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Algal symbionts of the freshwater sponge *Ephydatia muelleri*

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Abstract

The freshwater sponge, *Ephydatia muelleri*, is an emerging model system for studying animal:microbe symbioses. Intracellular green microalgae are one of the more common symbionts that live in a facultative mutualism with *E. muelleri*. While these symbioses have long been known, the identity of the algal symbionts in *E. muelleri* cells has not been studied in detail. Here, we isolate and characterize endosymbiotic algae from *E. muelleri* collected from different geographic locations. We find that the algae can be transmitted through asexually produced gemmules and importantly that they can form symbioses with different, differentiated sponge cell types in the adult sponge. Our findings indicate that at least two algal lineages form endosymbioses with *E. muelleri*. One of the lineages includes species commonly found in samples from two locations in Canada and one in the United States (clade 1: closely related to *Auxenochlorella pyrenoidosa*). The other clade includes algae found in sponges from one site in Maine, USA, and *Lewinosphaera symbiontica*, which is a strain isolated in 1956 from the freshwater sponge *Spongilla*. We compared microbiomes found in cultures of microalgae as well as the original sponge hosts, and found that very similar bacterial microbiomes associate with both clades (91 orders of Bacteria are shared among the samples we compared). The microbiomes found in the cultures resemble, with a high degree of overlap, the microbiome associated with the sponge host.

Keywords Green microalgae · Freshwater sponge · Endosymbiosis

1 Introduction

Sponges are among the earliest branching animal phyla (Telford et al. 2016) and form associations with a large variety of microbial partners (Thomas et al. 2016; Pita et al. 2018). The holobiont (i.e., host plus associated symbionts) performs key functions in the ecosystem, and holobiont partners generate reciprocal selective pressures that result in interesting evolutionary outcomes (e.g., Taylor et al. 2007; Webster

& Taylor 2012). Photosynthetic symbionts account for an important part of the holobiont both in marine and freshwater sponges. These symbionts provide photosynthates, secondary metabolites, and possibly are involved in nitrogen fixation to the sponge host (Wilkinson and Fay 1979; Wilkinson 1983; Arillo et al. 1993). Photosynthetic symbionts include cyanobacteria, dinoflagellates, rhodophytes, chlorophytes and diatoms (Scott et al. 1984; Rützler 1985; Taylor et al. 2007). Cyanobacteria are a common and well-described group of photosynthetic endosymbionts that can reside in marine and freshwater sponges (e.g., Thacker and Freeman 2012; Webster and Taylor 2012; Gaikwad et al. 2016; Kulakova et al. 2014) where they can play important roles in host adaptation through expanded metabolic function via photoautotrophy (e.g., Zhang et al. 2015; Hudspeth et al. 2022). A small number of marine sponges, like some in the Clionidae, possess photosynthetic dinoflagellate endosymbionts (*Gerakladium*) that can enhance bioerosion and growth rates (Hill 1996; Weisz et al. 2010; Ramsby et al. 2017). Co-occurrences of sponge:rhodophyte associations have been described where the organisms grow attached to

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one another producing surface proliferations (Tronchin et al. 2006) and some of the most abundant diatom species of the Antarctic plankton communities have been observed in high densities inside the tissues of several sponge species (Cerrano et al. 2004). Last but not least, and the focus of this study, is the long-known relationship of chlorophytes, also known as green algae, that form endosymbioses with freshwater sponges (Brandt 1882).

Algal photobionts are abundant in freshwater sponges of the Demospongiae, but less is known about this intriguing partnership than of marine sponges. The symbiotic intracellular algae of freshwater sponges include a diverse set of chlorophytes from a variety of genera mostly within the class Trebouxiophyceae (Kenney et al. 2019; Ereskovsky et al. 2022). Many of the reports of endosymbiotic green algae have been from the freshwater sponge, *Spongilla lacustris* (e.g., Castro-Rodriguez 1930; Gilbert and Allen 1973; Williamson 1979; Reisser 1984; Saller 1989, 1991; Masuda 1990; Handa et al. 2006). Symbiotic algae in *S. lacustris* and in *Radiospongilla cerebellata* have been shown to affect germination rates of gemmules (Agnes and Brøndsted 1953; Okuda et al. 2002) and have a positive effect on the growth rate of sponges due to net gain from photosynthesis (Frost and Williamson 1980; Sand-Jensen and Pedersen 1994). Additionally, Cook (1983) demonstrated that algal symbionts benefit another freshwater sponge species, *Ephydatia fluviatilis*, by providing glucose to host sponge cells, and by providing up to 20% of the total fixed carbon for the host's metabolism.

For the majority of freshwater sponge symbioses involving green algae, the precise identity of the partners is unknown. *Chlorella sorokiniana* is known to be a species of algae that is associated with *Spongilla* (Reisser 1984), but most studies have not identified the species of algae using molecular markers. For example, freshwater sponges with documented green algae of the Trebouxiophyceae include *Radiospongilla* sp. (Masuda 1990; Handa et al. 2006), *Eunapius fragilis* (Handa et al. 2006), *Lubomirskia* sp. (Berner and Titlyanov 1992; Bil et al. 1999; Ereskovsky et al. 2016), *E. fluviatilis* (Wilkinson 1980; Gaino et al. 1995), and *E. muelleri* (Hall et al. 2021). One study showed that gemmules from *R. cerebellata* possessed two different endosymbiotic algal species (Handa et al. 2006). In most of the cases described above, morphology was used for taxonomic classification. However, more recent genetic sequence data has become available for algae found in *Lubomirskia* and *Baikalospongia* freshwater sponge species; the symbionts in these sponges were identified as belonging to the coccoid green algal genus, *Mychonastes* in the case of *Lubomirskia baicalensis* (Chernogor et al. 2013) and multiple lineages of *Choricystis* in the case of several *Lubomirskia* and *Baikalospongia* sponge species in Lake Baikal (Kulakova et al. 2020). Pröschold and Darienko (2020)

recently analyzed a symbiotic strain of *S. lacustris* isolated by Lewin (1966) in Massachusetts, USA and determined it to be a member of the Chlorellaceae representing a new species of the genus *Lewinosphaera* (*L. symbiontica*) using both morphological and DNA sequence analysis. In spite of abundance and importance of freshwater sponge:algal symbioses, relatively little is known about the *Chlorella*-like algae that form endosymbiotic relationships with freshwater sponges.

Recently, Ereskovsky et al. (2022) reviewed the literature on cytosymbiosis and algal symbionts in sponges. They discuss the evidence that adult stem cells in sponges (most frequently archaeocytes) host endosymbionts and that this arrangement is contrasted with the observation that intracellular symbionts commonly reside in more specialized or differentiated cells in other animal hosts. They make the argument that Porifera should be a model for studying host:symbiont coevolution because sponge holobionts are ancient symbiotic associations among the Metazoa and because adult stem cell cytosymbioses are rare. We agree that sponges, particularly the freshwater sponges, offer unique opportunities to study animal:algal endosymbioses. We recently showed that the freshwater sponge, *E. muelleri*, is a tractable model for studying sponge:algal endosymbiosis (Hall et al. 2021). *Ephydatia muelleri* is nearly panglobal, is globally abundant, and easy to culture in the lab with and without symbionts. This species of sponge has a well-annotated chromosomal level genome and a developmentally staged transcriptome (Kenny et al. 2020). *E. muelleri* has also been an important species for studying physiology, cell biology, and genetics of sponges, especially in the context of animal evolution (Elliott and Leys 2007, 2010; Windsor & Leys 2010; Ludeman et al. 2014; Schenkelaars et al. 2016; Windsor-Reid et al. 2018; Hall et al. 2019; Mitchell and Nichols 2019; Colgren and Nichols 2022). Several transcriptomes representing various stages of symbiosis (Hall et al. 2021; Geraghty et al. 2021) have been analyzed, which revealed a set of sponge genes involved in establishment and maintenance of the symbiosis. However, algal-specific gene expression or the taxonomic identity of the native *Chlorella*-like algal symbiont in *E. muelleri* has not been analyzed. In an effort to better understand these important symbioses, we present morphological and molecular data describing *E. muelleri* algal and microbial endosymbionts isolated from several regions in Canada and the USA.

2 Materials and methods

2.1 Algal strain isolation and culturing

Ephydatia muelleri tissue possessing microalgal symbionts was collected from several different locations in the USA and Canada and algal symbionts were isolated from green *E.*

muelleri freshwater adult sponges or gemmules (Table S1). As adult sponges can possess both endosymbionts and other associated algae, to ensure that only intracellular endosymbionts were isolated (not free-living algae), gemmules or hatched gemmules were used when available. In the case of algae that were isolated from adult sponges, to ensure strains were pure, algae were cultured on agar plates as follows. Algae were isolated by grinding tissue followed by differential centrifugation as described in Hill et al. (2020). Algal cultures, both xenic and axenic, were grown in both Bold 3N medium (UTEX Culture Collection of Algae) and Bold Modified Basal Freshwater Nutrient Solution (BBM, Sigma Aldrich). All liquid cultures were grown at 20–25 °C under a light:dark cycle of 12:12 h for as long as 6 weeks before periodic subculture. To remove unwanted microorganisms from the aqueous cultures and to ensure strains were pure, algae were streaked on BBM agar plates for several rounds until single strains were isolated. Frozen stocks (-80 °C) of axenic strains were made using the GeneArt Cryopreservation Kit for Algae (Invitrogen).

2.2 Freshwater sponge culturing

Ephydatia muelleri gemmules were collected, stored, and cultured as described in Leys et al. (2019). Gemmules are overwintering, asexually produced small cysts of sponge stem cells that are capable of developing into a juvenile sponge upon hatching. Gemmules possessing high amounts of algae inside the gemmule covering and among the stem cells were green and could be used for algal isolation or for hatching sponges with algal endosymbionts in further experiments. Alternatively, aposymbiotic sponges hatched from gemmules could be infected with algae to establish the symbiosis according to Hill et al. (2020). Sponges and associated algal symbionts were either harvested and stored at -20° C for later DNA isolation or grown on 35 mm glass bottom culture dishes (MatTek Life Sciences) and fixed in cold 4% paraformaldehyde (PFA) in 190 proof ethanol overnight for microscopy.

2.3 Microscopy

Algae and associated microorganisms grown on agar plates were imaged using a Leica M165C stereoscope with Leica MC170 HD camera. Algae in liquid cultures were concentrated by centrifugation at 3000 rpm for 10 min to obtain a pellet that could be resuspended in algal media and mounted in 100% glycerol for morphological analysis using bright-field microscopy with a Nikon 80i fluorescence upright microscope at 100×oil immersion objective. Using the same microscope and lens, images of sponge cells and associated microalgae from ruptured gemmules were obtained by placing a single gemmule on a microscope slide and either

squashing with a coverslip or tearing with forceps and then placing a small amount of glycerol on the expelled cellular contents followed by a coverslip.

Sponges with symbionts that were fixed in 4% paraformaldehyde (PFA)/ethanol were washed three times in phosphate buffered saline (PBS), treated with phosphate buffered saline with 1% Tween 20 (PBST) for 45 min, washed again in PBS, and stained with Alexa Fluor™ 488 Phalloidin (1:40 dilution, Invitrogen, Molecular Probes) and Hoechst 33342 (1:2000 dilution, Thermo Scientific). Samples were imaged using the Leica SP8 laser scanning confocal microscope equipped with white light laser (WLL) system, hybrid detectors and 63× water immersion objective.

2.4 DNA extraction, molecular barcoding, and phylogenetic analysis

Genomic DNA of algal cultures was extracted using CTAB reagent and mechanical disruption with 0.5 mm Zirconia/Silica beads and the Bead Ruptor 4 (Omni International). The SSU rRNA and chlorophyte genes were amplified using GoTaq PCR 2X MasterMix (Promega). Chlorophyte gene marker amplifications were subject to the following thermocycling conditions: initial denaturation at 95 °C for 5 min, followed by 34 cycles of 95 °C for 30 s, 53 °C for 30 s, 72 °C for 1 min 20 s, followed by 1 cycle of 72 °C for 5 min. SSU rRNA gene marker amplifications were subject to the following thermocycling conditions: initial denaturation at 95 °C for 3 min, followed by 34 cycles of 95 °C for 1 min, 55 °C for 1 min, 72 °C for 3 min, followed by 1 cycle of 72 °C for 5 min. SSU rRNA primers used included EAF3, E528F, 920F, GF, GR, and ITS055R (Marin et al. 1998, 2003) and newly designed primers as shown in Table S2. Chlorophyte gene markers used included TUFA, ITS, 16S and RBCL (Burja et al. 2001; Fama et al. 2002; Sun et al. 2009; Bock et al. 2010; Zou et al. 2016, Table S3). All PCR products were purified using QIAquick MinElute PCR Purification Kit (Qiagen) and sequenced. All sequences are provided in File S1.

As morphology was not informative for the taxonomic identification of the algal isolates, we implemented a molecular taxonomy approach to decipher whether individuals of *E. muelleri* from different geographical locations possessed the same or different endosymbiotic *Chlorella*-like algae. For each geographic isolate shown in Fig. 1 and for an isolate from Virginia that was originally described in Hall et al. (2021), we sequenced three commonly used molecular markers for algal taxonomy: two chloroplast markers—the elongation factor Tu gene (*tufA*) and the large subunit of the enzyme ribulose biphosphate carboxylase (RuBisCo) (*rbcL*)—and the nuclear marker ribosomal small subunit DNA (SSU rDNA) including the internal transcribed spacer (ITS) between the small and large-subunit ribosomal (rRNA).

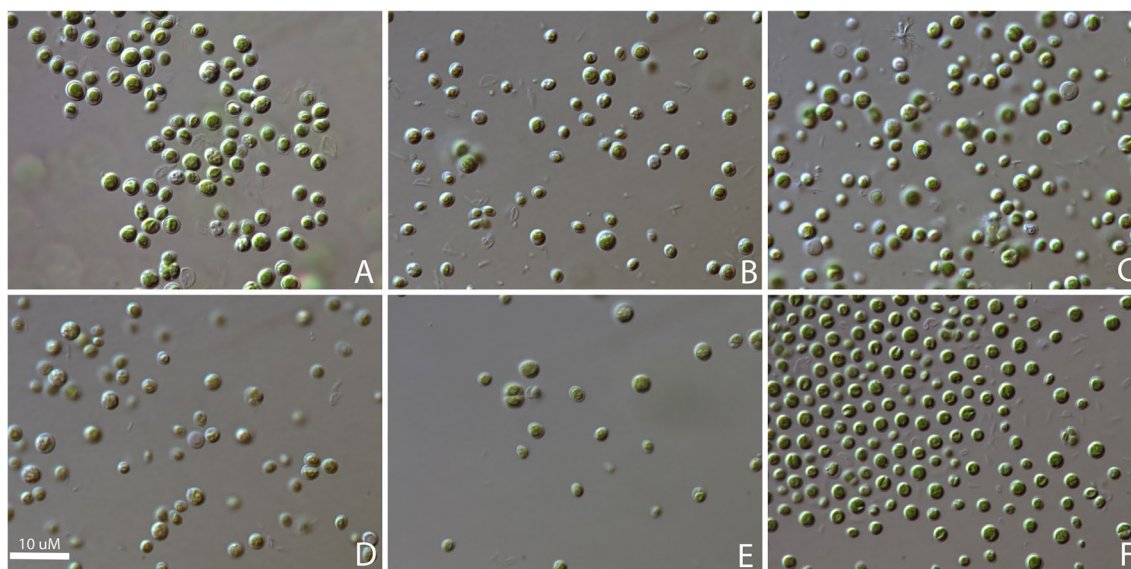


Fig. 1 Morphology of Symbiotic Algal Isolates of *Ephydatia muelleri*. **A.** O' Connor Lake, British Columbia, Canada (Emu_LOC). Isolated from gemmules. **B.** Saint Lawrence River, Montreal, Canada (Emu_MON). Isolated from adult tissue. **C.** Pemaquid River, Maine, U.S. (Emu_PR6). Isolated from adult tissue. **D.** Sooke Res-

ervoir, British Columbia, Canada (Emu_Sooke). Isolated from adult tissue. **E.** Dundee Pond, Maine, U.S. (Emu_DPAG). Isolated from hatched gemmules. **F.** Pemaquid River, Maine, U.S. (Emu_PRAG). Isolated from gemmules and hatched gemmules. Scale bar for all images = 10 µm

Known algal sequences were obtained from the NCBI database nucleotide collection (nr/nt) and aligned with our SSU rRNA and chlorophyte sequenced genes separately using MAFFT (Kato and Standley 2013). Phylogenetic trees were built with the chloroplast genes *tufA* and *rbcl* and the nuclear SSU-ITS concatenated using maximum likelihood in IQ-TREE (Nguyen et al. 2015; Minh et al. 2020) under the TIM2e + R3 model. All models were tested by ModelFinder (Kalyanamorthy et al. 2017) and the optimal was selected based on Bayesian Information Criterion (BIC) and Akaike Information Criterion (AIC) statistics Branch support was assessed using the ultrafast bootstrap (UF bootstrap) method (Minh et al. 2013).

2.5 Algal microbiome sequencing and analysis

Total genomic DNA of each of the xenic algal isolates was also used to complete a 16S rRNA gene microbiome analysis. A 464 bp hypervariable region of V3 (ONT_unipro341F) and V4 (ONT_unipro805R) of bacterial and archaeal DNA (Takahashi et al. 2014; Apprill et al. 2015; Parada et al. 2016) was amplified using LongAmp Taq 2X Master Mix (New England BioLabs). Thermocycling conditions for the first round of PCR included initial denaturation at 98 °C for 2 min, followed by 10 cycles of 95 °C for 20 s, 65 °C for 15 s, -1 °C/cycle, 65 °C for 30 s, followed by 25 cycles of 95 °C for 20 s, 55 °C for 15 s, and 65 °C for 1 min and a final extension at 1 cycle of 65 °C for 5 min. The PCR Barcoding Expansion Pack 1–96 (EXP-PBC096, Oxford

Nanoporetech, US) was used to barcode PCR libraries with thermocycling conditions of 16S rRNA gene amplicons with an initial denaturation at 95 °C for 3 min, followed by 15 cycles of 95 °C for 15 s, 62 °C for 15 s, 65 °C for 1 min, followed by 1 cycle of 65 °C for 5 min. The Ligation Sequencing Kit 1D (SQK-LSK109, Nanopore Technologies) and the NEBNext® Companion Module for Oxford Nanopore Technologies® Ligation Sequencing (E7180S, New England BioLabs) were used following the manufacturers' protocols. The barcoded gene libraries were pooled in equal amounts of DNA and sequenced in a MinION Flow Cell for ~5 h. The MinKNOW report containing the aggregated counts tables at different taxonomic levels was generated by the cloud-based Epi2ME software (Oxford Nanopore, USA). Following analyses were performed in R (version 4.2.1).

2.6 Sponge microbiome sequencing and analysis

DNA was extracted from about 20 mg of tissue using the DNeasy Blood and Tissue kit (Qiagen). The V4 hypervariable region was amplified with the same primers used for the algae samples, 515F-Y (Parada et al. 2016) and 806R (Apprill et al. 2015). DNA amplification was done in duplicates with the following conditions: 95 °C for 20 s, followed by 25 cycles of 95 °C for 10 s, 60 °C for 20 min, 72 °C for 30 min, and a final elongation at 72 °C for 5 min. Libraries were prepared with the Nextera XT DNA Library Preparation Kit (Illumina Inc.) and next generation paired-end sequencing was performed at the Natural History Museum

of London (<https://www.nhm.ac.uk/>) on an Illumina MiSeq device using v3 chemistry (2 × 300 bp). Read processing and taxonomic assignment followed the MiSeq SOP protocol (Kozich et al. 2013) in Mothur (v.1.41.3) inferring amplicon sequence variants (ASVs), allowing one mismatch per 100 bp. ASVs were classified using the Reference NCBI database (refseq_rna, update 12/01/2023, including 49,700,764 sequences), with a cutoff value of 80.

2.7 Statistical analyses of algal and sponge microbiomes

Measures of alpha diversity (Shannon index) were calculated using rarefied samples in R. These metrics were compared among clades or locations using analyses of variance (ANOVA), and Tukey's honest significant difference (HSD) for pairwise comparison. Beta-diversity among the microbiome samples was visualized using principal coordinate analysis (PCoA) on a Bray–Curtis dissimilarity matrix using "cmdscale" in vegan package v. 2.5-7 (Oksanen et al. 2018). Homogeneity of variance, which tests whether two or more groups are homogeneously dispersed in relation to their group centroid, was determined using the "betadisper" function in vegan. We compared distances among clades and locations by permutational multivariate analyses of variance (PERMANOVA) using "adonis" in vegan.

3 Results

3.1 Morphology and cellular location of *E. muelleri* algal symbionts

Both endosymbiotic and putatively ectosymbiotic algal strains were identified in *E. muelleri* tissues and gemmules.

In all cases, endosymbiotic algae were culturable in modified Bold's media and, regardless of geographic location, the predominant algae were unicellular zoochlorellae, small (4–10 μm), spherical, and without flagella (Fig. 1). As has been described in other work on green algal symbionts in freshwater sponges, these algae appear to be *Chlorella*-like in this facultative symbiosis. We did not observe significant morphological differences between the strains.

We considered whether algal symbionts transmitted to sponges through gemmules were present inside or outside of host thesocytes (sponge stem cells in the gemmules). We examined the cellular contents of *E. muelleri* gemmules to evaluate the location of algae prior to thesocyte differentiation (i.e., thesocytes are the resting state of archeocyte stem cells in the gemmule) and sponge development to determine where algal cells resided (Fig. 2). We observed that even 'yellow' gemmules (those without obvious symbionts) possessed some algal cells, mostly located outside of thesocytes (Fig. 2A). 'Green' *E. muelleri* gemmules possessed greater numbers of algae and while most of the algal cells were located outside of sponge cells, there were thesocytes that appeared to contain intracellular algae (Fig. 2B and C). Thus, it is possible that the algae persist in those thesocytes and are passed onto future cells through cell division. Gemmules were observed to contain diverse communities of bacteria.

We hatched *E. muelleri* gemmules to examine algal symbiont location in sponges developing from gemmules. Using confocal microscopy, we verified that all strains of algae isolated from *E. muelleri* gemmules or hatched gemmules (Table S1) were present in intracellular locations in juvenile sponges. *E. muelleri* hatched from 'green' gemmules collected from three of the geographical locations contained algae in intracellular spaces (Fig. 3A–C) in multiple cell types. Depending on the number of algae in the

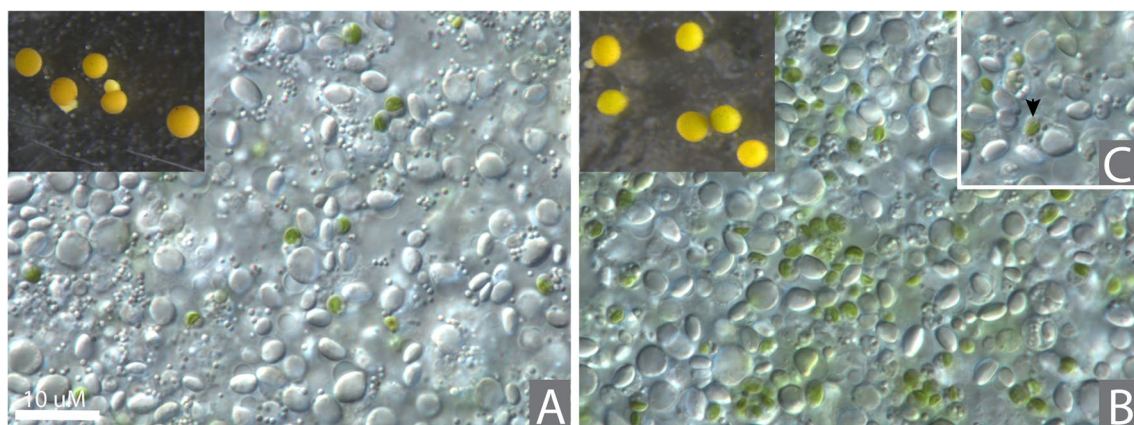


Fig. 2 Ruptured gemmules cellular content. **A.** Cells from a ruptured yellow *E. muelleri* gemmule with inset of yellow gemmules with two popped to show cellular contents. **B.** Cells from a ruptured green *E.*

muelleri gemmule with inset of green gemmules. **C.** Stem cell (thesocyte) with putative intracellular algae shown by arrowhead. Scale bar for gemmule cells = 10 μm

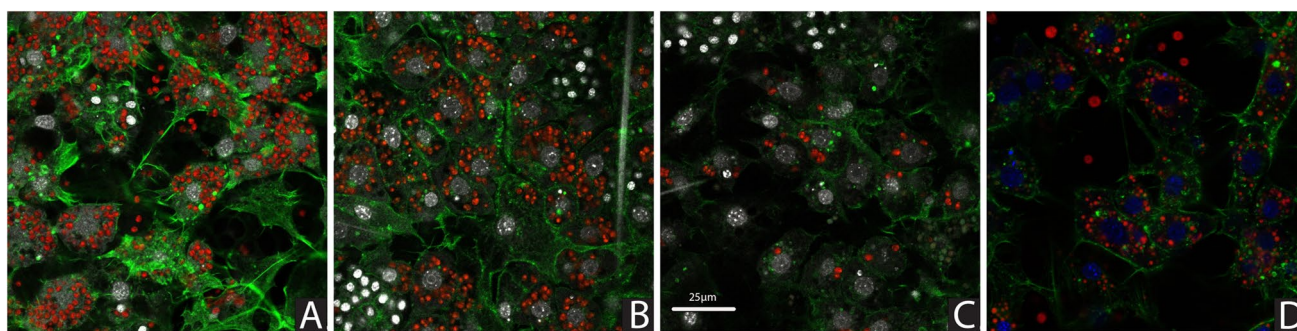


Fig. 3 *E. muelleri* cells with endosymbiotic algae. **A.** Panel **A–C** shows sponge tissue grown from green gemmules collected from different geographical locations. **A.** O' Connor Lake, British Columbia, Canada. **B.** Dundee Pond, Maine, U.S. **C.** Pemaquid River, Maine, U.S. Panel **D** shows *E. muelleri* tissue grown from “aposymbiotic” gemmules and infected with the algal endosymbiont strain isolated

from gemmules obtained from that geographic location. **D.** Pemaquid River, Maine, U.S. infected with native algal strain, 20 min post infection. Tissue stained with Phalloidin 488 for actin (green) and Hoescht for nuclei (white in panel **A–C.**, blue in panel **D.**) Autofluorescence of algae is observed (red). Scale bar for all images = 25 μm

gemmule and the length of time the sponge is grown with the algae, the numbers of algae per sponge cell varied, with dark green gemmules grown to adult sponges having numerous algae per cell (Fig. 3A and B). Sponges grown from gemmules possessing few algae (‘yellow’ gemmules) had fewer endosymbionts (Fig. 3C), but our earlier work showed that when sponges are grown under light, numbers of algae increase over time as the sponge and algal cells divide (Hall et al. 2021).

We have previously demonstrated (Hall et al. 2021) that *E. muelleri* hatched from aposymbiotic gemmules (or gemmules with very low numbers of algae) can be infected with symbiotic algal strains isolated from the same geographic location and that these endosymbioses can be maintained in the lab for as long as the sponges are viable. Here, we verified this finding using strains of algae and *E. muelleri* from new geographical locations. Figure 3D shows aposymbiotic *E. muelleri* hatched from gemmules from the Pemaquid River (Maine) infected with the Emu_PR6 native algal strain. While some algae remain in extracellular locations, 20 min after infection the majority of algae are intracellular.

3.2 Molecular barcoding and phylogenetic analysis of algal symbionts

To place our algal sequences within known algal clades, we collected the most similar sequences to those produced herein from the best hits resulting from BLAST analysis against the *nr* database from NCBI (File S1). All three strains from Canada (LOC, MON, SOOKE), one strain from Maine (DPAG), and the one from Virginia (VA SI) resulted in similar outputs. Using the nuclear marker SSU, the most similar sequences were those from *Pseudochlorella pringsheimii*. The chloroplast marker gene *tufA* for our strains blasted to *Auxenochlorella pyrenoidosa*. The

chloroplast marker *rbcL* was not particularly informative for the classification of these queries as it resulted in multiple best hits (*Auxenochlorella pyrenoidosa*, *Jaagichlorella luteoviridis*, *Chlorella vulgaris*, *Auxenochlorella protothecoides*, and *Chlorella pyrenoidosa*). The other two algal strains studied here, PR6 and PRAG, both isolated from individuals of *E. muelleri* from Pemaquid River (Maine), blasted against different algal strains than LOC, MON, SOOKE, DPAG and VA SI. The SSU queries for PR6 and PRAG were notably identical to the newly erected species of *Lewinosphaera symbiontica* that was recently described as an isolate from a freshwater sponge endosymbiont of *Spongilla* collected in Massachusetts in 1956 (Pröschold and Darienko 2020). The *tufA* marker revealed the closest similarity for these strains with *Chlorella* sp. and the *rbcL* marker with *Chlorella variabilis*.

For a more accurate taxonomic assignment and in order to determine possible evolutionary relationships between the algal isolates from this study and other described algal species, maximum likelihood phylogenetic trees were built for all the molecular markers. The nuclear markers SSU and ITS were concatenated and a phylogenetic tree was reconstructed based on the SSU-ITS tree reported in Pröschold and Darienko (2020). The alignment for SSU-ITS included 46 algal sequences with 2292 nucleotide positions, 456 distinct patterns, 334 parsimony-informative sequences, 220 singleton sites, and 1738 constant sites. The algal isolates from Virginia (VA SI), Canada (LOC, MON, SOOKE) and the strain DPAG from Maine (DPAG) clustered together in a clade with *Micractinium tetrahymenae* with the highest bootstrap support, while the other two strains from Maine (PR6 and PRAG) formed a highly supported clade with *L. symbiontica* (Fig. 4).

For the *tufA* phylogeny (Fig. S1), a dataset of 39 complete *tufA* sequences from different *Chlorella*-like algal

Fig. 4 Molecular phylogeny of Chlorellaceae based on the concatenated aligned sequences of the nuclear markers SSU and ITS. The tree is inferred by maximum likelihood and the nodal supports are ultrafast bootstraps. The reference sequences are based on the dataset from Pröschold and Darienko (2020). The algal isolates from this study are in bold. Note that only bootstrap values over 70 are shown



species was compiled, with 741 nucleotide positions, 319 distinct patterns, 292 parsimony-informative sequences, 35 singleton sites, and 414 constant sites. Here, our sequences for the *E. muelleri* algal strains were clustered together in similar clades as those recovered in the SSU-ITS phylogeny (Figs. 4, S1). While the strains VA SI, LOC, MON, SOOKE, and DPAG were placed in a clade with *A. pyrenoidosa* (clade 1), the strains PR6 and PRAG branched in clade 2 with two sequences of the *Chlorella* sp. strain 484 (Fig. S1).

Finally, to reconstruct the *rbcL* phylogeny (Fig. S2), we compiled a final dataset of 46 complete and partial sequences from different species of Chlorellaceae, with 1293 positions, 589 distinct patterns, 427 parsimony-informative, 94 singleton sites, and 772 constant sites. In the resulting maximum likelihood hypothesis, we observed the algal strains organized in two distinct phylogenetic clades (1 and 2) as with the previous markers with identical affinities. The strains PR6 and PRAG clustered here in a robustly supported clade (clade 2) with sequences from *Chlorella* sp. and *Chlorella variabilis* strains (Fig. S2). The strains VA SI, LOC, MON, SOOKE, and DPAG were recovered as a single clade (clade 1) with algal species from several genera including *Auxenochlorella*, *Jaagichlorella* and *Pseudochlorella* (Fig. S2).

3.3 Algal strain associated microbiomes

We found that the algae grew well, some strains for nearly a decade, in media or on plates with the microbes that were co-isolated from the sponges. Each of the strains of symbiotic algae isolated from either adult freshwater sponge tissue or from gemmules or hatched gemmules contained associated prokaryotic microorganisms (Fig. 5). We also found that these strains can be made axenic by colony isolation and multiple rounds of successive plating. The axenic strains also grow well in lab cultures, although most grow approximately 1.5 times slower than the xenic strains.

To better understand the diversity of bacteria we observed in algal cultures, we sequenced the microbiomes from endosymbiotic microalgal strains that were isolated from *E. muelleri* sponges and grown in culture for months to years with culture transfer approximately every 50 days (Table S1). Through long-read 16S rRNA gene amplicon sequencing we identified a total of 2,616 prokaryotic species (666 species with > 0.01 relative abundance). The identified species spanned 850 genera, 139 orders, 57 classes, and 26 phyla. We examined the bacterial community composition of the algal associated microbiomes using relative abundance plots at different taxonomic levels. At the phylum level, Cyanobacteria was the most

Fig. 5 Xenic strains of endosymbiotic algae isolated from *E. muelleri* gemmules. **A.** Pemaquid River, Maine, U.S. algal strain isolated from gemmules with associated bacterial colonies (arrowhead). **B.** Dundee Pond, Maine, U.S. algal strain isolated from hatched gemmules and associated bacteria (arrowhead)

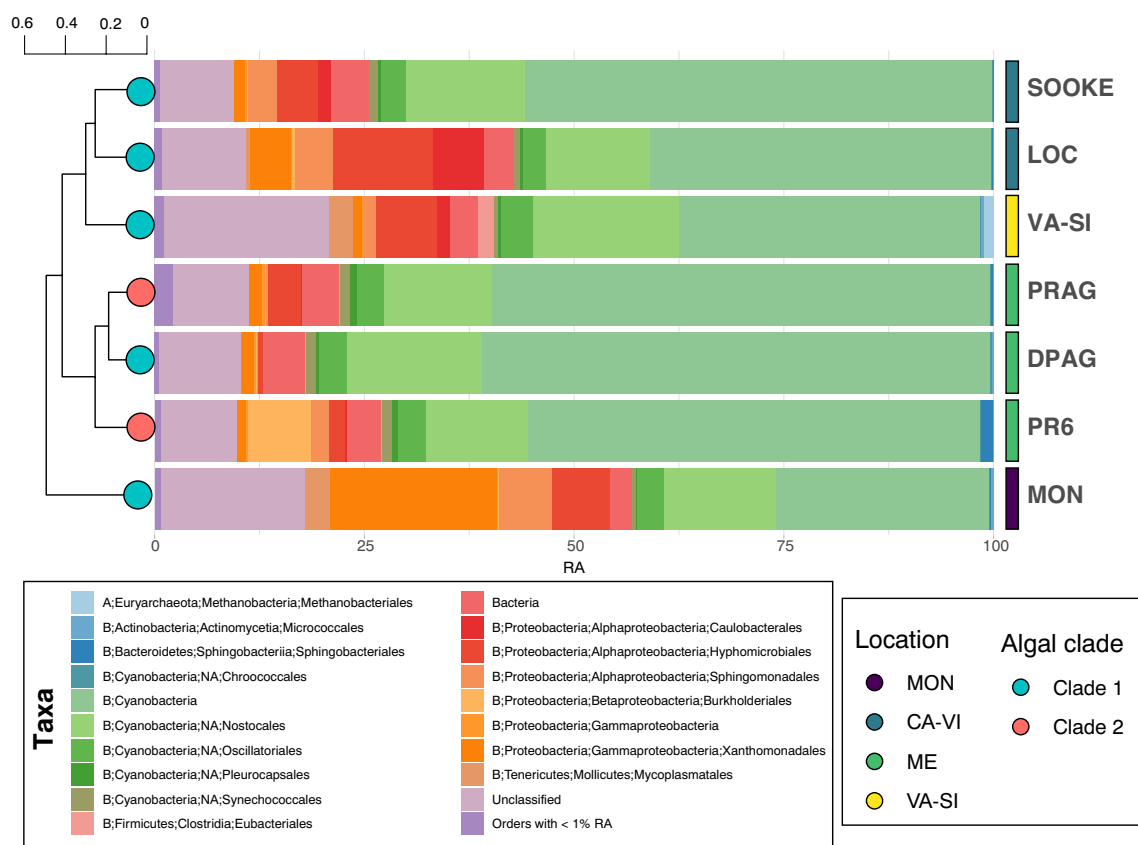
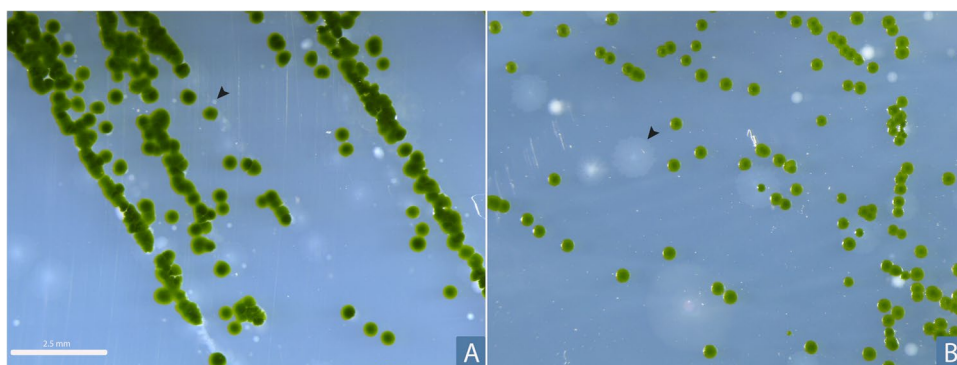


Fig. 6 Community composition of the microbiota in the algal isolates from *E. muelleri* at order taxonomic level represented as relative abundance in stacked bar charts and the dendrogram based in species level dissimilarity

abundant phylum in all samples followed by Proteobacteria (Fig. 6). These two phyla accounted for about 82% of the observations. Among Cyanobacteria, species from the orders Nostocales, Synechococcales, Oscillatoriales, and Pleurocapsales were present in all samples (Fig. 6). There were also four abundant Alphaproteobacteria orders and two Gammaproteobacteria orders shared among all samples (Fig. 6). Relatively high abundance of species from the orders Burkholderiales, Flavobacteriales, Mycoplasmatales, and Eubacteriales was observed in some but not

all samples (Fig. 6). The microbiomes of the algal cultures isolated from sponges collected in Maine (PR6, PRAG and DPAG) were more similar to each other than to the rest of the microbiomes, regardless of the algal strains belonging to both clades (Fig. 6).

After identifying the most abundant taxonomic groups in the algal associated microbiomes, we asked whether there were different diversity patterns among the samples. The alpha diversity (ShannonH) ranged from 1.7 to 2.9, but it was not significantly different between clades (p -value = 0.34)

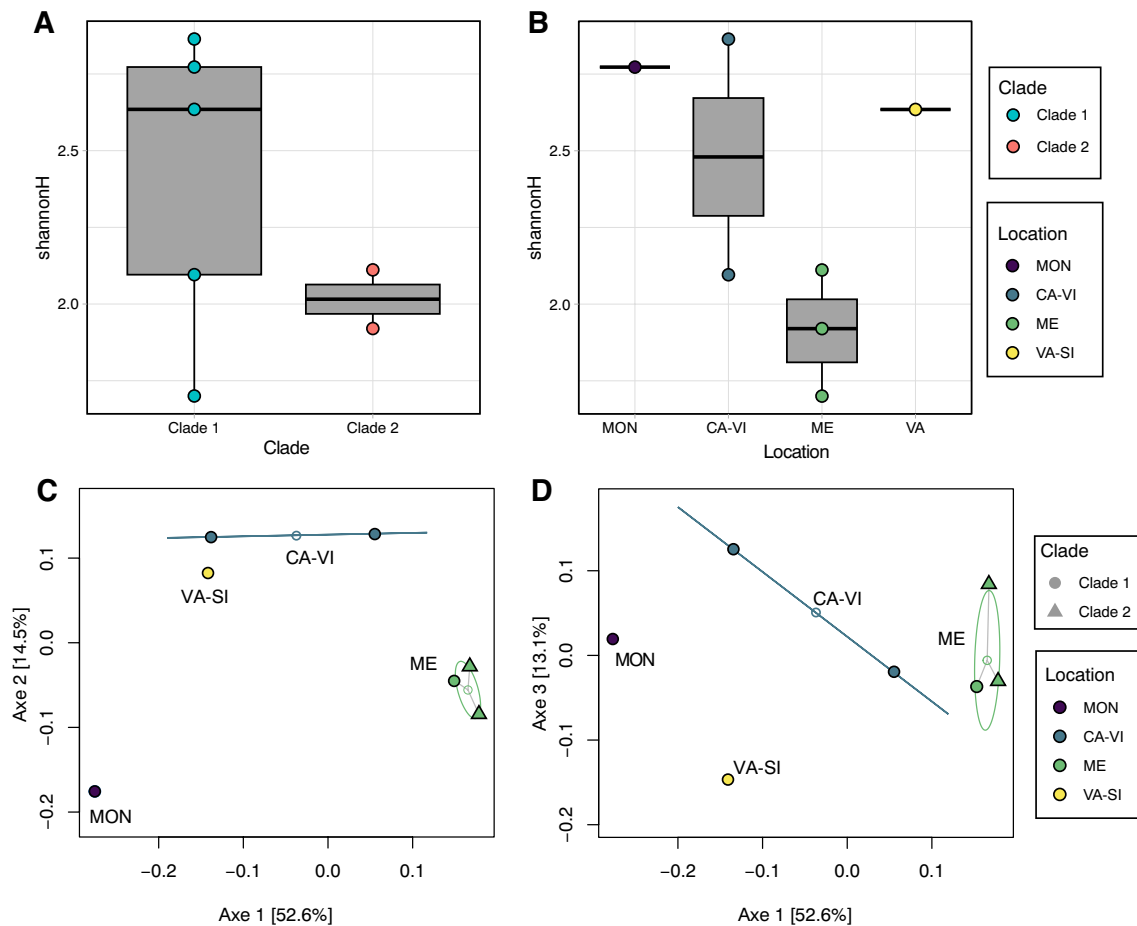


Fig. 7 Diversity and PCA Analysis. A-B. Shannon diversity for samples arranged by clade (A) and location (B). C-D. Principal Coordinate Analysis (PCoA) plots of beta diversity of the algal microbiomes based on Bray–Curtis dissimilarity at species level. Only the host

algal geographic location was a statistically significant grouping factor. Note that samples SOOKE and LOC are grouped in CA-VI, and PRAG, DPAG, and PR6 in ME

or location (p -value = 0.44) likely due to a large variability among the samples (Fig. 7A and B). Considering only CA (SOOKE and LOC) and ME (PRAG, DPAG, and PR6), there was marginally significant differences (p -value = 0.06).

To understand the structure in the composition of the microbial communities, we ordinated the samples using a principal coordinate analysis (PCoA) based on Bray Curtis dissimilarity (Fig. 7). The first two coordinate axes explain 73.6% of the total variance observed in our sampling (Fig. 7C and D). The ordination plot revealed that the samples have a tendency to group by geographic location, however, the phylogenetic relationship of the host (algal clades) does not have a strong influence in the microbial assemblage. PERMANOVA test showed significant clustering by location (p = 0.005), but not by sponge clade (p = 0.14). Assumption of homogeneity of dispersion within groups was not refuted (p -value > 0.1).

These algal culture samples were compared to the microbiome of the adult sponges originally collected from three

of the same locations as the algal isolates: CA-VI (which includes Sooke River (SOOKE) and Lake O'Connor (LOC)) and Maine (ME, Pemaquid River or PR). The sponge sample PR6_Em was the host from which the algal strain PR6 was sequenced. Sponge samples were sequenced with a different technology (File S1), therefore only a gross comparison at taxonomic levels of phylum and order was made. Similar to the algal samples, Cyanobacteria was the dominant group in the sponge microbiome (except for 3 samples in SOOKE), followed by Proteobacteria (Fig. 8). Many sponge samples, however, were characterized by high abundance of Bacteroidetes, that were not especially abundant in the algal microbiomes (Fig. 8). Similarly, sponge samples harbored medium abundances of Actinobacteria, Verrucomicrobia and Spirochaetes, that were decreased in the algal samples (Fig. 8A). Interestingly, similar to what occurs with the algal culture microbiomes (Figs. 6 and 8), sponge microbiomes from SOOKE were more similar to those collected in O'Connor Lake. At order level, the microbiomes of sponges

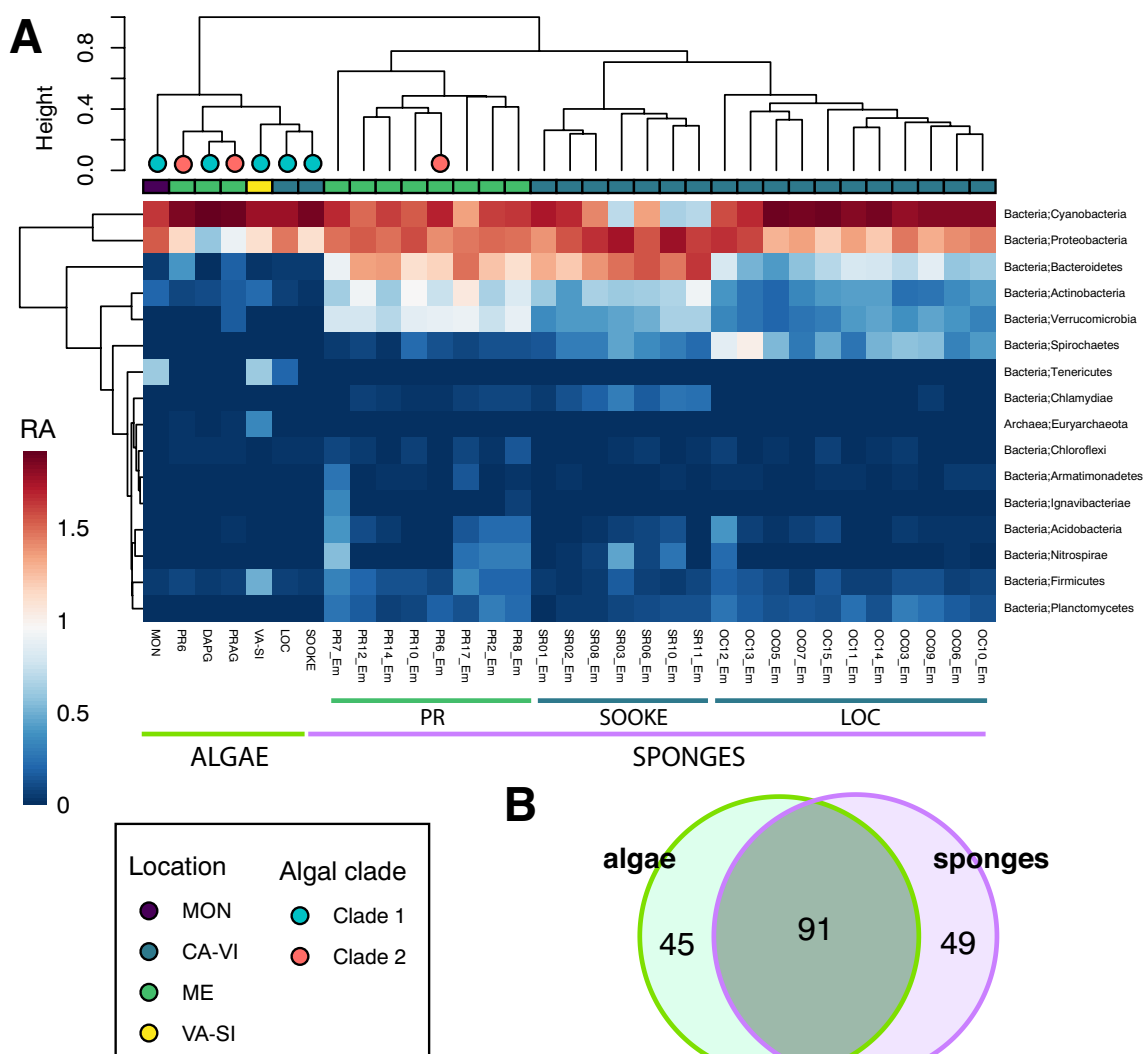


Fig. 8 Comparison of microbiota from algal isolates and sponges. **A.** Community composition comparison of the microbiota in the algal isolates and the sponge microbiome from *E. muelleri* at phylum taxonomic level represented as heatmap of log (relative abundance) and

dendrogram calculated over dissimilarities at species level. **B.** Venn diagram showing the number of bacterial and archaeal orders shared by the samples of algal cultures and sponge hosts compared

and algae shared 91 bacterial orders (Fig. 8B), that included Micrococcales (Actinobacteria), Chitinophagales and Flavobacteriales (Bacteroidetes), Pseudonabaenales, Nostocales, and Synechococcales (Cyanobacteria), Hyphomicrobiales, Rhodospirillales, Caulobacterales, Rhodobacterales, and Sphingomonadales (Alphaproteobacteria), and also Eubacteriales (Firmicutes).

4 Discussion

Endosymbionts found in *E. muelleri* from multiple locations in North America are green, coccoid, *Chlorella*-like microalgae (Fig. 1) that reside in a variety of sponge cells (Fig. 3). The transmission of algal symbionts in freshwater

sponges likely occurs both horizontally, through ingestion of algae from the environment, and vertically during fragmentation, budding, or gemmulation (recently reviewed in Ereskovsky et al. 2022). To our knowledge, algae have not been observed in eggs or larvae of freshwater sponges. Whether or not algal symbiont transmission from asexual gemmule to adult sponge occurs via arrested phagocytosis of algal cells in differentiating thesocytes (i.e., stem cells stored in overwintering gemmules) or from algal symbionts that are already endosymbiotic in thesocytes is unknown. We suspect that both could happen given the fact that some of the algal strains we isolated were from gemmules or hatched gemmules that had been washed in hydrogen peroxide to kill microbes on the gemmule surface, and we found algae in the gemmule cyst both inside and outside of thesocytes (Fig. 2).

While some algae in gemmules may be phagocytosed resulting in aposymbiotic sponges or sponges that do not show signs of algal division (e.g., Rasmont 1970; Simpson 1984), other algae avoid phagocytosis and persist in sponge cells (Fig. 3, Hall et al. 2021).

4.1 Algal identities

There is a moderate diversity of genera associated to freshwater sponges, which include *Choricystis*, *Lewinosphaera*, and *Chlorella*, all notoriously difficult to identify based on their morphology. In this sense, several authors have used molecular markers to identify endosymbiotic algal species coming from sponge tissues in the last decades (e.g., Annenkova et al. 2011; Pröschold et al. 2011; Kulakova et al. 2014, 2020; Pröschold and Darienko 2020). In *E. muelleri*, we identified at least two species of green coccoid algae that can form an endosymbiosis with host sponge cells. We found that both species belong to the Chlorellaceae with the predominant species found in Canada and the U.S. to be closely related to *Micractinium*, *Chlorella*, *Auxenochlorella*, and *Pseudochlorella* genera, with the definitive identity varying depending on the molecular marker and available data set used. The SSU-ITS phylogenies revealed that the closest relative to our isolated algal strains from Canada, Virginia and from Dundee Pond in Maine (clade 1 in all trees) is *Auxenochlorella pyrenoidosa*, a previously known algal endosymbiont. Although the genus *Auxenochlorella* has not been described to live within sponges before, a strain of *Auxenochlorella* was identified from *Hydra viridis* tissue (Darienko and Pröschold 2015). Based also on SSU-ITS data, we suggest that the algal strains belonging to clade 2, which were isolated from *E. muelleri* gemmules and an adult sponge from the Pemaquid River in Maine, are *Lewinosphaera symbiontica*, a species described by Pröschold and Darienko (2020) that was originally isolated in 1956 in Massachusetts in endosymbiosis with *Spongilla lacustris*. It is interesting to note that both PRAG and *L. symbiontica* have a 465 bp deletion in the 5' region of the SSU-ITS. While we do not have additional sequences from *L. symbiontica* for other genes we sequenced (TufA, rbcL, 16S), the sequences for PRAG and PR6 are identical for those markers (Figs. 4, S1, and S2).

We also isolated strains of green algae from *E. muelleri* adult tissue (and not gemmules) that are not likely endosymbionts, but algae associated with the sponge tissue or from the water surrounding and within the sponge. We did not find these algae in intracellular locations, but consistently found these strains associated with *E. muelleri* at two geographical locations in Maine (Dundee Pond and Pemaquid River). The SSU/ITS marker showed that each of these strains belong to the Scenedesmeaceae and are most closely related to *Scenedesmus*, *Desmodesmus*, and *Tetradesmus* algal species (File

S2). We do not know if these algae play roles in the sponge trophic biology, but they are consistently present in samples of *E. muelleri* sponges we collect from these locations.

Given that *E. muelleri* can live with and without their algal endosymbionts and that the algae can live inside the host cells or outside of the animal, we believe that this facultative symbiosis provides a good model for further study regarding the mechanisms that regulate the symbiotic relationship, the evolution of host:symbiont specificity, as well as the geographical factors that might influence sponge-microbe mutualisms.

4.2 Algal microbes

While it is well known that sponges have complex microbiomes, so far, the nature of the microbes that may be associated with their algal endosymbionts has not been explored. Algae and bacteria can influence each other's metabolism, physiology, and growth through nutrient exchange, and they can mutually influence aquatic ecosystems including relationships between symbiotic algae and their animal hosts (reviewed in Ramanan et al. 2016).

Microalgae are surrounded by a phycosphere, a region rich in organic material made by the alga that serves as an association/exchange network for other organisms, like bacteria, to interact (Cirri and Pohnert 2019). Microalgal-bacteria interactions that influence algal growth and biomass production are supported by an array of compounds produced by bacteria living in communities together with algae (Fuentes et al. 2016) and it has been suggested that the positive effects of algal-bacterial interactions on algal growth should influence future research to move beyond consideration of algal associated bacteria as only contaminants (Astafyeva et al. 2022). We observed that algal endosymbionts of *E. muelleri* grow well in cultures that include bacteria that are co-isolated along with the algae from either adult sponge cells or from gemmules or hatched gemmules. We also found that it was difficult to isolate some of the algae from their associated bacteria growing in cultures (Fig. 5) coming from poriferan sources (i.e., multiple rounds of colony isolation on agar plates were needed to get axenic cultures). In fact, we frequently observed bacterial/yeast colonies growing in predictive patterns with the algae (Fig. 5) and documented algae growing on top of the other microbes in culture.

4.3 Algal culture and sponge associated bacteria

Recent studies in *Hydra* indicate that interactions of the *Chlorella* algal symbiont and extracellularly located microbiota shape the host microbiome (Bathia et al. 2022). The possibility that associated bacteria are important to the algal symbionts or to the endosymbiosis within sponges led us to characterize the algal microbiomes of

each endosymbiont strain isolated from *E. muelleri* collected from different geographical locations (Fig. 6). To our knowledge, our study is the first to report the microbiota of endosymbiotic *Chlorella*-like algae isolated directly from environmental samples. A recent study has characterized the prokaryotic microbial community structure in *Chlorella vulgaris* SAG 211–12 cultures under different cultivation conditions in biotechnological systems (Haberkorn et al. 2020). In Haberkorn et al. (2020), the described bacterial community has originated from non-sterile handling of the pre-culture thus represents contamination that does not affect the algal growth in biotechnological reactors under different cultivation conditions. On the contrary, the algae prokaryotic community described here is representative of the microbial community that endosymbiotic *Chlorella*-like algae harbor in the wild.

In our study, each algal isolate possessed a unique microbiome signature, although there were some microbial families that were common to all microbiomes studied, and even to the microbiome of the host sponges (Figs. 6 and 8). The main phyla associated with the microbial communities present in endosymbiotic algal cultures isolated from *E. muelleri* were Cyanobacteria and Proteobacteria; though Bacteroidetes, Actinobacteria, Firmicutes, Tenericutes, and Euryarchaeota among a few others were also predominantly observed, but at lower frequencies. Some of the main families found in these microbiomes (i.e., Comamonadaceae, Caulobacteraceae, Chitinophagaceae, and Sphingomonadaceae) were also revealed from 16S sequencing of cultured strains of *Chlorella saccharophila* (Krohn-Molt et al. 2017). We acknowledge that long term cultivation of some of our strains (i.e., VA-SI, MON) may have resulted in inadvertent selection of distinct bacterial communities, but we believe that this baseline microbiome data provides information about microbes that can be associated with these endosymbiotic strains. The diversity patterns of the microbiomes of the algal cultures were independent of the identity of the algae, and they seem to be closely associated to either the geographical site and/or the host from which the algae were isolated (Fig. 7 and 8). This could indicate a strong influence of the environmental prokaryotes from which the host select their microbiomes, since these are Low Microbial Abundance sponges, with their microbiomes highly influenced by their horizontal acquisition strategy (Díez-Vives et al. 2022).

In summary, the fact that the microbiomes of the algal cultures and the sponge host microbiomes are so similar, sharing a high proportion of their taxa, strongly points to a crucial role of the prokaryotic symbionts in the growth and metabolic stability of both the algae and sponges. Freshwater sponges and their algal partners offer many opportunities to examine important questions in host:symbiont interactions from molecular, genetic, cellular, developmental, ecological, and evolutionary perspectives.

Supplementary Information The online version contains supplementary material available at <https://doi.org/10.1007/s13199-023-00934-8>.

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Data Availability All data is provided in figures and supplemental.

Declarations

Competing interests The authors have no competing interests to declare that are relevant to the content of this article.

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