



# Algae from *Aiptasia egesta* are robust representations of Symbiodiniaceae in the free-living state

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## ABSTRACT

Many cnidarians rely on their dinoflagellate partners from the family Symbiodiniaceae for their ecological success. Symbiotic species of Symbiodiniaceae have two distinct life stages: inside the host, *in hospite*, and outside the host, *ex hospite*. Several aspects of cnidarian-algal symbiosis can be understood by comparing these two life stages. Most commonly, algae in culture are used in comparative studies to represent the *ex hospite* life stage, however, nutrition becomes a confounding variable for this comparison because algal culture media is nutrient rich, while algae *in hospite* are sampled from hosts maintained in oligotrophic seawater. In contrast to cultured algae, expelled algae may be a more robust representation of the *ex hospite* state, as the host and expelled algae are in the same seawater environment, removing differences in culture media as a confounding variable. Here, we studied the physiology of algae released from the sea anemone *Exaiptasia diaphana* (commonly called *Aiptasia*), a model system for the study of coral-algal symbiosis. In *Aiptasia*, algae are released in distinct pellets, referred to as *egesta*, and we explored its potential as an experimental system to represent Symbiodiniaceae in the *ex hospite* state. Observation under confocal and differential interference contrast microscopy revealed that *egesta* contained discharged nematocysts, host tissue, and were populated by a diversity of microbes, including protists and cyanobacteria. Further experiments revealed that *egesta* were released at night. In addition, algae in *egesta* had a higher mitotic index than algae *in hospite*, were photosynthetically viable for at least 48 hrs after expulsion, and could competently establish symbiosis with aposymbiotic *Aiptasia*. We then studied the gene expression of nutrient-related genes and studied their expression using qPCR. From the genes tested, we found that algae from *egesta* closely mirrored gene expression profiles of algae *in hospite* and were dissimilar to those of cultured algae, suggesting that algae from *egesta* are in a nutritional environment that is similar to their *in hospite* counterparts. Altogether, evidence is provided that algae from *Aiptasia egesta* are a robust representation of Symbiodiniaceae in the *ex hospite* state and their use in experiments can improve our understanding of cnidarian-algal symbiosis.

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## INTRODUCTION

The family Symbiodiniaceae is a diverse group of dinoflagellates, many of which form symbioses with a variety of cnidarian hosts, including corals and sea anemones. In this symbiosis, the algal symbiont is housed within host gastrodermal cells where they provide photosynthate to the host and in return, the host provides inorganic nutrients, protection and a high light environment (Davy, Allemand & Weis, 2012; Marcelino et al., 2013). The two partners must work cooperatively to maintain and regulate symbiosis. For example, the host utilizes carbonic anhydrase and bicarbonate transporters to supply inorganic carbon to the symbiont to maximize photosynthesis (Tansik, Fitt & Hopkinson, 2017; Koch, Verde & Weis, 2020). Reciprocally, the symbiont may be modulating the host's immune system to prevent immune destruction or expulsion by the host (Detournay et al., 2012; Mansfield et al., 2017; Jacobovitz et al., 2021; Jinkerson et al., 2022). However, our understanding of the cellular mechanisms involved in maintaining symbiosis remains limited.

Studies often compare the biology of Symbiodiniaceae between its two life stages: inside the host, *in hospite* and outside the host, *ex hospite*. In this comparison, algae that are freshly isolated from the host are used to represent algae *in hospite*, while cultured algae are often used to represent algae *ex hospite*. Studies have found that compared to algae in culture, algae *in hospite* have lower cell division rates (Smith & Muscatine, 1999; Tivey, Parkinson & Weis, 2020) and differential expression of genes that are tightly linked to nutrient limitation (Maor-Landaw, Van Oppen & McFadden, 2020; Xiang et al., 2020; Cui et al., 2022). Based on these data, it is commonly hypothesized that the host limits nutrient transfer to the symbiont as a mechanism to maintain a stable population of symbionts (Falkowski et al., 1993; Smith & Muscatine, 1999; Xiang et al., 2020; Tivey, Parkinson & Weis, 2020; Cui et al., 2022). This comparison, however, introduces nutrition as a confounding variable because while algae *in hospite* are sampled from hosts maintained in oligotrophic seawater, algae *ex hospite* are sampled from cultures grown in nutrient-rich media. The nutrient-depleted phenotype found in algae *in hospite* could therefore not be caused by host-inhibitory mechanisms and instead may simply reflect the low nutrient availability in the water column surrounding hosts (for in-depth discussion, see Maruyama & Weis, 2021). In fact, studies have found that supplementing nutrients to the host, by feeding or by the addition of inorganic nutrients to the water column, will return the algae to a nutrient-enriched phenotype as measured by cell division rates or gene expression profiles, suggesting that the host readily supplies nutrients to their symbionts when available (Hoegh-Guldberg & Smith, 1989; Stambler et al., 1991; Falkowski et al., 1993; Hoegh-Guldberg, 1994; Muller-Parker et al., 1994; Smith & Muscatine, 1999; Ferrier-Pagès et al., 2001; Houlbrèque et al., 2004; Rosset, D'Angelo & Wiedenmann, 2015; Xiang et al., 2020; Tivey, Parkinson & Weis, 2020; Cui et al., 2022). Further work is necessary to tease apart the effects of symbiosis and nutrient availability on algal phenotypes to determine whether the host is limiting nutrient transfer to its symbionts.

Multiple transcriptomic studies have found that nutrient-related genes are differentially expressed between algae *in hospite* and those in culture (Maor-Landaw, Van Oppen & McFadden, 2020; Xiang et al., 2020; Cui et al., 2022). Examples include ammonium

transporter *AMT*, nitrate transporter *NRT*, purine nucleoside permease *NUP*, and nitrate reductase *NR* that are upregulated and glutamine synthetase *GS* that is downregulated in algae *in hospite* compared to those in culture. This gene expression pattern is indicative of nutrient limitation in Symbiodiniaceae and other marine algae (Hildebrand, 2005; Kang et al., 2007; Xiang et al., 2020). Some differentially expressed genes, however, are probably not linked to nutrition, such as the sugar transporter *SWEET1*, that is hypothesized to play a role in transferring photosynthate to the host. Sampling a more robust representation of Symbiodiniaceae in the *ex hospite* state that controls for nutrient availability could confirm whether these differentially regulated genes are reflective of nutrition, the symbiotic state, or both.

In cnidarian-algal symbiosis, the host regularly expels viable algae as a homeostatic mechanism to maintain a stable population of symbionts (Steele, 1975; Hoegh-Guldberg, McCloskey & Muscatine, 1987; Hill & Ralph, 2007; Thornhill et al., 2017). In this study, we hypothesized that sampling expelled algae as a representation of the *ex hospite* state in Symbiodiniaceae for comparison with their *in hospite* counterparts could remove nutrition as a confounding variable, as both the host and the expelled algae are in the same oligotrophic seawater environment. Therefore, physiological and/or gene expression differences found between algae *in hospite* and expelled algae in this comparison would be modulated by the host, and not be masked by the high-nutrient state of cultured algae as in the traditional comparison. In the sea anemone *Exaiptasia diaphana* (commonly called Aiptasia), a model system for the study of cnidarian-algal symbiosis, algae are released in distinct, brown pellets, herein referred to as egesta. The brown egesta primarily consist of algae, and are released separately from pellets containing digested food (Steele, 1975). The discrete nature of egesta, with their high abundance of algae, made them a promising experimental system for sampling purposes.

To begin the practice of sampling expelled algae in symbiosis studies, we first explored the basic biology of algae from Aiptasia egesta and characterized egesta appearance, timing of egesta expulsion, and characteristics of the algae including mitotic index, photophysiology, and ability to initiate symbiosis with new hosts. Then, we studied the gene expression of several nutrient-related genes to determine the nutrient status of algae from egesta. Finally, we discuss several practical considerations of the use of algae from Aiptasia egesta as an experimental model system to represent Symbiodiniaceae in the *ex hospite* state.

## MATERIALS & METHODS

### Animal and algal culture maintenance

Aiptasia (*Exaiptasia diaphana*, clone ID: H2) symbiotic with *Breviolum minutum* (culture ID: SSB01) and SSB01 cultures were used in this study. Animals were maintained in Coralife Instant Ocean (Tempe, AZ, USA) artificial seawater (ASW), and algae were maintained in F/2 media made in filtered artificial seawater (FSW). Animals and algae were housed at 25 °C in a Percival AL-41L4 incubator (Perry, IA, USA) with Zoo Med Laboratories (San Luis Obispo, CA, USA) 10,000 k fluorescent lamps set to 40  $\mu\text{mol photons/m}^2/\text{s}$  using shade cloth on a 12 h:12 h light:dark cycle. Aposymbiotic anemones were originally bleached

using menthol ([Matthews et al., 2016](#)) and maintained symbiont-free by incubation in the dark for several months before use. Unless otherwise stated, animals were fed three times a week *ad libitum* with freshly hatched brine shrimp and the water was changed three times a week with ASW.

### Confocal imagery and light microscopy

Live egesta samples were imaged on a Zeiss LSM 780 NLO confocal microscope system (Zeiss, Oberkochen, Germany). Images were taken on three separate channels, with excitation from a 405 nm Diode, Argon (488), and 633 nm HeNe. Emission was detected between 410–470 nm, 491–572 nm, and 647–722 nm, respectively. Simultaneously, using laser illumination, a transmitted light detector (T-PMT) acquired transmitted light images in bright field. Confocal z-stack images were processed for pseudocoloring and merged to create a maximum intensity projection using ZEN Black (Carl Zeiss AG) and ImageJ software. Separately, differential interference contrast (DIC) images were obtained using an Olympus Vanox-T AH2 microscope to image nematocysts.

### Time-lapse of egesta release

Sixteen to twenty *Aiptasia* were transferred to twenty cm diameter glass Carolina Biological (Burlington, NC, USA) culture dishes with one L of ASW. Animals were allowed to settle for at least 24 hrs prior to time-lapse photography. Immediately before imaging began, the culture dish was cleaned with a cotton swab and the water was changed. For time-lapse imaging, a Canon 5D Mark II DSLR with external flash was set to photograph the entire dish hourly for 24 hrs. To control for any effect of cleaning on timing of egesta release, two time-lapses (Experiment 1 & 2) began at 16:30 and two other time-lapses (Experiment 3 & 4) began at 22:30. Following their original light schedule, lights were set to come on at 7:00 and off at 19:00. Animals were not fed during the experiment. Each image was analyzed with ImageJ using the Cell Counter plugin, and each dark particle in an image were counted as egesta. The number of egesta in each time point was subtracted from the total number of egesta in the previous time point to calculate the quantity of egesta released in the hour. The quantity of egesta was then normalized to anemone number per container to obtain the number of egesta per anemone at each time point.

### Mitotic index

Thirty-two *Aiptasia* were housed individually in two mL FSW in 24-well plates and allowed to settle for 24 hrs. After settlement, each well was cleaned with a cotton swab and the water was changed. Egesta were collected after an additional 24 hrs. Of the 32 original anemones, eight released egesta large enough to be imaged. The eight anemones that released usable egesta were homogenized with a microcentrifuge pestle in 300  $\mu$ L FSW, spun down at  $800\times g$  for five min, and resuspended in twenty  $\mu$ L FSW. Resuspended algae and egesta were then imaged under DIC light microscopy.

Images were analyzed using ImageJ with the Cell Counter plugin by students enrolled in Oregon State University's Spring 2021 Z362 Invertebrate Biology laboratory. Following the method of [Baghdasarian & Muscatine \(2000\)](#), individual cells in each image were scored based on presence of a cell wall division plate, and at least 700 cells were counted

per sample. Students were tasked to save their cell counter files, and their counts were manually verified by loading the counter files onto the images. The mitotic index was calculated as the percent of cells in the entire population that were undergoing division and paired *t*-tests were performed for statistical analysis.

### **Algal photophysiology and cell viability**

Egesta were collected from Aiptasia less than 24 hrs after last cleaning, to ensure collection of freshly expelled egesta. All collected egesta were pooled together and pelleted at  $800\times g$  for five min, the supernatant was removed and the cells resuspended in one mL FSW. The pooled sample was divided into two 500  $\mu\text{L}$  aliquots. One was kept intact, while the other was homogenized using a motorized microcentrifuge pestle. From the homogenized sample, twenty  $\mu\text{L}$  was used to quantify cell number using a Thermo Fisher Scientific (Waltham, MA, USA) Countess II automated cell counter. Homogenized and intact egesta samples were then dispensed into twelve well plates in 95  $\mu\text{L}$  (20,000 cells per sample,  $n = 5$ ) and FSW was added up to two mL. Cultured algae (SSB01) was also collected, pelleted at  $800\times g$  for five min and resuspended in FSW. Five culture samples of 20,000 cells were then dispensed in twelve well plates and FSW added up to two mL. All plates were then returned to their original incubator.

Maximum quantum yield of photosystem II (Fv/Fm) was obtained using a Light Induced Fluorescence Transient Fast Repetition Rate fluorometer (LIFT-FRRf; Soliense Inc, Santa Cruz, CA). Fv/Fm was measured once a day at 13:00 following dark acclimation for 30 min to ensure measurement of maximum quantum yield. To measure Fv/Fm, excitation was delivered at 475 nm wavelength in four phases and fluorescence detected at 685 nm. The first phase was a saturating sequence of 100 flashlets, each lasting 0.7  $\mu\text{s}$  with a 1.5  $\mu\text{s}$  gap. The second phase was a relaxation phase of 80 flashlets beginning with a gap of twenty  $\mu\text{s}$ , increasing exponentially until the end of the sequence. The third phase was a sequence of 1,600 flashlets lasting two  $\mu\text{s}$  each with a 40  $\mu\text{s}$  gap. The final relaxation phase was identical to the second phase. Data were fitted using LIFT software with the three-component exponential model (SEQ\_3).

To test the ratio of live and dead cells, we used an Evans Blue cell viability dye (*Morera & Villanueva, 2009*). Approximately ten pellets of egesta less than 24 hrs old were collected and homogenized together using a motorized microcentrifuge pestle in 500  $\mu\text{L}$  of ASW. Cells were then pelleted at  $800\times g$  for five min and the pellet resuspended in 100  $\mu\text{L}$  of FSW. Cells were then stained with 20  $\mu\text{L}$  of Evans Blue dye and incubated for ten min. The sample was imaged under light microscopy and algal cells that internalized the blue dye were counted as dead while cells that were not stained were counted as alive.

### **Inoculation of aiptasia with algae**

Aposymbiotic anemones ( $n = 9$ ) were plated into 24 well plates in two mL FSW. Plates were placed in their original incubators set on a 12 h:12 h light:dark cycle for 72 h. After incubation in the light, animals were manually checked under fluorescence microscopy to verify aposymbiotic status.

Egesta were collected from Aiptasia less than 24 hrs after last cleaning. Samples of both egesta and algae in culture were pelleted at  $800\times g$  for five min and resuspended in 300  $\mu\text{L}$

FSW. To obtain freshly isolated symbionts, one symbiotic anemone was homogenized using a motorized microcentrifuge pestle in 300  $\mu$ L of FSW. Egesta and cultured algae were also homogenized using a motorized microcentrifuge pestle. All samples were then washed twice by centrifugation at  $800\times g$  for five min and resuspension in two mL FSW. Algal samples were counted using a Countess II automated cell counter, and samples were diluted to  $1 \times 10^6$  algal cells/mL in FSW.

To inoculate aposymbiotic *Aiptasia*,  $1 \times 10^5$  algal cells in a 100  $\mu$ L volume were gently pipetted over anemone mouths. Immediately after the addition of algae, twenty  $\mu$ L of brine shrimp extract was pipetted to anemones to induce a feeding response. Negative controls were fed the brine shrimp extract with no algae added. Anemones were allowed to take up algae for 24 hrs. The water was then changed, and anemones were moved to clean wells of a 24 well plate in two mL FSW. After an additional 24 hrs, the anemones were imaged under fluorescence microscopy. Briefly, anemones were relaxed in 0.18 M  $MgCl_2$  dissolved in FSW, and images of three tentacles per anemone were taken in several focal depths under brightfield and under the red Filter Set 15 (Carl Zeiss) to visualize algal cell auto-fluorescence using a Zeiss Axio Observer inverted microscope. Each manually constructed z-stack was merged into a single image using Adobe Photoshop. Symbiont density was determined from each image using ImageJ as the number of fluorescent algal cells normalized to two-dimensional tentacle area.

### Primer design

Target genes were selected based on their function and differential expression between algae in culture and *in hospite* from [Xiang et al. \(2020\)](#). Primer and transcript sequences were obtained from [Xiang et al. \(2015\)](#) and [Xiang et al. \(2020\)](#) for an ammonium transporter (*AMT*, transcript ID: s6\_38207), a nitrate transporter (*NRT*, s6\_422), purine nucleoside permease (*NUP*, s6\_27864), and nitrate reductase (*NR*, s6\_34). New primers were designed for glutamine synthetase (*GS*, s6\_5551), photosystem II protein D1 (*psbA*, s6\_1009), and a sugar transporter (*SWEET1*, s6\_35311). We chose to study the expression of *psbA* because its expression was consistent between algae in culture and *in hospite* ([Xiang et al., 2020](#)). Cyclophilin was chosen as an algal housekeeping gene ([Rosic et al., 2011](#)) and primers were designed to be specific for *B. minutum* and not amplify *Aiptasia* cyclophilin. Primer pairs were verified for amplification efficiency of at least 90% by assessing the slope of standard curves generated from quantitative PCR reactions of serial dilutions of template cDNA. Primers sequences are available in the supplementary data.

### Quantitative PCR

To quantify gene expression, 54 anemones were placed in three separate containers with 500 mL ASW (eighteen anemones per container). The average oral disk diameter of the anemones was 5.4 mm (data not shown). Anemones were fed every other day *ad libitum*. Egesta containing algae were collected 24 hrs after feeding, and the container cleaned after egesta were collected. We did not sample egesta containing digested food and they are easy to distinguish due to the pink color of the carotenoids from digested *Artemia* sp. nauplii. To obtain enough material for gene expression studies (approximately  $1 \times 10^6$  cells), egesta

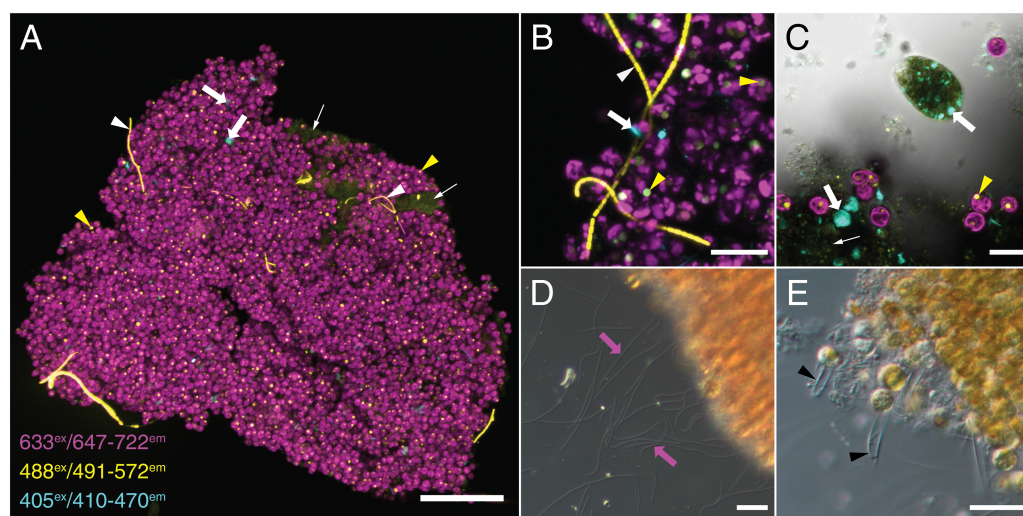
were collected a total of six times from each container. Immediately after collection, egesta were processed for RNA extraction as follows. First, egesta were homogenized in cold FSW with a motorized microcentrifuge pestle. To completely isolate algae from host cells, the sample was then further homogenized by passing the sample seven times through a three mL syringe fitted with a 23 gauge needle. Following protocols from [Xiang et al. \(2020\)](#), the homogenate was then loaded on a 50% isotonic Percoll solution made in FSW and spun for twenty min at  $9,000\times g$ . The supernatant containing host material was carefully removed, and the algae were resuspended in one mL cold FSW. The sample was then pelleted at  $3,000\times g$  for five min and the supernatant removed. The pellet was then frozen at  $-80\text{ }^{\circ}\text{C}$  until further processing.

To obtain algae *in hospite*, four random anemones from each container were sampled immediately after the last collection of egesta and processed for RNA extraction following the same procedure as those for egesta, and the algal pellets frozen for at least ten min. Three independent, but identical cultures of SSB01 grown in F/2 culture media were made simultaneously and allowed to grow for three weeks. These cultures were also sampled and followed the same procedure for RNA processing and the algal pellets were frozen for at least ten min.

After the final sample of egesta was collected, the frozen samples from each repeated collection were pooled together corresponding to their respective *Aiptasia* containers. Each frozen freshly isolated, culture, and egesta sample was then resuspended in  $300\text{ }\mu\text{L}$  Trizol. Sterilized glass beads were added to the sample, and the sample was homogenized with a bead beater for two min set at 4,000 RPM. The beads were then removed, the samples were centrifuged at  $16,000\times g$  for one min, and the supernatant containing RNA was extracted. The RNA was purified using a Zymo Direct-zol kit following the manufacturer's instructions. RNA samples were further processed using a Turbo DNA-free kit to remove genomic DNA contaminants. Samples were then purified again using a New England Biolabs (Ipswich, MA, USA) Monarch RNA purification kit, followed by a One-Step PCR inhibitor removal kit to remove carbohydrate contaminants. The RNA samples were analyzed for purity and concentration using a Nanodrop.

All samples were then diluted to ten ng/ $\mu\text{L}$  and cDNA was synthesized from nine  $\mu\text{L}$  (90 ng total) of RNA in  $30\text{ }\mu\text{L}$  reactions using the New England Biolabs Protoscript II cDNA synthesis kit with oligo d(T)<sub>23</sub> VN primers following the manufacturer's instructions. Quantitative PCR reactions were performed in twenty  $\mu\text{L}$  volumes using 1X Power SYBR Green PCR Mastermix,  $0.5\text{ }\mu\text{M}$  of each primer pair, and one  $\mu\text{L}$  of template cDNA. A BioRad CFX96 Real-Time System was used to carry out reactions using a two-step amplification phase ( $95\text{ }^{\circ}\text{C}/\text{ten min}$ , followed by 40 cycles of  $95\text{ }^{\circ}\text{C}/\text{ten s}$  and  $58\text{ }^{\circ}\text{C}/30\text{ s}$ ) and then a melting-curve analysis performed to confirm the presence of single amplicons. No-reverse transcription, no-primer, and no-template controls were included as negative controls.

For each sample and target gene,  $\Delta\text{Ct}$  values were calculated by subtracting the Ct value of the target gene from the Ct value of the housekeeping gene (cyclophilin). Then, the  $\Delta\text{Ct}$  value was subtracted from the mean  $\Delta\text{Ct}$  value of corresponding genes from cultured algae samples to calculate relative expression  $\Delta\Delta\text{Ct}$ . Fold gene expression was then calculated



**Figure 1** Confocal and DIC images of whole *Aiptasia egesta*. (A–C) Confocal images of whole *Aiptasia egesta*. Magenta: *Breviolum minutum* chlorophyll is detected with excitation at 633 nm and emission at 647–722 nm. Yellow: Cyanobacteria phycobillin (white arrowhead), host tissue (small white arrow), and *B. minutum* pyrenoids (yellow arrowhead) are detected with excitation at 488 nm and emission at 491–572 nm. Cyan: Pigment from an unknown protist (white arrow) is detected with excitation at 405 nm and emission at 410–470 nm. (D–E) Fired nematocysts (magenta arrows) and empty capsules (black arrowheads) are seen on the superficial surface of egesta under DIC microscopy. Scale bars: (A) 100  $\mu\text{m}$  and (B–E) 20  $\mu\text{m}$ .

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as  $2^{-\Delta\Delta\text{Ct}}$  for each gene from each sample. Statistical analyses were conducted on  $\Delta\Delta\text{Ct}$  values for each gene by ANOVA and *post-hoc* Tukey's HSD.

## RESULTS

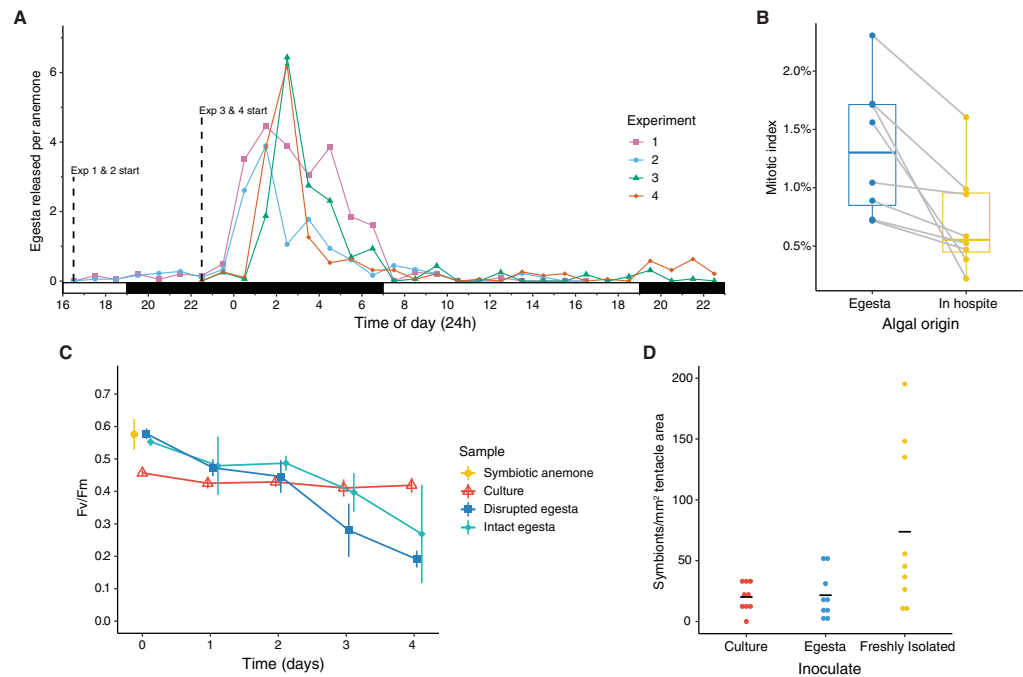
### Physical characteristics of egesta

Egesta observed under confocal microscopy revealed several autofluorescent features (Fig. 1). Chlorophyll autofluorescence was readily apparent from *B. minutum* under 633 nm excitation. Under 488 nm excitation, autofluorescence of putative pyrenoids from *B. minutum*, phycobillins from cyanobacteria, and host tissue were detected (Fig. 1A). A signal was also detected under 405 nm excitation within egesta (Fig. 1A), and we hypothesize that its origin comes from dead remains of an unidentified ciliate protist that autofluoresced under the same excitation wavelength (Fig. 1C). Differential interference contrast microscopy revealed that the surface of egesta contained discharged nematocysts (Fig. 1D) and the remains of nematocyst capsules (Fig. 1E). Egesta were also colonized by several kinds of unidentified protists and microbes (data not shown). None of the protists were observed to phagocytose *B. minutum*.

### Algal physiology

Egesta were released at night, beginning five to six hrs after lights off and peaking at seven to eight hrs after lights off (Fig. 2A). The rate of release slowly decreased until lights were turned on, and release decreased to background levels at the onset of light. The timing of



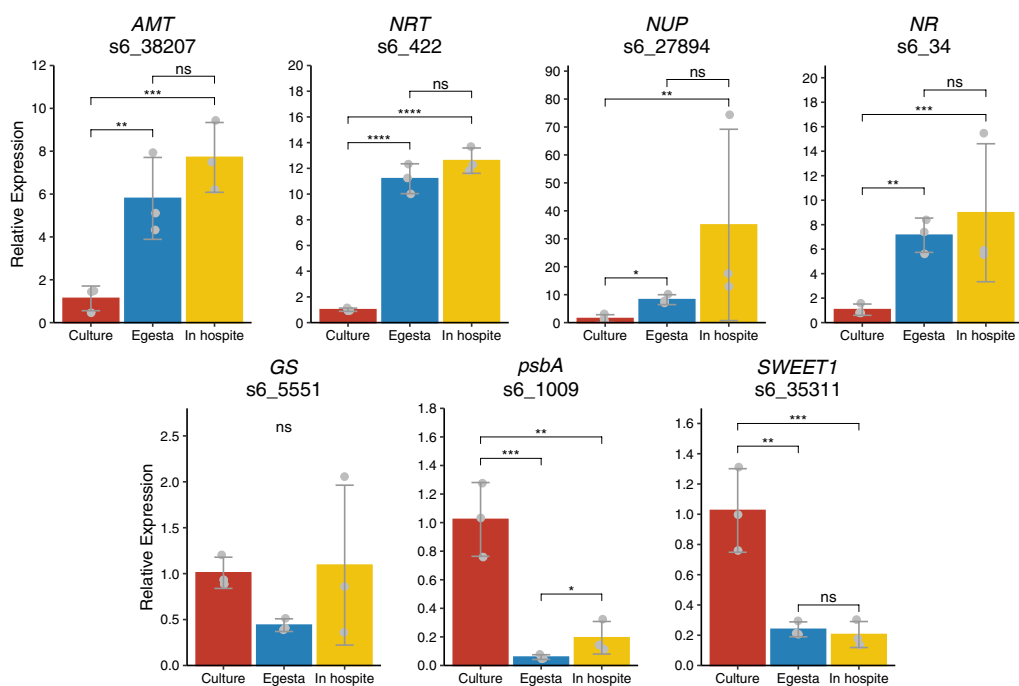


**Figure 2** The physiology of algae from egesta. (A) Amount of egesta released per anemone every hour over 24 hours. Experiments 1 and 2 began at 17:00 and experiments 3 and 4 began at 23:00. Black and white bars on *x*-axis depict dark and light periods, respectively. (B) The mitotic index of algae in egesta and in the host. Dots indicate individual samples and lines connecting dots indicate paired samples. The difference was statistically significant ( $p = 0.01$ , Paired *t*-test). (C) The maximum quantum yield of photosystem II of algae over time as measured by FRRf. Vertical lines indicate standard deviations around the mean. (D) Symbiont density in aposymbiotic *Aiptasia* inoculated with algae in culture, from egesta, or freshly isolated from hosts. Horizontal lines indicate mean symbiont density. Differences were not statistically significant ( $p = 0.06$ , Kruskal-Wallis test).

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dish cleaning prior to time-lapse imaging had a minor effect on timing of egesta release by delaying the time of peak egesta release by one hr (Fig. 2A). We noticed that the size of egesta was not uniform (data not shown)—presumably containing different amounts of algae, but we were unable to quantify the size of egesta and the number of symbionts per egestum in our experiments. Therefore, the time series data reflects only the number of discrete pellets that were released by anemones and is a proxy for the true number of expelled algae.

The released algae were alive and biologically active. The mitotic index of algae from egesta was higher than that of algae *in hospite* (Fig. 2B;  $p = 0.01$ , paired *t*-test), and the Fv/Fm of algae in egesta, disrupted and intact, at time zero was the same as those in symbiotic anemones (Fig. 2C). Algae from egesta then declined in Fv/Fm over time, with Fv/Fm remaining relatively high for two days before declining rapidly, with chlorophyll fluorescence becoming unmeasurable by day five. Mechanically disrupting the egesta did not have a significant impact on algal photophysiology. Algae from culture started at a lower Fv/Fm than those from egesta or anemones, but Fv/Fm did not decline over time. Cell staining with Evans Blue dye found that 1.5% (17 out of 1,132 total cells) of algae from



**Figure 3** Gene expression of nutrient-related, photosynthesis, and sugar transporting genes between cultured algae, algae from egesta, and algae *in hospite*. With the exception of GS, Expression of genes was similar in algae from egesta and algae *in hospite* compared to that of cultured algae. Bars indicate standard deviations around the mean. Transcript IDs are listed under gene abbreviations based on [Xiang et al. \(2015\)](#). Stars indicate statistically significant differences calculated on  $\Delta\Delta Ct$  values as determined by ANOVA and post-hoc Tukey's tests. \*  $p < 0.05$ , \*\*  $p > 0.01$ , \*\*\*  $p < .0001$ , \*\*\*\*  $p < 0.00001$ .

Full-size [DOI: 10.7717/peerj.13796/fig-3](https://doi.org/10.7717/peerj.13796/fig-3)

egesta were dead. Algae from egesta were capable of initiating symbiosis with aposymbiotic *Aiptasia*, reaching symbiont densities just as high as *Aiptasia* inoculated with cultured and freshly isolated algae (Fig. 2D). Negative controls remained aposymbiotic for the duration of the experiment (data not shown). While freshly isolated symbionts reached the highest density of symbionts, the difference was not significant ( $p = 0.06$ , Kruskal–Wallis test).

### Gene expression

Nitrogen transporter and nitrogen metabolism genes, *AMT*, *NRT*, *NUP*, and *NR*, were significantly upregulated in algae from egesta and algae *in hospite* compared to culture (Fig. 3). Expression levels of these genes were not significantly different between algae from egesta and algae *in hospite*. Glutamine synthetase, *GS*, gene expression showed no significant difference between algal sources. Expression levels of *psbA* and sugar transporter, *SWEET1*, were highest in cultured algae compared to algae from egesta and algae *in hospite*. Overall, with the exception of glutamine synthetase, expression levels of the tested genes were similar between algae from egesta and algae *in hospite* compared to algae in culture.

## DISCUSSION

### Egesta are sticky communities of microbes

Egesta primarily consisted of *B. minutum* as previously reported (Steele, 1975) and these egesta were released separately from digested food. In addition, egesta contained host material and a variety of microbes (Fig. 1). In sample handling, we noted that egesta were sticky and tended to adhere to plastic surfaces. The stickiness is possibly caused by the presence of fired nematocysts that were present on the egesta surface (Figs. 1D and 1E). The presence of nematocysts, both fired and unfired, were previously described in algae-containing egesta from the sebae anemone, *Heteractis crispa* (Alan Verde, Cleveland & Lee, 2015). The authors noted that egesta released by *H. crispa* were consumed by symbiotic anemonefish (Alan Verde, Cleveland & Lee, 2015). It is not known if other animals consume Aiptasia egesta in nature, but if so, they may aid in algal dispersal, as algae often survive digestion (Parker, 1984; Grupstra et al., 2021; Grupstra et al., 2022).

### Evidence for the preferential expulsion of dividing algae

The mitotic index of algae was higher in egesta than *in hospite*, a result that agrees with previous findings in Aiptasia and several corals (Baghdasarian & Muscatine, 2000). It was hypothesized by Baghdasarian & Muscatine (2000) that the host preferentially expels dividing algae. This hypothesis is well-supported from a temporal perspective because the egesta were released at night, and several studies have shown that peak cell division occurs at night in cultured Symbiodiniaceae (Fitt & Trench, 1983; Smith & Muscatine, 1999; Yamashita & Koike, 2016; Fujise et al., 2018). In contrast, algal cell division was not elevated at night *in hospite* in Aiptasia (Smith & Muscatine, 1999). In that study, however, the dividing algae may have been expelled by the host prior to sampling, leading to the detection of unchanged cell division at night (Smith & Muscatine, 1999). Measuring the mitotic index of algae from egesta that are released over a diurnal cycle may help complete the picture of algal population dynamics *in hospite*.

Nutrition may also explain the higher division rates in algae from egesta compared to those *in hospite*. As nutrient availability has been shown to strongly predict cell division rates in Symbiodiniaceae (Smith & Muscatine, 1999; Karako-Lampert et al., 2005; Tivey, Parkinson & Weis, 2020), the host may be preferentially expelling algae that sequester more nutrients for cell division than their more cooperative counterparts that remain undivided *in hospite*. Indeed, in competitive inoculation experiments, the symbiont *Durusdinium trenchii*, known to sequester more nitrogen than *B. minutum* in Aiptasia (Sproles et al., 2020), failed to proliferate in Aiptasia in the presence of the homologous *B. minutum* (Gabay et al., 2019). The preferential expulsion of dividing algae could be a mechanism for the removal of uncooperative symbionts. Future experiments using methods such as stable isotope analysis and NanoSIMS could investigate whether expelled algae sequester more nutrients than their counterparts *in hospite* and whether they continue to sequester more nutrients when establishing symbiosis with new hosts.

### Timing of algal release varies depending on host taxa

The night-time release of algae in egesta by *Aiptasia* is not shared by other hosts. In several corals, including *Pocillopora damicornis* and *Acropora digitifera*, expulsion rates increased with the onset of light and peaked in the middle of the day (Stimson & Kinzie, 1991; Koike et al., 2007). Other studies in the coral *Stylophora pistillata*, the soft coral *Xenia macrospiculata*, and the giant clam *Tridacna crocea* found no clear diel pattern for algal release (Hoegh-Guldberg, McCloskey & Muscatine, 1987; Umeki et al., 2020). Some corals released algae at night, with peak expulsion rates occurring at night for the corals *Millepora dichotoma* and *Heteroxenis fuscescens* (Hoegh-Guldberg, McCloskey & Muscatine, 1987). These contrasting patterns may reflect the actual differences in patterns of algal release between species, or it could reflect differences in sampling methodology. As peak algal motility occurs at midday for Symbiodiniaceae (Fitt & Trench, 1983; Yamashita & Koike, 2016), some sampling approaches could miss subpopulations of algae in the vessel depending on time of sampling.

### Algae from egesta are competent symbionts, but are short-lived *ex hospite*

Algae from egesta were short-lived *ex hospite* compared to their cultured counterparts, possibly due to nutrient limitation and/or the unknown effects of being in a microbial community (Fig. 2C). This supports the hypothesis that expelled algae do not form stable populations *ex hospite* (Thornhill et al., 2017). However, freshly expelled algae from egesta had high viability and were fully capable of initiating symbiosis with aposymbiotic hosts (Fig. 2D). In nature, larval and juvenile recruits probably rely on the continuous release of algae from adult hosts for a source of symbionts (Thornhill et al., 2017). This is supported by studies that found that the presence of adult coral colonies, presumably releasing algae, aided in symbiont acquisition by juvenile corals in aquarium experiments (Nitschke, Davy & Ward, 2016; Ali et al., 2019). In addition, a study found that a giant clam harboring Symbiodiniaceae released viable symbionts that were able to establish symbiosis with *Acropora tenuis* larvae (Umeki et al., 2020). Egesta are also potential vectors for the transmission of the microbiome between adults and recruits. In corals, there is evidence that *Acropora tenuis* and *Pocillopora damicornis* expel beneficial microbes into the seawater after spawning (Ceh, Van Keulen & Bourne, 2013). Additional experiments are required to test these hypotheses, particularly in coral recruits.

### Evidence that algae from egesta have a similar nutritional status to algae in hospite

In general, in our study, gene expression profiles of algae from egesta were more like those of algae *in hospite* than those of algae in culture. Several of the tested genes were nitrogen transporters or metabolizers. Transcriptional regulation of these genes is rapid (less than a day) in *B. minutum* and other marine algae in response to nutrient levels (Hildebrand, 2005; Kang et al., 2007; Xiang et al., 2020). As the algae from sampled egesta were 1–2 days old, changes in transcript levels for these genes should have occurred in this experiment if nutrient levels had significantly changed once algae were expelled by their hosts. Our data suggest that algae from egesta have a similar nutritional status to algae

*in hospite*, challenging the hypothesis that the host is actively limiting nutrient transport to the symbiont (Maor-Landaw, Van Oppen & McFadden, 2020; Xiang et al., 2020; Cui et al., 2022). The identification of nutrient-related genes as differentially expressed in symbiosis studies that used cultured algae to represent the *ex hospite* state may instead reflect the nutrient-rich state of algae in culture (Maor-Landaw, Van Oppen & McFadden, 2020; Xiang et al., 2020; Cui et al., 2022). In addition, we found that the sugar transporter *SWEET1* (s6\_35311), was not differentially regulated with symbiosis, challenging the hypothesis that this gene copy of *SWEET1* functions to transfer photosynthate from the symbiont to the host (Xiang et al., 2020). This current study, however, did not measure algal growth rates in culture—therefore, we were unable to determine the growth phase of the algae, which can have significant effects on algal physiology and gene expression (Droop, 1975; Mansour, Volkman & Blackburn, 2003; Xiang et al., 2020). Nevertheless, we found that algae in culture had the lowest expression rates of nutrient transporters and nutrient metabolizers, indicating a nutrient-replete phenotype compared to algae from egesta and algae *in hospite*. Altogether, these results warrant further testing of gene expression in algae from egesta using high throughput methods to develop a better understanding of symbiosis.

### Practical considerations for the use of *Aiptasia egesta* in experiments

Based on findings in this study, we argue that it is critical to further explore the biology of expelled algae as a representation of the *ex hospite* state of Symbiodiniaceae to better understand cnidarian-algal symbiosis. To facilitate their use by researchers, we discuss below practical considerations for the use of *Aiptasia egesta* in experiments.

We found that while egesta are primarily composed of Symbiodiniacean algae, they are also populated with microbes, including several protists and cyanobacteria. In our experience, processing of egesta by homogenization and separation by density centrifugation yielded samples of high purity. Collecting ample biological material for use in qPCR and inoculation experiments was difficult. To obtain  $1 \times 10^6$  algal cells from egesta, pellets had to be collected six times every other day from approximately twenty anemones with an average oral disk size of 5.4 mm. We also found that collection was variable between days and yield was not consistent (data not shown). For collection of egesta, if *Aiptasia* are housed without water flow, we recommend using a glass pasteur pipet, as the egesta tended to stick to plastic pipets. Egesta can also be mistaken for pedal lacerates, so it is recommended that a dissection microscope be used to aid in identification. It is possible that various factors, such as anemone size, light levels, and feeding, all influence the rate of egesta release, but these variables were not empirically tested. Another consideration is the induction of algal release by mild stress to increase sample biomass, however, the health of the algae may be affected, so this method will require further experimentation.

## CONCLUSIONS

Overall, this study establishes the foundation for using *Aiptasia egesta* in experiments as a representation of algae in the *ex hospite* state. We found that egesta were released at night, and that algae from egesta had a higher mitotic index than algae *in hospite*, were

photosynthetically active but short lived, and could establish symbiosis with new hosts. Finally, we found that algae from egesta had a similar gene expression profile to that of algae *in hospite* compared to algae in culture, warranting future high-throughput studies to determine symbiosis-specific genes in this comparison. Although there are limitations to the use of algae from egesta, their use in experiments alongside algae in culture and algae *in hospite* can provide valuable insight into cnidarian-algal symbiosis.

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## ADDITIONAL INFORMATION AND DECLARATIONS

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The authors declare there are no competing interests.

### Author Contributions

- Shumpei Maruyama conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the article, and approved the final draft.
- Julia R. Unsworth performed the experiments, analyzed the data, authored or reviewed drafts of the article, and approved the final draft.
- Valeri Sawiccy conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the article, and approved the final draft.

- Virginia M. Weis conceived and designed the experiments, authored or reviewed drafts of the article, and approved the final draft.

### Data Availability

The following information was supplied regarding data availability:

The raw data for egesta release timing, mitotic index, Fv/Fm, symbiont densities, and qPCR are available in the [Supplemental Files](#).

### Supplemental Information

Supplemental information for this article can be found online at <http://dx.doi.org/10.7717/peerj.13796#supplemental-information>.

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