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Cryptic Species: A Mismatch between Genetics and Morphology in *Millepora*

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**Abstract** Millepore morphology is highly variable and shows signs of phenotypic plasticity. Two species of *Millepora* are present around the islands of the Bahamas: one exhibiting a strong, blade-like structure, *M. complanata*, and the other having a delicate branch-like structure, *M. alcicornis*. The phylogenetic relationship of these corals has been under considerable debate for many years. The existence of a range of intermediate growth forms exhibiting characteristics of both recognized species has led to the re-examination of this species complex. Several methods were employed to examine the taxonomic relationship including ecological abundance surveys, morphological thin-section analysis, and sequencing of rDNA internal transcribed spacer (ITS) regions. Abundance surveys showed a demarcation of growth forms by depth at two sites but an intermingling of growth forms at a third site. Morphometric analysis resulted in discrimination between *M. alcicornis*, *M. complanata* and the intermediate growth forms. However, rDNA sequence differences revealed the presence of two distinct clades, each containing members of the two currently recognized species as well as intermediate growth forms. The sequence analysis suggests the presence of two, phenotypically plastic cryptic species. Although limited in scope, our results indicate that caution should be exercised when describing species based on morphology alone and that multiple characters, including genetic information, should be used when describing species relationships.

**Keywords** *Millepora*, Phenotypic Plasticity, Morphometric, Cryptic Species, rDNA, ITS

1. Introduction

The genus *Millepora* (family Milleporidae, class Hydrozoa, phylum Cnidaria), commonly referred to as “fire-coral,” is an integral part of reef communities[1]. Fire corals serve as important framework builders, second only to scleractinian corals[2]. This framework is supported by an algal zooxanthellate symbiont, which aids in light-enhanced calcification[3]. Millepores are distributed worldwide in tropical seas and typically range in depth from less than 1m to approximately 40m[4]. The habitats they live in can range, depending mostly on growth form, from strong turbulent shallow waters to sheltered deeper waters[5]. The morphology of the millepores is highly variable and is believed to show phenotypic plasticity[1,6].

Phenotypic plasticity is believed to be molded by selection, such that certain phenotypes are better able to exploit a given environment[7]. Hence, a given species may exhibit different phenotypes depending on the environment. Plasticity has been described in many marine taxa including corals[8-9], sponges[10], fish[11], barnacles[12], and mollusks[13]. The presence and range of taxa shown to exhibit phenotypic plasticity must be taken into consideration when applying morphological characters to define species.

The various growth forms of *Millepora* in the Caribbean (Figure 1) range from thinly encrusting sheets and delicate dendroid branches, *M. alcicornis*, to thicker, rigid bladed forms, *M. complanata*[1]. It is this variation in morphology that has led to constant controversy about Millepore classification. Debate about taxonomy is not restricted to the millepores; morphological plasticity has also been documented in many scleractinian corals[14].

Since *Millepora* was first recognized by Linnaeus in 1758, many naturalists have worked on the millepores and their research has resulted in numerous and widely varied classification schemes[6]. When first described, millepores were primarily classified using morphological characters such as texture of the surface of the coral, size and shape of pores, stinging properties, and visual appearance[15]. Early investigators[16] found *Millepora* to be quite diverse and recognized 22 different species from the Caribbean.

Later, Hickson[17] suggested that all recognized species were environmentally controlled growth forms of a single
Millepora species. Boschma[4] recognized ten species of Millepora, three from the Atlantic Ocean and seven from the Indian and Pacific Oceans. Although 50 species of millepores have been described by Vernon[18], currently there are 17 recognized extant Millepora species; 11 from the Pacific and six from the Atlantic[19-20]. Wide variation in growth form of all species and a lack of diagnostic morphological characters presents serious problems for correct identification at the species level.

De Weerdt[15] examined the importance of morphological characters when distinguishing between species of Millepora. Multiple surveys of M. alcicornis and M. complanata in the Caribbean showed that the growth forms are widely overlapping in the environments they inhabit. This wide range implied that the differences in morphology could not be attributed solely to plasticity, and that there is a genetic component controlling growth. Transplantation experiments conducted in the Caribbean, showed that bladed forms (M. complanata) developed finger-like projections when relocated to deeper depths, which seems to be optimal for the branching form (M. alcicornis), and that the branching form became more robust when relocated to shallower depths, which seems to be optimal for the bladed form[15]. De Weerdt[15] concluded that morphological characters can change depending on environment and thus are not conclusive indicators of species.

The recent use of rDNA sequence and the development of coral-specific primers have made more accurate taxonomic classification possible. The inherent uncertainty, due to phenotypic plasticity, in using morphological characters as a way to classify species, can be aided by determining genetic distance, or relatedness, between closely related growth forms using rDNA sequences[21]. Previous studies on a wide range of organisms have suggested that the internal transcribed spacer (ITS) regions of ribosomal DNA (rDNA) are highly variable and thus suitable for comparative genetic studies of closely related species and populations[22-23]. In eukaryotes, the nuclear ribosomal small subunit gene, 18S, is separated from the 5.8S gene by an internal transcribed spacer(ITS-1) region and the 5.8S gene is separated from the 28S gene by ITS-2. Ribosomal genes and their spacers evolve at different evolutionary rates[21] making this gene family an ideal candidate for untangling species taxonomic relationships. The 18S, 5.8S and 28S genes are highly conserved, but following transcription of these genes, the spacer regions are excised prior to the incorporation of the rRNA into ribosomes. Since the ITS regions do not contribute to formation of the ribosomes, it is somewhat free to accumulate mutations. This leads to higher rates of evolution because the ITS spacers have fewer functional constraints compared to the ribosomal genes (18S, 5.8S and 28S).

Takabayashi et al.[24] examined the ITS regions encompassed by the coral specific primer A18S[25] and the universal primer ITS 4 for seven different coral species. Takabayashi et al.[24] compared the relatedness of samples from Acropora, Seriatopora, Goniopora, Portites, Heliofungia, and Stylophora to each other, as well as to replicates within the same species. Amplified fragment size and sequence data were used to distinguish between the six different genera mentioned above, and between eight different samples of Acropora longicyathus. Takabayashi et al.[24] reported that the ITS region varied from 2 to 31% in different coral species making this region ideal for comparative analyses between populations. The ability to distinguish samples between and within species made this fragment a powerful tool for the phylogenetic study of corals.

Meroz-Fine et al.[26] utilized a combination of morphological characters and DNA sequence from the ITS region of the Red Sea fire-coral, M. dichotoma, to show that the currently recognized single species with two growth forms (blade and branching) was in fact two distinct species. They reported that the average ITS sequence divergence between growth forms was 11.9% while the average divergence within a growth form was between 3.7 to 4.5%. Meroz-Fine et al.[26] also reported that they could distinguish between the two growth forms based on the size of the amplified ITS region. The bladed form was composed of 900 base pairs while the branching form was composed of 800 base pairs. While much of the work conducted on Millepora has been done on M. dichotoma, little work has focused on the two prevalent millepores found in the Caribbean.

Initial abundance surveys on Millepora conducted in early 2003 at various sites around San Salvador Island, Bahamas revealed the presence of a wide range of morphologies that did not easily fit into the current classification scheme of the
two recognized species. The new morphologies showed characteristics of both *M. alcicornis* and *M. complanata* (Figure 2). These new growth forms were termed intermediates and the present study is an attempt to explain the phylogenetic relationship of these intermediate growth forms to the other two recognized species.

The purpose of our research is to determine whether the colony morphologies represented by the described species of *Millepora* are matched by genetic isolation. We hope to distinguish among four hypotheses: 1) Millepores of the Bahamas are heterogeneous assemblages of genetically distinct forms. 2) The described "species" are a spectrum of colony growth forms reflecting ecological conditions rather than genetic isolation. 3) The range of growth forms observed is the result of extensive hybridization. 4) Millepores are reproductively isolated cryptic species and that traditional macro- and microskeletal features used for classification cannot distinguish them.

**Figure 2.** Photographs depicting examples of intermediate growth forms of *Millepora* found around San Salvador Island, Bahamas

## 2. Materials and Methods

### 2.1. Collection Sites

Millepores used in this study were collected from reefs surrounding the island of San Salvador, Bahamas (Figure 3). San Salvador is located on the eastern edge of the Bahamas Island chain and is characterized by its karst and hypersaline lakes. Patch reefs included for collection were Lindsay Reef (24°00'32"N, 74°31'59"W), Rocky Point Reef (24°06'25"N, 74°31'17"W), and French Bay (23°56'59"N, 74°32'50"W) (Figure 3). Lindsay Reef and Rocky Point Reef are shallow reefs with maximum depths of approximately 5m. French Bay has both shallow (1-5m) and deep (5-10m) patch reefs.

**Figure 3.** Satellite image San Salvador, Bahamas showing the three collection sites[27]

### 2.2. Ecological Abundance Surveys

Twenty meter line transects were laid down at random points at both French Bay and Lindsay Reef. Transects 1-3m in depth were surveyed by snorkeling, and transects deeper than 3m were conducted using SCUBA. All *Millepora* colonies within one meter of either side of an outstretched tape measure were recorded giving a total area surveyed for each transect of 40m². Only colonies with the classic finely branched morphology were considered to be *M. alcicornis* and only bladed colonies with a complete absence of branches were considered to be *M. complanata*. *Millepora* colonies not meeting these specifications were recorded as intermediate growth forms.

### 2.3. Sample Collections

Coral samples were randomly collected from each of the aforementioned reefs by removing a small piece, approximately 4 sq. cm in size. Samples of *M. alcicornis*, *M. complanata*, and the intermediate morphology were transported from the collection sites to the Gerace Research Centre (San Salvador, Bahamas) in buckets containing seawater and held in a flow-through seawater tank for no more than two days until coral DNA was isolated.

### 2.4. Morphometric Analysis

Twelve specimens of *Millepora* were subjected to morphometric analysis following the method outlined by Amaral et al.[28]. Specimens included five individuals assigned to *M. complanata*, three individuals of *M. alcicornis* and four individuals that exhibited intermediate growth forms.

Specimens were cut and mounted onto standard microscope slides, ground to standard thin sections (30µ) and analyzed using a petrographic microscope. A series of 25 mm² grids was superimposed on each thin section such that replicate, non-overlapping grids could be analyzed. Each
grid was photographed using a SONY ExwaveHAD digital video camera and imported into Photoshop for analysis.

A variety of features of the skeletal microstructure were quantified using the Analysis Tools module in Photoshop. These included diameter and surface area of all gastropores and dactylopores, distance between gastropores, dactylopores, and their density (# pores/grid). The number of dactylopores associated with each gastropore also was recorded. Variable numbers of individual gastropores (8-23) and dactylopores (27-129) were measured from each grid. Total number of grids measured was determined by the surface area of the hydrozoan present on each thin section. N = 3-5 grids per coral (8-129/grid, see above). Within- and between-grid values were averaged. Since no difference was observed using pooled vs. unpoled data, pooled data from each specimen are presented here. A Q-mode distance matrix was generated from the average values obtained from each specimen using the Euclidean Distance Coefficient. The matrix was subjected to Cluster Analysis (UPGMA algorithm). An Analysis of Similarity (ANOSIM)[29] was performed to assess significance of the differences observed between samples (both procedures in Primer v. 6.15). The Q-mode dendrogram was compared to the results of molecular genetic analysis.

2.5. DNA Techniques

Genomic DNA was isolated using a procedure modified from Rowan and Powers[30] and Lopez et al.[31]. Coral tissue was removed by repeatedly blasting the skeleton with a 50cc syringe containing L buffer (100mM EDTA, 10mM Tris, pH 7.6). Coral tissue was centrifuged at 3500rpm for 10 minutes; the resulting pellet was washed in 10mL of L buffer and re-centrifuged. The tissue pellet was resuspended in 900µL of L buffer and macerated manually with a tissue homogenizer. The homogenate was centrifuged twice at 500rpm for 10 minutes in order to separate the coral tissue from the liberated zooxanthellae. Following the addition of 1% (w/v) SDS to the supernatant, the lysate was incubated at 65°C for 30-60 minutes. Pro K (0.5 mg/mL) was added and the lysate was incubated at 37°C for at least 6 hours. NaCl (0.8M) and CTAB (1% w/v) were added and the sample was incubated at 65°C for 30 minutes. Nucleic acids were precipitated twice in 70% (v/v) ethanol and 3M sodium acetate (pH 5.2) and immediately centrifuged. Following resuspension of the pellet in dH2O, the DNA was briefly centrifuged and the supernatant was retained.

ITS rDNA PCR amplification was performed using 100-300ng of template, 60pmol of the coral specific primer A18S (5'GATCGAAAAGGTATTTATGTAGGG-3') and 60pmol of the universal primer ITS 4 (5'-TCTCCGGCTTTGATATGC-3')[25], 10X Tfl PCR buffer (Promega, Madison, WI), 2.0mM MgSO4, 0.1mM dNTP and 1U of Tfl polymerase. The PCR profile was: 1 cycle of 94°C for 2 minutes; 30 cycles of 94°C for 1 minute, 55°C for 2 minutes, and 72°C for 3 minutes; and 1 cycle of 72°C for 5 minutes[26]. Amplified PCR products were run on 1.2% (w/v) low melting agarose gels. Discreet, prominent bands were excised and purified using the Wizard SV Gel and PCR Clean-Up System (Promega). Purified PCR products were ligated into pGEM-T vectors following the manufacturer’s protocol (Promega) and transformed into competent DH5α E. coli host cells. Following blue-white selection, positive colonies were harvested and plasmid DNA was isolated using the Zippy Plasmid Miniprep Kit (Zymo Research, Orange, CA). Plasmids containing ITS rDNA inserts were sequenced in both directions with 3.0 pmol of FM13 forward and reverse primers. Reagents and reaction conditions for sequencing were as specified by the USB Thermo Sequenase Cycling Sequencing Kit (USB, Cleveland, OH). PCR products were run in 5.5% KB Plus Gel Matrix acrylamide using a LI-COR 4300 DNA Analyzer (LI-COR, Lincoln, NE). Sequence reaction products were analyzed using e-Seq V3.0 (LI-COR).

2.6. Phylogenetic Analysis

Maximum likelihood trees were produced using MEGA 4.0.1. The sequence of Millepora exesa[32] was used for the outgroup (GenBank, accession no. U65484) and 1000 bootstrap replicates were performed. Nucleotide percent substitution was also calculated from sequence data and compared within and between morphologies using MEGA 4[33] (The Biodesign Institute, Tempe, AZ).

3. Results

3.1. Abundance Surveys

![Image](327x182 to 538x356)

Figure 4. Population density of millopones separated by depth and specific reef. Error bars represent standard error of the mean. Shallow = 1-3m and deep > 3m. Data shown are from French Bay and Lindsay Reef. N (number of transects) = 19 for French Bay deep (total of 370 colonies were counted; 299 alcockinis, 0 complanata and 71 intermediates), N = 24 for French Bay shallow, (total of 557 colonies were counted; 66 alcockinis, 463 complanata and 28 intermediates) and N = 10 for Lindsay Reef (total of 63 colonies were counted; 24 alcockinis, 17 complanata and 22 intermediates).

Yearly reef surveys were conducted between 2003 and 2009. Year to year variability was minimal and the same trends were exhibited on individual reefs. Representative
data are shown in Figure 4 for French Bay and Lindsay Reef. The two locations surveyed exhibited different assemblages of *Millepora*. Lindsay Reef is a shallow reef that contained similar, low densities of *M. alcicornis*, *M. complanata* and intermediate growth forms. In contrast, shallow reefs in French Bay included both species and intermediates, but *M. complanata* colonies were far more abundant than either *M. alcicornis* or the intermediate growth forms. Deep reefs in French Bay contained only *M. alcicornis* and the intermediate growth forms; no *M. complanata* colonies were observed in these transects.

### 3.2. Morphometric Analysis

Morphometric data were able to discriminate between the two standard Milleporid taxa (*M. complanata* and *M. alcicornis*) as well as those specimens exhibiting an intermediate growth form (Figure 5). Specimens of *M. alcicornis* are “tacked on” to the cluster containing the intermediate growth forms at relatively low similarity values.

![Figure 5. Dendrogram obtained from reduction of a Q-mode Euclidean Distance Matrix. Differences between specimens are significantly different (p < 0.05) across the three growth forms (*M. complanata*, *M. alcicornis* and Intermediate)](image)

### 3.3. PCR Amplification of rDNA

Fragments of the ITS rDNA region (18S rDNA, ITS-1, 5.8S rDNA, ITS-2, and 28S rDNA) were amplified from bladed, branching and intermediate growth forms and a single PCR amplification product of approximately 825 base pairs was obtained from every sample (Figure 6).

![Figure 6. PCR amplification of rDNA using the A18S/ITS4 primers. The single band corresponds to a size of approximately 825 base pairs. Ma = *M. alcicornis* (branched growth form), I = intermediate and Mc = *M. complanata* (bladed growth form). The number following the sample name denotes the sample number. NT = no template controls and a 100 base pair ladder is shown](image)

### 3.4. DNA Sequence Analysis

A total of 36 samples were sequenced (17 of *M. complanata*, 12 of *M. alcicornis*, 7 intermediate growth forms) each yielding a sequence length of approximately 825 base pairs. The aligned sequences including gaps were 843 sites long and contained 778 conserved sites, 50 variable sites, and 10 parsimony informative sites.

Genetic variation, as nucleotide percent substitution, both within and between morphologies of *M. alcicornis*, *M. complanata* and intermediate growth forms was relatively low ranging from 0.6% to 0.9% (Table 1). Nucleotide substitution rates were higher when *M. complanata* was compared to *M. exaesa* from the Red Sea[32].

<table>
<thead>
<tr>
<th>Morphology</th>
<th>M. complanata</th>
<th>M. alcicornis</th>
<th>Intermediate</th>
</tr>
</thead>
<tbody>
<tr>
<td>M. complanata</td>
<td>0.6%</td>
<td>0.8%</td>
<td>0.6%</td>
</tr>
<tr>
<td>M. alcicornis</td>
<td>0.9%</td>
<td>0.6%</td>
<td>0.6%</td>
</tr>
<tr>
<td>Intermediate</td>
<td>0.7%</td>
<td>0.7%</td>
<td>0.7%</td>
</tr>
<tr>
<td>M. exaesa</td>
<td>12.2%</td>
<td>---</td>
<td>---</td>
</tr>
</tbody>
</table>

Table 1. Nucleotide Percent Substitution in rDNA Sequence between and within the Three Morphologies of *Millepora* using A18S and ITS4 Primers. A Comparison of the rDNA Sequence between *M. complanata* and *M. exaesa* is also Included

Maximum likelihood (ML) analysis of ITS rDNA sequences produced consensus topologies with two major clades (Figure 7). The two clades each contained members of all three morphologies (*M. complanata*, *M. alcicornis*, and intermediate growth forms). Bootstrap values for the two main clades of the tree are represented as a percentage and were 75% for both clades (Figure 7).
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**Figure 7.** Maximum likelihood tree showing bootstrap values for 36 samples of *Millepora* from the full range of phenotypes. 03, 05 or 06 denote the year the sample was collected (2003, 2005 or 2006). MA = *M. alcicornis*, MC = *M. complanata*, I = intermediate growth form. Numbers after the species designations represent the sample number. FB = French Bay, LR = Lindsay Reef and RP = Rocky Point. Bootstrap values are listed on the branchpoints. *M. exaeoa* was used as the outgroup.

**Figure 8.** Representatives of *Millepora* samples that are classified as clade 1 and 2. Collection sites are FB = French Bay reef, LR = Lindsay Reef, and RP = Rocky Point reef.

**Table 2.** Nucleotide Position within the rDNA Sequence, Nucleotide Polymorphism and Clade Identification of the Conserved Pattern of SNPs Found in all *Millepora* Samples Sequenced

<table>
<thead>
<tr>
<th>Nucleotide Position</th>
<th>Clade</th>
<th>Nucleotide Polymorphism</th>
<th>Region</th>
</tr>
</thead>
<tbody>
<tr>
<td>238</td>
<td>1</td>
<td>C</td>
<td>ITS-1</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>T</td>
<td></td>
</tr>
<tr>
<td>284</td>
<td>1</td>
<td>A</td>
<td>ITS-1</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>T</td>
<td></td>
</tr>
<tr>
<td>310</td>
<td>1</td>
<td>T</td>
<td>ITS-1</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>A</td>
<td></td>
</tr>
<tr>
<td>713</td>
<td>1</td>
<td>T</td>
<td>ITS-2</td>
</tr>
<tr>
<td></td>
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<tr>
<td>733</td>
<td>1</td>
<td>C</td>
<td>ITS-2</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>A</td>
<td></td>
</tr>
</tbody>
</table>

Upon closer examination of the rDNA sequence an interesting pattern emerged. The presence of five single nucleotide polymorphisms (SNPs) were found to be generally conserved across all samples sequenced (Table 2). This set of conserved SNPs directly corresponded with the two different clades formed by the phylogenetic analysis (Figure 7). Members of each morphology, from the full range of phenotypes, were observed in each clade (Figure 8).

### 4. Discussion

The taxonomy of the millepores is currently based on morphological characters and does not take into account genetic differences that may be present[15]. Taken alone, our morphometric results corroborate earlier work that distinguishes between the millepores based on morphology. The fact that *M. alcicornis* was not as clearly differentiated by our cluster analysis may in part be the result of a smaller sample size of this species in thin section (two-dimensional planes obtained from branching hydrozoan colonies are quite small). A more interesting result is the contrast between the outcomes produced by genetic and morphometric data, which suggests that the standard, morphologically based, taxonomy may be incorrect; two species of *Millepora* may exist around San Salvador that cannot be distinguished based upon morphology alone. This hypothesis suggests that two phenotypically plastic cryptic species are present and appear to be reproductively isolated from one another.

Both currently recognized *Millepora* species are morphologically plastic and their relative abundance was found to be different between study sites. Abundance surveys at French Bay suggest the two morphologies may be utilizing different habitats since one is predominantly found in shallower waters, while the other is found almost exclusively in deeper waters (Figure 4). At first glance, this data seems to support the current taxonomy, but when phenotypic plasticity is incorporated, the various growth
forms present may simply be the result of the different environments in which they are found. Additionally, the occurrence of all growth forms in mutual proximity on Lindsay Reef supports Stem and Riding's[1] contention that morphological variation in the Millepores is not primarily a response of a single species to environmental differences. Results from abundance surveys at Lindsay Reef suggest that genetic differences may exist between the growth forms.

Results from thin section analysis support the abundance data but suggests a different taxonomic relationship than the sequence data. Morphometric results indicate that pore size and arrangement can be used as a diagnostic tool to distinguish the two species of Millepora (Figure 5). A possible explanation for this discrepancy with the genetic data is again phenotypic plasticity. While the environment has been shown to alter macro-morphological characteristics via plasticity, it is not much of a stretch to suggest that the environment can alter micro-morphological characters, such as pore size, as well. While more work needs to be done in this area, it is our assertion that the environment plays a large role in determining the macro and micro morphologies of these corals.

Results from the ITS rDNA sequence comparison of the two purported species of Millepora and the intermediate growth forms show that the three morphologies are very closely related (Table 1). However, the rDNA ITS region exhibits considerable divergence when compared to the reproductively isolated M. exesa found in the Red Sea.

Takabayashi et al.[24] have shown that the size of the PCR amplified ITS rDNA fragment may be used as a diagnostic marker to distinguish between closely related species. Meroz-Fine et al.[26] reported that two growth forms of M. dichotoma that were classified as a single species contained ITS rDNA regions that were quite different in size and sequence leading them to conclude the two growth forms were different species. Regardless of the growth form sequenced, all of our millepore samples had a fragment length of approximately 825 base pairs suggesting that the millepores may be one plastic species. However, the phylogenetic tree (Figure 7) generated from these sequences demonstrated the presence of two clades, each containing members of all three morphologies (Figure 8), which further suggests the current taxonomy of Millepora may not be accurate. The conservation of the SNP pattern (Table 2) suggests that these two clades are reproductively isolated and the intermediate growth forms are not a result of extensive hybridization.

However, basing species-level phylogenetic reconstructions on ITS regions is sometimes problematic due to intragenomic sequence variation in the rDNA tandem repeats[34]. Variant rDNA copies can arise spontaneously in a single generation from point mutations. LaJeunesse and Pinzon[35] maintain that the dominant rDNA sequence in the genome can be used for phylogenetic reconstructions. Sequencing rDNA ITS regions following cloning, as was done in this analysis, sometimes leads to the detection of rare variants in the repeated rDNA sequences[35]. In order to determine whether the SNP pattern we have uncovered is a diagnostic species identifier or is due to the detection of rare cloning variants, we have begun an analysis using PCR-denaturing gradient gel electrophoresis in which rare rDNA variants from a single sample can be isolated and sequenced[36].

5. Conclusions

Our results indicate the possibility that two reproductively isolated cryptic species that are independent of growth form, depth and reef location may exist off the coast of San Salvador, Bahamas. These results have important implications for using the paleontological record for investigating taxonomic and evolutionary relationships between closely related hydrozoan taxa. Inasmuch as macro- and microskeletal features are the only recourse for elucidating such relationships for fossil hydrozoans, we submit that a close examination of the facies in which specimens are preserved accompany their identification. In this fashion, patterns of ecophenotypic plasticity may be explored in tandem with taxonomic analysis.

Many new Caribbean coral species and species complexes have been recognized by integrating molecular genetic and morphometric analyses[37-38] and these have been extended to include fossil corals[39]. However, the morphometric component of this work is carried out using landmark analyses of scleractinian coral skeletal microstructure. Hydrozoan skeletons are much less complex than those featured by scleractinians (for example no septa, much less septal ornamentation). It is possible that microstructural analyses of hydrozoans simply cannot yield results that mirror molecular genetic analyses.

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