INTRODUCTION

Dimethylsulfiniopropionate (DMSP) is an algal metabolite that is produced by a wide variety of species at different intracellular concentrations (Keller et al. 1989). Marine dinoflagellates, including the coral symbiotic algae of the genus *Symbiodinium*, produce relatively high levels of DMSP (Keller & Korjeff-Bellows 1996, Yoch 2002, Van Alstyne et al. 2006). DMSP and its enzymatic cleavage products have multiple proposed functions at the cellular level, including those of herbivory deterrent (Wolfe et al. 1997), algal osmolyte (Kirst 1996), antioxidant (Sunda et al. 2002), antiviral defense mechanism (Evans et al. 2006), overflow mechanism for excess reduced sulfur (Stefels 2000), methyl donor (Ishida 1968), sulfur storage compound (van Diggelen et al. 1986), foraging cue (DeBose et al. 2008) and cryoprotectant in polar algae (Kirst et al. 1991, Karsten et al. 1996). Dimethylsulfide (DMS) is a significant degradation product of DMSP, and is a major source of sulfur to the atmosphere (Kettle & Andreae 2000). DMS is also reputed to affect ocean cloud cover and the radiative climate through formation of aerosols (Charlson et al. 1987). The enzyme DMSP lyase (dimethylpropiothetin dethiomethylase, EC 4.4.1.3; DL) is responsible for DMSP conversion, producing DMS and other products (Johnston et al. 2008). This enzyme has been reported in phytoplankton (Steinke et al. 1998), bacteria (de Souza &
Yoch 1995) and fungi (Bacic et al. 1998). As demonstrated by Sunda et al. (2002), DMSP is reactive toward hydroxyl radicals (·OH), but its enzymatic cleavage products, acrylate and DMS, are ~20 and ~60 times more reactive towards ·OH, respectively. Overall, the enzymatic conversion of DMSP forms several potential antioxidant scavengers (Sunda et al. 2002). Though many algal DMSP producers have DL capabilities (DMSP lyase potential activity [DLA]), this is not true for all algal species (Steinke et al. 1996, Niki et al. 2000, van Bergeijk & Stal 2001, Sunda et al. 2002). Since DMSP and DLA potentially mitigate stress, investigations of how these parameters vary across Symbiodinium clades may improve understanding of the symbiont–host relationship.

*Symbiodinium* are the most prominent dinoflagellates in symbioses with marine invertebrates and protists and are commonly found with members of the phyla Cnidaria (i.e. corals, anemones), Platyhelminthes, Mollusca, Porifera and Foraminifera (Trench 1979, Pawlowski et al. 2001). Many symbiont host species have been shown to contain DMSP, including corals (Jones et al. 1994, Hill et al. 1995, Broadbent et al. 2002), anemones (van Alstyne et al. 2006), flatworms (van Bergeijk & Stal 2001) and clams (Jones et al. 1994, Hill et al. 2000, 2004). Evidence suggests that algal symbionts are responsible for DMSP production in cnidarian species due to a positive correlation between DMSP concentration and symbiotic algal densities (cell number) in cnidian hosts, although partitioning between host corals and algal symbionts is unknown (Broadbent et al. 2002, Van Alstyne et al. 2006). Significant amounts of DMSP in reef corals suggest that these ecosystems could be significant sources of DMS to the atmosphere (Broadbent & Jones 2004, Jones & Trevena 2005), but it is currently unknown whether *Symbiodinium* have the enzymatic ability to convert DMSP into DMS. With evidence for DMSP variation in *Symbiodinium* from different coral species and bleached versus healthy corals (Hill et al. 1995, Broadbent et al. 2002, Van Alstyne et al. 2006, Jones et al. 2007), but no reports of DLA in *Symbiodinium* or their coral hosts, a characterization of baseline DMSP and DLA levels within and across algal and animal host species is warranted.

The present study investigated 5 *Symbiodinium microadriaticum* strains to determine (1) whether DLA was detectable in *S. microadriaticum* cultures, (2) whether DLA was primarily associated with the algal fraction of non-axenic cultures and (3) whether *S. microadriaticum* DLA and DMSP were significantly different between the algal strains investigated. *Emiliania huxleyi* strains were analyzed for comparison.

**MATERIALS AND METHODS**

**Algal cultures.** All experiments were conducted with strains of the dinoflagellate *Symbiodinium microadriaticum* and *Emiliania huxleyi*, purchased from the Provasoli-Guillard Center for the Culture of Marine Phytoplankton (CCMP; Bigelow Laboratory, Maine, USA). Algal strains included CCMP 373 and 374 (*E. huxleyi*; axenic) and *S. microadriaticum* CCMP 421, 828, 829, 830 and 1633 (Table 1). The cultures were maintained using sterile techniques and grown in sterile K, L1 or f/2-Si media (pH 8.0) according to preference (Bigelow Laboratory). *S. microadriaticum* cultures were not treated with antibiotics to obtain axenicity as our preliminary trials showed that antibiotic treatment negatively affected growth (data not shown). Thus, measures were taken to assess bacterial abundance and potential interference with DLA assays (detailed in ‘Bacterial analyses’ below). Algal cultures were grown at 26°C with a 12 h light:12 h dark cycle, without agitation (Rogers & Davis 2006). In common with other studies (Matrai & Keller 1994), our preliminary trials demonstrated that DMSP levels were

<table>
<thead>
<tr>
<th>CCMP strain</th>
<th>Species</th>
<th>Collection location (very approximate)</th>
<th>Ocean</th>
<th>Isolated from</th>
<th>Clade</th>
</tr>
</thead>
<tbody>
<tr>
<td>373</td>
<td><em>E. huxleyi</em></td>
<td>Sargasso Sea</td>
<td>North Atlantic</td>
<td>Sea water</td>
<td>–</td>
</tr>
<tr>
<td>374</td>
<td><em>E. huxleyi</em></td>
<td>Gulf of ME, USA</td>
<td>North Atlantic</td>
<td>Sea water</td>
<td>–</td>
</tr>
<tr>
<td>421</td>
<td><em>S. microadriaticum</em></td>
<td>Wellington, NZ</td>
<td>South Pacific</td>
<td>Sea water</td>
<td>E</td>
</tr>
<tr>
<td>828</td>
<td><em>S. microadriaticum</em></td>
<td>Florida Keys, FL, USA</td>
<td>North Atlantic</td>
<td>Sea water</td>
<td>A</td>
</tr>
<tr>
<td>829</td>
<td><em>S. microadriaticum</em></td>
<td>Great Barrier Reef, Australia</td>
<td>South Pacific</td>
<td><em>Tridacna crocea</em> (bivalve)</td>
<td>A</td>
</tr>
<tr>
<td>830</td>
<td><em>S. microadriaticum</em></td>
<td>Bermuda Biological Station, Bermuda</td>
<td>North Atlantic</td>
<td><em>Aiptasia pallida</em> (sea anemone)</td>
<td>B</td>
</tr>
<tr>
<td>1633</td>
<td><em>S. microadriaticum</em></td>
<td>Hawaii, USA</td>
<td>North Pacific</td>
<td><em>Aiptasia puchella</em> (sea anemone)</td>
<td>B</td>
</tr>
</tbody>
</table>
dependent upon the growth phase; therefore, all algal cultures were sampled during their exponential growth phase, at an average density of $1 \times 10^5$ cells ml$^{-1}$. Each strain was grown in semi-continuous batch culture in 50 ml conical flasks with 30 ml of culture in each, under cool-white fluorescent bulbs (100 µE m$^{-2}$ s$^{-1}$). All cultures were sampled 2 h ($\pm$1 h) into their light periods as DMSP (and DLA) concentrations varied with diel cycle in *S. microadriaticum* (authors' pers. obs.) and have been reported to vary in other algal species as well (Bucciarelli et al. 2007, Jones et al. 2007, Stefels et al. 2007, Sunda et al. 2007). *S. microadriaticum* cultures were genetically verified for clade type (see 'Algal analyses'). Each of the strain replicates was grown separately and analyzed individually.

**DMSP and DLA analyses. DMS analysis and calibra-
tion:** All samples were analyzed with a Hewlett-Packard 5890 Series II gas chromatograph (GC) fitted with a Chromosil 330 packed column (Supelco) and a flame photometric detector (FPD). System temperature settings were 150°C for the injector, 60°C for the column oven and 175°C for the detector. Nitrogen gas was the carrier (60 cm$^3$ min$^{-1}$), and air (60 cm$^3$ min$^{-1}$) were the flame gases. Data were collected and analyzed using HP ChemStation (Hewlett-Packard). Quantifications were made by headspace analysis following DMSP conversion to DMS by alkaline hydrolysis. Known concentrations of DMSP (purchased from Research Plus) were diluted in sterile water to give working solutions, which were frozen in small aliquots at –80°C. A standard curve for serial dilutions of DMSP was used to construct a calibration curve (using the square-root values of the peak area), and this linear regression served to convert peak areas from GC headspace measurements to DMS concentrations. Standards and controls (n = 5 each) were prepared in parallel to experimental samples by using the same Tris buffer for DLA, incubating at 30°C for 20 min and using the same total liquid volume (1 ml) in headspace vials. Addition of 5 N NaOH to standards (final volume of 1 ml; final concentration of 5 N NaOH) occurred prior to heating, and all vials were placed in the dark for at least 4 h for equilibration prior to sampling. Analytical replicates were used only for interstrain assessment of repeatability. The precision of the DMS analysis was <5%, and headspace storage trials showed no DMS losses with the analytical methods employed. The detection limit of the GC was 1 nmol DMSP l$^{-1}$.

**Total and particulate DMSP analyses:** Total DMSP was determined using a simple acidification/storage procedure (5 µl 50% H$_2$SO$_4$ ml$^{-1}$ of culture) followed by the total DMSP method (Kiene & Slezak 2006). Appropriate controls (n = 5) were prepared in parallel to sample preparation.

**DLA analyses:** Our DLA methods were modified from procedures described by Harada et al. (2004) and Steinke et al. (2000) and optimized (pH, exogenous DMSP concentration) for *Symbiodinium microadriaticum*. Briefly, DLA was determined by measuring the production rate of DMS prior to and after the addition of exogenous saturating levels of DMSP to permeabilized cells (using a Tris buffer, 200 mmol l$^{-1}$ Tris containing 500 mmol l$^{-1}$ NaCl, pH 8). Algal cell permeabilization is necessary to allow exogenous DMSP into the cells in order to detect an enzymatic response to saturating DMSP levels. Tests showed that DLA increased in samples with Tris buffer compared to those without, and Tris buffer produced higher DLA than other means of cell disruption (homogenization, varied Tris buffer strength). We found that Tris buffer at pH 8 yielded DLA in *S. microadriaticum* samples greater than those in pH 6 Tris buffer. Appropriate controls (biotic and abiotic in parallel to sample preparation) and standards were run in tandem. Controls consisted of 0.5 ml Tris buffer and 0.5 ml culture medium amended with 5 µl of 1 mol l$^{-1}$ DMSP at $t_0$. Spot checks of Tris buffer and culture pH before and after DLA analysis were conducted and never deviated significantly from pH 8 (stable at 7.98; pH above 8 results in increased abiotic conversion [any conversion of DMSP not attributed to biological enzymes]). Headspace vials were 6 ml in volume and sealed with polytetrafluoroethylene (PTFE)/rubber septa (National Scientific).

Algal cells in culture were concentrated by centrifugation at 1310 $\times g$ for 5 min prior to placement in headspace vials. Concentrating the cells was necessary as the amount of DLA in some cultures was too small and needed to be adjusted according to the limitations of our analytical system (Steinke et al. 2000). Centrifugation has also been used previously for concentrating algal cells in DLA and DMSP/DMS experiments (Steinke et al. 1998, Broadbent et al. 2002). We did not use filters (e.g. GF/F) to capture cells because the filters could not be fully submerged in the small liquid volume of the headspace vials. After centrifugation, algal cells were resuspended in their native media to a final volume of 0.5 ml. Tris buffer was added to samples to permeabilize the cells. Preliminary trials in native media versus those in Tris buffer verified that the buffer was necessary for optimal DLA measurement. Tris buffer (0.5 ml) was added to 0.5 ml algal culture, sealed, incubated in a 30°C water bath for 20 min and vortexed vigorously for 3 s before sampling at each time point (at
5 min intervals from 10 min prior to, through 30 min after, exogenous DMSP addition.

The temperature chosen for these experiments (30°C) was previously recommended to enable comparisons among samples and studies (Steinke et al. 2000). At t₀, 5 µl DMSP stock solution (1 mol l⁻¹) was added to give a final concentration of 5 mmol l⁻¹. This amount of DMSP was found to be saturating for lyase-catalyzed DMS production in this system according to preliminary tests. At each time point, 50 µl headspace samples were removed with an Agilent gas-tight syringe (same volume injected for all samples and standards) and injected into the GC for DMS measurement. At least 8 headspace samples (~10 min through 30 min) were measured for each vial to yield a rate of DMS increase with time. DMS production was linear for all samples taken, and DLAs for all samples were corrected for abiotic conversion of DMS by subtraction of DMS production rates measured in control vials. DLA is defined as nanomoles of DMS per minute and is also reported as DLA:chl a (nmol DMS · min⁻¹ · µg⁻¹) and DLA:DMSPₚ (nmol DMS · min⁻¹ · fmol⁻¹ DMSP). On a per cell basis, DMSP is reported as femtomoles per cell.

**Bacterial analyses. DLA in culture sample filtrates:** Algal cultures were sub-sampled prior to centrifugation to determine bacterial contribution to overall DMS production. Culture samples were filtered using Isopore membrane filters (Millipore; 25 mm diameter, 2 µm pore size) and gentle filtration (gravity or <50 mmHg vacuum; Steinke et al. 2000) to obtain an algal-free culture fraction while allowing unattached bacteria into the filtrate. Filtrate (0.5 ml) was immediately placed in a 6 ml headspace vial, 0.5 ml of Tris buffer was added and DLA was assessed as described above. Filtrate samples without algae were scaled up to represent the amount of bacteria present in unfiltered (same volume of whole culture as the filtrate) samples to calculate the maximum bacterial DLA contribution to the reported algal DLA measurements. Specifically, the contribution to total potential DLA by bacteria (bacterial contribution) was calculated as follows for each replicate: (1) filtered bacterial counts were divided by unfiltered bacterial counts to determine the ratio of bacteria in the filtered versus unfiltered culture samples, and (2) filtered DLA quantities were divided by unfiltered DLA quantities at t = 30 min; each DLA measure was corrected for abiotic conversion by subtracting the amount of DMS measured in controls. Bacterial contributions to total observed DLA (%) were therefore calculated by dividing the above DLA ratio (Point 2) by the above bacterial count ratio (Point 1) and multiplying by 100.

**Bacteria enumeration:** Prior to centrifugation, culture samples and culture filtrate sub-samples (0.5 ml each) were preserved (Sherr & Sherr 1993), stained with 4’-6-diamidino-2-phenylindole (DAPI, final concentration 20 µg ml⁻¹) and filtered onto 0.8 µm polycarbonate filters (after diluting for cell density) for bacterial enumeration. Cells were enumerated by counting 10 to 20 bacteria grid field⁻¹ in 30 random fields filter⁻¹ (1000-fold magnification) with an epifluorescence microscope (Kemp et al. 1993). Cell numbers in individual grids were averaged, and the numbers of cells per milliliter of culture was calculated.

**Algal analyses. Symbiodinium microadriaticum** were enumerated by hemocytometer using an epifluorescence microscope prior to and after centrifugation. Ten grid squares were counted for each sample and averaged to calculate the total number of algal cells per milliliter. Algal cell sizes were determined using a microscope, hemocytometer and eyepiece graticule. Chl a concentrations were measured fluorometrically with a Trilogy Laboratory Fluorometer (Turner Designs). Briefly, 1 ml aliquots of unconcentrated culture were filtered through Whatman GF/F glass fiber filters and extracted in 90% acetone for 24 h at 4°C (Parsons et al. 1984). *S. microadriaticum* genetic diversity was assessed using standard RFLP methods to verify *Symbiodinium* clades (Table 1; Rowan & Powers 1991). Algal DNA was extracted using the CTAB/phenol extraction methods as detailed by Coffroth et al. (1992) and Goulet & Coffroth (1997) and was amplified with PCR using a ‘universal’ primer (ss5) and the zooxanthella-biased primer ss3Z (Rowan & Powers 1991). Samples were subsequently digested with *Taql* following the protocol of Goulet & Coffroth (2004) and visualized by ultraviolet light after ethidium bromide staining of the product in a 2% agarose gel.

**Statistical analyses.** Prior to analyses, assumptions of normality and homogeneity were tested and data were transformed as necessary. Regression analyses were used to assess DLA among strains and the relationship between DMS concentration and cell number. Analysis of variance (ANOVA) was used to assess whether DLA:chl a measures differed among strains, filtrates, controls, or over time. ANOVA was also used to assess DMSPₚ and total DMSP (particulate and dissolved; DMSPₚ) values per cell. All statistical analyses were conducted using Minitab V. 10 (Minitab, Ver. 2000), with α = 0.05 for all tests.

**RESULTS**

**Algal DMSP**

For each of the *Symbiodinium microadriaticum* cultures investigated, DMSPₚ closely paralleled DMSPₚ measurements and were not significantly different (p > 0.05; Table 2). *S. microadriaticum* DMSPₚ per cell and DMSPₚ per cell varied according to strain with CCMP...
Table 2. *Symbiodinium microadriaticum* and *Emiliania huxleyi*. Comparison of DMSP<sub>p</sub> and DMSP<sub>t</sub>, (particulate and total dimethylsulfiniopropionate; fmol cell<sup>−1</sup>) and DMSP lyase potential activity (DLA; nmol min<sup>−1</sup> fmol DMSP<sup>−1</sup>; nmol min<sup>−1</sup> µg chl <i>a</i><sup>−1</sup>) in cultured strains of *E. huxleyi* (CCMP 373 and 374) and *S. microadriaticum* (CCMP 421, 828, 829, 830 and 1633). Bacterial DLA in *S. microadriaticum* is also shown. Averages ± SD are presented. Sample number is indicated in parentheses. ND: not detected; -: not tested.

<table>
<thead>
<tr>
<th>CCMP strain</th>
<th>DMSP&lt;sub&gt;p&lt;/sub&gt; (fmol cell&lt;sup&gt;−1&lt;/sup&gt;)</th>
<th>DMSP&lt;sub&gt;t&lt;/sub&gt; (fmol cell&lt;sup&gt;−1&lt;/sup&gt;)</th>
<th>DLA:DMSP&lt;sub&gt;p&lt;/sub&gt; (nmol min&lt;sup&gt;−1&lt;/sup&gt; fmol DMSP&lt;sup&gt;−1&lt;/sup&gt;)</th>
<th>DLA:chl a (nmol min&lt;sup&gt;−1&lt;/sup&gt; µg chl &lt;i&gt;a&lt;/i&gt;&lt;sup&gt;−1&lt;/sup&gt;)</th>
<th>Bacterial DLA (%) of total DLA</th>
</tr>
</thead>
<tbody>
<tr>
<td>373</td>
<td>7.6 ± 2.3 (5)</td>
<td>–</td>
<td>0.1 ± 0.01 (5)</td>
<td>0.2 ± 0.1 (4)</td>
<td>–</td>
</tr>
<tr>
<td>374</td>
<td>4.7 ± 2.1 (6)</td>
<td>–</td>
<td>0.007 ± 0.0002 (2)</td>
<td>0.006 ± 0.005 (2)</td>
<td>–</td>
</tr>
<tr>
<td>421</td>
<td>201.0 ± 138.9 (3)</td>
<td>199.0 ± 151.4 (3)</td>
<td>3.2 ± 2.7 (5)</td>
<td>0.6 ± 0.5 (5)</td>
<td>15.2 ± 6.7 (4)</td>
</tr>
<tr>
<td>828</td>
<td>122.6 ± 60.7 (3)</td>
<td>105.5 ± 51.6 (4)</td>
<td>4.7 ± 1.6 (4)</td>
<td>0.8 ± 0.3 (4)</td>
<td>1.4 ± 0.8 (3)</td>
</tr>
<tr>
<td>829</td>
<td>81.4 ± 42.9 (3)</td>
<td>85.4 ± 38.4 (4)</td>
<td>27.3 ± 16.7 (5)</td>
<td>5.3 ± 0.9 (5)</td>
<td>2.2 ± 0.5 (3)</td>
</tr>
<tr>
<td>830</td>
<td>33.8 (1)</td>
<td>43.0 ± 14.5 (4)</td>
<td>ND (4)</td>
<td>ND (4)</td>
<td>–</td>
</tr>
<tr>
<td>1633</td>
<td>329.9 ± 193.2 (3)</td>
<td>347.9 ± 201.9 (4)</td>
<td>6.5 ± 2.7 (4)</td>
<td>0.7 ± 0.4 (4)</td>
<td>1.0 (1)</td>
</tr>
</tbody>
</table>

1633 having the greatest and CCMP 830 the least DMSP per cell. DMSP<sub>p</sub> and DMSP<sub>t</sub> per cell values for CCMP 1633 were significantly different from those for *Emiliania huxleyi* CCMP 373 and 374 (<i>p</i> < 0.05), but not from those for other *S. microadriaticum* investigated. Average cell diameters (µm) for *S. microadriaticum* strains were (average ± SD; n = 30 strain<sup>−1</sup>) as follows: CCMP 421 (10.28 ± 1.08), 828 (10.69 ± 1.11), 829 (10.67 ± 1.22), 830 (10.74 ± 1.12) and 1633 (9.72 ± 0.82); and for *E. huxleyi* strains were as follows: CCMP 373 (5.08 ± 0.63) and 374 (5.03 ± 0.66). Only CCMP 1633 was statistically different in cell size (<i>p</i> < 0.05) among the various *S. microadriaticum* strains. Both *E. huxleyi* cultures were found to have less DMSP<sub>p</sub> when compared with *S. microadriaticum* cultures.

**Algal DLA**

In all strains, DMS evolution prior to exogenous DMSP addition (Fig. 1) was not statistically different from that in controls (<i>p</i> > 0.05). DLA was greatest in strain CCMP 829 and was undetectable in CCMP 830. Additionally, when normalized to cell number or chl <i>a</i> (fmol cell<sup>−1</sup>, nmol · min<sup>−1</sup> · µg<sup>−1</sup>), DLA for strains CCMP 373, 421, 828 and 1633 were markedly different from strain CCMP 829 (<i>p</i> < 0.01), but not from each other (<i>p</i> > 0.05), and CCMP 374 was statistically different from all other strains (<i>p</i> < 0.01) (Fig. 2, Table 2). DLA:chl <i>a</i> for *Emiliania huxleyi* CCMP 373 was greater than that of CCMP 374, but less than all *Symbiodinium microadriaticum* investigated (except CCMP 830; Table 2). DLA:DMSP<sub>p</sub> per cell averages were highest in strains 829 and 373 (0.06 and 0.02 nmol DMS · min<sup>−1</sup> · fmol DMSP<sup>−1</sup>, respectively) compared to other strains (Fig. 2). DLA:DMSP<sub>p</sub>...
per cell for strain CCMP 829 was statistically higher than that for all other strains in Fig. 2 (p < 0.01). CCMP 374 DLA measures (n = 2) are not presented in Fig. 2 because few replicates had measurable activity. Overall, CCMP 829 demonstrated the greatest DLA, whereas CCMP 421, 828 and 1633 had intermediate rates, with CCMP 830 having no detectable activity. Within strains, DMSPp concentrations were correlated with DLA. For each CCMP strain, r² values (in parentheses) were as follows: 373 (0.90), 374 (0.86), 421 (0.87), 828 (0.66), 829 (0.53) and 1633 (0.72). However, DMSPp concentrations were not correlated with DLA between strains (those strains with high DMSP did not necessarily have high DLA).

### Discussion

Our results demonstrate that 4 of the 5 symbiotic dinoflagellate strains of the genus *Symbiodinium* examined in the present study are capable of performing the enzymatic lysis of DMSP to DMS, indicating DLA in these algae. *S. microadriaticum* strains demonstrated a range of DLA when grown in culture and exposed to exogenous DMSP addition. One of the *S. microadriaticum* strains in the present study (CCMP 830) did not demonstrate DLA, suggesting that DL is not a universal enzyme in this species or that DLA was not detected within our analytical capabilities. Furthermore, DLA was primarily associated with the algae (not bacteria) in the non-axenic cultures investigated.

*Symbiodinium microadriaticum* strains exhibiting DLA averaged DMS production rates of 0.6 to 5.3 nmol · min⁻¹ · µg⁻¹ chl a. The highest average DLA:chl a DLA in this study, 5.3 nmol · min⁻¹ · µg⁻¹, occurred in CCMP 829. This strain was originally isolated from South Pacific clams, and DMSP concentrations in certain tissues of *Tridacna* sp. are some of the highest recorded in animal tissues to date (Hill et al. 2000). DLA for all algal lines closely parallel those reported by Harada et al. (2004) for particle-associated DLA in or near waters from the Gulf of Maine (DLA:chl a = 0.5 to 7.9 nmol · min⁻¹ · µg⁻¹). In their study, DLA:chl a rates ranged from <5 to 53 nmol · min⁻¹ · µg⁻¹, with the highest rate (53) occurring at an oligotrophic sampling site in the Sargasso Sea dominated by prymnesiophytes and dinoflagellates. It appears that differences in DLA normalized to chl a may be due to several factors, including species composition. Other studies have shown a relationship between high DMSP:chl a and DLA:chl a and have attributed this, in part, to nutrient limitation (Sunda et al. 2007). It is not likely that nutrient limitation played a role in our experiments, though nutrient quantities were not specifically addressed. More detailed experiments addressing the role of nutrient limitation and enzyme kinetics in *Symbiodinium* are needed.

DLA differences were detected between the strains investigated, though we did not specifically address enzyme turnover rates. Our finding that DLA was not detectable in CCMP 830 is consistent with the observa-
tions of Niki et al. (2000), who reported no DLA for 2 DMSP-producing Prymnesio phyceae species. The genus *Symbiodinium* encompasses 8 divergent clades (A to H) (for example, see Pochon et al. 2004, Coffroth & Santos 2005), and it is generally accepted that these clades are composed of several lineages representing species complexes (Santos 2004). *S. microadriaticum* strains (CCMP 830 and 1633) are the same algal clade (i.e. Clade B), but were isolated from different host origins (Atlantic and Pacific Oceans, respectively) and have very different DLA potentials, suggesting intrACLade DLA variation within a species (Table 2). *Symbiodinium* clades are known to have different susceptibilities to light and thermal stress (Rowan 2004), which may, in part, explain a corals’ sensitivity to bleaching. We hypothesize that if DMS has an antioxidant role in *Symbiodinium*, DMSP production and DLA in these algae might be expected to correlate with one of the primary mechanisms involved in coral bleaching, namely, damage to Photosystem II in the symbionts (Iglesias-Prieto et al. 1992, Lesser & Farrell 2004). Susceptibility to Photosystem II damage may be mitigated if DMSP and its enzymatic cleavage products serve to alleviate conditions of oxidative stress by scavenging harmful reactive oxygen species (ROS; Sunda et al. 2002).

Because bacteria contributed <5 to 20% of the total DLA, the whole culture DLA values can be primarily attributed to the algal cells. Steinke et al. (2002a) reached a similar conclusion, finding that >95% of DLA was found in the particle fraction >2 µm. Bacterial DLA was greatest for CCMP 421, and this result is unexplained by the number of bacteria present in the filtrate. All cultures had similar ratios of bacteria to algae (approximately 5:1 on a per cell basis, respectively), and the amount of bacteria in the filtrate versus that in the total culture sample was also similar among strains (76.9 ± 16.6%). Bacterial DLA may in fact be overestimated due to algal cell rupture via filtration through polycarbonate filters, allowing algal contents into the experimental filtrate.

While the use of axenic cultures allows a more direct analysis of DLA in the algal component alone, we found that *Symbiodinium microadriaticum* treated with antibiotics had lower growth rates (data not shown). Additionally, checking cultures for axenicity by using standard plating techniques will miss a substantial portion of bacteria (Kogure et al. 1979). Though the bacteria in the present study were not genetically characterized, DLA could be over- or underestimated if a significant fraction of DMSP-utilizing bacteria (e.g. *Roseobacter*) were present in cultures. This is due to the finding that bacteria, in addition to the DL pathway, can also metabolize DMSP via the demethylation/demethiolation pathway, which would reduce the DMS quantities evolved (Taylor & Gilchrist 1991). We do not suspect that demethylation/demethiolation played a substantial role in our experiments, as no methanethiol was detected. Alternatively, bacterial DLA may increase the quantity of DMS detected, an important factor that we have addressed in the present study and estimate to be small. It is also recognized that the possibility of algal-attached bacteria cannot be excluded and may in fact be partially responsible for some of the observed DLA. However, during bacterial enumeration of whole culture samples, bacteria were not observed to be more numerous in close proximity to algal cells.

Our data demonstrate DLA variability within and across *Symbiodinium microadriaticum* strains, providing only the first step in an effort to further elucidate DLA regulation in the algae involved in numerous symbioses. DLA variability within strains could be attributed to several factors. According to Steinke et al. (2007) in their study of *Emiliania huxleyi*, in vivo DLA may vary during the course of a day due to enzyme turnover, and individual strains of algae may respond differently to exogenous substrate (DMSP) additions. Our finding that CCMP 373 has greater DLA than CCMP 374 is in agreement with other reports (Steinke et al. 1998), though differences in experimental conditions preclude direct comparison with previously reported rates (Steinke et al. 1998, 2000, 2007). Additionally, *E. huxleyi* DLAs in the present study were used to demonstrate the validity of the given assay for the detection and optimization of DLA in *S. microadriaticum*; the assay was not optimized for *E. huxleyi* DLA measurements. While several assay parameters differed when compared to those in previous studies, our *E. huxleyi* data are in agreement with published DLA measurements when calculations incorporate differences in activity associated with pH. For example, using data presented by Steinke et al. (1998), the calculated DLA:DMSP<sub>p</sub> (nmol DMS·min<sup>−1</sup>·fmol<sup>−1</sup> DMSP) for CCMP 373 is 0.27 at an optimal pH of 6, but, at pH 8, it would be approximately 10% of that value based on the reported pH trials. Additional sources of variability may include shifts in enzyme affinity during growth, culture conditions and individual variability associated with enzyme assay parameters. Given these observations, the present study demonstrates that some *S. microadriaticum* have DL capabilities and that DLAs are distinguishable between algal strains at the given concentration of exogenous DMSP addition.

*Symbiodinium* are known to contain substantial quantities of DMSP (Jones et al. 1994, Hill et al. 1995, Broadbent et al. 2002, Broadbent & Jones 2004), with cultured zooxanthellae having 179 fmol cell<sup>−1</sup> DMSP (Keller et al. 1989). Our DMSP<sub>p</sub> measurements for *S. microadriaticum* strains (34 to 330 fmol cell<sup>−1</sup>) are in
agreement with those from previous studies (Jones et al. 1994, Hill et al. 1995, Broadbent et al. 2002). Several studies support the hypothesis that DMSP and DLA are produced at varying levels by different organisms and that DMSP and DLA vary between strains of the same species (Steinke & Kirst 1996, Steinke et al. 1996, 2002a,b, Wolfe et al. 1997, Niki et al. 2000). Our comparisons of per cell DMSP between the 5 strains of *S. microadriaticum* tested revealed differences, so further investigations of DMSP levels within *Symbiodinium* are warranted. We found no apparent correlation between intracellular DMSP concentrations and DLA between strains, though strain CCMP 830 had no detectable DLA and also had the lowest DMSP concentrations of the *S. microadriaticum* strains investigated.

Corals (host and algal cells) may be exposed to conditions of elevated oxidative stress when harmful ROS are not scavenged or detoxified by antioxidants. Because corals contain photosynthetic, oxygen-producing algae, high levels of antioxidant enzymes (and free radical scavengers) are found in host (and algal) tissues (Lesser & Shick 1989, Dykens et al. 1992, Downs et al. 2002). With proposed antioxidant functions, DMSP, and more importantly its enzymatic cleavage products via DLA, may play significant roles in alleviating conditions of oxidative stress on reefs. To investigate this potential antioxidant role we are exploring the effects of various oxidative stressors on DMSP levels and DLA in isolated coral symbiotic algae. Our studies herein have shown that coral symbiotic algae contain DLA, and further studies are directed towards determining if cnidian hosts, in common with other symbiotic host species, contain DLA.

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