Depth-dependent detritus production in the sponge, *Halisarca caerulea*

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Abstract

Sponges are an important ecological and functional components of coral reef ecosystems. Recently, a new hypothesis about the functional ecology of sponges in organic matter recycling pathways, the sponge-loop hypothesis, in which dissolved and particulate organic matter is taken up by sponges and shunted to higher trophic levels as detritus, has been proposed and demonstrated for shallow (< 30 m) cryptic species. However, support for this hypothesis at mesophotic depths (≈ 30–150 m) is lacking. Here, we examined detritus production, a prerequisite of the sponge loop pathway, in a reciprocal transplant experiment, using *Halisarca caerulea* from water depths of 10 and 50 m. Detritus production was significantly lower in mesophotic sponges compared to shallow samples of *H. caerulea*. Additionally, detritus production rates in transplanted sponges moved in the direction of rates observed for resident conspecifics. The microbiome of these sponge populations was also significantly different between shallow and mesophotic depths, and the microbial communities of the transplanted sponges also shifted in the direction of their new depth in 10 d largely driven by changes in *Oxyphotobacteria*, *Acidimicrobia*, *Nitrososphaeria*, *Nitrospira*, *Deltaproteobacteria*, and *Dadabacteriia*. This occurred in an environment where the availability of both dissolved and particulate trophic resources changed significantly across the shallow to mesophotic depth gradient where these sponge populations were found. These results suggest that changes in sponge detritus production are primarily driven by differential quality and quantity of trophic resources, as well as their utilization by the sponge host, and its microbiome, along the shallow to mesophotic depth gradient.

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coral reefs. Sponges consume large amounts of DOM, both dissolved organic carbon (DOC) and dissolved organic nitrogen (DON) (Yahel et al. 2003; de Goeij et al. 2008a, 2013; Mueller et al. 2014a; McMurray et al. 2016; Hoer et al. 2018), contributing between 56% and 97% of their daily organic carbon intake (see table 1 in de Goeij et al. 2017). Sponge consumption of large amounts of DOM can result in the release of cellular debris, primarily choanocytes, that fuel a “sponge loop” detrital pathway of significant importance to higher trophic levels on coral reefs (de Goeij et al., 2013, 2017; Rix et al. 2016, 2018). We now know that the release of DOC is light dependent (Mueller et al. 2014b), and that consumption of either particulate organic carbon (POC) and/or DOC can produce sponge detritus (Maldonado 2015). One of the uncertainties regarding the sponge-loop hypothesis had been whether the production of detritus occurs in open reef sponges rather than just cryptic species. McMurray et al. (2018) examined the production of detritus in nine species of open reef sponges, both high microbial abundance (HMA) and low microbial abundance (LMA) species, and found that while HMA sponges consumed primarily DOC and LMA sponges consumed primarily POC in the form of picoplankton, none of the species examined produced significant amounts of detritus. However, other studies have shown net detritus production in tropical open reef sponges (Reiswig 1971; Yahel et al. 2003), using a similar methodology. Net detritus production is also not confined to tropical sponges, both cryptic and open reef, but found in temperate (Alexander et al. 2014; Maldonado 2015) and cold-water ecosystems (see table 2 in de Goeij et al. 2017). Another unresolved question related to the sponge-loop hypothesis is whether sponges found at mesophotic depths on coral reefs (~30–150 m), where sponges are a dominant component of these communities (Lesser and Slattery 2018), participate in the sponge-loop pathway (de Goeij et al. 2017; Lesser et al. 2018).

Mesophotic coral ecosystems are deep-reef communities whose community structure and function change with increasing depth based on the availability of light and trophic resources (Lesser et al. 2018). In particular, benthic primary producers, such as corals and macroalgae decrease in abundance with increasing depth due to light limitation, while sponges increase in abundance, measured as either sponge density or percent cover, along a shallow (<30 m) to mesophotic depth gradient in many coral reef ecosystems (Lesser and Slattery 2018). This shift in benthic community composition is likely to also affect the availability of DOM, since benthic primary producers are considered to be an important source of bioavailable DOM on coral reefs (Haas et al. 2011; Mueller et al. 2014b). Furthermore, decreasing light levels reduce DOM release by benthic primary producers (Mueller et al. 2014b, 2016), which supports observations that DOM concentrations decrease with increasing depth (Torrèton et al. 1997; Slattery and Lesser 2015; Mueller et al. 2017; Lesser et al. 2019). In contrast, we also know that POC increases with increasing depth into the mesophotic zone (e.g., Lesser 2006). Given these opposing gradients in DOM and POM across a depth gradient, the relative importance of DOM in sponge nutrition may decrease with depth while POM importance will increase. In this regard, Slattery and Lesser (2015) showed a shift in the relative DOC:POC composition of the diet of Agelas confitena (=A. tubulata). Sponges on the shallow reef (23 m) consumed primarily DOM to meet their carbon demand (97.4% ± 0.4% [SE] DOC vs. 2.6% ± 0.6 [SE] POC), whereas sponges at the upper mesophotic depth of 46 m consumed DOM and POC in similar amounts (43.9% ± 8.0% [SE] DOC vs. 56.1% ± 4.6 [SE] POC). However, no measurements of detritus production were made on these sponges. Given the lower C : N ratio of POM compared with DOM, one hypothesis is that the balance between new sponge growth, which includes the production of choanocyte chambers, and cell turnover might be shifted in favor of new sponge growth at deeper depths, resulting in mesophotic sponges producing less detritus (Hadas et al. 2009; de Goeij et al. 2017). Simultaneously, knowing that the symbiotic microbes of these sponges play an important role in the consumption and processing of DOM within the holobiont, the composition as well as any functional differences in these microbial communities may shift with food availability along a gradient from shallow to mesophotic depths.

As the production of detritus is a prerequisite for the existence of sponge loop pathway as originally described, we compared the detritus production of the sponge Halisarca caerulea on shallow and mesophotic reefs. We assessed several factors that change across a depth gradient, that might affect detritus production, including food availability and composition, temperature, light, and changes in the symbiotic microbial communities. We further conducted a reciprocal transplant experiment to compare the detritus production of H. caerulea from shallow (10 m) and upper mesophotic (50 m) depths. Additionally, we quantified the availability of potential food sources (DOM and POM) along a depth gradient from 0 to 61 m, and the quality (C : N ratio) of those resources specifically at 10 and 50 m. Bulk stable isotope analysis of both sponge tissue and detritus was performed to elucidate the primary food sources for individual sponges. Lastly, the microbiomes of shallow and mesophotic sponges were compared to assess potential differences in their contribution to sponge nutrition.

Materials and methods

Site characterization

Field work was conducted in November 2017 at the Caribbean Research and Management of Biodiversity (CARMABI) marine field station, and sponge samples were collected from shallow (~10 m) and mesophotic (~50 m) depths near Buoy 1 (12°07'28.65"N, 68°58'23.23"W), a fringing reef on the leeward side of Curaçao. The study site is characterized by an
~ 100 m wide sandy reef terrace with patchy coral communities, gradually sloping toward a drop-off at 10–12 m depth (van Duyl 1985). The fore-reef slopes at an angle of ~30–40° until a sand flat is reached at a depth of 50–60 m. Temperature (°C) and downwelling irradiance ($E_d$) of photosynthetically active radiation were taken every minute over several days using HOBO Pendant Temperature/Light data loggers at each site ($n = 3$ for each depth). Measurements of irradiance were in lumens m$^{-2}$, and only used to express the percentage difference in downwelling irradiance ($E_d$) between sites. Depth profiles of photosynthetically active radiation (PAR: 400–700 nm) irradiance ($E_d$) with depth were taken using a single channel PAR recorder (RBRsoloPAR; RBR) fitted with a calibrated LiCor cosine-corrected, planar, sensor (LI 192SA).

**Detritus production by *H. caerulea***

*H. caerulea* is an encrusting LMA sponge found in both cryptic (i.e., reef interstices, reef depressions, and caves) and open reef environments and has previously been shown to be involved in the sponge-loop detrital pathway on shallow coral reefs (de Goeij et al. 2013). Individual *H. caerulea* were sampled for a transplant experiment from the shallow reef terrace (10–12 m depth; $n = 7–9$ replicates) and the mesophotic fore-reef slope (45–50 m depth; $n = 7–9$ replicates) at the Buoy 1 station. Sponges originating from the shallow reef terrace are hereafter referred to as “shallow,” whereas those sampled at the mesophotic fore-reef slope are referred to as “deep.” Each individual sponge was collected from the reef framework using a hammer and chisel and transported to the experimental location at 10 m. Subsequently, the attached coral rock substrate was cleared of epibionts and the sponge specimens were trimmed to an average size ($\pm$ SD) of 9.3 $\pm$ 3.5 cm$^2$ (de Goeij et al. 2008b). Six identical experimental platforms were constructed from PVC pipes; three were installed at 10 m depth, and three platforms were installed at 50 m depth (Fig. S1A, B). The platforms were placed ~ 2 m apart from each other along the 10 and 50 m iso-baths, respectively, and were designed to suspend sponges, facing downward, above a glass funnel (diameter: 7.5 cm) resting in a 15 mL collection tube (Fig. S1A). This positioning facilitated the collection of detritus while preventing the sponges from becoming smothered in sediment and/or self-produced detritus. The trimmed *H. caerulea* specimens were secured in harnesses made of cable ties and fishing line, and were hung above an individual funnel/collection tube (Fig. S1A). Sponges were suspended ~ 0.5 cm above the funnels to maximize the probability of catching released detritus, while still allowing sufficient water exchange to ensure food supply for the sponges. Each platform held five experimental samples (= sponges) and two similar sized pieces of sun-dried coral rock serving as controls for the experimental sponge detrital production (Fig. S1B).

Following a reciprocal transplant design, sponge treatments were designated as deep to deep (DD) controls, shallow to deep (SD) transplants, deep to shallow (DS) transplants, and shallow to shallow (SS) controls. To simulate reduced light conditions experienced by *H. caerulea* growing in cavities and overhangs and/or at mesophotic depths, green PVC roofing sheets were installed on top of the three shallow platforms (Fig. S1A). Placement of the green roofing sheets reduced the light intensity by > 90% to irradiances more typically experienced in cavities and overhangs (Scheffers et al. 2010; Alexander et al. 2015). Transparent roofing sheets were installed at mesophotic depths to simulate any alterations in hydrodynamic conditions experienced by shallow experimental sponges, and to maintain irradiances previously observed at these mesophotic depths (Lesser et al. 2018). Sponges were left to acclimatize for 1 week above glass funnels to ensure full recovery from collection and trimming, transplantation, and adjustment to new conditions (Alexander et al. 2015). All work carried out on the shallow reef terrace was conducted using SCUBA, whereas closed circuit rebreather diving was used at mesophotic depths. Collections and experiments were performed under the research permit (#2012/48584) issued by the Curaçaoan Ministry of Health, Environment and Nature (GMN) to the CARMABI foundation.

On 13 November 2017, all platforms including glass funnels, coral rock controls, and experimental sponges that had been acclimatized after handling were cleaned thoroughly with soft brushes to minimize contamination with detritus accumulated over the previous week. New collection tubes were placed underneath the funnels, and sponges were left undisturbed for an initial collection period of 24 h. Divers collected the individual samples of experimental sponge/rock control carefully by tapping each funnel which promoted all detritus on the glass surface to fall into the funnel below. Subsequently, the experimental sponge/rock control was held with one hand away from the funnel and with the other hand the glass funnel was delicately tapped and turned in circles to transfer detritus adhering to the glass into the collection tube. Once all detritus from the funnel was in the collection tube, the funnel was slowly removed and the collection tube was sealed with a screw cap. After these collections were concluded, platforms were cleaned, and new tubes were placed as described above. Collection tubes with detritus were transported to the nearby CARMABI research station and processed within 3 h. This initial collection of detritus was found to be compromised by divers kicking sediment into the funnels. Subsequently, a second collection period of 48 h was initiated and detritus was collected for both experimental and control samples and processed as described above. Following the acclimatization period and the detrital sample collection periods, all sponge specimens were removed from the transplant platforms and placed into individual plastic collection bags for transport to CARMABI Research Station for processing. Rates of detritus production were based on a 48 h collection after the sponges had acclimatized.

In the laboratory, collection tubes were centrifuged (7 min, rcf = 3825 × g, 6000 rpm) until detritus formed a solid pellet at the bottom of the tubes. Overlying seawater was decanted and
the detrital pellet was transferred into a pre-weighed 1.5 mL microcentrifuge tube. Ultrapure water was added and the pellet was resuspended to desalinate the sample. Subsequently, sample tubes were centrifuged (7 min, rcf = 14,000 × g, 12,000 rpm) and overlying water was removed with a transfer pipette. Subsequently, samples tubes were covered with parafilm and freeze-dried for 24 h before weighing. Rates of detritus production were calculated based on surface area normalized total production, minus control samples, and then divided by the number of days the experiment was conducted. Sponge tissue samples were also collected for 16s rRNA metagenetic, and bulk stable isotope analyses. Each individual specimen was dipped in filtered (0.2 um) seawater to remove any particulate material from the tissue surface. Strips of tissue were then excised from the sponge using sterile scalpels, taking care to avoid coral rock or crustose coralline algae found below the sponge tissues. Pieces of tissue for 16s rRNA metagenetic analyses were collected and placed in a DNA preservation buffer (Seutin et al. 1991) and frozen at −20°C. Tissue samples for bulk stable isotope analysis were freeze-dried for 24 h. Samples were transferred on ice to the University of New Hampshire (UNH) for subsequent analyses (see below).

Differences in abiotic factors between 10 and 50 m were assessed using a two-tailed t-test, whereas analyses conducted throughout the entire depth range were analyzed using type II regression and ANOVA. All experimental results were assessed for treatment effects using ANOVA after checking for normality and heteroscedasticity. Any values not meeting these requirements were transformed before analysis. Ratios are a priori not normally distributed and were transformed prior to analysis. Any significant treatment effects were then followed by post hoc pairwise Tukey’s HSD to detect treatment effects.

Environmental analyses

Seawater samples (1 L) from the surface, 10, 15, 30, 46, 50, and 61 m (n = 3 per depth) were collected from the site and returned to the laboratory where they were each immediately filtered onto GF/F filters. Each GF/F filter was placed in a 2 mL cryovial with DNA preservation buffer (Seutin et al. 1991) and stored at −20°C for subsequent DNA extraction. The seawater filtrate was frozen, transported to UNH, and analyzed for dissolved inorganic nitrogen (DIN) consisting of nitrate/nitrite (NOx: μmol L⁻¹) and ammonium (NH₄: μmol L⁻¹) using high throughput colorimetric assays on a Smartchem Analyzer. The relative differences in laboratory duplicates averaged 4.9% for NOx and 7.8% for NH₄, with average Certified Reference Material (CRM) recoveries of 102.6% for NOx and 95% for NH₄. These same samples were analyzed for DOC (μmol C L⁻¹) and DON (μmol N L⁻¹) as total dissolved nitrogen (TDN) minus DIN, using high-temperature catalytic oxidation and high-temperature oxidation with chemiluminescent detection, respectively, at the UNH Water Quality Analysis Laboratory. The relative differences in laboratory duplicates averaged 4.9% for DOC and 7.8% for TDN, with average CRM recoveries of 97% for DOC and 102.1% for TDN.

For live POC, we analyzed aliquots (3 mL) of seawater from samples (100–200 mL) collected at the surface, 10, 15, 30, 46, and 61 m (n = 3 per depth). Acid washed and deionized water-rinsed 100 mL syringes were filled by divers ~1–2 m from the benthos, regardless of substrate angle. These samples were returned to the laboratory and fixed at a final concentration of 0.5% electron microscopy grade paraformaldehyde in filtered (0.2 μm) seawater, and frozen at ~80°C. Samples were shipped frozen and then maintained in liquid nitrogen until analysis by flow cytometry (Lesser et al. 1992; Lesser 2006). Each sample was analyzed for cell abundances using a Becton Dickinson FACScan flow cytometer equipped with a 15 mW, 488 nm, air-cooled Argon ion laser. Simultaneous measurements of forward light scatter (FSC, relative size), 90° light scatter (SSC), chlorophyll fluorescence (> 650 nm), and phycoerythrin fluorescence (560–590 nm) were made on all samples as described previously (Lesser et al. 1992; Lesser 2006). Differentiation of cyanobacteria from prochlorophytes was based on the presence of phycoerythrin fluorescence. The volume of sample analyzed by the FACScan was determined gravimetrically, where the difference in mg was proportional to the volume of sample analyzed in mL. The abundance of heterotrophic bacteria was determined using SYBRGreen (Molecular Probes), a dsDNA specific dye, which stains all prokaryotes and picoeukaryotes (emission fluorescence: 515–525 nm). Subtraction of photosynthetic prokaryotes from the total prokaryotes provided the concentration of the heterotrophic bacterial component of the community, including numerous Archaea. Gating populations, based on size, allowed for the separation and enumeration of the picoeukaryotic fraction. All flow cytometry measurements included the analysis of fluorescent-calibrated beads of varying sizes to calculate the equivalent spherical diameter (ESD), and biovolume for cell populations, as needed. All enumerated cells were converted to carbon and nitrogen equivalents using the following conversions; Synechococcus: 470 fg C cell⁻¹ (Campbell et al. 1994), Prochlorococcus: 53 fg C cell⁻¹ (Morel et al. 1993), and heterotrophic bacteria: 20 fg C cell⁻¹ (Ducklow et al. 1993); and Synechococcus: 35 fg N cell⁻¹ (Bertilsson et al. 2003), Prochlorococcus: 9.4 fg N cell⁻¹ (Bertilsson et al. 2003), and heterotrophic bacteria: 3.3 fg N cell⁻¹ (FaggerBakke et al. 1996). For eukaryotic phytoplankton, the carbon and nitrogen contents were not calculated as the volume measurements required for C and N conversions were not available. However, the number of eukaryotic phytoplankton available in the plankton, and often consumed by sponges, was low (Lesser 2006).

Stable isotope analyses

Samples from each control and for all experimental sponges, and the detritus they produced (as described above), were sent to the Stable Isotope Laboratory at the Marine Biological Laboratory, and acidified using 1 N HCl to remove any inorganic carbon (sensu Jaschinski et al. 2008) for the bulk...
analysis of particulate C and N, as well as the ratio of the natural abundance of the stable isotopes δ15N and δ13C. Samples were analyzed using a Europa ANCA-SL elemental analyzer-gas chromatograph attached to a continuous-flow Europa 20-20 gas source stable isotope ratio mass spectrometer. The carbon isotope results were reported relative to Vienna Pee Dee Belthemite, and nitrogen isotope results were reported relative to atmospheric air; both were expressed using the delta (δ) notation in units per mil (%o). The analytical precision of the instrument was ±0.1‰, and the mean precision of sample replicates for δ13C was ±0.4 ‰, and for δ15N was ±0.2 ‰. The δ13C and δ15N tissue data were also analyzed in a mixing model using the R package SIAR as described by van Duyl et al. (2018) to determine the proportional contribution of algal vs. coral DOM, as well as POM and detritus for sponges in each treatment. We used data from different sites in the Caribbean that are consistent with the known variation in these values. These inputs into the model included δ13C and δ15N values for algal DOM (δ13C = -15.42 ‰; δ15N 0.80, average of Dictyota sp. and Lobophora sp., table 3 in van Duyl et al. 2018), and based on the assumption that autotrophic algal DOM is a reflection of isotopic value of its source, coral DOM (δ13C = -17.19 ‰; δ15N 0.26, average of all coral species table 2 in van Duyl et al. 2011), and based on the assumption that autotrophic coral DOM is a reflection of the isotopic value of its source, POM (δ13C =-24.91 ‰; δ15N 5.62, van Duyl et al. 2018) and detritus (δ13C = -19.54 ‰; δ15N 2.56, from 10 and 50 m controls [no sponges]). A trophic enrichment factor of 0.5 ± 0.5% (SD) for δ13C and 3.0 ± 0.5% (SD) for δ15N was used (van Duyl et al. 2018).

DNA extraction and metagenetics

DNA was extracted from GF/F-filtered seawater and the preserved tissue and detrital pellets using the DNaseasy PowerSoil DNA isolation kit (Qiagen), using manufacturer’s instructions with modifications to cell lysis as described by Sunagawa et al. (2010), followed by two rounds of 2 min bead beating using a QiaGen QuickLyser set at 50 MHz. All microbial gDNA was checked for quality and concentration using a Nanodrop spectrophotometer 2000c.

Microbial samples were amplified using PCR amplify the COI and CR1 gene regions targeting the universal bacterial/archaeal 16S rRNA gene. Samples were amplified with new degenerate primers targeting the hypervariable region V3–V4, consisting of the forward primer 515F (5′-GTGYCAGCMGCCGCGGTAA; Parada et al. 2016) and the reverse primer 806R (5′-GGACTACNVGGGTWTCTAAT; Apprill et al. 2015). Fluidigm linker sequences CS1 (5′-ACACTGACGACATGGTTCTACA) and CS2 (5′-TACGACTGAGACATTGTTCTCA) were added to the 5′ end of both forward and reverse primers to facilitate Illumina MiSeq. The 16S rRNA gene PCR consisted of a 25 μL reaction with 12.5 μL Amplitaq Gold 360 Master Mix (Applied Biosystems), 1.0 μL GC-enhancer, 0.5 μL 515F (10 μmol L−1) and 0.5 μL 806R (10 μmol L−1), 2.0 μL of DNA template (40–60 ng), and 8.5 μL nuclease free water (Integrated DNA Technologies). Reactions were performed using the following protocol: initial denaturation for 10 min at 95°C, 30 cycles of 95°C for 45 s, 50°C for 60 s, and 72°C for 90 s, followed by a 10 min extension at 72°C. The PCR products were then electrophoresed on a 1% agarose gel.

Sequencing and bioinformatics

The 16S PCR amplicons containing Fluidigm linkers were sequenced on an Illumina MiSeq System using the Mid-Output reagent kit (2 x 150 bp reads, ~20–40,000 reads per sample) at the University of Illinois at Chicago Research Resources Center’s Sequencing Core. Amplicon sequence variants (ASVs) were inferred and tabulated across samples using DADA2 (Callahan et al. 2016). Briefly, raw reads were trimmed of the initial 20 bp to remove residual primer and then truncated beyond the first instance of quality scores below 3 (truncQ = 2). The maximum expected error during denoising was 2 (maxEE = 2). The error model was built from the first 100 M bases and inspected using “plotErrors.” Denoised reads were then merged and chimeric contigs discarded using mergePairs. Taxonomic ranks were assigned to the inferred ASVs using the SILVA ribosomal reference database release 128, using the DADA2 function “assignTaxonomy.”

Analyses of the sponge, detrital, and seawater microbial communities were accomplished using PhyloSeq functions in R (McMurdie and Holmes 2013). The ASV count table was first filtered to discard samples with fewer than 10,000 counts, and then filtered to include ASVs detected in more than 2 samples, and accounting for at least 10 occurrences across samples. Samples were then rarefied to normalize for sequencing effort. Alpha diversities were estimated using Shannon and Chao1 richness indices. Samples were ordinated based on Bray-Curtis distance using nonmetric multidimensional scaling (method = NMDS).

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Metagenomic functional abundances were predicted from 16S abundances using PICRUSt2 v2.1.0-b (https://github.com/picrust/picrust2/wiki). Briefly, 16S ASVs inferred by DADA2 were aligned with HMMER (Eddy 2008) and placed in the provided reference tree using EPA-ng and GAPP (Barbera et al. 2019). Gene family copy numbers were predicted for 16S as well as KEGG functions (i.e., EC and KO accessions) using Hidden State Prediction (castor; Louca and Doebeli 2018) based on
16S ASV abundances and phylogenetic proximity to reference taxa with available genomes. Abundances of biological pathways encoded by microbiomes were then predicted using MinPath (Ye and Doak 2009). Abundances of KEGG accessions corresponding to metabolic processes of interest were tested for treatment effects using ANOVA (Table S2).

Results

Environmental variability along the shallow to mesophotic depth gradient

Environmental differences, where they existed, between experimental sites were assumed to reflect their depth specific variability. At 10 m, the average temperature during the deployment was 28.6°C ± 0.9°C (mean ± SD) and at 50 m it was 28.1°C ± 0.9°C, which was not significantly different between depths (two-tailed t-test, t(14538) = 0.412, p = 0.680). From the profile data ambient $E_d$ at 10 m was $\sim 740 \mu$mol quanta m$^{-2}$ s$^{-1}$, and at 50 m it was $\sim 72 \mu$mol quanta m$^{-2}$ s$^{-1}$. The complete profiles of irradiance with depth can be seen in Lesser et al. (2018). Using a 90% and 10% decrease due to the green and transparent roofing sheets, the sponges were exposed to $\sim 74 \mu$mol quanta m$^{-2}$ s$^{-1}$ at 10 m, and $\sim 65 \mu$mol quanta m$^{-2}$ s$^{-1}$ at 50 m water depth. However, sponges at both depths were hanging upside down and receiving primarily upwelling irradiances ($E_u$) reflected off the sandy bottom. Given that the reflectance of these substrates can be affected by sediment type, presence of microphytobenthic organisms, and organic content, the reflectance of $E_u$ can be anywhere from 30% to 55% across the visible spectrum (Louchard et al. 2003). Thus, we applied a 40% reduction in $E_u$ to obtain values of $E_d$, or upwelling irradiance, as an estimate of what the experimental sponges were seeing during the acclimatization and experimental periods. This resulted in estimated $E_d$ irradiances of $\sim 44 \mu$mol quanta m$^{-2}$ s$^{-1}$ at 10 m and $39 \mu$mol quanta m$^{-2}$ s$^{-1}$ at 50 m. The combined nitrate and nitrite (NO$_x$) for 10 m was $1.35 \mu$mol L$^{-1}$ ± 0.03 (mean ± SD), and at 50 m it was $1.48 \mu$mol L$^{-1}$ ± 0.09. (mean ± SD). These changes were not significantly different between depths (two-tailed t-test, t(4) = 2.56, p = 0.063).

The trophic resources available to sponges varied significantly with increasing depth at the beginning of the experiment. For the dissolved fractions; both DOC (ANOVA: $F_{1,4} = 15.15$, $p = 0.03$) and DON (ANOVA: $F_{1,4} = 14.2$, $p = 0.033$) decreased significantly with increasing depth (Fig. 1A,B). At the end of the experiment, additional measurements of DOC and DON from the experimental sites revealed that DOC concentration at 10 m was $77.44 \mu$mol C L$^{-1}$ ± 1.85 (mean ± SD), and at 50 m it was 68.56 $\mu$mol C L$^{-1}$ ± 0.65 (mean ± SD), which was significantly different between depths (two-tailed t-test, t(4) = 7.83, $p = 0.001$). The DON concentration at 10 m was 5.38 $\mu$mol N L$^{-1}$ ± 0.55 (mean ± SD) and at 50 m it was 3.24 $\mu$mol N L$^{-1}$ ± 0.99, (mean ± SD) which was also significantly different between depths (two-tailed t-test, t(4) = 3.27, p = 0.031). The C : N ratios for DOC:

\[ y=84.41-0.74x, R^2=0.835, p=0.03 \]

\[ y=5.92-0.055x, R^2=0.826, p=0.033 \]

Fig. 1. (A) Water column DOC (mean ± SD) as a function of depth (n = 3 samples per depth) in Curaçao. A significant regression (ANOVA: $F_{1,4} = 15.15$, $p = 0.03$) was observed for DOC as a function of depth; (B) water column DON (mean ± SD) as a function of depth (n = 3 samples per depth) in Curaçao. A significant regression (ANOVA: $F_{1,4} = 14.2$, $p = 0.033$) was observed for DON as a function of depth.

DON averaged $14.39 \pm 0.65$ (SE) at 10 m and $21.17 \pm 4.05$ at 50 m depth and were not significantly different from each other (two-tailed t-test, t(4) = 1.97, p = 0.119).
As was the case for the dissolved fractions, the particulate fractions varied with depth. *Synechococcus* populations were a component of the picoplankton and declined significantly (ANOVA: $F_{1,5} = 22.82$, $p = 0.009$) with increasing depth (Fig. 2A). In contrast prochlorophytes, another component of the picoplankton, increased significantly (ANOVA: $F_{1,5} = 56.32$, $p = 0.002$) (Fig. 2B). Eukaryotic phytoplankton also declined significantly (ANOVA: $F_{1,5} = 17.71$, $p = 0.017$) with increasing depth (Fig. 2C). Heterotrophic bacteria increased significantly (ANOVA: $F_{1,5} = 43.46$, $p = 0.003$) (Fig. 2D) in Curacao as a function of depth.

![Fig. 2](image-url)
5.16, p = 0.0003 (SE) at 10 m, and 5.01 ± 0.498 at 50 m depth, and were not significantly different from each other (two-tailed t-test, t(4) = 308, p = 0.773).

**Detritus production by H. caerulea**

The treatment effects on the corrected (i.e., sponge detritus minus control detritus) rates of detritus production for *H. caerulea* were significant (ANOVA: F[3,20] = 6.54, p = 0.0029). Post hoc analysis revealed that SS produced significantly more detritus (1.358 ± 0.326 [SE] mg cm⁻²) than DD sponges (0.284 ± 0.075 [SE] mg cm⁻²) (Tukey’s HSD, p < 0.05; Fig. 4) while the SD sponges (0.436 ± 0.102 [SE] mg cm⁻²) were not significantly different from DD samples (Tukey’s HSD, p > 0.05) and DS sponges (0.839 ± 0.137 [SE] mg cm⁻²) showed intermediate rates of detritus production that were not significantly different from either SS or DD samples (Tukey’s HSD, p > 0.05; Fig. 4).

**Stable isotope signatures of H. caerulea tissues and detritus**

At the end of the 10 d, experiment treatment effects for the δ¹³C stable isotope values of *H. caerulea* tissue (Fig. 5) were not significantly different (ANOVA: F[3,28] = 0.817, p = 0.495) whereas the δ¹⁵N of the same tissues (Fig. 5) were significantly different (ANOVA: F[3,28] = 4.059, p = 0.016). The post hoc Tukey’s HSD analysis showed that for the tissue, δ¹⁵N SS and DS were significantly lower compared to SD, with DD treatments indistinguishable from all other treatments (Tukey’s HSD, p < 0.05). There was no effect of treatment (ANOVA: F[3,28] = 0.991, p = 0.412) on tissue C : N ratios, which varied from 5.2 to 6.2. For the sponge-produced detritus samples, treatment effects for the δ¹³C stable isotope values (Fig. 5) were significantly different (ANOVA: F[3,24] = 38.06, p < 0.0001) with SS and DS treatments significantly enriched in δ¹³C compared to DD and SD treatments (Tukey’s HSD, p > 0.05), while the δ¹⁵N values of sponge detritus (Fig. 5) were not significantly different from each other (ANOVA: F[3,24] = 1.854, p = 0.164). There was no effect of treatment (ANOVA: F[3,24] = 0.726, p = 0.546) on C : N ratios which ranged from 7.9 to 9.2. The SIAR mixing model, which calculates the percentage of the sponge diet in each treatment that comes from different trophic sources in the water column, showed a significant effect of treatment for the percentage contribution of algal DOM (ANOVA: F[3,28] = 7.478, p = 0.0008) with the only significant (Tukey’s HSD, p < 0.05) pairwise comparison being the SD treatment compared to all other treatments, which were not significantly different (Tukey’s HSD, p > 0.05) from each other (Fig. 6). For coral DOM, there was also a significant effect of treatment (ANOVA: F[3,28] = 14.439, p < 0.0001) with the SS and DS treatment groups consuming significantly more coral DOM (Tukey’s HSD, p > 0.05) compared to DD and SD, which were not significantly different from each other (Fig. 6). For live POM (i.e., plankton), there was a significant effect of treatment (ANOVA: F[3,28] = 14.094,
p < 0.0001) with the SS and DS treatment groups consuming significantly less POM (Tukey’s HSD, p > 0.05) compared to DD and SD, which were not significantly different from each other (Fig. 6). Lastly, there was no significant effect of treatment on the percentage of detrital POM contributing to the diet of experimental sponges (ANOVA: F_{3,28} = 2.043, p = 0.0131) (Fig. 6).

Microbial composition and function of sponge microbiomes

For the microbiome community, a multidimensional analysis of the Beta diversity estimates using Bray–Curtis distance matrices at the class level (Fig. 7A) for the treatment, ambient seawater and detritus samples showed a significant effect of treatment (PERMANOVA: F_{3,24} = 1.98, p = 0.002). All treatment samples were significantly different in their microbial composition from their surrounding seawater and detritus, and seawater and detritus were significantly different from each other (Fig. 7A). Post hoc PERMANOVA on the different treatment groups revealed significant differences between the SD and DD treatments (PERMANOVA: F_{1,12} = 2.38, Bonferroni-corrected p = 0.02), the SD and DS treatments (PERMANOVA: F_{3,24} = 1.98, Bonferroni-corrected p = 0.024), and the SS and DD treatments (PERMANOVA: F_{3,24} = 1.98, Bonferroni-corrected p = 0.018). The composition of the H. caerulea tissue consisted of 54 class level assignments among all treatment groups. Working with those classes that consisted of greater than 1% of the relative abundance (13 classes; Fig. 7B), significant differences were found at the class level (Table S1), where significant post hoc comparisons occurred for the SS vs. DD treatments for the Oxyphotobacteria, which included the genera Cyanobium, Prochlorococcus, and Synechococcus, where the SS treatment had a significantly greater abundance (Tukey’s HSD, p < 0.05; Table S1).
and Fig. 7B). For the DD samples Acidimicrobiia, Nitrososphaeria, Nitrospira, Deltaproteobacteria, and Dadabacteriia were significantly higher (Tukey’s HSD, *p* < 0.05; Table S1) in abundance (Fig. 7B). In addition, for Acidimicrobiia and Nitrospira, the DS vs. DD and SD vs. DD pairwise comparisons were also significantly different (Tukey’s HSD, *p* < 0.05; Table S1).

For *H. caerulea*, those genes involved in carbon metabolism (KEGG pathway 01200) showed no significant effects when all treatments were considered. Subsequently, the analysis was collapsed to analyze only shallow (SS and DS) vs. deep (DD and SD) samples. Genes for carbon fixation (ribulose-1,5-bisphosphate carboxylase/oxygenase [RuBisco], small chain, KO1602) were significantly enriched (ANOVA: *F*₁,₂₆ = 12.8, FDR corrected *p* = 0.014) in shallow vs. deep samples, whereas chitin degradation (chitin disaccharide deacetylase, K03478) was significantly enriched (ANOVA: *F*₁,₂₆ = 18.6, FDR corrected *p* = 0.019) in deep vs. shallow samples. No genes for nitrogen metabolism (KEGG pathway 00910) were significantly different between shallow and deep samples.

**Discussion**

Most of the available substrate on a coral reef is found in the three-dimensional framework of its cryptic spaces and marine caves, estimated to be 75–90% of total reef volume (Ginsburg 1983), and dominated by sponges (e.g., Jackson and Winston 1982; Richter et al. 2001; van Duyl et al. 2006; Slattery et al. 2013). It has been shown that the majority of organic carbon removal by the reef framework, > 95% for both Indo-Pacific and Caribbean regions, is in the form of DOC (de Goeij and van Duyl 2007). Many of the sponges commonly found in, but not restricted to, cryptic environments, including *H. caerulea*, have been shown to utilize DOC (de Goeij et al. 2008a), and they release a significant amount of carbon as detritus (de Goeij et al. 2013). This led to the formulation of the sponge-loop hypothesis, where DOM taken up by sponges is subsequently released as detritus, which then enters the benthic detrital food chain (de Goeij et al. 2013). While the ubiquity of the sponge loop is gaining support in different tropical (i.e., Red Sea) and even deep-sea coral reef habitats (Rix et al. 2016, 2018), some of the early uncertainties, such as the influence of depth and food availability on sponge-loop fluxes, and the difference between cryptic and open reef sponges in detritus production (Slattery and Lesser 2015; Pawlik et al. 2015a, b), remain unanswered. Here, we have shown that for *H. caerulea*, the rate of detritus production at mesophotic depths (i.e., 50 m) was significantly lower than conspecifics at shallow depths (i.e., 10 m). This is significant because it suggests that populations of *H. caerulea* at mesophotic depths may: (1) exhibit depth-specific differences in the microbiome that may influence detritus production due to differences in DOM composition, bioavailability, and utilization (e.g., coral vs. algal-derived DOM) and/or (2) shift energy from cell turnover (= maintenance costs) to increased net biomass...
(= costs of growth) which includes new choanocyte chambers, and potentially less detritus production, because of better quality food (i.e., greater assimilation efficiencies of more abundant, low C : N, POM vs. DOM), especially when combined with increased clearance rates at mesophotic depths.

Detritus production occurs whether POM or DOM is consumed (Maldonado 2015), and tracer studies have shown that algal-derived DOM is incorporated at a higher rate by the microbiome whereas the incorporation of coral-derived DOM occurs at a higher rate by the sponge host (Rix et al. 2017). For the samples in this experiment, the sponge tissue and sponge-derived detritus samples were completely separated from each other based on their bulk stable isotopic values (Fig. 5). Compared to previous work on *H. caerulea* from the same location, the \( \delta^{13}C \) tissue values are similar (van Duyl et al. 2011), while the \( \delta^{13}C \) stable isotope values for sponge tissues from this study are enriched in \( ^{13}C \) by \( \approx 2.5 \) relative to DOM suggesting the broad use of DOM as a food source (de Goeij et al. 2008b), and macroalgal-derived DOM in particular. Additionally, van Duyl et al. (2011) used the same mixing model and estimated that 74.8% of the diet of *H. caerulea* originates from coral-derived mucus in shallow, cryptic environments, from which up to 80% enters the DOM pool (Wild et al. 2004). However, van Duyl et al. (2011) did not include macroalgal-derived DOM sensu stricto (i.e., crustose coralline algae were used as a proxy) in their analysis, which may contribute a significant amount of the available DOM on open coral reefs, and sponges have been shown to utilize it as a food source (van Duyl et al. 2018). The \( \delta^{13}C \) stable isotope values of sponge-derived detritus signatures, however, were enriched in \( ^{13}C \) by \( \approx 2.0-4.0\% \) relative to POM suggesting that the isotopic value of sponge-derived detritus is influenced by the consumption of POM.

Using these SIA values, the isotope mixing model results for *H. caerulea* show that all sponges in these experiments use \( \approx 1.5-2.0 \) times as much algal-derived DOM relative to coral-derived DOM suggesting that the microbiome may have had a greater impact on the assimilation of DOM (Rix et al. 2017). However, while it has been shown that detritus production for algal-derived DOM occurs at higher rates than for coral-derived DOM (Rix et al. 2017), rates of detritus production for mesophotic *H. caerulea* are lower despite a DOM resource primarily from macroalgae, and cyanobacterial mats (Brocke et al. 2015; de Bakker et al. 2017). This could imply that the overall lower concentration of DOM at mesophotic depths caused the decrease in detritus production, or alternatively, shallow-water sponges acquire more algal-derived DOM from internal sources such as the translocation of DOM from the photosynthetic cyanobacteria in their microbiome. While the irradiiances that these sponges were exposed to are equivalent such that rates of cyanobacterial productivity might not differ, the cyanobacteria in the microbiome of *H. caerulea* are distinct from the water column (Shannon–Weaver alpha-diversity index, PERMANOVA: \( F_{1,30} = 7.56, p = 0.001 \), with sponge cyanobacteria comprised of seven genera while seawater from both 10 and 50 m is dominated by two genera (Fig. S2), and are present in greater abundances in SS vs. DD treatments. Taken together, the bulk \( \delta^{13}C \) values of *H. caerulea* tissues and detritus represents a mixed diet of both DOM (derived from macroalgae and coral) and POM, and based on the mixing model of the relative consumption of different trophic resources, POM consumption increases with increasing depth consistent with its increased availability as depth increases (Lesser 2006; Lesser and Slattery 2013; Slattery and Lesser 2015).

Mesophotic sponges have increased amounts of POM available and decreased amounts of DOM compared with their shallow-water counterparts (Figs. 1, 2), but what role would the modest amounts of increased POM have in detritus production for these sponges? The isotopic signatures of the detritus produced by these sponges shows lower \( \delta^{13}C \) and \( \delta^{15}N \) values which likely represent both sponge-derived detritus, as well as water column detrital POM not produced by sponges, given that these sources of detritus could not be separated in these analyses (i.e., controls were subtracted for calculating rates of detritus production but could not be subtracted for other analyses). The colonization of the detrital POM and sponge-derived detritus by bacteria, both bacterioplankton and bacteria from the microbiome, and eukaryote detritivores from the surrounding seawater and from sponge-associated detritivores is also likely to have occurred. The C : N ratios of sponge-derived detritus (8.46 ± 0.33 [SE]) across all treatments were significantly (two-tailed t-test, \( t(58) = 7.62, p < 0.0001 \)) higher than the sponge tissues (5.48 ± 0.23 [SE]), which is consistent with detritus produced during laboratory studies (Rix et al. 2016). This is not that surprising given the presence, albeit in low concentrations for an LMA sponge, of low C : N bacteria in the microbiome and their potential influence on the C : N ratio of the sponge holobiont. However, the confounding signals from these bacteria, detritivores and bacterivores colonizing the detritus make interpreting the detritus isotopic values, relative to the sponge tissue, difficult at best.

Based on the total organic carbon (i.e., both POC and DOC) available at shallow and mesophotic depths, the sponges used in this study do not appear to be limited by the availability of carbon. In addition, the C : N ratios of sponge tissues from both shallow and mesophotic depths, ranging from 5.2 to 6.2, strongly suggest carbon sufficiency. A number of studies, however, have suggested that sponges may be limited by the availability of nitrogen (Hadas et al. 2009; Rix et al. 2016; de Goeij et al. 2017; Lesser et al. 2018). This is because DOM in the open, oligotrophic waters of coral reefs has a high C : N ratio (>10: Ogawa and Tanoue 2003; Tanaka et al. 2011; Rix et al. 2016), such that one would hypothesize that if sponges depend on DOM for a large proportion of their carbon this would result in nitrogen limitation and the need for other sources of nitrogen such as POM (Hadas et al. 2009), or potentially through internal recycling of nitrogen between the sponge and it is microbiome (Hoffmann et al. 2009). In this study, the concentration of DOM at mesophotic depths was
lower than shallow depths, but the C : N ratio of mesophotic DOM (~ 21) was greater than that of shallow DOM (~ 14), initially suggesting a depletion of N with depth. However, only ~ 20% of DOC is bioavailable in shallow reef waters (de Goeij and van Duyl 2007; Lønborg and Álvarez-Salgado 2012; Lønborg et al. 2017) and potentially even lower at mesophotic depths. If we assume 10% bioavailability for the deeper depths, and we also assume the bioavailability of DON tracks DOC, then the C : N ratios for shallow (~ 14) and mesophotic (~ 21) DOM are still sufficient to support metabolism in terms of the availability of carbon. But if 40–70% or more of DOC uptake is used to fulfill respiratory requirements in H. caerulea (de Goeij et al. 2008a; Hadas et al. 2009; Morganti et al. 2017; Hoer et al. 2018), then the remaining carbon, along with the available nitrogen, should also be sufficient to support balanced growth as it results in a lower mean C : N for DOM of ~ 8.64 and 4.32 for shallow sponges at 40% and 70% of carbon respired and ~ 6.35 and 3.18 for upper mesophotic sponges (Table S3). In addition, the live POM C : N ratios are low, and do not differ significantly across the shallow to mesophotic depth gradient (~ 5). As a result, we have no reason to suspect any carbon or nitrogen limitation from shallow to mesophotic depths relative to growth in H. caerulea. Our calculations suggest that had this been one of the ~ 50% of LMA sponges that do not utilize DOM (de Goeij et al. 2017; Morganti et al., 2017), the DOM available to sponges, after satisfying standard metabolic requirements (i.e., 70% of DOC is respired), would have been significantly nitrogen limited in shallow waters (C : N ~ 30.55) compared to mesophotic depths (C : N ~ 14.12). As a result, the POM fraction, while small, is an increasingly important source of nitrogen to offset decreases in DON with increasing depth for growth of the sponge host, as the sponge microbiome is not likely to ever be nitrogen limited due to the abundant in hospite concentrations of DIN.

Most sponges have been found to consume food, both POM and DOM, in amounts proportional to the abundance of these resources (Pile 1997; Lesser 2006; de Goeij et al. 2008a). Despite suggestions to the contrary that sponges are actively selecting different food resources (McMurray et al. 2016), most sponges appear to passively select particles based on the functional morphology of their feeding apparatus and the size-selective constraints imposed by the choanocytes (Hadas et al. 2009; Maldonado et al. 2012). If DOM, as described above, is sufficient to support the energetic and biosynthetic requirements for maintenance metabolism, then the additional low C : N POM can be used for sponge host growth, especially if consumed in proportion to its availability. Sponges increase in percent cover and abundance with increasing depth (Lesser and Slattery 2018), as DOM resources decrease with depth and POM resources increase. This observation has led to suggestions that sponges may be nitrogen limited in regards to the total sponge biomass that can be supported (de Goeij et al. 2017; Lesser et al. 2018). But in this study, the C : N ratio of both DOM and POM (see above) does not support either qualitative, or quantitative, carbon or nitrogen limitation across the shallow to mesophotic gradient. Given this, the primary restriction to increased sponge growth and biomass accumulation at increasing depths would be the amount of DOM and POM consumed to support the growth, biomass, and greater abundance of sponges from shallow to mesophotic depths (Ribes et al. 2003, 2005; Lesser 2006; Lesser and Slattery 2018). Consuming a more stoichiometrically balanced (i.e., lower C : N) diet across the shallow to mesophotic depth gradient is essential, but in order to support the faster growth and greater biomass of mesophotic sponges consuming more food with increasing depth appears to be required. Given that a proportional increase in the uptake of DOM and POM occurs as resources increase (e.g., Morganti et al. 2017), the rate of sponge pumping would also be important. Rates of sponge pumping (i.e., Q values) do increase with increasing depth, at least down to 30 m (Fig. S3), and if true into the mesophotic zone more food would be consumed with increasing depth. When food is qualitatively, or quantitatively, limiting most marine invertebrates conserve maintenance metabolism at the expense of growth, and when food is not limiting, then both higher rates of growth and biomass accumulation can be supported (Fraser and Roger 2007; Sololova et al. 2012). The efficient assimilation of low C : N resources into energetically more expensive biomass production vs. maintenance (= cell turnover and detritus production) might subsequently decrease the amount of detritus produced as depth increases.

Sponges contain diverse microbial communities within their tissues (Hentschel et al. 2002, 2012; Taylor et al. 2007) that are, in part, known to be responsible for the ability of sponges to uptake and process DOM (de Goeij et al. 2008b; Rix et al. 2016, 2017). Additionally, sponge microbiomes can be structured by environmental factors such as light or nutrients that vary with depth (e.g., Morrow et al. 2016). The microbiome communities of H. caerulea are structured by their native depths, the effect of which is not significantly diminished by short-term experimental transplantation. However, while transplant treatments (i.e., SD and DS) do not differ from either SS or DD treatments, they do differ from each other suggesting that microbial composition has begun transitioning toward communities more characteristic of the transplanted depth over a period of 10 d. The microbiome of H. caerulea is dominated by 13 classes of prokaryotes (Fig. S5) for which significant treatment effects were only observed for the Oxyphotobacteria, Nitrososphaeria, Acidimicrobiia, Nitrospira, Deltabacteria, and Deltaproteobacteria (Table S1). Members of the Oxyphotobacteria (= Cyanobacteria) are found in significantly greater abundances in shallow treatments. Photosynthetic cyanobacteria have been reported to be the most abundant group within the microbiome of many sponge species (Steindler et al. 2005; Usher, 2008; Thacker and Freeman 2012), and in previous studies the transfer of carbon rich organic material (e.g., glycerol) from cyanobacterial
symbionts to their sponge hosts has been shown (Wilkinson 1979). Since the productivity potential of cyanobacteria is related to light levels, which are equivalent between the shallow and mesophotic treatments, it is reasonable to assume that shallow populations of H. caerulea are receiving more translocated DOM from their symbionts as a result of their greater abundance in the shallow treatments, as well as what they uptake from their environment, and where they also produce more detritus. However, Nitrososphaeria, Acidimicrobia, Nitrospira, Dadabacteriia, and Deltaproteobacteria were all significantly more abundant in DD treatments. These taxa are involved in ammonia oxidation, organic matter decomposition, heterotrophy/autotrophy/nitrite oxidation, heterotrophy/nitrite reduction, and sulfate/sulfur reduction, respectively. These metabolic capabilities include many steps in the mineralization of organic material, as well as both the nitrogen and sulfur cycles. While a linkage between differences in metabolic capabilities have been suggested for HMA vs. LMA sponges (Weisz et al. 2008), similar differences along environmental gradients, such as shifts in the abundance of specific groups within a common microbiome, have not yet been fully explored. Here, the differential abundance of specific microbial taxa in the host microbiome with changing depth predicts that these sponges can alter their metabolic phenotype to either increased primary productivity (e.g., increased abundance of RuBiso genes in shallow sponge microbiome) or enhanced metabolic capabilities to process organic material (e.g., increased abundance of genes to degrade chitin in mesophotic sponge populations).

We have shown that the sponge H. caerulea produces significantly less detritus at mesophotic depths than conspecifics at shallow depths, and that transplanting these populations initiates immediate changes in detritus production, trophic ecology, and shifts in their microbiome in response to their environment and availability of trophic resources. Sponges increase in abundance with increasing depth into the lower mesophotic (> 60 m) zone (Lesser and Slattery 2018). Simultaneously, DOM continues to decrease while POC increases with increasing depth into the lower mesophotic zone (Lesser et al. 2019), making it essential to understand the differential uptake, assimilation and incorporation of DOM and POM, and the potential that increased growth efficiencies are responsible for decreases in detritus production and increased sponge biomass with increasing depth.

**References**


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Conflict of Interest
None declared.