



# Using Seawater to Document Coral-Zoothanthea Diversity: A New Approach to Coral Reef Monitoring Using Environmental DNA

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### Specialty section:

This article was submitted to  
Marine Molecular Biology and Ecology,  
a section of the journal  
Frontiers in Marine Science

**Received:** 09 September 2017

**Accepted:** 19 January 2018

**Published:** 20 February 2018

### Citation:

Shinzato C, Zayasu Y, Kanda M,  
Kawamitsu M, Satoh N, Yamashita H  
and Suzuki G (2018) Using Seawater  
to Document Coral-Zoothanthea  
Diversity: A New Approach to Coral  
Reef Monitoring Using Environmental  
DNA. *Front. Mar. Sci.* 5:28.  
doi: 10.3389/fmars.2018.00028

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Frequent, high-density coral monitoring is essential to understand coral reef ecosystems. For this purpose, we developed a novel method for simultaneous monitoring of *Acropora* corals and their symbiont, *Symbiodinium*, from environmental DNA (eDNA) in seawater using next generation sequencing technology (NGS). We performed a tank experiment with running seawater using 19 *Acropora* species. Complete mitochondrial genomes of all the *Acropora* species were assembled to create a database and major types of their *Symbiodinium* symbionts were identified. Then eDNA was isolated by filtering inlet and outlet seawater from the tanks. *Acropora* and *Symbiodinium* DNA were amplified by PCR and sequenced. We detected all of the tested *Acropora* types from eDNA samples. Proportions and numbers of DNA sequences were both positively correlated with masses of corals in the tanks. In this trial, we detected DNA sequences from as little as 0.04 kg of *Acropora* colony, suggesting that existence of at least one adult *Acropora* colony (~30 cm diameter = 1 kg) per m<sup>2</sup> at depths <10 m could be detected using eDNA in the field. In addition, we detected major types of *Symbiodinium* within host corals from seawater, suggesting that it should be possible to detect major coral symbiont types if *Acropora* corals exist nearby, and possible free-living state *Symbiodinium* cells from eDNA in seawater. eDNA abundance of *Symbiodinium* types did not correlate well with frequencies of major *Symbiodinium* types in the corals, suggesting that quantification of *Symbiodinium* is difficult at this stage. Although this is the initial attempt to detect coral and *Symbiodinium* simultaneously from eDNA in seawater, this method may allow us to perform high-frequency, high-density coral reef monitoring of both corals and their symbionts in the near future.

**Keywords:** scleractinian coral, *Acropora*, *Symbiodinium*, environmental DNA, illumina next-generation sequencing

## INTRODUCTION

Coral reefs are estimated to harbor about one-third of all described marine species (Knowlton et al., 2010); however, they face a range of anthropogenic challenges, including ocean acidification and increasing seawater temperatures (Hoegh-Guldberg et al., 2007). It has been reported that tropical storms, predation by crown-of-thorns starfish, and bleaching are major causes of coral reef decline (De'ath et al., 2012). Frequent high-density coral monitoring is essential to understand dramatically changing coral reef environments. In addition, it is also important to develop strategies against many threats and policies for quick rehabilitation and conservation. However, actual coral reef monitoring is generally performed infrequently because it requires direct observation by scuba diving and sampling by hand.

DNA that originates from various sources, such as mucus, metabolic waste, and damaged tissues from multicellular organisms exists in seawater and is called environmental DNA (eDNA) (Bohmann et al., 2014; Kelly et al., 2014). Recently eDNA has begun to be employed in aquatic environmental research (Rees et al., 2014, 2015), and it is also being used to monitor marine biodiversity (Miya et al., 2015; Yamamoto et al., 2017). However, we are unaware of any reports that have attempted to use eDNA from seawater to monitor coral reefs. In this study, we performed tank experiments with running seawater as an initial proof of concept.

The structure of coral reefs is formed by calcium deposition by anthozoan cnidarians known as scleractinian corals. We examined the genus *Acropora* (Scleractinia, Acroporidae) in this study because it is a keystone reef taxon globally, ranging from the Red Sea through the Indo-Pacific Ocean to the Caribbean. It is also the most speciose coral genus with more than 100 described species (Wallace, 1999). Their widespread distribution notwithstanding, *Acropora* species are among the most sensitive to increasing water temperatures (Loya et al., 2001); hence, they are expected to decline in the near future (Alvarez-Filip et al., 2013).

Scleractinian corals, including *Acropora*, form obligate endosymbioses with photosynthetic dinoflagellates of the genus *Symbiodinium* (Yellowlees et al., 2008; Bourne et al., 2009; Shinzato et al., 2014a). The genus *Symbiodinium* displays tremendous taxonomic and physiological diversity (Lajeunesse, 2001; Stat et al., 2008). Nine divergent lineages, clades A–I, have been described, based on nuclear ribosomal DNA (rDNA) and chloroplast 23S rDNA (Pochon and Gates, 2010). It has been reported that stress tolerance of *Symbiodinium* differs among clades or types (Tchernov et al., 2004; Sampayo et al., 2008; Hawkins and Davy, 2012) and *Symbiodinium* genetic composition may play important roles in coral health and stress tolerance (Jones et al., 2008; Stat et al., 2008; Lajeunesse et al., 2010; Fisher et al., 2012; Keshavmurthy et al., 2012, 2014). Thus, simultaneous investigation of corals and *Symbiodinium* is probably important to understand coral reef ecosystems.

Corals release massive amounts of soluble mucus, which transfers large amounts of energy and nutrients to the reef substrate (Wild et al., 2004). Due to the symbiont natural

rates of increase, corals steadily release *Symbiodinium* cells into the surrounding environment (Yamashita et al., 2011), suggesting that close to reefs, seawater should contain detectable quantities of DNA from both corals and *Symbiodinium*. Recently, whole genome sequences of an *Acropora* coral (Shinzato et al., 2011) and *Symbiodinium* (Shoguchi et al., 2013; Lin et al., 2015; Aranda et al., 2016) have been published, and next-generation sequencing (NGS) technologies have been used to investigate coral reef biodiversity (Shinzato et al., 2014b, 2015; Combosch and Vollmer, 2015; Bongaerts et al., 2017). In the genus *Symbiodinium*, each clade contains multiple genetic types, and identification has been performed using ribosomal, mitochondrial, plastid, and nuclear DNA markers (Rowan and Powers, 1991; Wilcox, 1998; Lajeunesse, 2001; Santos et al., 2002; Takabayashi et al., 2004). Recently internal-transcribed spacer 2 (ITS-2) regions have been widely used as genetic markers (Lajeunesse, 2001, 2002, 2005; Pochon et al., 2007; Stat et al., 2011) and databases of *Symbiodinium* DNA sequences, such as SYM-BLAST (<https://www.auburn.edu/~santosr/symbblast.htm>), *Symbio*GBR (Tonk et al., 2013), and GeoSymbio (Franklin et al., 2012), are currently available. In this laboratory study, we developed a technique that would enable simultaneous detection and identification of coral and *Symbiodinium* eDNA in reefs, in order to assess distributions of corals and their symbiont types.

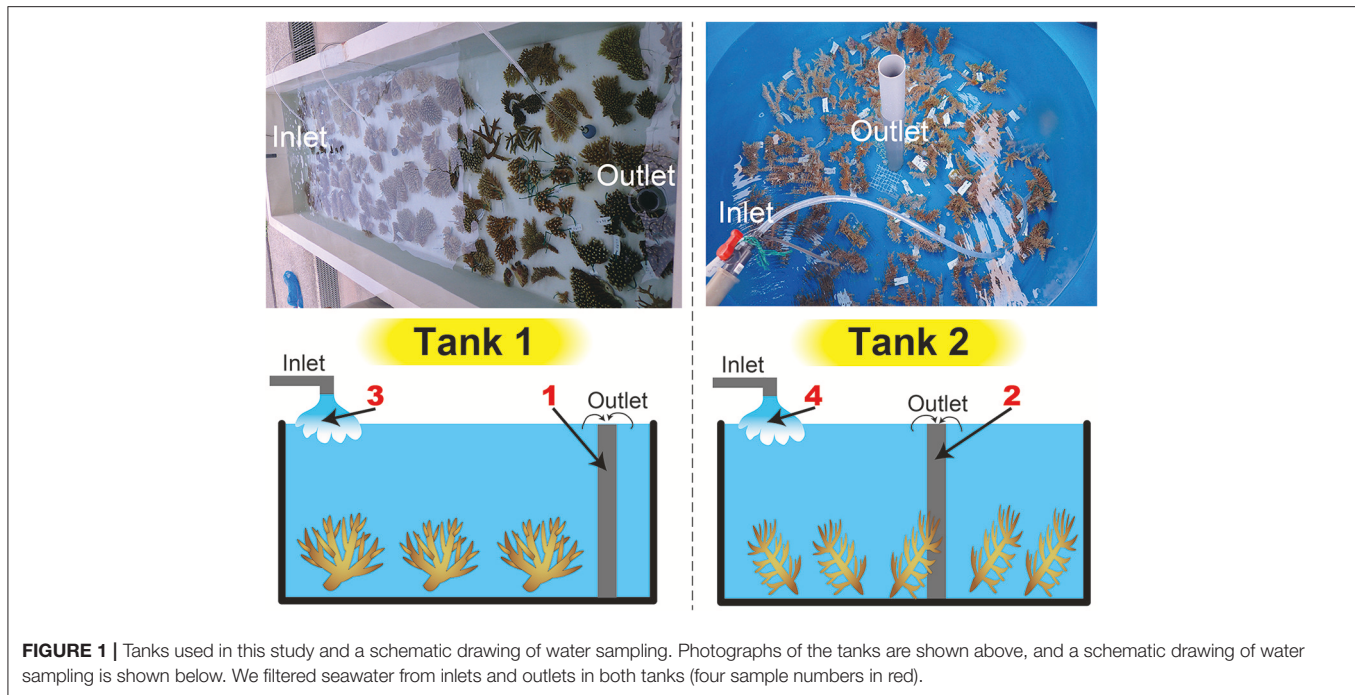
## MATERIALS AND METHODS

### Coral Sampling and Culturing

*Acropora* corals from Sekisei Lagoon, Ishigaki Island, Okinawa Prefecture, Japan were identified and ~20-cm-diameter fragments were collected in May 2016. In an exception, sampling of corals was permitted by the Okinawa Prefectural Government for research use (No. 27-73). In total, we collected 19 species (63 colonies) and put 16 shallow-water (5–10 m) *Acropora* species in Tank 1 (*A. acuminata*, *A. austera*, *A. cytherea*, *A. carduus*, *A. digitifera*, *A. florida*, *A. grandis*, *A. hyacinthus*, *A. intermedia*, *Acropora* sp., *A. microphthalmal*, *A. muricata*, *A. nasta*, *A. selago*, *A. tenuis*) and three deeper-water species ( $\geq 15$  m) in Tank 2 (*A. echinata*, *Acropora* sp., *A. awi*) respectively at Seikai National Fisheries Research Institute at Ishigaki Island, and kept for 2 days with running seawater (Figure 1, Table 1). Numbers of colonies and coral masses determined in seawater differed among species in order to test detection limits of eDNA in seawater (Table 1). Tank sizes and water flow of the two overflow tanks are as follows; Tank 1: 300 cm length  $\times$  80 cm width  $\times$  50 cm height (water depth: ~20 cm) and a flow rate of 35 L of seawater/min, Tank 2: 200 cm diameter and 100 cm depth (water depth: ~80 cm) and 10 L/min. Small fragments of each coral colony were isolated and stored in the guanidinium reagent, CHAOS (Fukami et al., 2000), for DNA extraction.

### Reconstruction of Coral Mitochondrial Genomes

DNA from each colony was isolated from CHAOS solution using a Maxwell 16 Automated Purification System (Promega). We used 200 ng of DNA from each sample for PCR-free library preparation with KAPA HyperPlus Kits (NIPPON Genetics



Co, Ltd) following the manufacturer's instructions. All libraries were multiplexed and sequenced (250-bp paired-end, using a HiSeq 2500 in Rapid mode; Illumina). Raw sequence data were submitted to the DNA Databank of Japan (DDBJ) Sequence Read Archive (DRA) under accession number DRA005695 (Table S1, BioProject PRJDB5633). Sequencing data from each library were assembled with IDBA\_UD assembler version 1.1.1 (Peng et al., 2012) with different kmer lengths (60, 80, and 100). Identification of complete circular mitochondrial genomes from assembled contigs was performed by (1) comparing them with the *Acropora tenuis* complete mitochondrial genome (NCBI accession: AF338425.1) (BLASTN  $e\text{-value} \leq 1e^{-100}$ ), and (2) confirming that 100 bp of both head and tail DNA sequences of a contig were identical, indicating that the sequence was circular.

### eDNA Extraction and PCR Amplification

After 2 days of incubation, 3 L of raw seawater were collected from outlet pipes of the two tanks (Figure 1) and were filtered through a 0.8  $\mu\text{m}$  pore size polycarbonate filter (47-mm $\Phi$ , Advantec Toyo Kaisha, Ltd., Tokyo Japan). In addition, seawater from inlet pipes was also collected to provide control samples (before the seawater was passed over the corals) (Figure 1). Filters were frozen at  $-20^{\circ}\text{C}$  for several days, and then boiled in 500  $\mu\text{L}$  of TE buffer. Details of the TE boiling method for DNA extraction are provided in Koike et al. (2007) and Yamashita et al. (2011). A part of the *Acropora* mitochondrial control region (mtCR,  $\sim 800\text{--}1,000$  bp in length) has been widely used as a molecular marker for *Acropora* species identification (van Oppen et al., 2002; Suzuki et al., 2008). Thus, eDNA was subjected to PCR amplification using RNS2 (5'-CAGAGTAAGTCGTAACATAG-3') and GR (5'-AATTCGGGTGTGTGTTCTCT-3') primers for

amplifying mtCR (van Oppen et al., 2002; Suzuki et al., 2008) together with genomic DNA isolated from *Acropora echinata* as a positive PCR control (expected amplicon size: 935 bp). PCR cycling conditions were 15 min at  $95^{\circ}\text{C}$ , followed by 32 cycles of 30 s at  $94^{\circ}\text{C}$ , 90 s at  $58^{\circ}\text{C}$ , and 60 s at  $72^{\circ}\text{C}$ , with an extension of 30 min at  $60^{\circ}\text{C}$  in the final cycle. To amplify  $\sim 600\text{--}700$  bp of *Symbiodinium* internal-transcribed spacer regions (both ITS-1 and ITS-2) r18Sf primer (5'-GAAAGTTTCATGAACCTTAT-3'), which binds to an 18S region, and Sym28Sr primer (5'-CTTGTRTGACTTCATGCTA-3'), which binds to the ITS2-flanking 28S region, were used (Yamashita and Koike, 2013), together with genomic DNA isolated from clade C *Symbiodinium* culture strain CCMP2466 (clade C, type C1; purchased from the Provasoli-Guillard National Center for Culture of Marine Algae and Microbiota, East Boothbay, Maine, USA) as a positive PCR control (expected amplicon size: 676 bp). PCR cycling conditions were 35 cycles of 45 s at  $94^{\circ}\text{C}$ , 45 s at  $51^{\circ}\text{C}$ , and 60 s at  $72^{\circ}\text{C}$ . To check major symbiotic *Symbiodinium* types in each *Acropora* colony, DNAs isolated from each colony were also used for PCR amplification using r18Sf and Sym28Sr primers under the same PCR conditions mentioned above. All PCR products were cleaned with a QIAquick PCR Purification Kit (Qiagen).

### PCR Amplicon Sequencing and Bioinformatics Analysis

Sequencing libraries of cleaned PCR products (ten samples, DNA amounts used from each sample are shown in Figure S1) were prepared using a KAPA Hyper Prep Kit (NIPPON Genetics Co, Ltd) without fragmentation. Libraries were multiplexed and 300-bp paired-end reads were sequenced with a MiSeq platform (Illumina) with PhiX control (Illumina). Raw sequence data



**TABLE 1** | *Acropora* corals used in the study.

Tank	Species	Number of colonies	IDs for each colony	Weight in water (kg)
1	<i>A. tenuis</i>	15	ten1 ~ 15	11.2
	<i>A. digitifera</i>	5	dig1 ~ 5	1.5
	<i>A. hyacinthus</i>	4	hya1 ~ 4	0.23
	<i>A. cytherea</i>	3	cyt1 ~ 3	0.31
	<i>A. selago</i>	2	sel1 ~ 2	0.15
	<i>A. nasuta</i>	2	nas1 ~ 2	0.29
	<i>A. acuminata</i>	1	acu	0.07
	<i>A. austera</i>	1	aus	0.11
	<i>A. florida</i>	1	flo	0.17
	<i>A. grandis</i>	1	gra	0.11
	<i>A. intermedia</i>	1	int	0.1
	<i>Acropora</i> sp.	1	sp	0.05
	<i>A. muricata</i>	1	mur	0.07
	<i>A. microphthalma</i>	1	mic	0.06
	<i>A. microphthalma</i> deeper-morph	1	micD	0.04
<i>A. carduus</i>	2	car1 ~ 2	0.17	
2	<i>A. echinata</i>	7	ech1 ~ 7	1.1
	<i>A. awi</i>	7	awi1 ~ 7	1.3
	<i>Acropora</i> sp.	7	sp1 ~ 7	1.4

*Acropora* sp. is unidentified species similar to *A. echinata*.

were submitted to the DDBJ DRA under accession number DRA005680 (BioProject PRJDB5633). Read pairs that both PCR primers of *Acropora* (RNS2 and GR) or *Symbiodinium* (r18Sf and Sym28Sr) recognized were selected using custom perl scripts (Supplementary Materials) and subjected to quality filtering. Low quality bases (Phred quality score <30) were trimmed using SolexaQA (Cox et al., 2010) and high-quality sequences longer than 200 bp were retained for subsequent analyses.

To identify *Acropora* species from eDNA, we used mtCR sequences based on the complete, assembled, mitochondrial genomes of corals used in this study (see above) and on *A. digitifera* sequencing data from five individuals previously collected at Kabira, Ishigaki, Japan (Shinzato et al., 2015). Although mtCR is highly variable within *Acropora* mitochondrial genomes (Figure S2), percent identities of mtCR sequences between species are high (88.5–100%, Table S2). Lengths of mtCR exceeding paired-end sequencing length (300-bp paired-end) and 300-bp sequences flanking the RNS2 primer are more variable than those of the GR primer (Tables S3, S4). Accordingly, 300 bp flanking the RNS2 primer sequence in the mtCR from each colony were extracted and exactly the same sequences of clustered using CDHIT-EST (Li and Godzik, 2006), and the clustered sequences (14 types) comprised a mtCR database in this study (Table S3). Quality trimmed reads having the RNS2 primer sequence were identified and chimera filtering based on the mtCR database was performed using the UCHIME2 algorithm (Edgar, in review). We aligned these filtered sequences to the database using BLASTN (Camacho et al., 2009) with

e-value cutoff of  $1e^{-100}$ . Output files were stringently filtered and numbers of valid alignments that ensure all of the following criteria were counted (details are available on Supplementary Materials); (1) percent identity = 100% as mitochondrial genome sequences of all colonies were in the database; (2) the BLASTN bit score of the best alignment must be greater than the second-best alignment. In order to determine the correlation between coral wet masses and the number of valid alignments, regression analysis was performed using R, version 3.1.3 (R Core Team, 2015).

To develop a *Symbiodinium* ITS-2 database, we first clustered exactly the same DNA sequences in a database of ITS-2 gene sequences of most reported *Symbiodinium* types, GeoSymbio (410 ITS-2 sequences) (Franklin et al., 2012) using CDHIT-EST (Li and Godzik, 2006) with option “-c 1.” This resulted in 391 sequences (different ITS-2 sequences sharing same DNA sequences in the original GeoSymbio database were annotated using a concatenation of the original GeoSymbio names with “\_”). Then we downloaded *Symbiodinium* DNA sequences containing the term “internal transcribed spacer 2” from NCBI in October 2017, and Blasted (Camacho et al., 2009) these against the clustered GeoSymbio (BLASTN, e-value <  $1e^{-5}$ ). Aligned sequences longer than 200 bp were extracted and annotated with the names of BLASTN top-hit sequences in the clustered GeoSymbio together with their NCBI accession numbers. Finally, exactly the same DNA sequences were clustered using CDHIT-EST (Li and Godzik, 2006) with option “-c 1.” For identification of *Symbiodinium* types from eDNA and each *Acropora* colony, ITS-2 gene sequences flanking the Sym28Sr primer were subjected to chimera filtering (Edgar, in review) using the database and aligned to the *Symbiodinium* ITS-2 database using BLASTN (Camacho et al., 2009) with an e-value cutoff of  $1e^{-100}$ . Output files were stringently filtered and the number of valid alignments that ensure all of the following criteria were counted (details are available on Supplementary Materials); (1) percent identity  $\geq 98\%$  as there might be *Symbiodinium* types that are not included in the NCBI database; (2) the BLASTN bit score of the best alignment must be greater than the second-best alignment.

## RESULTS

### Assembling the Complete Mitochondrial Genomes of *Acropora* Corals and Identifying Major Types of Symbiotic *Symbiodinium*

We isolated genomic DNA from tissues taken from each of 63 colonies, containing DNA from both *Acropora* and *Symbiodinium*, except for *A. digitifera*, from which we failed to isolate useful DNA. We successfully assembled complete (circular) mitochondrial genomes of all species (accession numbers: LC201813–LC201870) (Table S1). We identified putative mtCR in the assembled mitochondrial genomes and used these to create a database comprised 14 mtCR types (Table S3).

For development of a *Symbiodinium* ITS-2 database, 5,962 DNA sequences of possible *Symbiodinium* ITS-2 were downloaded from NCBI, annotated using GeoSymbio database, and exactly the same DNA sequences were clustered. Then, we prepared a *Symbiodinium* ITS-2 database containing 2,352 non-redundant sequences. Using the latter database, we identified major types of *Symbiodinium* from each colony (Figure 2). Most colonies exclusively harbored clade C (99.96% of total valid alignments, 4533872 out of 4535626, Figure 2 and Supplementary Material), but clade D was also a major symbiont in some colonies (e.g., D1 and D1a in *A. tenuis* ten3 and ten12 colonies, Figure 2). Interestingly, C50 is the most major in most colonies in shallow-water *Acropora* in Tank 1, although C3d\_C21 occurred in half of Tank 2 *Acropora* colonies (Figure 2).

## Identification of *Acropora* Corals from eDNA in Seawater

After culturing corals for 2 days, we isolated eDNA from both inlet and outlet seawater. We performed PCR to amplify *Acropora* mtCR and the *Symbiodinium* ITS-2 region (Figure S1). Although all PCR products using *Symbiodinium* primers could be identified, some PCR products of *Acropora* mtCR were not visible by electrophoresis. However, we did isolate DNA from all *Acropora* mtCR PCR products (Tank 1 outlet: 8.4 ng/ $\mu$ L, Tank 2 outlet: 4.3 ng/ $\mu$ L, Tank 1 inlet: 4.6 ng/ $\mu$ L, Tank 2 inlet: 4.5 ng/ $\mu$ L, respectively) and successfully prepared Illumina sequencing libraries for all samples (Figure S1). These ten amplicon samples, including two positive controls (*A. echinata* and *Symbiodinium* culture strain CCMP2466), were multiplexed and sequenced (Table 2).

As mtCR sequences are highly conserved among *Acropora* corals (Tables S2–S4), most sequences mapped to more than two sequences in the databases. Thus, we subsequently filtered mapping output so as to count only valid alignments to the database. In the positive control of *A. echinata* DNA PCR amplicon, 99.1% of valid alignments (35339 of 35662) were correctly assigned to *Acropora* sp. (Figure 3), indicating that our filtering method worked properly. As expected, few valid alignments from inlet water of Tank 1 and Tank 2 were detected (Table S5). We identified all 14 mtCR types in the database, although only a single valid alignment for car1 (*A. carduus*) was detected (Figure 3A). The most abundant mtCR types in Tank 1 included *A. tenuis* (Figure 3A), probably reflecting 15 *A. tenuis* colonies in the tank (Table 1). In Tank 2, mtCR sequences from *Acropora* sp. and *A. echinata* were detected more frequently than *A. awi* (Figure 3). However, although coral species in Tank 1 and Tank 2 were different, a small proportion of alignments to some mtCR types of coral species that did not exist in each Tank were detected in outlet water of both tanks (Tank 1: 0.05%; 40 out of 67971, Tank 2: 0.8%; 440 out of 52449, Figure 3 in red colors). In inlet water of Tanks 1 and 2, few valid alignments were observed (Figure 3). The proportions of valid alignments of each mtCR types to the total numbers of valid alignments and proportions of weights of corals (Figure 3B, Table S5) were positively correlated in both tanks (regression analysis,  $R^2 = 0.7793$ ,  $p$ -value <

0.001), in Tank 1 (regression analysis,  $R^2 = 0.9807$ ,  $p$ -value < 0.001) and in Tank 2 (regression analysis,  $R^2 = 0.5985$ ,  $p$ -value < 0.01), respectively, suggesting that proportions of DNA sequences reflect the proportions of coral types in surrounding water. In addition, the number of sequences and weights of corals (Figure 3C, Table S5) were positively, but not strongly correlated in both tanks (regression analysis,  $R^2 = 0.6371$ ,  $p$ -value < 0.01), in Tank 1 (regression analysis,  $R^2 = 0.6923$ ,  $p$ -value < 0.01), and in Tank 2 (regression analysis,  $R^2 = 0.7109$ ,  $p$ -value < 0.01), respectively. We detected coral colonies heavier than 0.04 kg (e.g., *A. microphthalmal* deeper-morph: 0.04 kg in Tank 1, Table 1). Overall, the calculated average number of valid alignments per kg (Table S5) was  $6145 \pm 7058$  ( $R^2 = 0.7962$ ,  $p < 0.001$ ), while for *Acropora* species in Tank 1 it was  $3389 \pm 2895$  ( $R^2 = 0.987$ ,  $p < 0.001$ ) and in Tank 2 it was  $13492 \pm 10411$  ( $R^2 = 0.5658$ ,  $p < 0.05$ ).

## Identification of *Symbiodinium* Types from eDNA in Seawater

In the positive control of *Symbiodinium* CCMP2466 (clade C, type C1) PCR amplicon, 97.23% of valid alignments were correctly assigned to C1 types (C1 and C1ca\_C1b\_C1e, Figure 4), indicating that our bioinformatics filtering worked properly for *Symbiodinium* as well. Clade C, the major symbiotic *Symbiodinium* clade in the corals used (Figure 2), was detected in both tanks (Figure 4). The most abundant clade C type detected was C50 in Tank 1 and C3d\_C21 in Tank 2, respectively, possibly reflecting abundant clade C types of the *Acropora* corals in each tank (Tank 1: C50, Tank 2: C3d\_C21, Figure 2). We were not able to detect some clade C types that were identified from the corals; however, the total number of valid alignments of these types comprised only ~1% of total alignments (46314 out of 4535626) of symbiotic *Symbiodinium* in the *Acropora* corals (Figure 2, Supplementary Materials), indicating that these were minor types of clade C symbiont, which could not be detected in seawater. Although clade C types were exclusively detected from *Acropora* colonies, those in outlet water were low (Tank 1: 35.6%; 44178 out of 36748, Tank 2: 89.3%; 111107 out of 124375). We detected considerable number of clade D types in outlet water of both Tanks 1 and 2 (Figure 4). Within clade D, D1 was the most abundant in both tanks, possibly reflecting more D1 symbionts in the corals (Figure 2). We also detected other clade D and clade A types, which were not detected in the *Acropora* corals used in this study (Figure 2), from both outlet and inlet seawater. In outlet water, only small numbers of valid alignments originated from ITS-2 types that were not detected in *Acropora* corals (Figure 2 and Supplementary Materials, Tank 1: 1%; 1003 out of 124223, Tank 2: 3.2%; 4026 out of 124375), albeit in inlet water, large proportions of valid alignments originated from ITS-2 types that were not detected in the corals (Figure 2 and Supplementary Materials, Tank 1: 51.7%; 19005 out of 36748, Tank 2: 99.8%; 38821 out of 38918). In inlet water detected ITS-2 types were considerably different between Tanks 1 and 2 (Figure 4). For example, some abundant types in Tank 1 inlet water (A2-ISS-C2-Sy and D5) were not detected in Tank 2 inlet water, whereas A3 and F4.4b were abundant types in Tank 2 inlet water (Figure 4).





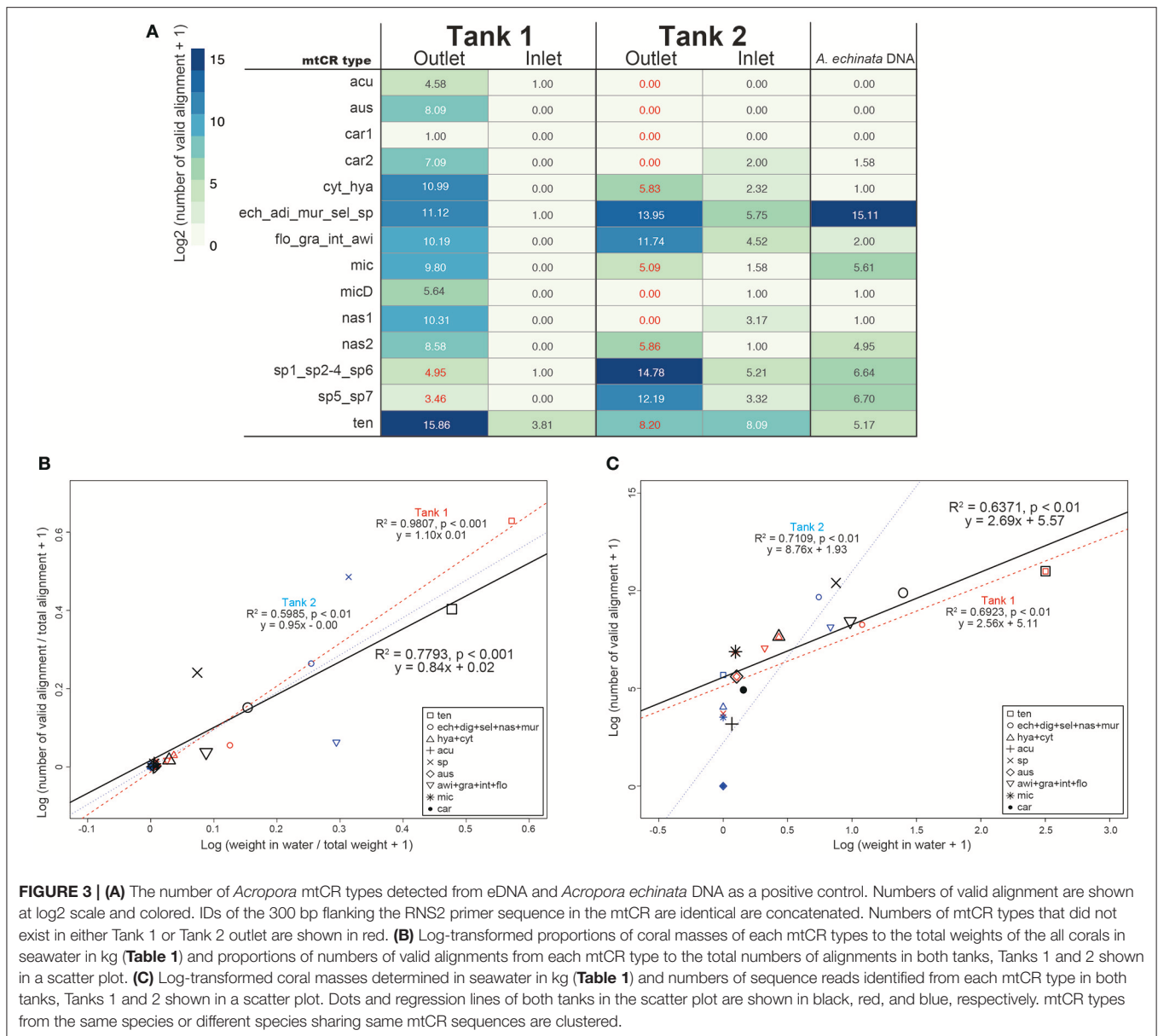
**TABLE 2** | Summary of sequencing of PCR amplicons from eDNA in culturing tanks (Tank 1 and Tank 2) and positive controls for *Acropora* and *Symbiodinium* (isolated DNAs of *Acropora echinata* and cultured clade C *Symbiodinium* CCMP2466, respectively).

Sample name	Number of raw sequence pairs (basepair)	Number of quality trimmed and chimera removal sequences	Number of aligned sequences to the database (BLASTN, e-value < 1e <sup>-100</sup> )	Number of valid alignment
<i>Acropora echinata</i> DNA	611664 (376582569 bp)	248162	245331	35662
<i>Acropora</i> -1 (Tank 1 outlet)	575413 (352350985 bp)	213996	212207	67971
<i>Acropora</i> -2 (Tank 2 outlet)	743548 (454706122 bp)	266882	263520	52449
<i>Acropora</i> -3 (Tank 1 inlet)	645400 (312578543 bp)	600	111	16
<i>Acropora</i> -4 (Tank 2 inlet)	658164 (334083049 bp)	2888	1642	411
Clade C <i>Symbiodinium</i> (CCMP2466)	954817 (586923250 bp)	251453	423560	309108
<i>Symbiodinium</i> -1 (Tank 1 outlet)	582922 (358869403 bp)	257035	217861	124223
<i>Symbiodinium</i> -2 (Tank 2 outlet)	586197 (360725736 bp)	202786	228327	124375
<i>Symbiodinium</i> -3 (Tank 1 inlet)	682968 (396295801 bp)	206604	66946	36748
<i>Symbiodinium</i> -4 (Tank 2 inlet)	688667 (415307055 bp)	436168	53358	38918

(Suzuki et al., 2012a), but were seriously damaged due to an outbreak of crown-of-thorns starfish, *Acanthaster planci*, in 2012 (Suzuki et al., 2012b). Therefore, the amount of *Acropora* eDNA contained in flowing seawater could be small. *Acropora* corals observed around the marine station were *A. selago*, *A. muricata*, and *A. nasuta* and their coverage was less than 1% (Suzuki et al., 2012a). Although some of the minor reads mapping to the mtCR database from inlet pipes might have originated from minor contaminants or artifacts from other samples, all of the observed *Acropora* species naturally occurring near the research institute were detected from inlet pipes (Figure 3), suggesting that these reads possibly originated from outer reefs and this technology could detect small amounts of *Acropora* DNA in seawater. There are still some difficulties in applying eDNA techniques to monitoring coral reefs, e.g., lack of suitable DNA markers for corals and reference DNA databases. In this study, it was not clear that we could distinguish DNA sequences from all colonies due to the high conservation and slow evolution of mitochondrial DNA in anthozoan cnidarians, including scleractinian corals (Shearer et al., 2002). For example, mtCR sequences of *A. grandis*, *A. intermedia*, and *A. awi* are same (Table S2). Thus, additional markers for nuclear DNA will be needed to reliably identify *Acropora* species using eDNA, specifically markers that are more variable than mitochondrial DNA (Ladner and Palumbi, 2012; Suzuki et al., 2016), although nuclear DNA of marine organisms is usually present at low concentrations and is rapidly degraded in seawater (Rees et al., 2014). However, the success of using a nuclear DNA marker (e.g., internal transcribed spacer 1) to detect common carp from eDNA has recently been reported (Minamoto et al., 2016), implying that it might be possible to employ nuclear markers for identifying corals from seawater. An additional problem is that species that are not in the database cannot be detected. Species identification from eDNA depends upon completeness of the reference database and enhancement of a coral DNA database will be required for more precise coral species detection and identification.

In a previous study, detection of *Symbiodinium* DNA from seawater was complicated by the low primer specificity (Pochon et al., 2010). In this study, we used r18Sf and Sym28Sr primers (Yamashita and Koike, 2013) that were used for amplifying *Symbiodinium* DNA from environmental samples (Yamashita et al., 2013, 2014). This primer set could be used not only for analyzing *Symbiodinium* within host animals, but also for detecting free-living state *Symbiodinium* in coral reef environments (Yamashita and Koike, 2013; Yamashita et al., 2013, 2014). As hoped, major types of *Symbiodinium* symbionts in corals used in this study were detected from eDNA in outlet water. Most valid alignments in outlet water were assigned to ITS-2 types detected from *Acropora* corals (Figures 2, 4, Supplementary Materials, Tank 1: 99%, Tank 2: 96.7%), suggesting that it would be possible to detect most coral symbiont types from seawater, if *Acropora* corals exist nearby. However, eDNA abundance did not correlate well with frequencies of the major *Symbiodinium* types of the corals used in this study. For example, clade C types were exclusively detected from *Acropora* colonies, but not from outlet water of both Tanks 1 and 2, indicating that quantification of *Symbiodinium* types is presently difficult. It might be due to a failure to detect minor symbiotic *Symbiodinium* types in *Acropora* corals (Figure 2) or to significant copy number differences of the ITS-2 gene among *Symbiodinium* types. Indeed, it is reported that there are thousands of copies of ITS regions in *Symbiodinium* genomes and copy numbers of ITS1 in clade D are ~3x higher than in clade C (Mieog et al., 2007). These might be major factors accounting for the inconsistency between sequence reads and major *Symbiodinium* types.

The primer set r18Sf and Sym28Sr can detect even relatively rare *Symbiodinium* clades in coral reef environments (Yamashita and Koike, 2013; Yamashita et al., 2013, 2014). Thus, the primer set was powerful enough to amplify the ITS-2 gene from low concentrations of *Symbiodinium* cells. In our experiments, clade A was detected in both inlets and clade D types that were not

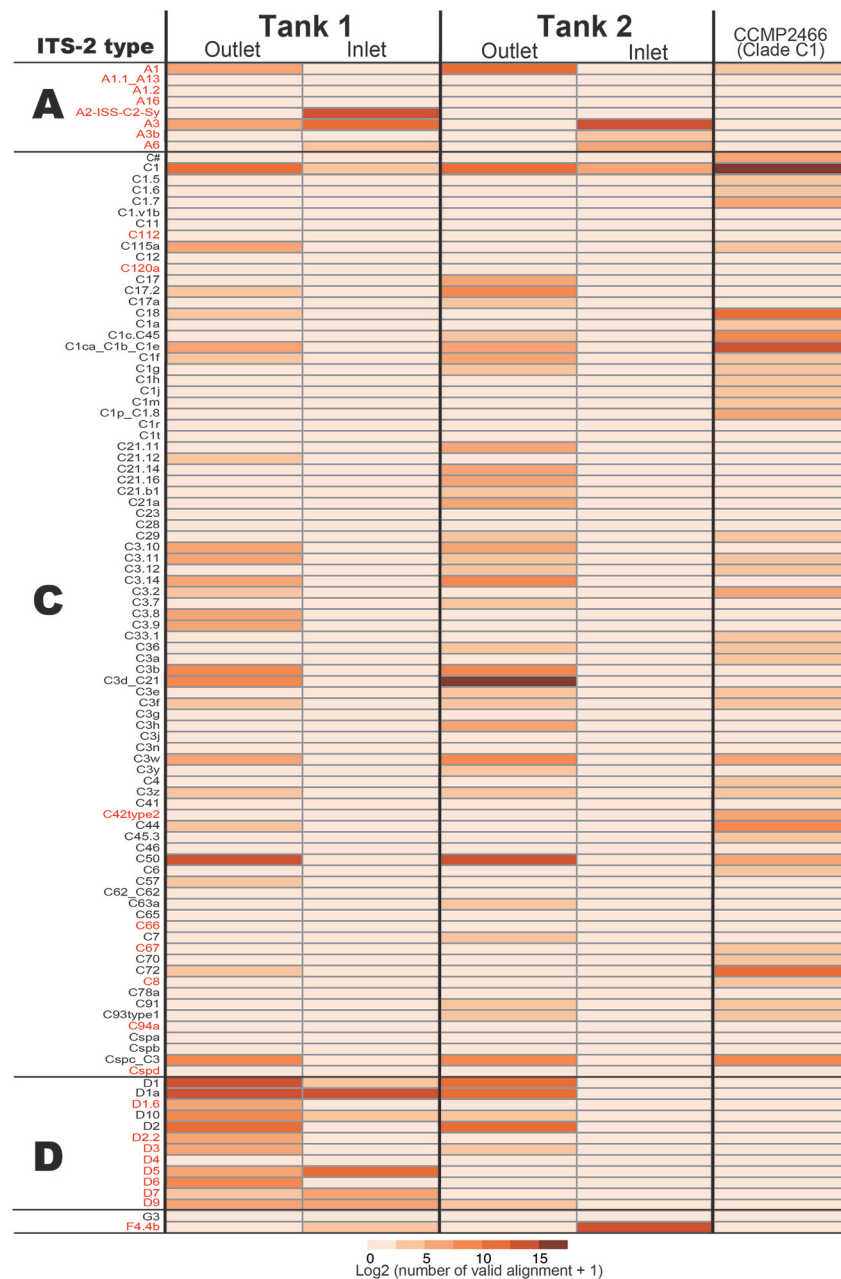


major coral symbionts in the coral used in this study (Figure 2) were identified in the Tank 1 inlet. Likewise, clade F was detected in the Tank 2 inlet (Figure 4). Clade F is a common foraminiferan symbiont in the tropical and subtropical Indo-Pacific and Caribbean Oceans (Pochon and Pawlowski, 2006), but was previously detected in corals (Lien et al., 2012). The detection of different *Symbiodinium* types in inlet water for the two tanks probably stems from different cell types between two inlets and the great sensitivity of the technique. Chance events involving the distribution of individual *Symbiodinium* cells to the two tanks were potentially able to bias the results. Therefore, it should be taken into account that both free-living state *Symbiodinium*, as well as coral symbionts are thoroughly mixed in the water column and regular, continuous observation of symbiotic *Symbiodinium*

in corals under different environmental conditions would be necessary for assessing diverse coral reef environments.

In this study, we demonstrate that detection of nearby coral species and their symbiotic algae is feasible in coral reef environments using eDNA isolated from seawater. With high-quality molecular markers and an extensive database, we could perform more precise, detailed monitoring of coral reef ecosystems, including corals, symbionts, and other organisms. Coral damage and recovery after bleaching or severe storms varies even at small spatial scales, depending on reef environmental conditions, such as reef topology and seawater temperature. It is expected that high-frequency, high-density monitoring will enable us to establish effective conservation strategies. We hope that eDNA technology may become a





**FIGURE 4 |** The number of *Symbiodinium* ITS-2 types detected by eDNA PCR amplicon sequencing and CCMP2466 as a positive control. ITS-2 types that share the same GeoSymbio names in the database were summed (raw numbers of valid alignments are available in Supplementary Materials). ITS-2 types belonging to the same clade (A, C, or D) are separated and ITS-2 types in red indicate that these were not detected in the *Acropora* colonies used in this study (Figure 2).

powerful tool for conservation management of coral reefs in the future.

## AUTHOR CONTRIBUTIONS

CS, HY, and GS: conceptualization and commencement of the study. YZ and GS: Animal collection and *Acropora* species identification. HY and GS: Coral culturing, environmental DNA isolation, and PCR amplification. MiK and MaK: Preparation of

sequencing libraries and producing sequencing data. NS provide sequencing reagents and facilities. CS analyzed the data and wrote the main manuscript text, and all authors edited and approved the manuscript.

## ACKNOWLEDGMENTS

This study was supported by JSPS KAKENHI grants (17KT0027, 17K07949 and 26290065 for CS, 17K15179

for YZ, 26291094 for HY, 15H04538 for GS). We thank Dr. Steven D. Aird (OIST) for editing the manuscript.

## SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fmars.2018.00028/full#supplementary-material>

Supplementary Materials: Five files and directories (Supplementary\_Fig-Table.pdf, Perl\_scripts, Database, Commands.doc and Sym\_num-valid-alignments.xlsx) are included.

Explanation of files:

Supplementary\_Fig-Table.pdf: Supplementary Figures and Tables shown in PDF format.

Commands.doc: bioinformatics commands that we used in this study.

Sym\_num-valid-alignments.xlsx: Actual numbers of valid alignments used in Figures 2, 4.

Perl\_scripts: custom perl scripts used in this study.

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- Database: Fasta files used for BLASTN databases (*Acropora*: RNS2\_acropora.fa, *Symbiodinium*: sym-ITS-2\_NCBI-GeoSym.fa) in this study.
- Figure S1** | Electrophoresis of PCR amplicons from eDNA. These were used for library preparation (total amounts of DNA used are shown) and sequenced with a MiSeq (300-bp paired-end libraries). A 100-bp DNA Ladder (TaKaRa, code 3422) was used to determine DNA marker sizes. Amplicon sizes of positive controls (*Acropora ehinata* and cultured clade C *Symbiodinium* CCMP2466) are 935 and 676 bp, respectively.
- Figure S2** | Rate of identical nucleotides in 60-bp sliding window of aligned mitochondrial genomes of *Acropora* corals used in this study. mtCR and RNS2 and GR primer flanking sequences are shown.
- Table S1** | Shotgun sequencing data (HiSeq2500, 250-bp paired-end libraries) from coral colonies used in this study. Abbreviations are same as **Table 1**.
- Table S2** | Percent identity of mtCR sequences to the most similar sequence within the database used in this study. Abbreviations are same as **Table 1**.
- Table S3** | Percent identities of 300 bp flanking the RNS2 primer to the most similar mtCR sequence in the corals used in this study.
- Table S4** | Percent identities of 300 bp flanking the GR primer to the most similar mtCR sequence in the corals used in this study.
- Table S5** | Weight in water and number of valid alignments of *Acropora* species used in this study.
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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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