Evidence of nitrification and denitrification in high and low microbial abundance sponges

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Abstract Aerobic and anaerobic microbial key processes were quantified and compared to microbial numbers and morphological structure in Mediterranean sponges. Direct counts on histological sections stained with DAPI showed that sponges with high microbial abundances (HMA sponges) have a denser morphological structure with a reduced aquiferous system compared to low microbial abundance (LMA) sponges. In Dysidea avara, the LMA sponge, rates of nitrification and denitrification were higher than in the HMA sponge Chondrosia reniformis, while anaerobic ammonium oxidation and sulfate reduction were below detection in both species. This study shows that LMA sponges may host physiologically similar microbes with comparable or even higher metabolic rates than HMA sponges, and that anaerobic processes such as denitrification can be found both in HMA and LMA sponges. A higher concentration of microorganisms in the mesohyl of HMA compared to LMA sponges may indicate a stronger retention of and, hence, a possible benefit from associated microbes.

Introduction

Sponges are evolutionary ancient Metazoa, the first multicellular organisms on the tree of life. They can harbor great amounts and a large variety of microbes (see Taylor et al. 2007 for review). Microscope studies showed that, in some species, 33% of the space in the sponge tissue is occupied by microbes (Vacelet and Donadey 1977). These species have been termed ‘bacteriosponge’ (Reiswig 1981) or ‘high microbial abundance’ (HMA) sponges (Hentschel et al. 2003) and can have as many as $10^8$–$10^{10}$ cells g$^{-1}$ of sponge wet weight, which is 2–4 orders of magnitude higher than the microbial concentration in seawater. The microorganisms in HMA species are sponge-specific and differ from those in the water column in number and nature (Hentschel et al. 2003). In contrast, ‘low microbial abundance’ (LMA) sponges (Hentschel et al. 2003) and can have as many as $10^8$–$10^{10}$ cells g$^{-1}$ of sponge wet weight (Hentschel et al. 2003). Microorganisms can be present in the sponge matrix (mesohyl) between sponge cells, within sponge cells (Vacelet and Donadey 1977) and even within the nucleus (Friedrich et al. 1999; Vacelet 1975).

Molecular studies revealed the presence of a large variety of microbes associated to sponges, which belong to 14 different phyla (Taylor et al. 2007). Sponge-associated microorganisms are very similar across sponge species and within species across latitudinal gradients (Hentschel et al. 2002). Unfortunately, the ever-increasing knowledge of the taxonomic nature of sponge-associated microbes infers little about their metabolic function, activity and benefit to the sponge. Thus, microbial mediated biochemical processes must be uncovered if the nature of the sponge-microbe association is to be understood. To date, several microbial processes have been found in sponges: nitrification (Bayer et al. 2007; Bayer et al. 2008; Corredor et al. 1988; Diaz and Ward 1997; Jiménez and Ribes 2007; Southwell et al. 2008), nitrogen fixation...
(Wilkinson and Fay 1979; Mohamed et al. 2008), photosynthesis (Arillo et al. 1993), methane oxidation (Vacelet et al. 1996) and sulfate reduction (Hoffmann et al. 2005a). Until recently, sponges were assumed to have an exclusively aerobic metabolism, similarly to most invertebrates. Their enormous capacity to filtrate water (e.g. Reiswig 1971) was thought to provide adequate oxygenation across the whole sponge at all times. The discovery of anaerobic microbes in Porifera led to the proposition of anoxic niches within sponges (Webster et al. 2001). When oxygen concentrations were measured with microelectrodes inside sponges, it became evident that sponges that do not ventilate become quickly anoxic. This is the case for Aplysina aerophoba (Hoffmann et al. 2008), Geodia barretti explants (Hoffmann et al. 2005a), Dysidea avara (Schläppy et al. 2007), Chondrosia reniformis (Schläppy et al. in press) and Cliona orientalis (Schönberg et al. 2004). The spatial and/or temporal anoxic microniches within the tissues of those species could result in the activation of the anaerobic microbes present in the sponge. The link between anoxia in sponges and anaerobic microbial processes was made by Hoffmann et al. (2005b) who reported the simultaneous occurrence of anoxic zones in sponge tissues and of sulfate reduction, an anaerobic microbial process, in Geodia barretti, a cold-water sponge.

Vacelet and Donadey (1977) observed that sponges with a dense mesohyl and a reduced aquiferous system host large amount of microorganisms, while species with a looser mesohyl and well developed aquiferous system host comparatively less microorganisms. Recently, Weisz et al. (2008) provided supporting evidence for this original observation and established that HMA sponges are denser (heavier) and have a lower pumping rate as the LMA species. In this study, we investigate whether dense tissue and a high number of associated microbes lead to more diverse microbial processes, including anaerobic ones. We present a new method of quantification, counting microbes directly on tissue sections, to determine both microbial numbers and tissue density. We chose a LMA sponge, Dysidea avara, and a HMA sponge, Chondrosia reniformis. Additionally, we included Aplysina aerophoba for which microbial cell counts using the classical sponge slurry approach were already made (Friedrich et al. 1999) and for which microbial processes have already been described (Bayer et al. 2008; Jiménez and Ribes 2007).

Materials and methods

Microbial counts and quantification of the aquiferous system

The Dysidea avara specimens came from two different locations and dates: (1) Adriatic Sea, Limski Canal, Croatia close to Rovinj, Croatia (44°67.50’N 13°37.0’ Sec.), April 2005 and (2) Northern Adriatic Sea Muntanya de Montgó, Punta del Romani at Cala Illa Mateuca in the township of L’Escala, Girona, Spain (42°06.863’N 03°10.116’E), Northern Mediterranean in March 2006. Aplysina aerophoba and Chondrosia reniformis specimens only came from the Croatian site. Sponges were immediately fixed in 2% formalin after collection, dehydrated in a 30, 50, 70% ethanol series and stored in 70% ethanol. Three tissue blocks were cut from each individual, at different zones of the sponge (apex, middle and basis). Each block contained the continuum from sponge surface to sponge core. After saturation with liquid cryomedium (Jung Tissue Freeze Medium®, Leica Microsystems, Nussloch) for 12 h at 4°C, the blocks were trimmed into 1 x 0.5 cm pieces. They were embedded in base molds with fresh cryomedium and left to harden for 12 h at −80°C. For histological analysis, 5 μm-longitudinal sections were made using a cryostat microtome (HM 505E, Microm, Walldorf, Germany) at −35°C. All sponge sections were mounted on gelatinized glass slides and prepared for microscopic analysis by using 0.2% 4, 6-diamino-2-phenyindole (DAPI, Sigma). Since auto-fluorescence was expected to occur on the sections stained with DAPI and thus give a spurious DAPI signal, we also used a second staining method, fluorescence in situ hybridization (FISH), to confirm that the DAPI signals really represented microbial DNA. FISH was performed with sections of each species using the Cy3-labeled oligonucleotide probe mix EUB I-III (Daims et al. 1999). Formamide concentrations in the hybridization solution and washing buffer were 35% v:v. After rinsing and air-drying, the sections were mounted in Citiflour.

Microscopical analysis was first carried out (at 100× magnification) with a Zeiss Axiophot microscope equipped with Zeiss filters (for DAPI and Cy3) on the entire tissue section to obtain an overview and train visual perception. Color micrographs were taken using a Zeiss AxioImager M1 microscope with an AxioCam MRC camera system. Digital image processing was performed using AxioVision 4.4 software. An ocular with a 122 μm × 122 μm-counting grid and scale bar was used for determining the proportion of different tissue types (e.g. cortex, mesohyl) and for counting of DAPI- or FISH-stained microbial cells (at 1,000× magnification). To determine microbial abundance, thirty visual fields were inspected per specimen on a transect from the sponge’s surface to its core by counting 150 grid-cells per tissue block. With three tissue blocks per specimen, this reveals n = 450 grid-cell counts for determination of microorganism numbers per specimen. A percentage estimation of the aquiferous system (proportion of choanocyte chamber and canals to mesohyl in %) was carried out for each grid and averaged for each species.
Microbial processes in *Dysidea avara* and *Chondrosia reniformis*

Twelve specimens of *D. avara* and *C. reniformis* were collected by SCUBA diving in 10–15 m water depth in Muntanya de Montgó, Punta del Romani at Cala Illa Mateua in the township of L’Escala, Girona, Spain (42°06.863’N 03°10.116’E). The specimens were allowed to acclimatize for 1 day before the start of the experiments in a 20-l aquarium with freshly sampled seawater, an air stone and re-circulating water at 15°C. Prior to starting the denitrification and anammox incubation experiments, three specimens of each species were cut into explants of 0.2–0.5 g⁻¹ and used directly for the experiment. All specimens looked healthy at the start of the experiments.

Nitrification

The presence of nitrification (the process of NH₄⁺ transformation to NO₂⁻ and subsequently to NO₃⁻) was tested by conducting an incubation experiment in which three specimens of *C. reniformis* and three of *D. avara* were kept in 1-l beakers with fresh marine seawater, with an air stone and a magnetic stirrer. An additional three beakers were set up identically but without sponges as controls. The water was kept at constant temperature and in the dark. Additional ammonium (10 μM) was added to each beaker. Water aliquots (10 ml) were taken over the course of 24 h and immediately frozen at −20°C until the start of the analyses. Ammonium was processed in an auto-analyzer while nitrite and nitrate were determined with a NO–NO₂–NO₃ analyzer (Thermo Environmental Instruments Inc., USA). Aliquots of 200 μl were injected in a glass chamber that contained a reduction solution (nitrite: 7.5 g Na I mixed with 2.5 g H₃PO₄ 99%; NO₃: 0.1 M Va (III) Cl). The reduction solutions were made anew for each day of measurement. After passing through 2 M NaOH, the NO gas was freed from the sample, and NO₃ was measured by passing through the ozonator. The integrated area under the curve (using the ECW 2000 Integration Package) and not the peak was used to calculate the NO₃ and NO₂⁻ content of the samples. Nitrates were calculated: NO₃⁻ − NO₂⁻ = NO₂⁻. The rate of production or consumption was calculated as the average of all three sponge replicates minus the average of the three controls.

Denitrification and anammox

Denitrification occurs through the combination of 2 moles of NO₃⁻ to 1 mol of N₂. Depending on the residual ¹⁴N in the system (either through the seawater or due to sponge microbe nitrification) and when the ¹⁵NO₃⁻ tracer is added, denitrification can produce N₂ of any of the three combinations through random pairing: ¹⁴N¹⁴N, ¹⁴N¹⁵N or ¹⁵N¹⁵N (Nielsen 1992). Anammox (anaerobic ammonium oxidation) combines 1 mol of NO₂⁻ with 1 mol of NH₄⁺ ion to N₂. We specifically checked for anammox by using ¹⁵NH₄⁺ and ¹⁴NO₂⁻ as labels. Thus, anammox can yield ¹⁴N¹⁴N or ¹⁴N¹⁵N but not ¹⁵N¹⁵N when the incubation is done with ¹⁴NO₂⁻ (Thamdrup and Dalsgaard 2002). Sponge explants of *C. reniformis* (0.3–0.9 g wet weight) and *D. avara* (0.02–0.4 g wet weight) were incubated in 12.5-ml gas-tight glass vials (Labco Exetainer, Labco Limited, UK) filled with natural seawater spiked with 10 μM ¹⁴NO₃⁻ for the denitrification experiment and with 10 μM ¹⁵NH₄⁺ and 2 μM ¹⁴NO₂⁻ for the anammox experiment. The exetainers were free of air bubbles. We estimated a respiration rate of 9 μmol O₂ cm⁻³ sponge day⁻¹ based on Hoffmann et al. (2005b) and estimated that 90% of the oxygen in the exetainers should be depleted within 24 h (but residual oxygen may have been present). Microbial processes were stopped at 0, 2, 3, 6, 12 and 24 h for denitrification and 0, 1, 3, 6, 12 and 24 h for anammox by injecting 150 μl of saturated HgCl₂ solution in the exetainers. Those were gently shaken for good mixing of the chemical and stored up-side-down (to avoid gas leakage) after a head space of 2 ml of He gas was added to trap the produced N₂. Samples were stored at room temperature until analyzed. Gas chromatography-isotopic ratio mass spectrometry was carried out to determine the isotope ratio (¹⁴N¹⁴N, ¹⁴N¹⁵N, and ¹⁵N¹⁵N) of the N₂ gas trapped in the exetainers’ head space by injecting 250 μl. The concentration of the different isotopic combinations was determined, and the N₂ production rates were calculated from the slope of the N₂ increase over time (Nielsen 1992).

Sulfate reduction

Both sulfide microelectrodes and radioactive isotopes were used in this study to detect sulfate reduction in *D. avara* and *C. reniformis*. The presence of sulfide was tested by the use of a sulfide microsensor with a 20-micron tip and a 1-μM detection limit. Additionally, an incubation experiment of sponge explants was performed with ³⁵SO₄²⁻ using a modified method of the whole core method (Jørgensen 1978). Incubation of sponge pieces was carried out in 5-ml syringes filled with 4.5-ml seawater for 0, 10, 30, and 60 min. The short incubation times were chosen as immediate re-oxidation of the tracer masked sulfate reduction rates in the tissue of other sponges (Hoffmann et al. 2005b). The incubations were performed with air (oxic), with argon (anoxic), and with argon and an extra carbon source that was made of a mixture of acetate, lactate, propionate and butyrate. A volume of 50 μl of a 25-mM stock solution of ³⁵SO₄²⁻ of 600 KBq adjusted to pH 8 with NaOH and was used to inject into the syringe containing the sponge. Sodium sulfate (10 mM) was also
added as carrier for the tracer (Fossing and Jørgensen 1989). The reaction was stopped after 0, 10, 30, and 60 min by adding 20% zinc acetate. Reduced sulfur species were analyzed with the chromium reduction method (Fossing and Jørgensen 1989), with the two-step method.

**Results**

**Microbial counts and prevalence of the aquiferous system**

Visual training allowed the clear distinction between microbial or sponge cells. Microorganisms were visible as $<$2-μm “speckles” at 1,000× magnification and as characteristic FISH signals, while nuclei of the sponge cells were $\sim 5$ μm. Choanocyte chambers were characterized by the arrangement of the choanocyte nuclei lining a lacuna, whereas canals were areas devoid of cells or any other structures (Figs. 1, 2, 3). *Chondrosia reniformis*’ extracellular matrix was heterogeneous and very dense (Table 1) with few visible canals (Fig. 1a). Two microbial niches were found: the collagen and the mesohyl (Fig. 1b). The microbes were more abundant in the mesohyl and exhibited distinct clusters or chains of microorganisms of 4–16 μm in size (Fig. 1c, d), which were absent in the collagen. Thus, microorganism counts were carried out separately for the mesohyl ($2.7 \times 10^{10}$ cells cm$^{-3}$ sponge) and collagen ($0.97 \times 10^{10}$ cells cm$^{-3}$ sponge) and proved to be significantly different (chi-square, $P = 0.04$, $\hat{x} = 0.05$, $df = 1$, with Yate’s correction). Water canals were absent in the collagen and only in low numbers in the choanosome. *Aplysina aerophoba* showed homogenous tissue composition and high tissue density (Fig. 2a, b; Table 1). Sponge cell nuclei were evenly distributed in the mesohyl but less so in the cortex and in inclusions. The mesohyl had canals of various sizes, the choanocyte chambers (4–20 μm across, average 9–10 μm; Fig. 3c) and sponge cells were uniformly scattered over the whole area. Sponge-associated microorganisms were densely packed in the mesohyl (Fig. 2d). *Dysidea avara* had a loose tissue composition (Fig. 3a). The choanosome was composed of choanocytes forming a giant complex of associated chambers (Fig. 3b). The microorganisms were $<$1 μm, had a variety of morphologies and were arranged in clumps (Fig. 3c, d). The aquiferous system was most developed in *D. avara* followed by *A. aerophoba* and finally *C. reniformis*, which had densest body structure (Table 1). When the prevalence of the aquiferous system (‘porosity’) of the sponges was related to the number of microbes in the whole sponges (averages in Table 1), *C. reniformis* and *A. aerophoba* clustered separately but yet in the vicinity of each other.

![Fig. 1](image)

*Fig. 1* *Chondrosia reniformis* cryo-sections a general overview, $\times 3.2$, b mesohyl and collagen overview, DAPI, $\times 100$, c mesohyl detail with choanocyte chamber, DAPI, $\times 1,000$ and d mesohyl detail with sponge-associated microbes and sponge cell nuclei, DAPI, $\times 1,000$. Col collagen, SS sponge surface, Mes mesohyl, CC choanocyte chamber, SC sponge cell nucleus, M microbe
Fig. 2 *Aplysina aerophoba* cryo-sections  
**a** general overview, ×2.5,  
**b** overview of the cortex region and of the mesohyl, DAPI, ×100,  
**c** detail of the mesohyl with sponge-associated microbes, sponge cell nuclei and canals, DAPI, ×1,000 and  
**d** detail of the mesohyl with choanocyte chambers, DAPI, ×1,000.  
Col collagen, SS sponge surface, Mes mesohyl, CC choanocyte chamber, Ca canal, SC sponge cell nucleus, M microbe

Fig. 3 *Dysidea avara* cryo-sections  
**a** general overview, ×3.2,  
**b** choanosome with sedimentary inclusions, DAPI, ×100,  
**c** choanosome detail with choanocyte chamber, DAPI, ×1,000 and  
**d** mesohyl detail with sponge-associated microbes and sponge cell nuclei.  
Col collagen, SS sponge surface, CC choanocyte chamber, SC sponge cell nucleus, M microbe, SI sedimentary inclusion
Table 1  Average aquiferous system in % of the whole sponge and average number of microbes × 10^{10} cm^{-3} sponge + SEM

|                | A. aerophoba | C. reniformis | D. avara |
|----------------|--------------|---------------|----------|
| n              | 3            | 2             | 4        |
| Aquiferous system \% of whole sponge ± SEM  | 21 ± 0.7     | 6 ± 1         | 47 ± 1   |
| Average number of microbes × 10^{10} cm^{-3} sponge ± SEM | 3.3 ± 0.09   | 2.4 ± 0.003   | 0.9 ± 0.05 |

Microbial processes

The microbial processes of nitrification and denitrification were found in both D. avara and C. reniformis (Table 2), while anammox and sulfate reduction were not detected. Ammonium, which fueled nitrification, was produced in both study species at very similar rates (Table 2). At the same time, nitrite was excreted by both sponge species but at very low rates (Table 2 and Fig. 5), which corresponded to only 0.3 and 0.4% of the ammonium excreted. Nitrate was produced in both species (Table 2 and Fig. 5) indicating net nitrification. Nitrate production rates were about ten times higher than nitrite production rates (Table 2) and corresponded to 4.2 and 5.1% of the ammonium excreted in C. reniformis and D. avara, respectively. Anammox being undetectable, a valid way of calculating denitrification rates was to take the slope of the N\(_2\) increase over time of the \(^{15}\text{NO}_3^-\) incubation experiment (Nielsen 1992). Denitrification occurred in both study species and two types of N\(_2\) were produced: \(^{29}\text{N}_2\), resulting from the pairing of \(^{14}\text{N} - ^{15}\text{N}\), and \(^{30}\text{N}_2\), resulting from the pairing of \(^{15}\text{N} - ^{15}\text{N}\) (Fig. 6). The rates of production of \(^{29}\text{N}_2\) and \(^{30}\text{N}_2\) are shown in Table 2. Using random isotope pairing and taking both denitrification and combined nitrification–denitrification into account, the total denitrification rate was 10 and 14.9 nmol N\(_2\) g\(^{-1}\) wet weight sponge h\(^{-1}\) in C. reniformis and D. avara, respectively. The net production of N\(_2\) mostly occurred after 4 h of incubation (Fig. 6).

Discussion

Relation between microbial abundance and aquiferous system

The extracellular matrix of C. reniformis had two clearly visible components: collagen and mesohyl that differed enough in their microbial composition to be analyzed separately. Chondrosia reniformis’ collagen and mesohyl had significantly different amounts of microbes pointing clearly to the existence of different microbial niches within this species. This distinction did not arise in A. aerophoba and D. avara where the whole extracellular matrix was one single microbial habitat. The occurrence of microbes in the collagen of C. reniformis was unexpected, and the presence of these two well-defined sponge environments holds various implications. From a purely methodological point of view, it is clear that the use of ‘slurries’ to assess microbial abundance in sponges may obscure tissue heterogeneity and would have, in the case of C. reniformis, yielded lower estimates. Though we do not know about any possible origin or function of microbes in the collagen fibers, we assume that the different tissue types may harbor different
microbial communities due to the variability in the availability of nutrients, oxygen and space. In *C. reniformis*, we observed differences in the size and the morphotype of microbes in the collagen (0.2 μm, rods, vibrio-shaped) compared to those in the choanosome (rods, 2 μm), which suggests strongly that the two different tissue types have different abilities in the uptake and maintenance of sponge-associated microbes. In view of our results (and using visual training with both FISH and DAPI stains to recognize typical microbes), we feel confident that our method not only yields representative counts of microbes in sponges but also allows the detection of potential microbial niches (i.e. collagen versus mesohyl) within sponge tissues.

Our new method of assessing microbial abundance yields substantially higher counts than the often used ‘slurry’ method. Friedrich et al. (2001) reported microbial abundance of 6.4 ± 4.6 x 10^8 g^-1 sponge tissue in *Aplysina aerophoba*, while we found an average of 3.3 ± 0.09 SEM x 10^10. Further, *D. avara* has been qualified as being ‘virtually free of microorganism symbionts’ in a study done with a transmission electron microscope (Turon et al. 1997) but our results show *D. avara* as having microbial numbers three orders of magnitude higher (10^9 cm^-3 sponge) than seawater (~ 10^6) and are therefore in line with sediments (4 x 10^9 cm^-3, Llobet-Brossa et al. 1998).

The inspection of tissue sections of three Mediterranean sponges, *A. aerophoba*, *C. reniformis* and *D. avara* revealed that high-microbial abundance (HMA) sponges (*A. aerophoba* and *C. reniformis*) have a considerably smaller aquiferous system than low microbial abundance (LMA) sponges and also denser tissues. *D. avara* (a LMA sponge), in contrast, had a highly developed aquiferous system and far fewer microbes than the two other species (Fig. 4a). These findings are consistent with previous observations and support the idea that two different life strategies exist in sponges (e.g. Vacelet and Donadey 1977; Weisz et al. 2008). Our data, moreover, clearly shows that the reason for the high abundance of microbes is not only related to habitat availability (i.e. more mesohyl volume per sponge volume in species where the aquiferous system is reduced) but that the habitat present (spoon mesohyl) is also more densely populated in sponges with a reduced aquiferous system. Within the HMA sponges, however, the relationship is not entirely consistent because *A. aerophoba* had the larger aquiferous system of the two HMA and the most microbes in the mesohyl. Other unknown factors must be playing a role in determining the number of microbes than the sponge tissue density alone.

### Table 2 Microbial processes and rates in *Chondrosia reniformis* and *Dysidea avara*

|                  | NH₄⁺ | NO₂⁻ | NO₃⁻ | Total net nitrification | ²⁹N₂ | ³⁰N₂ | Total net denitrification |
|------------------|------|------|------|-------------------------|------|------|--------------------------|
| *C. reniformis*  | 195.8| 0.8  | 8.0  | 8.8                     | 4.9  | 4.9  | 10                       |
| *D. avara*       | 241.7| 0.8  | 13.9 | 14.7                    | 4.9  | 10   | 14.9                     |

Fig. 5 Incubation of *C. reniformis* and *D. avara* with 10 μM ^15NO₃⁻: production of nitrite and nitrate per cm³ sponge as average water concentrations in three replicate incubations minus concentration in control incubations divided by sponge volume at a given point in time. Nitrite was produced in very low rates and nitrate in higher rates for both sponges, and revealed total nitrification rates of 14.7 and 8.8 nmol cm⁻³ sponge day⁻¹ for *D. avara* and *C. reniformis*, respectively.

Fig. 6 Production of ^14N¹⁵N and ^¹⁵N¹⁵N after addition of 10 μM ^¹⁵NO₃ (with standard error of the mean) indicates denitrification in *C. reniformis* and *D. avara*.
Anoxic conditions are known to occur in both *C. reniformis* (Schla¨ ppy et al. in press) and in (Table 2). The absence of sulfate reduction was surprising. Production were highest in *D. avara* Furthermore, nitrification, denitrification and ammonium while anammox and sulfate reduction were not detected. Assuming 1.2 g wet weight = Original net nitrification rates converted into other units for comparison purposes assuming dry weight = 1 cm−1.

| Target spp. | This study (μmol g−1 wet weight sponge day−1) | Bayer et al. (2007) (μmol g−1 wet weight sponge day−1) | This study (nmol g−1 dry weight h−1) | Jiménez and Ribes (2007) (nmol g−1 dry weight h−1) | Bayer et al. (2008) (nmol g−1 dry weight h−1) |
|-------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|
| *C. reniformis* | 0.176 | 0 | 73.33 | 702 | A. aerophoba 89–344 |
| *D. avara* | 0.294 | ~0.3 | 122.5 | 0 |

The link between anatomy and microbial load has been attributed to the nutritional strategy of the sponge (Vacelet and Donadey 1977). Species with a well developed aquiferous system (Vacelet and Donadey 1977), large choanocyte chambers (Boury-Esnault et al. 1990) and higher pumping rates (Weisz et al. 2008) are thought to rely predominantly on capture of particulate organic matter, while those with a less pronounced aquiferous system are thought to host more microbes and subsequently use them as a food source (“microbial farming”). Our observation of higher microbial densities cm−3 mesohyl in HMA species supports this idea; we have, however, no proof yet if our target sponges also consume their associated microbes.

Microbial processes

Our aim was to investigate whether dense tissue and a high number of associated microbes may lead to more diverse microbial processes and higher rates. In this study, we found nitrification and denitrification rates in both *C. reniformis* (a HMA species) and *D. avara* (a LMA species), while anammox and sulfate reduction were not detected. Furthermore, nitrification, denitrification and ammonium production were highest in *D. avara*, the LMA species (Table 2). The absence of sulfate reduction was surprising. Anoxic conditions are known to occur in both *C. reniformis* (Schläppy et al. in press) and in *D. avara* (Schläppy et al. 2007) and sulfate reducers were found through the use of a specific FISH probe in *C. reniformis* (Schumann-Kindel et al. 1997). One possible explanation is the rapid re-oxidation of the reduced sulfide that has been shown to occur in another sponge species (Hoffmann et al. 2005b). Alternatively, sulfate reduction may be inhibited through the production of nitrite and nitrate by the nitrifiers, which fuel high rates of denitrification (as measured in this study) and out-compete sulfate reduction. Finally, it is possible that some sponges can specifically inhibit sulfate reduction to prevent accumulation of toxic sulfide, the end product of microbial sulfate reduction.

Nitrite and nitrate were produced in both *C. reniformis* and *D. avara* indicating that the nitrifiers were active. The two nitrification steps (NH4− to NO2− to NO3−) were identified and therefore reveal the presence and activity of both ammonium oxidizing and nitrite oxidizing microbes (only very little nitrite was detected). Nitrification has been shown by Bayer et al. (2007) for *D. avara* and *C. reniformis* but denitrification is shown here for the first time in Mediterranean sponges. The only other evidence of denitrification in sponges is in *Geodia barretti*, a cold-water species (Hoffmann et al. 2009), with 92 nmol N cm−3 day, equals 3.8 nmol N cm−3 h−1. Our rates are more than double of those found in *G. barretti*, with 10 and 14.9 nmol N cm−3 h−1.

The total net nitrification rates reported in this study are in the range of those reported by other authors for our target species although they are highly variable (Table 3). Those results do not take into account the anaerobic loss of nitrate through denitrification, hence, they are only indicative of net nitrification and definitely represent an underestimate of total nitrification in these sponges. The presence of denitrification may explain some of the variability in the nitrification results. Our rates of denitrification are very close to the nitrification rate, which indicates that approximately half the nitrate produced by the sponge is re-used for denitrification. The nitrifiers remove only a small portion (about 5%) of the ammonium produced, the rest being presumably expelled through ventilation by the sponge. The rather large ammonium production may not only be a product of sponge respiration but also of fermentation processes in times of anoxia. The fact that both nitrification and denitrification occur simultaneously was apparent in the comparatively high rate of 29N2 produced. It could only stem from unlabeled NO3−, the source of which can either be the seawater in which the sponge were incubated in or, more likely, the nitrification activity of sponge-associated microbes. This co-occurrence of two microbial processes may be similar in the other species of sponges for which nitrification rates have been reported (Southwell et al. 2008). Temporal and spatial occurrence of oxic and anoxic zones within the sponge would favor both the nitrifiers and the denitrifiers, in turn. The occurrence of nitrification and denitrification suggests strongly that the microbial community inside our study species is able to deal with shifts from aerobic to anaerobic conditions inside the sponge. Furthermore, the co-occurrence of nitrification and denitrification might explain the variations in NO3− production seen in some species (Bayer et al. 2008).
We conclude that both HMA and LMA sponges host an active community of microbes with nitrification and denitrification. Total rates of microbial processes in LMA sponge can be even higher than in HMA sponges. The main benefit to the sponge of having such a guild of microbes may reside in the fact that they get rid of ammonium and nitrite (which could be deleterious to the sponge) and that microbial denitrification may out-compete microbial sulfate reduction, which yields toxic sulfide. The sponge, by controlling its pumping activity, has a means of controlling the activity of both the nitrifiers and denitrifiers populations. It is also possible that the presence of autotrophic nitrifiers and heterotrophic denitrifiers provide the sponge with an optimized system for food utilization and storage under both aerobic and anaerobic conditions, in case it actually does feed on the associated microbes (something that is still to be proven). Our results show that both LMA and HMA sponges may host metabolically diverse and active microbial communities, and that anaerobic and aerobic processes can be found in both sponge types. Higher numbers of microbes per volume of mesohyl in HMA sponges may indicate a stronger retention of and, hence, a possible benefit from associated microbes.

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