studies did not directly measure DOC, but derived the dissolved organic carbon fraction from the total organic carbon fraction. Direct DOC measurements are challenging, as they are performed almost within detection limits of current analytical systems.

Both microbial abundance and growth form are suggested to affect the capability of sponges to utilize dissolved food sources. Higher DOM uptake is predicted for HMA sponges in comparison with LMA sponges, as microbes are considered to play an essential role in the processing of DOM (Reiswig 1974;

Freeman and Thacker 2011; Maldonado et al. 2012; Hoer et al. 2018). However, this distinction is not always clear, as the diet of some LMA sponges also consists mainly of DOM (e.g., de Goeij et al. 2008; Mueller et al. 2014). This is particularly found for sponges with an encrusting growth form (reviewed by de Goeij et al. 2017), since it is hypothesized that their high surface-to-volume ratio is advantageous for the uptake of DOM compared to lower surface-to-volume ratio of erect, massive (e.g., ball, cylinder) growth forms (Abelson et al. 1993; de Goeij et al. 2017).

To quantify the metabolic- and carbon removal rates of deep-sea sponges, we investigated the oxygen and dissolved and particulate organic carbon removal rates of six dominant North-Atlantic deep-sea sponges with different morphological traits (three massive HMA demosponges, two encrusting LMA demosponges, and one massive LMA hexactinellid) and composed carbon budgets using ex situ incubation experiments. Additionally, we determined different biomass metrics for the six targeted species (planar surface area, volume, wet weight, dry weight, ash-free dry weight, and organic carbon content) and provide species-specific conversion factors.

Materials and methods

Study areas, sponge collection and maintenance

We investigated the following dominant North-Atlantic deep-sea sponge species (Fig. 1 and Table S1): Vazella pourtalesii (Hexactinellidae; LMA; massive vase), Geodia barretti (Demospongiae; HMA; massive globular), Geodia atlantica (Demospongiae; HMA; massive bowl), Craniella zetlandica HMA; massive globular), Hymedesmia (Demospongiae: paupertas (Demospongiae; LMA; encrusting sheet), and Acantheurypon spinispinosum (Demospongiae; LMA; encrusting sheet). Sponge specimens were collected by ROV during four research cruises in 2016, 2017 (two cruises), and 2018 (Fig. 1). Whole V. pourtalesii individuals were collected in August 2016, attached to their rocky substrate at ~ 300 m depth, during the Hudson cruise 2016-019 (Kenchington et al. 2017) at the Emerald Basin on the Scotian Shelf, Canada (44°19'8.73"N 62°36′18.49″W). Sponges were kept in the dark in a 500-liter flow-through holding tank onboard the vessel. The temperature was maintained at 6°C, and surface (~ 5 m depth) seawater was added at a rate of 1.2 L min⁻¹. The sponges were transported without air exposure to the Bedford Institute of Oceanography, Dartmouth, Nova Scotia, Canada. In the lab, sponges were kept in the dark in a 500-liter flow-through holding tank. Seawater pumped from the Bedford Basin was passed through a sand filter and 20μ bag filters and maintained at 6°C. This water was continuously pumped into the sponge holding tank at 7 L h⁻¹. A small magnetic drive pump was added to the bottom of the holding tank to provide circulation and horizontal flow across the sponges. Whole C. zetlandica individuals were collected during the Kristine Bonnevie cruise 2017610 (April 2017) at 60°42′12.5″N

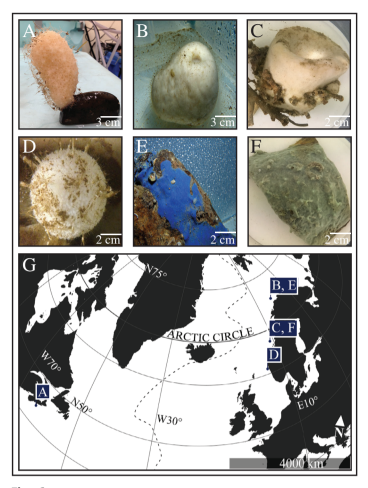


Fig. 1. Photographs of six dominant North-Atlantic deep-sea sponge species used in the study (top) and their North-Atlantic sampling area (bottom). (**A**) *V. pourtalesii,* (**B**) *G. barretti,* (**C**) *G. atlantica,* (**D**) *C. zetlandica* (courtesy of Erik Wurz), (**E**) *H. paupertas,* (**F**) *A. spinispinosum,* (**G**) location where individuals of each sponge species were collected during four different research cruises. The dotted line represents the Mid-Atlantic Ridge.

4°39′09.9″E in the province of Hordaland, Norway, and kept in 14-liter onboard flow-through aquaria with seawater pumped through at 120 L h⁻¹. Temperature was maintained at 8°C. Whole G. atlantica and A. spinispinosum individuals were collected attached to their rocky substrate during the G.O. Sars cruise 2017110 (August 2017) at the Sula reef (64°42'25.2"N 7°59'24.0"E) of the Northern Norwegian coast at depths of 250-400 m. During the same cruise, G. barretti individuals were collected at the Barents Sea (70°47′20.8″N 18°03′47.2″E), at a depth of 272 m. The latter three sponge species were kept on board the research vessel in the dark in 20-liter flow-through tanks in a climate room at 6°C. North-Atlantic seawater was pumped in from a depth of 6 m at 30 L h⁻¹. H. paupertas individuals were collected attached to rocky substrate during the G.O. Sars cruise 2018108 (August 2018) in the Barents Sea at 70°47′13.9″N 18°03′23.8″E. These sponges were kept on board the research vessel under similar

conditions as during the previous year. Ex situ experiments with G. atlantica, A. spinispinosum, and H. paupertas individuals were performed on board. The other individuals were transported without exposing them to air to the laboratory facilities at the University of Bergen, Norway, where the experiments took place. In Bergen, sponges were kept in a dark climate room (8°C) in multiple 20-liter flow-through aquarium systems. Each holding tank contained a maximum of five sponges. Flow originated from unfiltered water pumped from 200 m depth from the outer fjord near Bergen at ~ 50 L h $^{-1}$ with a temperature ranging from 6°C to 8°C. All sponges and substrate were cleared from epibionts prior to incubations.

Incubations, sample treatment, and analysis

All sponges were allowed to acclimatize for a minimum of 1 week prior to the incubation experiments. During experiments, individual sponges were enclosed in flow chambers (either 2, 3, or 6 liter depending on sponge biomass) for 2-8 h (see below) with magnetic stirring devices to ensure proper mixing (de Goeij et al. 2013). Chambers were acid-washed (0.4 mol L⁻¹ HCl) prior to the incubations and kept in a water bath to maintain a constant seawater temperature during the incubations (6–9°C depending on the incubation). Chambers were closed without trapping air in the system. The length of each individual incubation was determined during test incubations based on sponge size and oxygen removal (ideally timed to about > 10% to < 40% [O₂] decrease). At set time intervals depending on the incubation length ($t_{\text{sample}} = 0$, 30, 60, 90,120, 180, 240, 360, or 480 min), 85-100 mL water samples were taken with acid-washed 100-mL polycarbonate syringes. Sample water volume was replaced with aquarium water (drawn in by suction to maintain volume and eliminate air exposure). The concentration of bacterio- and phytoplankton in the water bath from which water is drawn in, is assumed to be the same as the concentration at t_0 of the incubation, and remains constant over time. Consequently, during each sampling, a known number of t_0 plankton with the same volume as the sample drawn is added again to the incubation water at that time interval. Subsequently, the concentration of plankton at the beginning of the next time interval was recalculated to correct for replacement water. These corrections were only performed for plankton, as the amount of DOC and O2 added with the drawn in water, was too small to have an effect on the total concentration of each substance within the incubation chamber.

Seawater incubations without sponges were performed as controls at each location according to the aforementioned protocol (Canada, n = 4; on board incubations in 2017, n = 4; on board incubations in 2018, n = 3; Bergen, n = 6).

Water samples were subdivided to analyze the concentrations of DOC and abundances of bacterio- and phytoplankton. Unfortunately, due to logistical issues (samples were lost during transport) and a malfunctioning autoanalyzer at the University of Amsterdam, we were not able to analyze all

DOC samples. For two species, C. zetlandica (n = 3) and H. paupertas (n = 3), no DOC samples could be measured, and we could only successfully analyze DOC samples (at GEOMAR Helmholtz Centre for Ocean Research, Kiel, Germany) for a subset of the total amount of individuals incubated for the other species; V. pourtalesii (n = 4/7), G. barretti (n = 3/12), G. atlantica (n = 4/6), and A. spinispinosum (n = 3/4).

Dissolved oxygen concentrations (O_2) were continuously measured during the incubations with OXY-4 mini optical oxygen sensors (*PreSens*, Germany). Sensors do not consume oxygen and due to their small dimensions (Ø 2 mm), flow and mass-transport inside the chambers are not disturbed. O_2 concentrations were recorded every 15 s (*OXY-4-v2_30FB software*).

Prior to DOC sampling, syringes, glassware, and pipette tips were rinsed three times with acid (8 mL, 0.4 mol L⁻¹ HCl), three times with Milli-Q (80 mL), and twice with sample water (10 mL). Twenty milliliter of sample water was filtered (< 20 kPa Hg suction pressure) over pre-combusted (4 h at 450°C) GF/F glass microfiber ($\sim 0.7 \mu m$ pore-size) filter and collected in pre-combusted (4 h at 450°C) amber glass EPA vials (40 mL). Samples were acidified with six drops of concentrated HCl (12 mol L⁻¹) to remove inorganic C, and stored in the dark at 4°C until analysis. DOC concentrations were analyzed using a total organic C analyzer and applying the hightemperature catalytic oxidation method Shimadzu) modified from Sugimura and Suzuki (1988). Every 8-10 d the instrument was calibrated by measuring standard solutions of 0, 42, 83, 125, 208, and 417 μ mol C L⁻¹, prepared from a potassium hydrogen phthalate standard (Merck 109017). Every measurement day, ultrapure (Milli-Q) water was used to determine the instrument blank (< 1 μ mol C L⁻¹). On every measurement day TOC analysis was validated with deep seawater reference (DSR) material provided by the Consensus Reference Materials Project of RSMAS (University of Miami) yielding values within the certified range of $42-45 \mu \text{mol C L}^{-1}$. Additionally, two internal standards were prepared each measurement day using a potassium hydrogen phthalate (Merck 109017) with DOC concentration within the samples range. DOC of each sample was determined from 5 to 8 injections. The precision was < 4% estimated as the standard deviation of replicate measurements divided by the mean.

Duplicate 1 mL samples for bacterio- and phytoplankton were fixed at a final concentration of 0.5% glutaraldehyde for 15–30 min at 4°C in the dark. After fixation, the samples were snap-frozen in liquid nitrogen and stored at -80° C until further analysis. Thawed samples were analyzed using a BD-FACSCalibur flow cytometer (*Becton Dickinson*, San Jose, California) with a 15-mW air-cooled argon laser (Brussaard 2004). Phytoplankton were enumerated for 10 min at 80 μ L min⁻¹ with the trigger on Chlorophyll a, red autofluorescence (Marie et al. 1999). Phycoerythrin containing cells (e.g., cyanobacterial *Synechococcus*) were discriminated by their

orange autofluorescence. Bacterioplankton samples were diluted (5x or 10x, to keep the event rate at 200–800 events s⁻¹) in sterile TE-buffer, pH 8.0 (10 mmol L⁻¹ Tris, *Roche Diagnostics*; 1 mmol L⁻¹ EDTA, *Sigma-Aldrich*) to avoid electronic coincidence, and stained with nucleic acid-specific SYBR Green I to a final concentration of 1×10^{-4} of the commercial stock (Marie et al. 1999; Brussaard 2004). Samples were corrected for blanks (TE-buffer with SYBR Green I) prepared and analyzed in a similar manner as the samples. Bacterioplankton samples were incubated in the dark for 15 min at room temperature after which samples were allowed to cool down at room temperature. Samples were analyzed for 1 min at 40 μ L min⁻¹. Listmode files were analyzed using CYTOWIN freeware (Vaulot et al. 1989).

Sponge biomass metrics

After the incubations, sponges were removed from their substrate and analyzed for volume (by water displacement) and (dripping) wet weight. Sponges were photographed, and planer surface area was calculated with ImageI on scaled pictures (Schneider et al. 2012). Then, all sponges were dried for 72 h in a drying oven at 60° C to determine dry weight. Randomly selected 1-cm³ cubes (n=6) of each massive sponge were transferred into a pre-weighed crucible and combusted at 450° C in a muffle furnace (4 h). Combusted samples were cooled to room temperature in a desiccator and weighed (ash weight). Subsequently, ash-free dry weight was calculated by subtracting ash weight from dry weight and normalized to total volume of the original sponge specimen. The rest of the dried sponges was crushed and ground up with mortar and pestle and stored in a desiccator until further analysis.

Samples for organic C content analysis were decalcified with HCl (4 mol L⁻¹) to ensure removal of inorganic C and subsequently lyophilized for 24 h in a *FD5515 Ilchin Biobase* freeze-drier. After freeze-drying, aliquots of approximately 10 mg were placed in tin-capsules and analyzed on an Elemental Analyzer (*Elementar Isotope cube, Elementar GmbH,* Langenselbold, Germany) coupled to a BioVision isotope ratio mass spectrometer (*Elementar Ltd,* Manchester, UK).

Oxygen and carbon removal rates

Kinetics for each component were described according to its best fitting mathematical model, integrated over the entire time frame of the incubation, to estimate and compare the most reliable initial fluxes of those components, based on the tangent of each model at t_0 .

To calculate changes in O_2 concentrations over time, a linear regression analysis was performed for each individual incubation. Resulting net O_2 removal rates were subsequently compared between sponge and seawater control incubations with a Welch's *t*-test for each species and a respective set of seawater controls. O_2 removal rates were corrected for background seawater respiration by subtracting the average respiration in the seawater controls from the sponge incubations.

Initial net live bacterio- and phytoplankton removal rates were calculated assuming exponential clearance of cells in incubations (Scheffers et al. 2004; de Goeij et al. 2008). The live planktonic fraction was dominated by two general cell types, heterotrophic bacteria and phytoplankton, the latter represented by *Synechococcus*-like cyanobacteria. To calculate net removal rates for each planktonic component, the average initial cell concentrations of all incubations were used as a starting point. Standardized data were fitted to an inverse exponential model to calculate final cell concentrations. Final concentrations were subtracted from the initial corrected concentrations and differences were compared between treatments (sponge vs. control incubations) using an unpaired *t*-test. Clearance rates (CR) were calculated according to Riisgård et al. (1993):

$$CR = V_w/t \times ln(C_0/C_t)$$

 V_w is the water volume in incubation chamber (in mL). t is the duration of incubation (in min). C_0 is the initial cell (bacterio- or phytoplankton) concentration (in cells mL⁻¹). C_t is the cell concentration at time point t (in cells mL⁻¹).

A conservative estimate of live particulate organic carbon (LPOC) removal was obtained using established conversion factors. Heterotrophic bacterial cells were converted using 30 fg C per bacterial cell (Fukuda et al. 1998; Leys et al. 2018) and phytoplankton using 470 fg C per *Synechococcus*-type cell (Bertilsson et al. 2003; Pile and Young 2006).

Initial net DOC removal rates were calculated by applying a 2G-model (de Goeij and van Duyl 2007; de Goeij et al. 2008). This is a simplified biexponential model to describe bioprocessing of DOC over time, assuming that the complex and heterogeneous DOC pool comprises two major fractions: a fast- (C_f) and slow-removable (C_s) fraction, for labile and refractory components of DOM, respectively (de Goeij and van Duyl 2007; de Goeij et al. 2008). In an assumed wellmixed system, the fast and slow removal fractions of DOC will be consumed according to their specific removal rate constants k_f and k_s , respectively. The sum of the individual removal rates is used here to describe total DOC removal.

$$\frac{dDOC}{dt} = -\left(k_f C_f + k_s C_s\right)$$

Integrating this equation yields the function that describes the concentration of DOC as a function of time:

$$DOC(t) = C_{f,0} \times e^{-kft} + C_{s,0} \times e^{-kst}$$

Experimental data is described with the model by estimating model variables $C_{\rm f}$ and $C_{\rm s}$ using a (10,000 iterations) minimalization routine (de Goeij et al. 2008). The initial DOC removal rate was calculated from the estimated values of these variables, based on the integration of the model over the

entire time frame of the incubation (tangent at t_0), and is given by

$$Flux_{DOC} = -\left(k_f C_{f,0} + k_s C_{s,0}\right)$$

Carbon mass balance

Total net organic carbon (TOC) removal rates were estimated as the sum of net initial LPOC and DOC removal rates. O_2 removal served as a proxy for respiration assuming a balanced molar ratio of carbon respiration to net O_2 removal (1 mol C respired equals 1 mol O_2 removed), yielding a respiratory quotient of 1 (Yahel et al. 2003; Hill et al. 2004).

$$RQ = \frac{\text{moles of C respired per unit time}}{\text{moles of O}_2 \text{ consumed per unit time}} = 1$$

To establish a mass balance for the different deep-sea sponge species, the quotient $\Delta O_2/\Delta TOC$ was calculated using the (bi)exponential removal rates for LPOC and DOC. This quotient describes how much of the carbon intake is used for respiration. Sponges do not meet their minimal respiratory demands at $\Delta O_2/\Delta TOC > 1$. At $\Delta O_2/\Delta TOC < 1$, sponges take up more carbon than their minimal respiratory demands, meaning they can invest excess carbon in processes such as growth and reproduction. Carbon budgets where only calculated for sponges of which we had complete sets of O_2 , LPOC, and DOC data (V. pourtalesii (n = 4), G. barretti (n = 3), G. atlantica (n = 4), and A. spinispinosum (n = 3)).

Results

Sponge biomass metrics

Sponge characteristics (phylogeny, growth form, abundance or associated microbes) and biomass metric conversion factors are given in Table 1. Average sponge biomass metrics; planar surface area (PSA), volume, wet weight (WW), dry weight (DW), ash-free dry weight (AFDW), and organic carbon (C) content are shown in Table S1. Encrusting sponges have a one- to two-orders of magnitude higher planar surface area: volume ratio (4.2–10.0) than massive sponges (0.2–0.3) and an order-of-magnitude higher volume: DW ratio (21.4–22.4 and 3.3–7.3, respectively). HMA sponges show a significantly higher organic C content than LMA sponges (t = -8.13, df = 27, p < 0.001; Table 1), with lowest values for the hexactinellid V. pourtalesii.

Oxygen removal rates

The concentration of O_2 in the incubation chambers linearly decreased with time for V. pourtalesii (t = 4.59, df = 7, p < 0.01), G. barretti (t = 3.69, df = 11, p < 0.01), G. atlantica (t = 5.11, df = 5, p < 0.01), G. zetlandica (t = 3.5, df = 3, p < 0.05), G. H. paupertas (G = 4.38, df = 2, G = 2, G = 2, G = 3, G = 3, G = 3, G = 3, G = 4.38, df = 2, G = 2, G = 3, G = 3, G = 3, G = 3, G = 4.38, df = 2, G = 4.38, df = 2, G = 4.38, df = 2, G = 4.38, df = 3, G = 4.38, G =

Table 1. Characteristics, biomass metrics and conversion factors for six dominant North-Atlantic deep-sea sponges. H = Hexactinellidae, D = Demospongiae, LMA = low microbial abundance, HMA = high microbial abundance. Planar surface area (PSA) is the surface area covered in a 2D top view, volumes are measured by water displacement in mL and the weight is given as g dry weight (DW). Conversion factors are based on average biomass metrics (planar surface area, volume, wet weight (WW), DW, ash-free dry weight (AFDW), organic carbon (C) content) for all individuals used in the experiments shown in Table S1.

Sponge species	Class	Growth form	LMA/HMA	PSA: Volume (cm²: mL)	Volume: Weight (mL: g DW)	Organic C content (% DW)
V. pourtalesii	Н	Massive, vase	LMA	0.3	5.2	5.5
G. barretti	D	Massive, globular	HMA	0.3	3.3	15.9
G. atlantica	D	Massive, bowl	HMA	0.3	7.3	20.3
C. zetlandica	D	Massive, globular	HMA	0.2	4.3	20.4
H. paupertas	D	Encrusting, sheet	LMA	10.0	21.4	12.6
A. spinispinosum	D	Encrusting, sheet	LMA	4.2	22.4	10.9

species are depicted in Table 2. Examples of O_2 concentration profiles during incubations for all species and controls are shown in Fig. S1. Initial O_2 removal rates for all sponges averaged $3.3 \pm 2.8~\mu \text{mol}~O_2$ g DW_{sponge} h⁻¹ (mean \pm SD throughout text unless stated otherwise), ranging from 1.0 (*C. zetlandica*) to 7.8 (*A. spinispinosum*).

Plankton (LPOC) removal rates

Bacterioplankton concentrations exponentially decreased in incubations with *G. barretti* (t = 2.44, df = 19, p < 0.05), *V. pourtalesii* (t = 5.91, df = 9, p < 0.001), *G. atlantica* (t = 6.62, df = 5, p < 0.01), *H. paupertas* (t = 2.81, df = 4, p < 0.05), and *C. zetlandica* (t = 4.25, df = 8, p < 0.01) compared to seawater control incubations (Fig. 2A–F). Incubations with *A. spinispinosum* showed no significant decrease in

bacterioplankton compared to control incubations (t=-0.72, df = 4, p=0.51) (Fig. 2F). Average bacterioplankton C removal and clearance rates per species are presented in Table 3. Initial bacterioplankton C removal rates averaged 0.25 \pm 0.35 μ mol C DW_{sponge} h⁻¹ for all species, ranging between 0.00 (A. spinispinosum) and 0.82 μ mol C DW_{sponge} h⁻¹ (V. pourtalesii) (Table 3). Bacterioplankton clearance rates averaged 0.69 \pm 1.06 mL mL_{sponge}⁻¹ min⁻¹ for all species, ranging from 0.00 (A. spinispinosum) to 2.22 mL mL_{sponge}⁻¹ min⁻¹ (V. pourtalesii).

Compared to control incubations, phytoplankton (i.e., *Synechococcus*-type cyanobacteria) concentrations decreased exponentially in incubations with *V. pourtalesii* (t = 5.34, df = 9, p < 0.001), *G. barretti* (t = 2.20, df = 11, p < 0.05), and *G. atlantica* (t = 11.92, df = 6, p < 0.001). Incubations with *C. zetlandica*

Table 2. Overview of oxygen removal rates by deep-sea sponge species (mean \pm SE). CA = Canada, NO = Norway. (1) Leys et al. (2011), (2) Leys et al. (2018), (3) Kutti et al. (2013).

Sponge species	O_2 removal (μ mol O_2 g DW _{sponge} ⁻¹ h ¹)	O_2 removal $(\mu \text{mol } O_2 \text{ mL}_{\text{sponge}}^{-1} \text{ h}^{-1})$	Original location	<i>T</i> (°C)	Method	Reference
V. pourtalesii (n = 7)	3.4 ± 0.7	0.7 ± 0.1	Emerald Basin (CA)	6.7	Incubation	This study
A. vastus $(n = 22)$	-	0.1	Fraser Ridge Reef (CA)	9.0	In-ex*	1
G. barretti (n = 12)	1.3 ± 0.2	0.5 ± 0.1	Barents Sea (NO)	9.0	Incubation	This study
G. barretti (n = 17)	1.4 ± 0.3	0.3 ± 0.1	Langenuen fjord (NO)	8.0–9.0	In-ex	2
G. barretti (n = 6)	1.5	-	Continental Shelf (NO)	6.9–7.4	Incubation	3
G. atlantica $(n = 6)$	5.8 ± 0.9	0.8 ± 0.1	Sula reef (NO)	6.0	Incubation	This study
C. zetlandica (n = 4)	1.0 ± 0.3	0.2 ± 0.1	Continental Shelf (NO)	9.0	Incubation	This study
H. paupertas $(n = 3)$	5.9 ± 1.5	1.5 ± 0.6	Barents Sea (NO)	6.0	Incubation	This study
A. spinispinosum (n = 4)	$\textbf{7.8} \pm \textbf{0.8}$	0.4 ± 0.0	Sula reef (NO)	6.3	Incubation	This study

^{*}Experiments performed in situ.

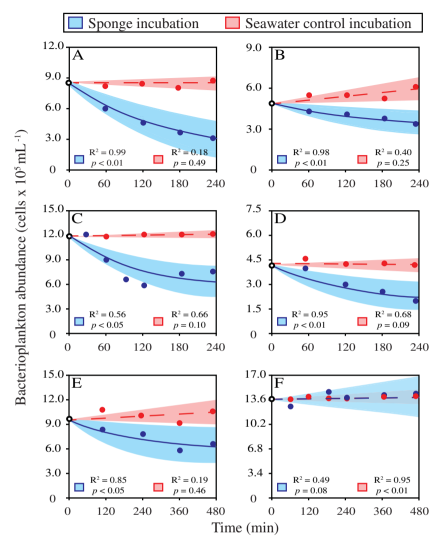


Fig. 2. Average abundances of bacterioplankton over time during incubations with six dominant North-Atlantic deep-sea sponge species (blue) in comparison to seawater control incubations (red). (**A**) *V. pourtalesii* (n = 7), (**B**) *G. barretti* (n = 12), (**C**) *G. atlantica* (n = 6), (**D**) *C. zetlandica* (n = 4), (**E**) *H. paupertas* (n = 3), and (**F**) *A. spinispinosum* (n = 4). Decrease of bacterial abundance over time during incubations is modeled with an exponential fit, shades depict 95% confidence intervals of the model. Note that x- and y-axis show different ranges per species.

 $(t=1.23, \, {\rm df}=3, \, p=0.31)$ and $A. \, spinispinosum \, (t=1.56, \, {\rm df}=7, \, p=0.16)$ showed no significant decrease compared to seawater control incubations. Average phytoplankton C removal and clearance rates per species are presented in Fig. S2 and Table 3. Initial phytoplankton C removal rates averaged $0.04\pm0.07~\mu{\rm mol}$ C g DW_{sponge} h⁻¹ for all species, ranging from $0.00~(A. \, spinispinosum/C. \, zetlandica)$ to $0.15~\mu{\rm mol}$ C DW_{sponge} h⁻¹ ($V. \, pourtalesii$) (Table 3). Phytoplankton clearance rates averaged $0.54\pm0.96~{\rm mL}~{\rm mL_{sponge}}^{-1}~{\rm min}^{-1}$ for all species, ranging between $0.00~(A. \, spinispinosum)$ and $1.77~\mu{\rm mol}~{\rm mL_{sponge}}^{-1}~{\rm min}^{-1}$ ($V. \, pourtalesii$).

Combined initial plankton removal rates amounted to total live particulate organic carbon (LPOC) removal rates of, on average, $0.30 \pm 0.39 \, \mu \text{mol C g DW}_{\text{sponge}} \, \, \text{h}^{-1}$, ranging from 0.00 (*A.* spinispinosum) to 0.97 $\mu \text{mol C g DW}_{\text{sponge}} \, \, \text{h}^{-1}$ (*V. pourtalesii*).

Dissolved organic carbon (DOC) removal rates

Changes in DOC concentration over time during incubations with four different species (V. pourtalesii (n=4), G. barretti (n=3), G. atlantica (n=4), and A. spinispinosum (n=3)) significantly fitted the biexponential 2G-model and thereby showed significant removal of DOC, while no DOC removal occurred in the seawater controls (Figs. 3 and S3). Initial DOC removal rates averaged $18.70 \pm 25.02 \, \mu \text{mol C g DW}_{sponge} \, h^{-1}$ for all sponges, ranging from 3.70 (G. barretti) to $56.07 \, \mu \text{mol C g DW}_{sponge} \, h^{-1}$ (A. spinispinosum) (Table 3).

Carbon mass balance

Carbon mass balances were only constructed for individuals for which a complete set of O₂, LPOC, and DOC was available. Note that initial removal rates depicted in Table 4

Table 3. Average initial (mean \pm SD) dissolved organic carbon (DOC), bacterio- and phytoplankton carbon (BC and PC, respectively) removal rates, and bacterio- and phytoplankton clearance rates of six dominant North-Atlantic deep-sea sponge species. Initial removal rates for bacterio- and phytoplankton are based on exponential uptake during incubations, whereas initial removal rates for DOC are based on a biexponential 2G-model uptake. NA: Not available.

Sponge species	DOC removal rate	BC removal rate	PC removal rate	Bacterioplankton clearance rate	Phytoplankton clearance rate
	(μ mol C g DW _{sponge} ⁻¹ h ⁻¹)			$(mL mL_{sponge}^{-1} min^{-1})$	
V. pourtalesii	$\textbf{9.17} \pm \textbf{2.69}$	$\textbf{0.82} \pm \textbf{0.43}$	0.15 ± 0.18	$\textbf{2.22} \pm \textbf{1.25}$	$\textbf{1.77} \pm \textbf{1.37}$
G. barretti	$\textbf{3.70} \pm \textbf{0.26}$	$\textbf{0.02} \pm \textbf{0.02}$	<0.01	0.15 ± 0.17	$\textbf{0.17} \pm \textbf{0.15}$
G. atlantica	5.85 ± 5.55	$\textbf{0.12} \pm \textbf{0.08}$	0.11 ± 0.13	0.08 ± 0.03	$\textbf{0.25} \pm \textbf{0.24}$
C. zetlandica	NA	$\textbf{0.02} \pm \textbf{0.02}$	0.00	0.06 ± 0.06	$\textbf{0.05} \pm \textbf{0.11}$
H. paupertas	NA	$\textbf{0.55} \pm \textbf{0.42}$	NA	0.15 ± 0.10	NA
A. spinispinosum	56.07 ± 19.92	0.00	0.00	0.00	0.00

are thus based on these measurements only and can deviate from those depicted in Tables 2 and 3. For the four assessed species, more than 90% of the average net total organic carbon (TOC) removal was accounted for by DOC (V. pourtalesii 92.0 \pm 5.5%, G. barretti 99.5 \pm 0.5%, G. atlantica 93.6 \pm 8.4%, A. spinispinosum 100%) (Table 4). Assuming a respiratory quotient of 1 in combination with exponential removal of LPOC and DOC during the incubations, all four species were found to match their minimal required C uptake ($\Delta O_2/\Delta TOC \leq 1.0$), but only when DOC was included in the mass balance. The HMA species show 2–5 times higher $\Delta O_2/\Delta TOC$ ratios than the two LMA species.

Discussion

In this study we show, for the first time, that multiple, dominant, North-Atlantic deep-sea sponge species, irrespective of taxonomic class, growth form, and abundance of microbial symbionts, are capable of consuming ambient DOC, and that this consumed DOC—representing more than 90% of the total organic carbon uptake—is essential to satisfy their minimal respiratory demands. We hypothesize that the combined pallet of dissolved and particulate food allows deep-sea sponges to thrive in otherwise food-limited environments.

Deep-sea sponges oxygen and carbon removal rates

Deep-sea sponge respiration rates (i.e., using O_2 uptake as proxy for respiration) show consistency throughout literature, as most are roughly within the same order of magnitude (Table 2), regardless of the experimental method used. When comparing respiration rates of deep-sea sponges to those reported for temperate (e.g., Thomassen and Riisgård 1995; Coma 2002) and tropical sponges (e.g., Reiswig 1974; Yahel et al. 2003), rates of deep-sea sponges are consistently one to two orders of magnitude lower (Fig. 4). Correspondingly, DOC and plankton carbon removal rates of deep-sea sponges are lower than those found for tropical species (e.g., de Goeij et al. 2008; Hoer et al. 2018; Fig. 4). Differences in O_2 and plankton carbon removal rates can be explained by the

positive effect of temperature on metabolism and physiological processes (see also Clarke and Fraser 2004). DOC removal rates seem to follow a similar trend, yet due to the very limited amount of data available, the relation with temperature was not found to be significant.

In general, O2, planktonic carbon and (specifically) DOC fluxes appear to be higher for encrusting sponges compared to massive growth forms (Fig. 4). For example, the deep-sea encrusting sponge A. spinispinosum has an order-of-magnitude than higher DOC flux massive deep-sea h^{-1} (56.1)µmol C g DW_{sponge} vs. 3.7-9.2- μ mol C g DW_{sponge} h⁻¹), as is the case for encrusting tropical species (218.3–253.3 μ mol C g DW_{sponge} h⁻¹, de Goeij et al. 2008) compared to massive tropical species (10.0--11.9 μ mol C g DW_{sponge} h⁻¹, Yahel et al. 2003; Hoer et al. 2018). This corroborates earlier suggestions that high surface-to-volume ratios enable encrusting sponges to have higher removal efficiencies compared to massive species (Abelson et al. 1993; Kötter et al. 2003; de Goeij et al. 2017).

In addition to morphology, higher net DOC removal rates are generally predicted for HMA sponges in comparison with LMA sponges, as microbes are considered to play an important role in the processing of DOM (e.g., Reiswig 1974; Hoer et al. 2018). However, both LMA species used in this study, A. spinispinosum and V. pourtalesii, showed high removal rates of DOC (56.1 and 9.2 μ mol C g DW_{sponge} h⁻¹, respectively), despite their different growth forms (encrusting vs. massive) and different phylogeny (demosponge vs. hexactinellid). Interestingly, other hexactinellids were previously not found to consume DOM (Yahel et al. 2007). However, Yahel et al. (2007) did not directly measure DOC, but derived it from TOC analysis, potentially resulting in an underestimation of actual DOC removal rates. Our results thereby add to the increasing body of evidence that also sponges with low microbial abundances are capable of consuming DOC (e.g., de Goeij et al. 2013; Rix et al. 2016; Morganti et al. 2017; Rix et al. 2020). However, we conclude that there are still too few available sponge carbon fluxes to fully understand the key

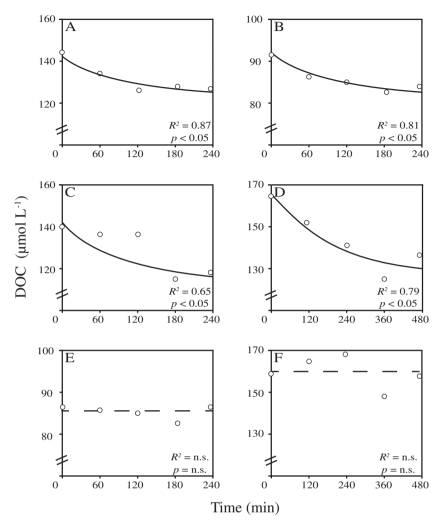


Fig. 3. DOC removal over time by four dominant North-Atlantic deep-sea sponge species compared to seawater controls in ex situ incubations. (**A**) *V. pourtalesii,* (**B**) *G. barretti,* (**C**) *G. atlantica,* (**D**) *A. spinispinosum,* and (**E** and **F**) seawater control. Trend lines are given by a 2G-model fit.

functional traits, such as abundance of microbial symbionts or morphology, that determine the strategy of sponges to process DOM (discussed in de Goeij et al. 2017).

Deep-sea sponge carbon budgets

The contribution of DOC to the total organic carbon removal of the investigated sponges (92–100%) is at the high

end of the range reported for shallow water sponges (56–97; see table 1 in de Goeij et al. 2017). Indirect measurements recently suggested that DOC accounts for 95% of the TOC removal of *G. barretti* (Leys et al. 2018), which is very close to the fluxes presented here. The minimal respiratory carbon demands for all species with a complete O₂, LPOC and DOC dataset were, by far, met (Table 4). Both HMA species show

Table 4. Mass balance based on minimal respiratory demands of four dominant North-Atlantic deep-sea sponge species. Note that mass balances were only based on individual sponges with a complete set of data for: oxygen (O_2) dissolved organic carbon (DOC), bacterio- and phytoplankton organic carbon (LPOC). Mass balances were constructed from initial rates based on linear O_2 , exponential LPOC and biexponential (i.e., 2G-model) DOC removal rates (average \pm SD). Total organic carbon (TOC) removal rates are calculated as the sum of initial LPOC and DOC removal rates.

Sponge species	O_2 (μ mol O_2 g DW_{sponge}^{-1} h^{-1})	TOC (μ mol C g DW _{sponge} ⁻¹ h ⁻¹)	Mass balance $\Delta O_2/\Delta TOC$
V. pourtalesii (n = 4)	$\textbf{3.19} \pm \textbf{1.96}$	10.00 ± 2.44	0.32
G. barretti (n = 3)	1.93 ± 1.09	$\textbf{3.72} \pm \textbf{0.24}$	0.52
G. atlantica $(n = 4)$	$\textbf{4.76} \pm \textbf{1.76}$	6.07 ± 5.55	0.78
A. spinispinosum $(n = 3)$	8.44 ± 1.04	56.07 ± 19.92	0.15

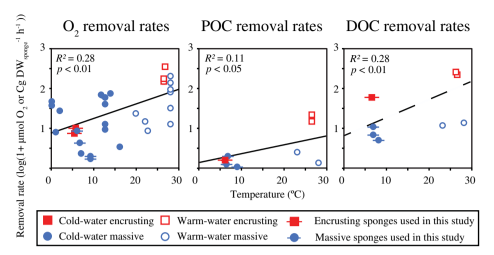


Fig. 4. Oxygen, particulate, and dissolved organic carbon (POC and DOC) removal rates for cold-water ($<15^{\circ}$ C) and warm-water ($>15^{\circ}$ C) marine sponges plotted against temperature. Removal rates are log transformed. Red squares depict encrusting sponges, blue circles depict massive sponges (i.e., all non-encrusting growth forms). R^2 values are based on the linear regression of all values (encrusting + massive). Regression lines are given by log $(1 + \mu \text{mol O}_2 \text{ g DW}_{\text{sponge}}^{-1} \text{ h}^{-1}) = 0.033 * \mathcal{T}(^{\circ}\text{C}) + 0.90$, $(\log(1 + \mu \text{mol POC g DW}_{\text{sponge}}^{-1} \text{ h}^{-1}) = 0.017 * \mathcal{T}(^{\circ}\text{C}) + 0.21$ and $(\log(1 + \mu \text{mol DOC g DW}_{\text{sponge}}^{-1} \text{ h}^{-1}) = 0.041 * \mathcal{T}(^{\circ}\text{C}) + 0.77$. All non-log-transformed rates are given in Table S2.

higher $\Delta O_2/\Delta TOC$ ratios than the two LMA species. These differences might be explained by aerobic microbial processes in HMA sponges, such as nitrification (Hoffmann et al. 2009) or ammonia oxidation (Mohamed et al. 2010), which require O_2 in addition to the O_2 demand based on carbon respiration. Moreover, the organic carbon uptake needed to balance respiration requirements of HMA sponges is potentially further reduced by sponge-associated chemoautotrophs using inorganic carbon sources, which are transferred to the sponge host (van Duyl et al. 2008; Pita et al. 2018; Shih et al. 2020). However, van Duyl et al. (2020) found inorganic carbon uptake to represent only 2–3% of deep-sea sponge carbon budgets.

Bacterio- and phytoplankton contributed only a small fraction (< 10%) to the TOC removal. However, these particulate food sources may contain valuable nutrients, such as vitamins, fatty acids, and amino acids (Pütter 1925; Phillips 1984), which are essential for anabolic processes that require organic carbon, such as growth, cell-turnover, and reproduction. Therefore, a complete carbon budget should include these processes. However, deep-sea sponges most likely grow slowly (Leys and Lauzon 1998), and we assume that within the short (2–8 h) timeframe of our incubations, growth is negligible. For several shallow water encrusting sponges, a rapid cell turnover and the subsequent release of "old" cells as detritus was shown (de Goeij et al. 2009, 2013; Alexander et al. 2014; Rix et al. 2016). This loss of carbon could have a major impact on carbon budgets. In fact, Rix et al. (2016) found that the deepsea encrusting sponge H. coriacea converted 39% of organic carbon derived from deep-sea coral mucus into detritus, and detritus production by deep-sea sponges has been argued to have a major contribution to the total sedimentation rate of the Greenland-Iceland-Norwegian seas (Witte et al. 1997).

However, work by McMurray et al. (2018) suggests that massive sponges that are not space-limited may not show rapid cell turnover and detritus production, and Leys et al. (2018) reported no production of new cells during experiments with *G. barretti*, suggesting minimal cell turnover in the investigated time frame. In conclusion, reports on deep-sea sponge detritus production and cell turnover are contradictive and still very limited, which does not warrant generalizations at this point. Likewise, only limited data is available on the reproduction of deep-sea sponges (Spetland et al. 2007) as well as seasonal changes in metabolic rates (Morley et al. 2016).

Interpretation of sponge metabolic rates at organism and ecosystem scale

Our understanding and interpretation of metabolic rates at both organism and ecosystem scale is currently hampered by two issues. Firstly, the use of a multitude of sponge biomass metrics (e.g., m², cm³, wet weight, dry weight, ash-free dry weight) in combination with a lack of conversion factors makes it almost impossible to compare metabolic rates between different sponge species, and to upscale fluxes from organism to ecosystem level. The use of a specific metric may depend on the context and the research question at hand. For example, when extrapolating individual fluxes to the ecosystem level, planar surface area is potentially the most practical standardization metric in use (read: fast and low-cost). Abundance data in deep sea, but also in shallow-water, habitats are usually collected via 2D video surveys or photo quadrants using ROVs (van Soest et al. 2007; Roberts et al. 2009). However, 2D planar surface area severely underestimates the volume and (organic) biomass of erect vs. flat organisms (e.g., massive vs. encrusting sponges; see also discussion in de Goeij et al. 2017). Arguably the best metric to standardize metabolic rate is organic biomass (i.e., ash-free dry weight) or organic carbon content, excluding ecologically inert hard constituents, such as silica spicules (Rützler 1978). However, an increase in inorganic spicule content requires additional energetic costs at the expense of organic material (McDonald et al. 2002; Carballo et al. 2006). Therefore, in ecological terms, volume, wet weight and dry weight provide alternatives. Volume and wet weight are compromised by effects of large variations in shape, form and tissue densities and compositions of sponges (Diaz and Rützler 2001). We therefore use dry weight here as comparative measure and suggest that future physiological studies on sponges best provide a combination of metrics and/or conversion factors between these metrics.

Secondly, the extrapolation of findings from ex situ studies to in situ conditions is challenging for a number of reasons. Ex situ experiments are usually performed on smaller sponges (~0.1 kg wet weight in this study), while, for example, G. barretti specimens of 24 kg wet weight have been found in situ (Klitgaard and Tendal 2004). Sponge metabolism per mass unit has been found to decreases with increasing mass (Frost 1980; Morganti et al. 2019). Although we did not find a significant size effect on O2 consumption (Fig. S4), both size and temperature are important factors to consider before fluxes can be upscaled to the ecosystem level. Furthermore, we incubated four of our six sponge species in surface water. These surface waters may contain (1) higher concentrations of plankton and DOC compared to in situ conditions and (2) a higher fraction of labile DOC that is bioavailable to metabolize by organisms. Both concentration and lability of DOC are known to potentially effect sponge carbon fluxes (Morganti et al. 2017; Wooster et al. 2019). However, for all tested deepsea sponges, regardless of differences in initial ambient plankton and DOC concentrations, DOC proved essential to meet, and exceed, their minimal respiratory demands. Additionally, the species that was incubated in deep-sea water (i.e., fjord water from 200 m), G. barretti, required a similarly high relative amount of DOC to meet its minimal respiratory demands as the other three tested sponges. Moreover, sponge holobionts might be capable of assimilating recalcitrant DOM. It is known that many sponges, including several LMA species, host specific Chloroflexi lineages (Schmitt et al. 2011; Radax et al. 2012; Thomas et al. 2016). Recently, it was suggested that a specific Chloroflexi lineage found in Geodia spp., SAR202, might oxidize recalcitrant organic matter (Landry et al. 2017) by degrading complex carbohydrates and aromatic compounds (Thrash et al. 2017; Colatriano et al. 2018). In addition, several other studies have now shown an important role of sponge cells in the assimilation of DOM (de Goeij et al. 2009; Achlatis et al. 2019; Rix et al. 2020).

In conclusion, in order to understand deep-sea spongemediated ecosystem processes, it is crucial that future studies investigate in situ organic matter fluxes, and the means or strategies by which various sponge hosts and their microbial symbionts metabolize and cycle DOM.

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Conflict of Interest

None declared.

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