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Molecular diversity of Symbiodinium spp. within six coral species in Larak Island, the Persian Gulf

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Abstract

Reef-building coral harbor communities of photosynthetic taxa of the genus Symbiodinium (zooxanthellae). The genus Symbiodinium is currently classified into nine genetic clades (A-I). Various corals harbor different Symbiodinium clades; some show specificity to a single strain. Coral and their zooxanthellae are sensitive to environmental stresses. In the Persian Gulf, coral reefs are subject to harsh environmental conditions including extreme temperatures and high salinity. This is the first study to use clade specific primers to clarify the diversity of Symbiodinium in each coral species of Larak Island. For this purpose six coral species were collected at two different locations in Larak Island. After DNA extraction, PCR amplification was performed using clade specific primers. The results showed that multiple Symbiodinium clades are hosted by most coral species. In addition, among thirteen obtained Symbiodinium sequences, the frequency of either tree clades, A, C and D was almost the same. Corals species may contain different clades of Symbiodinium depending on the region and on the tolerance characteristics of each clade. Thus, knowledge of zooxanthellae diversity associated with scleractinian can contribute to a better understanding of the sensitivity of corals to environmental conditions.

Keywords: Persian Gulf, *Symbiodinium*, Clade A, Clade C, Clade D.

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Introduction

Reef corals are hosts to a group of diverse exceptionally dinoflagellate symbionts of the genus Symbiodinium referred (commonly to and zooxanthellae) (Pochon Gates. 2010). The high productivity and diversity of coral reefs is largely due to this mutualistic symbiosis (Silverstein et al., 2012). The Symbiodinium transports newly fixed organic carbon to the coral and, in return, receive inorganic waste metabolites from host respiration and an environment free from predators (Davy, 2012).

Symbiodinium are morphologically and most species cryptic morphologically similar (Baker, 2003). Therefore, their identification must be done on molecular methods. It has been established that there are at least nine major clades of this dinoflagellate genus (Pochon and Gates, 2010). Numerous sub-clades and types have been identified within Symbiodinium clade, most commonly using the internal transcribed spacer-2 (ITS-2) marker (LaJeunesse, 2002).

Scleractinian coral and their endosymbiont are sensitive environmental stresses that include salinity (Reimer. 1971). high temperatures (Hoegh-Guldberg Smith, 1989), low temperatures (Steen and Muscatine, 1987), ultraviolet radiation (Gleason and Wellington, 1993) and turbidity (Trench, 1986). The physiological responses Symbiodinium vary greatly among phylogenetic types; for example, among

the known clades, Symbiodinium clade D has a higher thermal tolerance than other clades, and clade D increases the resistance of corals that harbor them to elevated sea surface temperatures (Rowan, 2004: Berkelmans and van Oppen. 2006). thermal The and physiological flexibility of Symbiodinium may provide mechanisms for scleractinian corals to survive under unfavorable conditions (Berkelmans and van Oppen, 2006).

The previous studies have shown that corals associate with different Symbiodinium clades or types depending on local environmental conditions (Baker, 2003). The Persian Gulf, located in the northwest of the Indian Ocean, is known as one of the most extreme environments for coral reefs with high temperature fluctuations. high salinity, high turbidity and low depth (Sheppard et al., 1992; Baker, 2004). Research conducted on northern Persian Gulf zooxanthellae since 2005 demonstrates the occurrence of Symbiodinium clades D, C and A in the shallow waters of the world's hottest sea (Mostafavi et al., 2005, 2014; Shahhosseiny et al., 2011). The present study aims to investigate the diversity of Symbiodinium in coral species of Larak Island, one of the seventeen islands in the northern Persian Gulf. Individual colonies can with associate several genetically simultaneously. symbionts distinct Corals of the same species may vary in depending their symbionts on environmental characteristics (van Oppen *et al.*, 2001). Therefore, the results of this survey will provide a valuable insight to the survival of Iranian coral reefs in the unfavorable environment of the Persian Gulf.

Material and methods

Sampling

Colonies of six species of scleractinian corals, Favia pallida (DANA, 1846); Leptastrea transversa (Klunzinger, 1879); Psammocora contigua (Esper, 1794); Stylophora pistillata (Esper, 1797); Pocillopora damicornis (Linnaeus, 1758) and Acropora

downingi (Wallace, 1999) were collected from two sites, S1 (latitude 26° 53' 22.39" N, longitude 56° 21' 10.44" E) and S2 (latitude 26° 52' 33.61"N, longitude 56° 20' 9.84" E), at depths 3 m (S1) and 6 m (S2) off Larak Island (Fig. 1) during March 2013. collection. After samples were preserved in DMSO buffer (20% DMSO, 0.25M EDTA, saturated with NaCl, pH=8) and transferred to the marine biology laboratory in Islamic Azad University, Science and Research Branch.

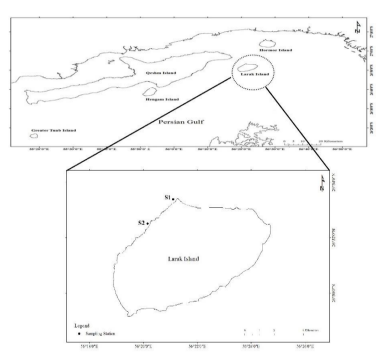


Figure 1: Map of the Larak Island showing the position of the sampling locations.

DNA extraction PCR amplification and sequencing

DNA was extracted using cetyl trimethylammonium bromide (CTAB)/Chloroform method (Baker, 1999). Polymerase Chain Reactions (PCR) assays were performed using three clade-specific primer sets

targeting the ITS1-5.8S-ITS2 (for clade A) (Correa *et al.*, 2009), domain 2 of the LSU (for clade D) (Correa *et al.*, 2009) and partial ITS 1 (for clade C) (Ulstrup and van Oppen, 2003) of the nuclear rDNA of *Symbiodinium*. Each PCR reaction comprised, 1.5 mM MgCl₂, 0.2 mM dNTPs Mix, 10 pmol

of each primer, 0.3 U Tag DNA polymerase and 1 ng DNA Template for a total volume of 25 µL. Amplification was performed using a Bio-Rad PCR Thermal Cycler with the following thermal profile: 30 cycles of 30s at 94°C, 30 s at 58°C, 56°C and 60°C (for clade A, C and D, respectively), 30s at 72°C and a final extension for 5 min at 72°C. The PCR products were analyzed by electrophoresis in 1.5% agarose gels. Thirteen PCR products obtained from this study were directly sequenced to confirm that the specific primers only recognize the target of interest. Finally, this was approved by constructing a phylogenetic tree based on sequences of each molecular marker.

Phylogenetic analyses

Sequences obtained from this study were deposited in a GenBank and their accession numbers are shown in Table 1. The new nucleotide sequences obtained during the present study were aligned with sequences available from Genbank using the software CLUSTALW (Thompson et al., 1994). Three alignment datasets generated (for clade A, C and D). The alignment datasets were analyzed using maximum likelihood (ML), maximum parsimony (MP) and Bayesian methods.

Table1: Host species, Symbiodinium clade and GenBank accession numbers.

Coral species	Symbiodinium	GenBank
	clade	accession numbers
Favia pallida	Clade A	KT069226
Leptastrea transversa	Clade A	KT069227
Stylophora pistillata	Clade A	KT069228
Pocillopora damicornis	Clade A	KT069229
Favia pallida	Clade C	KT069230
Leptastrea transversa	Clade C	KT069231
Psammocora contigua	Clade C	KT069232
Stylophora pistillata	Clade C	KT069233
Pocillopora damicornis	Clade C	KT069234
Favia pallida	Clade D	KT069235
Leptastrea transversa	Clade D	KT069236
Psammocora contigua	Clade D	KT069237
Acropora downingi	Clade D	KT069238

The most appropriate model selection for ML and Bayesian analyses was performed using Akaike Information Criterion (AIC) in MODELTEST 2.3 (Nylander, 2004). The general timereversible model (Rodriguez *et al.*, 1990) with gamma parameter (GTR+G)

gave the best fit to the data (clade A, C and D). ML and MP analyses were conducted using the MEGA6 (Tamura *et al.*, 2013) and PAUP beta version 4.0b10 (Swofford, 2002) respectively. ML and MP clades were assessed with 1,000 bootstrap replicates. The

Bayesian analysis was implemented in MrBayes 2.3 (Ronquist Huelsenbeck, 2003) and was based on the model selected by MODELTEST above. Starting from random trees, four Markov chains (with one cold and three heated chains) were run simultaneously to sample trees using the Markov Chain Monte Carlo (MCMC) principle which approximates the posterior probability (PP) of trees. After the burn-in phase (the first 5 million generations was discarded), every 100th tree out of 20⁶ was considered. The phylogenetic trees all analyses generated in were visualized using TREEVIEW (Page, 1996).

Results

The results showed that there are three clades of Symbiodinium from Larak Island; clade A, C and D. Symbiodinium sequences belonging to clade A were identified from F. pallida, L. S. pistillata transversa, and damicornis, clade C from F. pallida, L. transversa, S. pistillata, P. damicornis, P. contigua and clade D from F. pallida, L. transversa, P. contigua and A. downingi.

Figs. 2, 3 and 4 show the ML phylogenetic tree for the aligned sequences. As the tree topologies were similar in all analyses, the bootstrap values for MP and Bayesian posterior probabilities are shown on the ML tree (Figs. 2, 3 and 4).

The phylogenetic tree of Symbiodinium clade A sequences is shown in Fig. 2. The clade A Symbiodinium from four coral species strongly clustered (ML=MP=90%, PP=0.9) with subclade A1 Symbiodinium that is hosted Cassiopea xamachana in Jamaica (AF427466), Acropora sp. in Japan (AB849873-AB849875) and Zoanthus South sansibaricus in Africa (KM032592).

The phylogenetic tree ofSymbiodinium clade C ITS -1 sequence is shown in Fig. 3. The clade C Symbiodinium sequences from five coral species together with previously reported sequences, including some sequences from zoantharians Madagascar (KM032585), Singapore (EU333738) and South Africa (KM032562) formed a highly supported monophyly (ML= 99%, MP=100%, PP=1.0) within the Symbiodinium clade C radiation.

Phylogenetic results of large subunit ribosomal (LSU) RNA gene analyses are shown in Fig. 4. In the tree of clade D, all D-matching sequences from four coral species grouped with five previously reported sequences belonged to clade D AB778761, AB778758. AB778750 nudibranch and KF672733, KF030947 from stony corals in a well- supported monophyly (ML= 100%, MP=100%, PP=1.0).

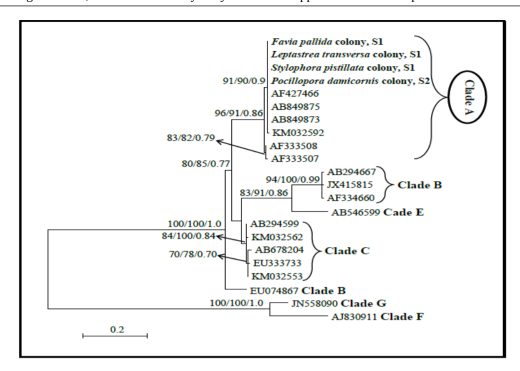


Figure 2: Maximum likelihood tree of ITS1-5.8S-ITS2 sequences for clade A. Values at branches represent maximum likelihood bootstrap percentages from 1000 trees/ maximum parsimony bootstrap percentages from 1000 trees/Bayesian posterior probabilities.

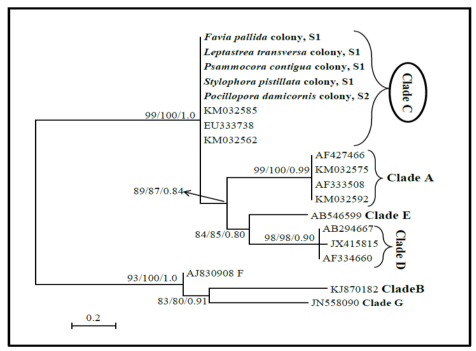


Figure 3: Maximum likelihood tree of partial ITS 1 for clade C. Values at branches represent maximum likelihood bootstrap percentages from 1000 trees/ maximum parsimony bootstrap percentages from 1000 trees/Bayesian posterior probabilities.

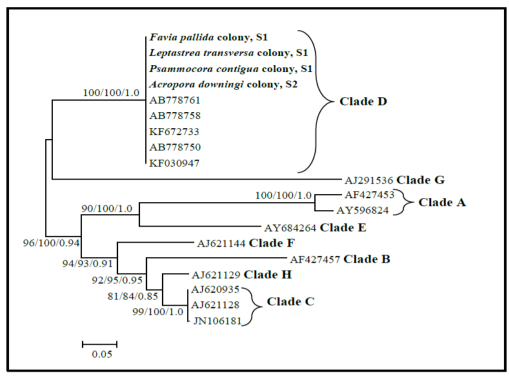


Figure 4: Maximum likelihood tree of LSU for clade D. Values at branches represent maximum likelihood bootstrap percentages from 1000 trees/ maximum parsimony bootstrap percentages from 1000 trees/Bayesian posterior probabilities

Discussion

In coral reefs off Larak Island, colonies of F. pallida and L. transversa, hosted Symbiodinium clades A, C and D simultaneously, S. pistillata and P. damicornis were associated with both Symbiodinium clades A and C, P. contigua harbored clade D and also clade C. Although the majority of coral contained species different Symbiodinium clades; downingi specificity show single Symbiodinium strain, clade D.

Several studies have shown that most coral colonies harbor multiple clade of *Symbiodinium* (Berkelmans and van Oppen , 2006; Mieog *et al.*, 2007). Therefore, in the present study PCR was performed with clade-specific primers. It appears that the occurrence

of *Symbiodinium* clade in coral species is depends on a combination of factors such as symbiont availability (van Oppen *et al.*, 2001). Previous research conducted on the zooxanthellae of the Persian Gulf have demonstrated the occurrence of *Symbiodinium* clades D, C and A in this region (Baker *et al.*, 2004; Mostafavi *et al.*, 2007, 2013; Shahhosseiny *et al.*, 2011; Hume *et al.*, 2015). Consequently, three pairs of primers belonging to clade A, C and D were used to amplify the selected marker in these clades.

Most previous studies on the zooxanthellae of the Persian Gulf have shown that clade D of *Symbiodinium*, is the most abundant of all coral symbionts (Baker *et al.*, 2004;

Mostafavi *et al.*, 2005, 2014; Shahhosseiny *et al.*, 2011).

Globally, clade D is common to corals from environments with high temperature and salinity (Fabricius et al., 2004), warm temperature and low light (Ulstrup and van Oppen, 2003) and high turbidity (van Oppen et al., 2001; Chen et al., 2003). Therefore, it has been concluded that predominance of clade D in the Persian Gulf is due to the unfavorable condition of this region (Mostafavi et al., 2007). However, among the coral species collected in this study, clade D was not the most abundant symbiont. The results of the present study have shown that four colonies of the six species harbored clade A and/or D, while five of specimens colonies six associated with clade C. The clade C dominance reported here agrees with another study of Symbiodinium from the southern Persian Gulf (Hume et al., 2015).

Although Clade C was more dominant *Symbiodinium*, other clades including A and D were commonly detected in corals off Larak Island.

Clade A of *Symbiodinium* have been found in corals inhabiting very shallow high-irradiance waters (Rowan *et al.*, 1997). It is known that only clade A of *Symbiodinium* are capable of producing considerable amounts of mycosporine-like amino acids (MAAs) (Banaszak *et al.*, 2000), compounds that help to protect against damage from UV radiation (Neale *et al.*, 1998). On the other hand, clade D of *Symbiodinium* is

thermally tolerant and increases the resistance of corals that harbor them to elevated sea surface temperature. Furthermore, the thermally resistance of clade C has been reported previously (Mostafavi *et al.*, 2013). It can therefore be concluded that the simultaneous presence of these clades in common coral species of the Persian Gulf may impart some protection from future stresses that lead to coral bleaching.

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