INTRODUCTION

Visual responses of soft and hard corals to various insults such as disease, predation and competition can appear similar in a particular host. Thus, coral diseases or syndromes are often described in terms of changes of color or outward morphological features of the host. Little is known about the pathogenesis of many of these syndromes. The etiology of only 5 coral diseases has been demonstrated using Koch’s postulates (reviewed by Sutherland et al. 2004, Weil et al. 2006, Harvell et al. 2007, Bourne et al. 2009). In the Caribbean sea fan coral Gorgonia ventalina, the appearance of purpled tissue is caused by an
increase in the number of purple sclerites and is a generalized response to invading pathogens or other biotic agents (Petes et al. 2003, Alker et al. 2004, Smith & Weil 2004, C. S. Couch et al. unpubl.). Necrotic lesions, surrounded by purple halos in *G. ventalina*, have been described as aspergilliosis (Smith et al. 1996, Nagelkerken et al. 1997, Kim & Harvell 2001). The terrestrial fungus *Aspergillus sydowii* was isolated from fans with these signs and Koch’s postulates fulfilled in controlled laboratory experiments (Smith et al. 1996, Geiser et al. 1998). More recently, culture-dependent sampling showed that fungal species other than *A. sydowii* can be associated with lesions (Toledo-Hernández et al. 2008). During field surveys in Mexico, Florida, and Puerto Rico, a new type of lesion was identified (Weil & Hooten 2009, Weil & Rogers 2011; our Fig. 1) that is characterized by small (1 to 3 mm in diameter), circular to oblong multi-focal purple spots (MFPS; description after Work & Aeby 2006).

Preliminary histological evaluation of these multi-focal purple lesions revealed the presence of a stramenophile (ovoid parasites distributed within a mucus net; Petes et al. 2003) similar in appearance to the quahog parasite unknown (QPX) parasite of clams (Ragone Calvo et al. 1998, Ragan et al. 2000) and *Labyrinthula zosterae* that infects marine grasses (Durako & Kuss 1994, Ralph & Short 2002). Labyrinthulomycetes (Labyrinthulomycota) produce a very distinctive, basophilic-staining ectoplasmic network matrix with continuity from the network matrix to the cell cytoplasm (Porter 1990). Although Labyrinthulomycetes are common in estuarine and marine habitats, only a few of the identified Labyrinthulomycetes appear pathogenic and have been reported to be the primary infectious agent implicated in damaging epizootics, especially in the sea grasses *Zostera marina* and *Thalassia testudinum* (Muehlstein et al. 1991, Durako & Kuss 1994, Ralph & Short 2002), octopus (Polglase 1980), nudibranch (McLean & Porter 1982), abalone (Bower 1987), and the quahog clam *Mercenaria mercenaria* (Ragone Calvo et al. 1998, Ragan et al. 2000, Lyons et al. 2007).

Given the impact of Labyrinthulomycetes on the health of other marine invertebrates, our goal was to explore the disease dynamics of MFPS in sea fans and to characterize the Labyrinthulomycetes-like microorganism isolated from purple sea fan lesions. We surveyed sea fan disease at 2 reefs in La Paraguera, Puerto Rico (PR) and collected apparently healthy and MFPS-affected *Gorgonia ventalina* from sites in Florida and Puerto Rico to (1) isolate the potential pathogen, (2) identify and characterize the
Labyrinthulomycetes using histological, molecular, and culture techniques, and (3) perform preliminary inoculation experiments to fulfill Koch’s postulates.

MATERIALS AND METHODS

Sea fan surveys

**Puerto Rico**

Sixteen 20 m² band transects were randomly laid (4 transects per depth) at 2 reefs in La Parguera, PR—Media Luna (17° 56.093' N, 67° 02.931' W) (at depths of 3 to 18 m) and Buoy (17° 53.38' N and 66° 59.09' W) (at depths of 18 to 25 m). Transects were laid annually in September or October from 2006 to 2010 at Media Luna and 2007 to 2010 at Buoy. Along each transect all apparently healthy *Gorgonia ventalina* colonies and colonies with lesions were enumerated. Any purple or abnormally colored areas on the sea fans were checked carefully. Lesions were also recorded and described as follows: large irregular purple spots (aspergillosis-like; full description Work & Aeby 2006), MFPS, predation, bleaching, and other abnormalities. Other abnormalities included growth anomalies, cyanobacteria, red band disease, etc. Signs of predation by snails (flamengo tongue) and by fireworms were noted. Prevalence (the total number of each abnormality divided by the total number of individuals) of each type of abnormality was calculated per transect, depth, and year.

MFPS prevalence data was arcsine transformed before data analysis to meet the assumptions of normality and equal variances (Zar 1999). An analysis of covariance (ANCOVA) was performed to compare depth over time (using time as the covariate) using IBM SPSS Statistics 17.0 (Insightful). Statistical differences among depths were identified using Fisher’s least significant difference (LSD) test; however, due to the differences in depth and number of individuals within each site, the sites could not be compared.

**Florida**

From 20 sea fan colonies (n = 8 healthy and n = 12 MFPS), pieces of apparently healthy (n = 8) and MFPS tissue (n = 12), ~6 × 9 cm, were collected from Big Pine Ledges (24° 33.207' N, 81° 22.731' W), Florida Keys in February 2010. Samples were wrapped individually in wet paper towels, placed in separate bags and shipped overnight to the University of Washington with heat packs to keep the colonies warm. Upon arrival, fans were split among 4 aquaria that each contained 32 ppt seawater maintained at 25°C; 2 aquaria held apparently healthy sea fans and 2 contained MFPS fans. Five of the MFPS sea fans were used for parasite isolation; pieces of 6 healthy and 6 MFPS sea fans were stored in 70% ethanol for DNA analysis (see ‘Molecular methods’).

Sea fan collection

**Florida**

Apparently healthy and MFPS sea fans (n = 15 each) were collected from Media Luna Reef, La Parguera, PR, in July 2010. Fragments from each fan were divided in half (~5 × 5 cm) and were flash frozen or fixed in 10% seawater buffered formalin for 24 h followed by storage in 70% ethanol.

Labyrinthulomycetes isolation and maintenance

Based on past descriptions of Labyrinthulomycota-like cells in sea fans located within the inner axis (Petes et al. 2003, C. S. Couch & L. Mydlarz pers. comm.), the inner axis of sea fans from Puerto Rico and Florida were targeted for isolation. A square piece of *Gorgonia ventalina* tissue including the lesion and 0.5 cm of surrounding tissue was excised and dipped twice into 70% ethanol and filtered sea water (FSW) containing 200 U ml⁻¹ penicillin, 0.2 mg ml⁻¹ streptomycin, and 1 µg ml⁻¹ fungizone (FSW+PSF). The tissue section was then placed into a fresh container of FSW+PSF, and 0.25 cm of tissue around the border was cut off. The remaining tissue (the lesion plus 0.25 cm of surrounding tissue) was used for pathogen isolation where the piece of sea fan was cut down the central axis and then chopped into smaller pieces. These pieces were placed into 4 different 25 cm² T-flasks (Falcon) containing FSW+PSF and held for 30 min at room temperature. The seawater was then replaced with either QPX broth (Kleinschuster et al. 1998) containing 100 U ml⁻¹ penicillin and 0.1 mg ml⁻¹ streptomycin (PS) or Nutrient Broth (Difco) containing PS (200 U ml⁻¹ penicillin, 0.2 mg ml⁻¹ streptomycin). Half of the culture flasks were placed at 25°C and half at 30°C (one of each media type per temperature). Cultures were examined daily at 100× and 200× magnification for possible growth of Labyrinthulomycetes using a Nikon
Eclipse TS100 inverted microscope equipped with an Olympus digital camera for taking photomicrographs. When a possible Labyrinthulomycota was identified, fresh QPX broth and PS were added to the flask; after approximately 5 d, cells were scraped from the original culture and sub-cultured into a new flask with QPX broth and PS. Henceforth, every 5 to 7 d, the cultures were sub-cultured, whereby 50 µl of cells were scraped from the bottom of the flask and placed into a new flask with QPX broth and PS. No other microorganisms were present based on observations at 200×; nor have any been observed in 2 yr of culture.

Characterization of a Labyrinthulomycete

In culture

Photomicrographs were taken of the isolated Labyrinthulomycota using an Olympus digital camera. To obtain size estimates of the Labyrinthulomycota-like cells, 50 µl aliquots of culture media containing the cells were added to a microscope slide placed in a petri dish and allowed to grow for 3 d at 25°C before staining with the Hema 3 kit (Protocol, Fisher Scientific). Photomicrographs were taken at 20× magnification, and Labyrinthulomycetes total cell length was measured using Image J 1.44 p software (NIH).

Histology

Sea fan sections, approximately 2 × 2 cm, were decalcified using sodium citrate buffered (5%) formic acid (15%), embedded in paraffin, and sectioned at the Cornell University’s Veterinary Histology Laboratory (Luna 1968). De-paraffinized 5 µm tissue sections from the 15 apparently healthy and 15 MFPS sea fans were stained with hematoxylin and eosin and viewed under a bright field Olympus BH-2 microscope with the Olympus DP-20 camera system. From 16 of these sea fans (8 MFPS, 8 healthy), a second tissue section was taken and stained for presence of melanin using Fontana-Mason (FM) silver stain protocol (Mydlarz et al. 2008). Stained tissue sections were examined for presence/absence of Labyrinthulomycetes cells (and cell types) or other parasites within the tissues or axial cortex, and any other tissue or cellular abnormalities including melanization. In sea fans where >10 Labyrinthulomycetes cells were identified, photomicrographs were taken at 20× magnification, and total cell length was measured using Image J 1.44 p software.

Molecular methods

Total genomic DNA was extracted from approximately 1 cm² of ethanol preserved or frozen sea fan tissue using a blood and tissue DNA extraction kit (Qiagen) following the manufacturer’s instructions. To better facilitate extraction, the ethanol-preserved tissue and skeleton were diced into fine pieces with a fresh razor blade prior to extractions, and the frozen tissue was ground into a fine powder in liquid nitrogen with a mortar and pestle.

Total DNA, ~50 to 100 ng, (2 to 4 µl) from 5 apparently healthy and 6 MFPS sea fan tissues from Florida, 15 healthy and 15 MFPS tissues from Puerto Rico or 1 µl of genomic DNA from the 2 independent cultures of the microorganism isolated as described in ‘Labyrinthulomycetes isolation and maintenance’ above was used as template for 25 µl PCR reactions containing 1 U of Taq DNA Polymerase (Invitrogen), 1× PCR buffer, 2 mM MgCl₂, 0.4 mM BSA, 0.2 mM dNTPs, and 25 µM of each primer. Labyrinthulomycota small-subunit ribosomal DNA (SSU rRNA) specific primers, Laby-A and Laby-Y, were used (Stokes et al. 2002). Cycling conditions were: 94°C for 30 s, 50°C for 30 s, 72°C for 1.5 min for 35 cycles followed by a final 5 min extension at 72°C. Five µl of each PCR reaction were separated on a 2% agarose/TAE gel for use in purifying the PCR products amplified from the 41 samples and 2 isolates. Those containing bands of the expected (~430 bp size) were purified using the QIAquick PCR kit (Qiagen) and sent to the Cornell University Life Sciences Core Laboratories for direct sequencing on an Applied Biosystems Automated 3730 DNA Analyzer. Although direct sequencing of all of the PCR products showed unambiguous DNA chromatographs throughout, the homogeneity of samples was additionally confirmed by cloning the PCR products into the TOPO-TA vector, using the TOPO-TA cloning kit (Invitrogen) for sequencing of 2 to 5 clones of each isolate. The basic local alignment search tool (BLAST) algorithm (Altschul et al.1990) was used to compare resulting sequences, including the overlapping 398 nucleotides (nt) from Florida samples and 292 nt from Puerto Rico samples, with those deposited in the National Center for Biotechnology Information (NCBI) GenBank database, and alignments were made using the EMBL-EBI Clustal W tool (Larkin et al. 2007, Goujon et al. 2010). Representative se-
sequences from the cultured microorganism and sea fan isolates were submitted to GenBank (accession numbers JQ248602 and JQ248603 to JQ248606, respectively).

**Temperature and growth optima**

Two types of assays were used in 2 trials to quantify Labyrinthulomycota cultures: cell counts using a hemocytometer and total protein concentration. In the first trial, a Labyrinthulomycota culture held at 22°C was divided into 18 sub-cultures that were incubated at 15, 20, 25, 30, and 32°C in triplicate for 3 d. Culture temperatures and incubation period were based on previous visual observations of Labyrinthulomycota growth, where over-growth of culture flasks occurred after 3 d at temperatures of 25°C and higher. In the second trial, 9 sub-cultures were incubated at 20, 25, and 30°C in triplicate for 3 d.

In the second trial, total protein was also assessed. After 3 d, the media was poured off, rinsed once with 3 ml of 0.22 µm-filtered artificial sea water, and replaced with 3 ml of 0.22 µm-filtered artificial sea water. With a sterile wooden dowel, the bottom of each culture was scraped until no Labyrinthulomycota growth was visible on the bottom of the culture. Seven hundred µl of each culture was placed in a bead beater and mixed at 300 rpm for 30 s; 400 µl was set aside for protein assays and 300 µl for cell counts and held on ice until use.

Prior to counting using a hemocytometer, cells were vortexed for about 20 s. In each culture, cells were counted in triplicate. In the first trial, only total cell counts were noted. In the second trial, cell types were distinguished as either trophic cells (solitary Labyrinthulomycetes cells) or sori (Labyrinthulomycetes cells containing spores) (Fig. 2). No other cell types (i.e. zoospores) were distinguishable or present.

Total protein was extracted from each sample by adding 400 µl of extraction buffer, 0.15 µg ml⁻¹ dithiothreitol (DTT) in Tris-HCl, to each tube. The contents of the tube were mixed and lysed for 2 min with the Fisherbrand disposable pestle grinder system and incubated for 45 min on ice for extractions. Protein was measured using the DC protein kit (Bio-Rad) and read in triplicate using the Synergy HT multi-Detection microplate reader with KC4 software (Biotek Instruments) at 750 nm.

Both cell counts and protein concentrations exhibited departures from normality due to high levels of variation and were log-transformed. Differences among temperature treatments and between trials (total cell counts only) were assessed using an analysis of variance (ANOVA) and identified with the Tukey post hoc test (Zar 1999). Statistical analyses were run using IBM SPSS Statistics 17.0 (Insightful).

**Inoculation experiments**

Pieces (6 × 9 cm) of 12 healthy Gorgonia ventalina individuals (n = 2 pieces per sea fan in Trial 1 and n = 3 pieces per sea fan in Trial 2) were collected at Laurel patch reef, La Parguera, PR (17°56.608’ N, 67°03.208’ W) in May and September 2010, and held in situ to heal on the reef for 3 d. Sea fans were then collected from the reef and moved into static 10 gallon (ca. 38 l) aquaria at the University of Puerto Rico, Isla Magueyes Laboratory in La Parguera. Fans were distributed equally, in a clonally replicated design, using the fragment described in ‘Sea fan collection’ above, and were acclimated for 2 d. Seawater was changed daily.

In the first trial, one fragment of each of the 12 sea fans was injected with the Labyrinthulomycota culture (~4000 cells per injection point) and the other fragment injected with the QPX media only. Specifically, the inoculum was injected into the central axis in 3 locations on the surface of each sea fan fragment, ~3 cm apart. The sea fans were then held for a week before collection for routine paraffin histology (as described in ‘Histology’) where areas adjacent to the...
infection points (n = 3) were chosen. In the second trial, similar methods were followed, except for an inclusion of a bath treatment for exposure, in addition to the injection method used in the first trial; also, the trial was extended to 14 d. In each aquarium in the bath treatment, sea fans were exposed to 5 ml (~400,000 cells) of Labyrinthulomycota culture for 24 h. Temperature was held at ambient 29.2°C (range: 28.6 to 32.5°C) in Trial 1, and 28.8°C (28.2 to 31.5°C) in Trial 2.

RESULTS

Sea fan surveys

Puerto Rico

An average of approximately 245 (range: 243 to 247) sea fan colonies were surveyed at Media Luna (survey years: 2006 to 2010) and 150 (138 to 161) at Buoy (survey years 2005 to 2010). Survey data (non-transformed) for all abnormalities are shown in Fig. 3A,B, and trends in MFPS are shown in Fig. 3C,D. At Media Luna, MFPS exhibited positive correlations ($R^2 = 0.674$) in time (p < 0.0001) from 2006 to 2010, with significant differences among depths (p < 0.0001), where pairwise comparisons indicate the regression line for each depth to be different (p < 0.05) (total ANCOVA: $F = 38.801, df = 4, p < 0.0001$). No correlation between MFPS and time was observed at BUOY (p > 0.05), and significant differences were only seen at depth where MFPS detected at 18 m was greater than that at 17, 20, or 25 m (p < 0.0001; total ANCOVA: $F = 10.04, df = 4, p < 0.0001$). No correlation was observed between large irregular lesions over time at Media Luna (p > 0.05), although depth was significant (p = 0.02) where fans in 6 m of water differed from those at 3, 10, and 18 m (total ANCOVA, $F = 2418.044, df = 4, p = 0.04$). No other variables exhibited statistically significant correlations over time or depth (p > 0.05). MFPS were not correlated with any other abnormalities.

Characterization of a new Labyrinthulomycetes

In culture

After incubating tissue and skeleton of sea fans collected from Florida in QPX medium for 3 d, Labyrinthulomycetes trophic cells were first observed in culture. After 5 d of growth, trophic cells, sori, and spores (Porter 1990) were observed (Fig. 2). Based
on published descriptions of morphology of the trophic cells and sori of the Labyrinthulomycota (Porter 1990), the cultured Labyrinthulomycetes best matched the genus *Aplanochytrium*, a member of the Family Thraustochytridae. Similarly, Labyrinthulomycetes cultured on glass slides and stained with Giemsa included trophic cells (length = 10.22 ± 0.15 µm, mean ± SE, range 4.41 to 16.12 µm, n = 150), spores (4.48 ± 0.14 µm, 2.36 to 7.31, n = 57), and sori (18.61 ± 0.89 µm, 12.77 to 31.43, n = 225).

**Histological**

In both sea fans exhibiting MFPS (n = 8; Table 1) and apparently healthy colonies (n = 7), tissue sections revealed ovoid to elongated trophic cells either solitary or undergoing binary fission and ectoplasmic networks (mucus) mainly within a space—apparently where the proteinaceous gorgonin had split apart—usually just beneath the cortex of the axis or sometimes within the central chord or core of the axis (Fig. 4A). Labyrinthulomycetes cells averaged 9.51 ± 0.47 µm (range 5.61 to 16.3) in sea fans with MFPS (n = 100 cells) and 8.49 ± 0.38 µm (range 4.9 to 13.4) in apparently healthy sea fans (n = 100 cells) (Table 1). The presence of Labyrinthulomycetes cells was generally limited to relatively small areas of the sea fan colonies, and the majority of sea fans did not display any obvious tissue changes or damage other than the longitudinal splitting of the gorgonin axis described above. Occasionally, cells with a mucus trail (ectoplasmic net) were seen in the coenenchyme or sea fan
polyps (Fig. 4B). In 2 of the sea fan colonies analyzed, infections were more systemic where large numbers of Labyrinthulomycetes cells were visualized and colonies were devoid of polyps. No apparent entry point or mode of entry was observed.

Invertebrates, including copepods and worms, were also observed in the coenenchyme within grossly purple MFPS tissue of some (6.7% of apparently healthy and 33% of MFPS, 20% of which were accompanied by Labyrinthulomycetes infections) of the sea fans. Fungal hyphae were also noted in the axis of 1 MFPS sea fan and 2 apparently healthy sea fans (n = 15 respectively), alongside infection with the Labyrinthulomycetes.

FM staining indicated a host melanization response only in areas where Labyrinthulomycetes or fungi were present (Fig. 4C), and in most fans examined (7 out of 8), melanization occurred with Labyrinthulomycetes infection. In the one fan where melanization was not detected, only 2 Labyrinthulomycetes cells were noted. Non-melanized gorgonin ranged in color from pink in the axis center (or clear) to light brown or tan (Fig. 4D).

**Molecular**

Detectable PCR products were amplified with primers targeting the phylum Labyrinthulomycota (Stokes et al. 2002) from the cultured cells and many of the field-collected samples as follows: 4 out of 5 healthy colonies from Florida, 3 out of 6 MFPS samples from Florida, 7 out of 15 healthy samples from Puerto Rico, and 5 out of 15 MFPS samples from Puerto Rico (Table 1).

Alignment of all of the Labyrinthulomycota SSU rRNA sequences derived from the cultured cells, healthy and MFPS sea fan tissue from Florida strongly suggests that the 3 sources contain the same microorganism. All of the sea fan derived sequences are identical to one another within this amplified SSU rRNA region. This sequence is also identical (100% similarity) to multiple SSU rRNA sequences in the NCBI database from the genus *Aplanochytrium/Labyrinthuloides* (synonymized by Leander & Porter 2000 as *Aplanochytrium*), suggesting that the sea fan isolate is very likely a member of this genus in the family Thraustochytridae.

Of the Puerto Rico samples that amplified with the Labyrinthulomycota SSU rRNA primers, 5 healthy and 4 MFPS were sequenced. Of these, 2 of the Labyrinthulomycetes sequences isolated from the apparently healthy sea fans matched the genus *Thraustochytrium* in the NCBI database based on identical sequences (100%) to multiple SSU rRNA sequences of the genus *Thraustochytrium*. The remaining healthy and MFPS-fan derived sequences appear to belong to the same *Aplanochytrium* genus as the Florida isolates (Table 1).

---

### Table 1. Gorgonia ventralina. Presence of Labyrinthulomycetes in sea fans (SF) with multifocal purple spots (MFPS) and healthy sea fans collected at Media Luna Reef, La Paraguera, Puerto Rico (July 2010) and summary data for sea fans collected at Big Pine Ledges, Florida (February 2010) based on histological and molecular evidence. Where positive PCR results were obtained using Labyrinthulomycetes-specific SSU rRNA primers (Stokes et al. 2002), sequence data indicate an *Aplanochytrium* sp. unless otherwise noted. nd: no data (Labyrinthulomycetes cells not measured); na: not applicable (no histological observations made).

<table>
<thead>
<tr>
<th>Specimen</th>
<th>PCR result</th>
<th>Presence</th>
<th>Histology</th>
<th>Mean ± SE length (µm) (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MFPS sea fans (Puerto Rico)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MFPS SF 1</td>
<td>+</td>
<td>+</td>
<td>nd</td>
<td></td>
</tr>
<tr>
<td>MFPS SF 2</td>
<td>+</td>
<td>−</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MFPS SF 3</td>
<td>+</td>
<td>+</td>
<td></td>
<td>10.66 ± 0.60 (13)</td>
</tr>
<tr>
<td>MFPS SF 4</td>
<td>+</td>
<td>+</td>
<td></td>
<td>8.38 ± 0.35 (16)</td>
</tr>
<tr>
<td>MFPS SF 5</td>
<td>+</td>
<td>+</td>
<td></td>
<td>10.23 ± 0.46 (15)</td>
</tr>
<tr>
<td>MFPS SF 6</td>
<td>−</td>
<td></td>
<td></td>
<td>8.75 ± 0.53 (11)</td>
</tr>
<tr>
<td>MFPS SF 7</td>
<td>−</td>
<td>−</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MFPS SF 8</td>
<td>−</td>
<td>−</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MFPS SF 9</td>
<td>−</td>
<td>−</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MFPS SF 10</td>
<td>−</td>
<td>+</td>
<td>nd</td>
<td></td>
</tr>
<tr>
<td>MFPS SF 11</td>
<td>−</td>
<td>−</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MFPS SF 12</td>
<td>−</td>
<td>−</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MFPS SF 13</td>
<td>−</td>
<td>+</td>
<td></td>
<td>9.49 ± 0.35 (19)</td>
</tr>
<tr>
<td>MFPS SF 14</td>
<td>−</td>
<td>+</td>
<td></td>
<td>9.56 ± 0.55 (28)</td>
</tr>
<tr>
<td>MFPS SF 15</td>
<td>−</td>
<td>−</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Summary</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Puerto Rico 5/15</td>
<td>8/15</td>
<td>9.51 ± 0.47</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Florida 3/6</td>
<td>na</td>
<td>na</td>
<td>na</td>
<td></td>
</tr>
</tbody>
</table>

| **Healthy sea fans (Puerto Rico)** |            |          |           |                          |
| Healthy SF 1         | −          | −        |           |                          |
| Healthy SF 2         | +          | +        |           | 9.39 ± 0.41 (19)         |
| Healthy SF 3         | +          | +        |           | 7.62 ± 0.62 (9)          |
| Healthy SF 4         | −          | −        |           |                          |
| Healthy SF 5         | −          | +        |           | 7.53 ± 0.07 (15)         |
| Healthy SF 6         | −          | −        |           |                          |
| Healthy SF 7         | +          | −        |           |                          |
| Healthy SF 8         | −          | +        |           | 8.73 ± 0.61 (10)         |
| Healthy SF 9         | −          | −        |           |                          |
| Healthy SF 10        | −          | −        |           | 7.54 ± 0.07 (15)         |
| Healthy SF 11        | +          |          |           |                          |
| Healthy SF 12        | +          |          |           | 9.77 ± 0.67 (7)          |
| Healthy SF 13        | +          |          |           | 9.86 ± 0.51 (16)         |
| Healthy SF 14        | −          |          |           |                          |
| Healthy SF 15        | +          |          |           | 7.54 ± 0.65 (8)          |
| **Summary**         |            |          |           |                          |
| Puerto Rico 7/15     | 7/15       | 8.49 ± 0.38 |                          |
| Florida 3/5          | na         | na      | na        |                          |

*Sequence data indicate a *Thraustochytrium* sp.

*No sequence data obtained*
Temperature and growth optima

Labyrinthulomycetes growth was quantified using cell counts in 2 trials. Total cell counts reported for Trial 1 indicated that growth was directly related to culture temperature (ANOVA, p < 0.0001), with the following statistical trend: 15°C < 20°C < 25°C < 32°C = 30°C (Tukey test, p < 0.05, Fig. 5A). A similar trend was observed in total cells for the second trial: 20°C < 25°C < 30°C (Tukey test, p < 0.05), although cell counts were different between trials (p = 0.037). In the second trial, where cell types were differentiated, counts differed significantly in total sori (p < 0.003) and total trophic cells between temperatures (ANOVA, p < 0.001): sori 20°C = 25°C < 30°C (Tukey test, p < 0.05), and trophic cells 20°C < 25°C < 30°C (Tukey test, p < 0.05; Fig. 5B). Total protein content in Trial 2 (Fig. 5C) was similar for cell cultures grown at 20°C (8.55 ± 0.89 mg protein culture⁻¹) and 25°C (8.96 ± 0.51 mg protein culture⁻¹), which were significantly lower than those held at 30°C (11.09 ± 0.84 mg protein culture⁻¹) (20°C = 25°C < 30°C ; ANOVA: p < 0.0001, Tukey test: p < 0.05).

Inoculation experiments

The inoculation experiments did not provide evidence of new Labyrinthulomycetes infections, as Labyrinthulomycetes cells were detected by histology in both control and inoculation treatments. In the first trial, Labyrinthulomycetes cells were visualized in 5 of the control replicates and 4 of the inoculated treatments (n = 12 colonies each; 2 colonies contained Labyrinthulomycetes cells in both control and treatment replicates). In the second trial, 1 lesion developed in each of 2 sea fans held in bath treatment with Labyrinthulomycetes cells: 1 sea fan contained Labyrinthulomycetes cells in association with the developed lesion. However, Labyrinthulomycetes cells were also detected in all of the control and bath treatment samples from these colonies.

DISCUSSION

The first step to validate disease etiology is pathogen isolation and characterization, and for the first time, Labyrinthulomycetes cells were isolated, cultured, and characterized from sea fans with MFPS. Identical sequence data were obtained from sea fans with MFPS from Florida and Puerto Rico, as well as most apparently healthy sea fans from both localities.
Labyrinthulomycetes from the genus *Aplanochytrium*. However, a second Labyrinthulomycetes sequence was detected in 2 healthy individuals collected in PR, a Labyrinthulomycetes of the genus *Thraustochytrium*. Observations including general pathology and morphology of cells (size) in sea fans from PR indicate that infections by Labyrinthulomycetes are localized within colonies (Table 1) and that *Aplanochytrium* and *Thraustochytrium* may be indistinguishable. The observation of Labyrinthulomycetes in both apparently healthy and MFPS sea fans implies an opportunistic, like many marine pathogens: often present in the marine environment but only pathogenic under some conditions. Therefore, additional work is needed to understand the pathogenesis of Labyrinthulomycetes in sea fans including whether MFPS are formed in response to Labyrinthulomycetes infections. Based on 5 yr of survey data collected off La Parguera, PR, incidence of MFPS is increasing in sea fan colonies, particularly MFPS at shallow depths. This indication of an emerging disease compels further characterization of the relationship between MFPS and biotic invaders.

Labyrinthulomycetes have been previously associated with apparently healthy cnidarian hosts (based on either light or electron microscopy or both) including hydroids (Raghukumar 1988) and scleractinian corals (Kramarsky-Winter et al. 2006, Siboni et al. 2010). Based on 18S rRNA sequences, Labyrinthulomycetes from the family Thraustochytriidae were associated with scleractinian corals: *Fungia granulosa* in the Gulf of Elat (/Aqaba) (genera unknown, Harel et al. 2008) and with *Favia* spp. from both the Gulf of Elat (/Aqaba) and the Great Barrier Reef (*Aplanochytrium* sp, *Thraustochytrium* sp, and *Labyrinthuloides* sp., Siboni et al. 2010). These results are similar to the molecular analyses of Labyrinthulomycetes present in sea fan colonies described in the present study. In scleractinian corals, Labyrinthulomycetes can form white patches on the host surface (Kramarsky-Winter et al. 2006) and have been described in the mucus, coral surface, and coral tissues (Raghukumar & Balasubramanian 1991, Kramarsky-Winter et al. 2006, Siboni et al. 2010). The Labyrinthulomycetes–scleractinian relationship may be mutualistic or commensal, with the thraustochytriids providing the host with nutrition and nutrient cycling in exchange for coral mucus as a carbon source (Siboni et al. 2010).

Unlike the mutualistic Labyrinthulomycetes relationships described in other cnidarians, our histological evidence suggests a parasitic relationship. In many infected sea fans, evidence of poor health included longitudinal splitting of the gorgonin axis, patches of Labyrinthulomycetes aggregations surrounded by melanized gorgonin and lack of polyps in colonies with the highest intensity infections. However, whether poor health is caused by Labyrinthulomycetes infection or compromised individuals allow the opportunistic Labyrinthulomycetes to invade is unknown. Melanin staining suggests a host response to the Labyrinthulomycetes, and similar melanin staining has been previously observed in purple tissue and axis of sea fans infected with fungus (Mydlarz et al. 2008) and/or Labyrinthulomycetes (Petes et al. 2003) as well as in pigmented *Porites* spp. (Palmer et al. 2008). Melanin encapsulation is a key component of the invertebrate immune response and is well characterized in insects and crustaceans (reviewed by Rowley 1996, Cerenius et al. 2010). Similarly, the melanization response plays a role in scleractinian wound healing (Palmer et al. 2011). Labyrinthulomycetes infections, similar to fungal infections of sea fans, appear to be opportunistic; both parasite types appear most frequently within the sea fan axis. It has been hypothesized that the skeleton may have lower levels of anti-bacterial (Kim et al. 2000) and anti-fungal compounds (Kim et al. 2000, Dube et al. 2002, Ward 2007, Ward et al. 2007) than are found in the coenenchyme (Ward et al. 2007), perhaps allowing initial infection. The melanization response by amoebocytes (Mydlarz et al. 2008) builds a thick wall over the fungus or Labyrinthulomycetes, in an effort to encapsulate or kill the parasites or perhaps to strengthen the axis. It appears that both parasites weaken the gorgonin by lateral spread of the hyphae or mucus net through the gorgonin or splitting it apart as the number of parasite cells and mucus increase, which weakens the gorgonin so the sea fan blade may be more susceptible to breakage from storm-driven wave and current damage.

Despite 2 inoculation trials, there was no evidence of cultured Labyrinthulomycetes infecting sea fans. In fact, histological evidence from these trials further corroborates that Labyrinthulomycetes are found in otherwise healthy sea fans. One possible explanation for this observation is that the sea fan colonies used in these experiments had been previously exposed and infected and may not be susceptible to re-infection, or that susceptible individuals were already infected, thus masking any *Aplanochytrium* cells acquired by exposure. Another explanation is that although both injection and bath challenges were attempted, possibly they did not replicate the Labyrinthulomycetes mode of entry into the sea fan.
tissues and axial skeleton or that the cultured cells were no longer infectious. Experimental inoculation trials were run within a temperature range that should be favorable for Aplanochytrium growth (28 to 32°C). The optimal growth-temperature was 30°C, similar to that of Aspergillus sydowii (Alker et al. 2004, Ward et al. 2007). In fact, fungal growth decreased at 31°C coincident with maximum sea fan host anti-fungal activity (Ward et al. 2007).

Pathogen-growth optima in culture may not translate directly to the dynamic seen in a host–pathogen relationship where host immune response and the pathogen growth have a complex interplay (Dahl et al. 2011, Perrigault et al. 2011). For example, although in vitro growth of the QPX, a thraustochytrid parasite of quahog clams, Mercenaria mercenaria, is greatest between 20°C and 23°C (Perrigault et al. 2010), both natural and experimentally produced infections in quahogs were more severe (based on prevalence and mortality) at 13°C (compared to 21°C and 27°C), where host immune responses were compromised (Perrigault et al. 2011). QPX infections are chronic, and QPX is regularly detected in healthy individuals, indicating an opportunistic parasite. In fact, both QPX strains and host genetics are important in the virulence of QPX disease (Dahl et al. 2008). This complex interplay of QPX disease in clams may provide clues to better understanding the sea fan–Labyrinthulomycetes relationship, as the same factors may be important for disease transmission and host immunity.

In conclusion, Labyrinthulomycetes of the family Thraustochytridae appear to be opportunistic parasites of sea fan corals. Although Labyrinthulomycetes were detected in association with MFPS, MFPS do not always indicate Labyrinthulomycetes infections and are likely a host response to biotic stressors. Additional time course sampling and surveys of natural populations may clarify the relationship between MFPS and Labyrinthulomycetes infection. The increase in purple spots may indicate compromised health in sea fans around La Parguera, PR. This hypothesis is supported by increases in disease in other corals species in this area (E. Weil pers. obs.) and in coral disease in the Caribbean and globally over the past 2 decades (Harvell et al. 2004, 2007).

Acknowledgements. This work was conducted under National Science Foundation grant number OCE0849776. Initial parasite isolation was supported, in part, by the School of Aquatic and Fishery Sciences at the University of Washington. Florida sea fan corals were collected by E. Bartels of the Mote Tropical Research Laboratory on Summerland Key under Florida Keys National Marine Sanctuary permit number FKNMS-2008-001. The authors thank C. S. Couch and E. Peters for editorial comments. The Department of Marine Sciences University of Puerto Rico provided logistical support for field work and lab experiments.

LITERATURE CITED


Editorial responsibility: Garriet Smith, Aiken, South Carolina, USA
Submitted: December 12, 2011; Accepted: May 16, 2012
Proofs received from author(s): September 12, 2012