Population Recovery and Differential Heat Shock Protein Expression for the Corals *Agaricia agaricites* and *A. tenuifolia* in Belize

Martha L. Robbart  
*Smith College*

Paulette M. Peckol  
*Smith College*

Stylianos P. Scordilis  
*Smith College*, sscordil@science.smith.edu

H. Allen Curran  
*Smith College*, acurran@smith.edu

Jocelyn Brown-Saracino  
*Smith College*

Follow this and additional works at: https://scholarworks.smith.edu/geo_facpubs

Part of the Geology Commons

**Recommended Citation**  
https://scholarworks.smith.edu/geo_facpubs/69

This Article has been accepted for inclusion in Geosciences: Faculty Publications by an authorized administrator of Smith ScholarWorks. For more information, please contact scholarworks@smith.edu
Population recovery and differential heat shock protein expression for the corals *Agaricia agaricites* and *A. tenuifolia* in Belize

Martha L. Robbart\(^1,3\), Paulette Peckol\(^1\)*, Stylianos P. Scordilis\(^1\), H. Allen Curran\(^2\), Jocelyn Brown-Saracino\(^1\)

\(^1\)Department of Biological Sciences, and \(^2\)Department of Geology, Smith College, Northampton, Massachusetts 01063, USA
\(^3\)PBS&J Environmental Services, 2001 Northwest 107th Avenue, Miami, Florida 33172, USA

**ABSTRACT:** Over recent decades, coral reefs worldwide have experienced severe sea-surface temperature (SST) anomalies. Associated with an El Niño-Southern Oscillation (ENSO) event of 1997–1998, nearly 100% mortality of the space-dominant coral *Agaricia tenuifolia* was reported at several shelf lagoonal sites of the Belize barrier reef system; a less abundant congener, *A. agaricites*, had lower mortality rates. We assessed *A. agaricites* and *A. tenuifolia* populations at coral reef ridges in the south-central sector of the Belize shelf lagoon and forereef sites to document recovery following the 1998 ENSO event and subsequent passage of Hurricane Mitch. To investigate the difference in heat stress tolerance between the 2 species, heat shock protein (HSP) expression was examined in the laboratory under ambient (28°C) and elevated (+6°C) temperatures. Populations of *A. agaricites* and *A. tenuifolia* surveyed at forereef sites in 1999 showed after effects from the 2 disturbances (partial colony mortality was ~23 and 30% for *A. agaricites* and *A. tenuifolia*, respectively), but partial mortality declined by 2001. At reef ridge sites, *A. tenuifolia* exhibited 75 to 95% partial colony mortality in 1999 compared to 18% in the less abundant *A. agaricites*. We measured a significant increase in percentage live cover at ridge sites for both *Agaricia* species from 1999 to 2001, except at Tunicate Ridge; at this site, which has restricted water flow, live *A. tenuifolia* cover remained low (~10%) 3.5 yr after the 1998 warming event, due in part to high sponge cover (>75%). Immunoblotting results indicated that *A. agaricites* had twice as much HSC 70 (16.9 µg cm\(^{-2}\)) as *A. tenuifolia* (8.7 µg cm\(^{-2}\)) at ambient temperatures and 6× as much under the +6°C treatment. In addition to the inducible response by *A. agaricites*, this species expressed HSP 90, whereas *A. tenuifolia* did not. The distinctive patterns of population recovery and HSP expression suggest that *A. tenuifolia* has a lesser ability to produce HSPs for protection against environmental stress than *A. agaricites*. Such differences in resilience to large-scale environmental disturbances such as intermittent ENSO episodes may drive a dramatic change in coral species abundance patterns.

**KEY WORDS:** *Agaricia* · Coral reefs · ENSO episodes · Heat shock proteins · Population dynamics

---

**INTRODUCTION**

Most reef-building scleractinian corals live near their upper thermal limits and may experience heat stress with an increase of only a few degrees centigrade (Jokiel & Coles 1977, Pitt et al. 2001). Widespread coral bleaching has been associated with thermal anomalies that occur with elevated sea-surface temperatures (SST) accompanying El Niño-Southern Oscillation (ENSO) events (Brown 1997, Wilkinson 2000). Bleaching also has been induced in laboratory heat shock experiments (Hayes & King 1995). Long-term studies of coral reefs affected by thermal anomalies with resultant coral mortality have revealed that recovery may take several decades (Glynn 1993, Eakin 1996).

Environmental stresses, such as increase or decrease in temperature or salinity, or exposure to heavy metals may deleteriously affect normal metabolic functioning at the cellular level. Proteins that are critical to the maintenance of cell function may become denatured as a result of prolonged or severe environmental stress. For example, heat stress may cause a breakdown of
enzymatic pathways with consequent biochemical and metabolic dysfunction (Cossins & Bowler 1987); effects from such a stressor may be exacerbated by influences from other environmental factors (i.e. irradiance, salinity, circulation) (Fitt 2001).

Often, such environmental challenges elicit a cellular response to protect an organism from permanent damage that may result from exposure to these extreme conditions (Craig 1985, Welch 1992). In response to thermal anomalies or UV radiation, Lesser et al. (1990) demonstrated increased activities of enzymes responsible for detoxifying active forms of oxygen in symbionts of the zoanthid *Palythoa caribaeorum*. Similarly, heat shock proteins (HSPs) serve a protective or reperative function and directly reduce the accumulation of denatured proteins caused by significant environmental change, including heat stress, heavy metals and other metabolic poisons (Craig 1985, Welch 1992, Sharp et al. 1997). Once exposed to non-lethal stress that elicits a response, namely, increased synthesis of HSPs, organisms often show a greater resistance to future, similar environmental disturbances. HSPs are thus vital to organisms in variable environments and are responsible for acclimation at the cellular level (Lindquist & Craig 1988, Laszlo 1992, Welch 1992).

The cosmopolitan distribution of HSPs across widely disparate taxa led researchers to focus on the role of these proteins in the physiological responses of scleractinian corals to thermal stress. Several HSPs have been documented in coral species, including constitutive (cognate) heat shock protein 70 (HSC 70), and inducible HSP 70, HSP 60 and HSP 25, under heat and UV radiation stress (Hayes & King 1995, Fang et al. 1997, Sharp et al. 1997, Branton et al. 1999, Downs et al. 2000). HSP 70 is the best-documented heat shock protein, largely because it is highly inducible. For example, field transplant experiments with the Pacific coral *Goniopora djiboutensis* demonstrated elevated levels of HSC/HSP 70 following heat shock (Sharp et al. 1997).

However, many studies identifying a 70 kDa stress protein did not differentiate between the constitutive (HSC 70) and inducible (HSP 70) forms (Welch 1992, Thompson et al. 2001). Differentiation between the constitutive and inducible forms is important because increased synthesis of HSC 70 may act to stabilize proteins at the onset of environmental stress prior to HSP 70 induction. Thompson & Scordillis (2001) documented elevated levels of HSC 70 in mice 15 min after exercise stress, suggesting this so-called ‘constitutive’ protein was acting as a ‘first defender’ following the stress. Therefore, distinguishing between HSC 70 and HSP 70 synthesis in organisms responding to environmental stress provides greater understanding of the mechanism of molecular stress response.

While bleaching events have affected the wider Caribbean since the early 1980s (Lasker et al. 1984, Williams et al. 1987), there were no records of mass bleaching of corals on the Belize (Mesoamerican) barrier reef system before the El Niño-Southern Oscillation (ENSO) episode of 1995 (Burke et al. 1996, McField 1999). Associated with this event, McField (1999) measured over 50% coral bleaching on the forereef region off Belize. Although most coral colonies recovered within 6 mo, ~10% experienced at least partial colony mortality. Again, in 1998, the Belize barrier reef was subjected to large-scale disturbances by a severe ENSO event (early August to October 1998) and Hurricane Mitch (October 25 to 31). Aronson et al. (2002a) reported positive (>1°C) SST anomalies during autumn 1998 for shelf lagoon and forereef regions off south-central Belize. SSTs exceeded the HotSpot threshold for over 2 wk and reached +4°C in late September. Hurricane Mitch, a category-5 storm on the Saffir/Simpson Hurricane Scale, passed directly over the Bay Islands of Honduras, south of Belize, and introduced enough fresh water to create a freshwater lens 3 m deep at Carrie Bow Caye in southern Belize (Aronson et al. 2002a). Following these major disturbances, Kramer & Kramer (2000) reported significant remnant bleaching at forereef sites off Belize 6 mo after the initial bleaching response. In contrast, Mumby (1999) found that although ~80% of coral colonies on the forereef region (0 to 10 m) of Glover's Atoll (Belize) appeared to be either fully or partially bleached, these colonies regained their usual coloration in subsequent months.

The 1998 ENSO episode had a particularly strong impact on the shallow shelf lagoonal region off south-central Belize. Mass bleaching was first observed in this area in early September 1998 (Nemecek 1999). In the Pelican Cayes, nearly 100% mortality was documented for the spacedominant coral *Agaricia tenuifolia* at lagoonal reef sites to a depth of 15 m (Aronson et al. 2000, 2002a, Peckol et al. 2003) following the massive and prolonged bleaching event. Patch reefs also experienced severe and lingering effects from bleaching, particularly for *Montastraea annularis*, a major reef-building coral (Peckol et al. 2003). A congener of *A. tenuifolia*, *A. agaricites*, showed lower percentage colony bleaching (Peckol et al. 2001).

Heat stress associated with the 1998 ENSO episode was hypothesized to be responsible for this mass mortality event of *Agaricia tenuifolia* in the lagoonal reefs of the Pelican Cayes, Belize (Aronson et al. 2000, 2002a). Fitt & Warner (1985) found that congener *A. lamarcki* was also sensitive to temperature stress. These lagoonal reefs are steep-sided ‘ponds’ with limited water flow (Shyka & Sebens 2000). Due to the enclosed nature and proximity of reefs to freshwater
incursions during storms, these coral populations may have different stress tolerance levels than populations in forereefs or other locations (Aronson et al. 2000). However, the widespread nature of the bleaching event, including forereef and lagoonal regions, the onset of bleaching prior to Hurricane Mitch and the depth to which coral mortality extended in the Pelican Cayes (15 m), suggest that prolonged heat stress was the primary cause of the mass coral mortality at this location.

These events led to the present study in which *Agaricia agaricites* and *A. tenuifolia* populations were surveyed at lagoonal and forereef sites to document the recovery following these major environmental disturbances. To investigate the effect of heat stress on *A. tenuifolia* and contrast the effects with a congener, *A. agaricites*, we measured heat shock protein levels in experimentally heat shocked *A. tenuifolia* and *A. agaricites*. Designed to look at a question of physiological responses, this study used immunoblotting for the heat shock cognate protein HSC 70 and inducible heat shock protein HSP 90. Thus, the present study was designed to address the hypothesis that variable HSP expression in *A. agaricites* and *A. tenuifolia* may have been important in their differential response to the ENSO warming event of 1998.

MATERIALS AND METHODS

Study organisms. *Agaricia* spp. are found throughout the western Atlantic at depths ranging from ~1 to >70 m (Humann & DeLoach 2002). *A. agaricites* is a hermaphroditic brooder, creating internally fertilized planulae larvae (Fadlallah 1983). These brooded planulae settle close to parent colonies and are able to quickly colonize disturbed reefs. Although no studies have been done on reproductive biology of *A. tenuifolia*, it is also believed to be a brooder as it exhibits similar population dynamics to *A. agaricites* and other brooding corals such as *Porites astreoides* (Gleason et al. 2001).

*Agaricia agaricites* and *A. tenuifolia* have somewhat different but overlapping, biogeographic and depth ranges. *A. agaricites* is abundant in Florida, the Bahamas and the Caribbean, while *A. tenuifolia* is common in the western Caribbean and rarer in the eastern part of the basin (Humann & DeLoach 2002). *A. tenuifolia* is found across a range of depths (1 to ~30 m) and flow regimes in Belize (Sebens et al. 2003). *A. agaricites* is reported to occur to greater depths than its congener, up to 73 m (Humann & DeLoach 2002).

Study sites. Our 3 forereef sites (Fig. 1), Tobacco, South Water and Curlew, located near associated cayes, showed well-developed spur and groove forma-

![Fig. 1. Belize barrier reef system showing locations of study areas](image-url)
protocol (Kramer & Lang 2003, Peckol et al. 2003). 10 m transect lines (~15 to 22 transects per site, except June 2002, \( n = 8 \)) were positioned haphazardly on the reef surface parallel to reef growth (8 to 10 m apart) along the reef ridges and forereef spurs (6 to 15 m depths). The following information was collected for all corals under the transect lines: genus and species of coral, water depth, maximum colony diameter, percentage of colony dead and percentage of colony bleached/diseased. We expressed mean partial colony mortality (coral colonies were replicates) for each species at each site. To express percentage live coral cover (overall) and by species (for Agaricia tenuifolia and A. agaricites), we considered each 10 m transect as a replicate. Because an estimate of 10% live cover of small versus large coral colonies can represent quite different amounts of live coral cover, we adjusted percentage live cover estimates for colony diameter (percentage of colony alive \( \times \) colony diameter). We then summed the adjusted percentage live cover estimates and expressed the sum as a percentage of the 10 m transect. Reported percentages thus reflect the overall (total) live coral cover, as well as A. tenuifolia and A. agaricites live cover, along 10 m transects.

Percentage cover data were arcsine-transformed prior to use of analysis of variance (ANOVA). We did not include January 1999 or June 2002 assessments in the statistical analyses because of more limited site monitoring or transect number.

**Heat shock experiments.** Agaricia tenuifolia and A. agaricites were collected at patch reefs in the lagoonal region off Belize in June 2001 to develop techniques used later on experimental corals (see Robbart 2002). Corals used in heat shock experiments were collected at 15 m on the forereef off Carrie Bow Caye in August 2001 (ambient water temperature = 28°C) and transported to the laboratory in coolers at ambient temperature. Fore reef corals were selected because their exposure to elevated summer temperatures was less likely than populations from shallower, patch reef sites, thus reducing the likelihood of pre-existing, background levels of HSPs related to thermal acclimation (Fitt et al. 2001) prior to the beginning of the experiment.

Coral fragments (3 to 5 cm²) were held in aerated 150 L aquaria at ambient (28°C) temperatures or heated to 34°C (+6°C). Direct irradiance was blocked with opaque screens. Aquaria were maintained at experimental temperatures for 2 h and then returned to ambient (28°C) for 2 h. Corals (\( n = 4 \), each species and treatment) were then removed and placed in RNA Later (Ambion) and frozen at -20°C. Coral samples were transported on dry ice.

**HSP determination.** HSPs were analyzed for coral samples from the heat shock experiment (ambient and +6°C treatments). While some degree of natural biological variation was seen in the initial coral samples (Robbart 2002). HSP levels did not differ significantly between initial samples and the ambient treatment for either Agaricia species. Samples were cleaned of macroscopic epibionts before processing. The area of living coral tissue was measured by flattening aluminum foil in a non-overlapping fashion over the surface of the coral sample (Marsh 1970) and tissue surface area was calculated using an area meter scanner (CID).

Coral fragments (1 to 3 cm² pieces) were heated in Laemmli SDS polyacrylamide gel sample buffer (Laemmli 1970) to separate coral tissue from the calcium carbonate. Tissue aliquots in the sample buffer were precipitated with 3 volumes of 100% acetone (HPLC grade), mixed and centrifuged for 1 min at 14,000 \( \times g \). The pellet was resuspended in cold, distilled H₂O equal to the original volume and acetone-precipitated again. After the third resuspension/precipitation the samples were frozen as a pellet at -80°C.

Laemmli sample buffer was added to the sample pellet and heated in a boiling water bath for 2 to 3 min to solubilize protein. Protein concentrations were estimated using the Lowry method (Lowry et al. 1951) using a standard curve of twice recrystallized bovine serum albumin.

Protein extracts were separated by 1-dimensional gel electrophoresis (Laemmli 1970). Samples were loaded onto 12% (HSC 70) or 10% (HSP 90) separating gels (5% stacking gels) using 5, 6 or 6.5 µg of extract for HSC 70 gels, and 20 or 25 µg for HSP 90 gels, along with 0.1 µg of HSC 70 standard (SPP-751, StressGen Biotech) and 0.15 or 0.025 µg of HSP 90 standard (SPP-770, StressGen Biotech.) (Thompson et al. 2003). Electrophoresis was performed on gels until the dye front was 0.5 cm from the bottom.

After electrophoresis, gels were transferred onto PVDF membranes (Millipore Corp.) (Towbin et al. 1979) and probed with the appropriate monoclonal antibodies (StressGen Biotech); either rat anti-hamster HSC 70 (1:3000; SPA-815) or rat anti-human HSP 90 monoclonal antibody (1:2000; SPA-840). After incubation overnight in primary antibody at 4°C, secondary antibody (HSC 70; 1:3000 and HSP 90; 1:2000, goat anti-mouse polyclonal-HRP-coupled, A-0412, Sigma) was applied to blots for 2 h at RT. A monoclonal antibody to plant HSC 70 (SPA-817) was tested to detect zooxanthellar HSPs. Although none were detected (Robbart 2002), the antibody was not specific for dinoflagellate HSPs. Blots were then developed in TMB peroxidase substrate (KPL Laboratories) and dried overnight before being scanned densitometrically for analysis.

Immunoblots were scanned using an HP Scanner at 1200 dpi and Adobe Photoshop software. Protein
bands were quantified with Scion Image (Scion Corp.) and areas were analyzed using JMP (SAS Institute).

RESULTS

Coral abundance, colony mortality and percentage live cover

Abundance patterns revealed species differences with depth and reef type for colonies of *Agaricia agaricites* and *A. tenuifolia* at forereef and reef ridge sites (Figs. 2 & 3). *A. tenuifolia* was generally more abundant at shallower depths, 4 to 9 m, while *A. agaricites* showed no clear depth preference (χ² analysis: forereef, χ² = 97.4, p < 0.001; ridges, χ² = 31.8, p < 0.001). Although *A. agaricites* and *A. tenuifolia* showed similar abundances at forereef sites, the latter species was 4 to 5 times more prevalent at the ridge sites.

Total live coral cover (adjusted for colony diameter along 10 m transect lines) at forereef sites (Fig. 4) was relatively stable between June 1999 and June 2001, and similar among sites (2-way ANOVA, site and time, p > 0.05). During the January 1999 census, coral cover at South Water Caye was somewhat (1-way ANOVA, p > 0.05) reduced (41%) compared with subsequent sampling periods (~52%), perhaps reflecting combined, immediate effects of the thermal anomaly and Hurricane Mitch.

Live coral cover was substantially lower at ridge sites compared with forereef areas (Figs. 4 & 5). Two-way ANOVA revealed a significant increase in coral cover from June 1999 to June 2001 at ridges (F = 4.45, p = 0.04); however, live coral at Tunicate remained lower than the other 2 ridge sites (F = 6.3, p = 0.003).

This site difference was still evident during the June 2002 census (Fig. 5).

Percentage live *Agaricia* cover (adjusted for colony diameter along transect lines) was relatively low at both forereef and ridge sites (Figs. 4 & 5). During the January 1999 survey at South Water forereef and Wee Wee ridge sites, live cover was ~3 and 1% for *A. tenuifolia* and *A. agaricites*, respectively. We found no significant change in percentage live cover between June 1999 and June 2001 for either species (2-way ANOVA, p > 0.05) at forereef sites; however, live cover of *A. tenuifolia* was higher (sites, $F = 8.6, p = 0.0004$) at Curlew than the other forereef sites. At ridge sites, we measured a significant increase in percentage cover for *A. agaricites* (2-way ANOVA, $F = 21.4, p = 1.4 \times 10^{-5}$) and *A. tenuifolia* ($F = 12.2, p = 0.0008$) from June 1999 to 2001 except at Tunicate, where live cover remained relatively low (for *A. agaricites*: site, $F = 3.7, p = 0.03$; *A. tenuifolia*: site, $F = 10.0, p = 0.0001$). By the June 2002 census (not included in the statistical analysis because a lower number [n = 8] of transects were measured), *A. agaricites* and *A. tenuifolia* had increased somewhat at Wee Wee and Peter Douglas ridges, but showed low percentage live cover at Tunicate (Fig. 5).

Expressing relative abundance (percentage live *Agaricia* relative to total live coral cover) of *Agaricia* spp. revealed distinctive species recovery patterns following the 1998 thermal anomaly and Hurricane Mitch. Except at Curlew where relative abundance remained stable at ~5%, *A. agaricites* demonstrated 2- to 3-fold increases in percentage live cover from June 1999 to 2001 (2-way ANOVA: forereefs, $F = 6.3, p = 0.01$; ridges, $F = 21.6, p = 1 \times 10^{-5}$). *A. tenuifolia* showed no significant increases in relative abundance at forereef or ridge sites (2-way ANOVA: time, $p > 0.05$) during this period.

Fig. 3. *Agaricia agaricites* and *A. tenuifolia*. Abundance (number of colonies) with depth (m) at Belize reef ridge sites, Wee Wee, Peter Douglas, and Tunicate. Data combined from June 1999 and 2001 surveys.
Monitoring percentage partial colony mortality provided an additional measure of change for *Agaricia agaricites* and *A. tenuifolia* (Table 1). During June 1999, partial colony mortalities were similar for both species at forereef sites, averaging ~30 and 23% for *A. tenuifolia* and *A. agaricites*, respectively, and declining somewhat by 2001 (2-way ANOVA, p > 0.05, both time and species). At reef ridge sites, *A. tenuifolia* showed high partial colony mortality (75 to 95%) during 1999, significantly higher than colony mortality at forereef sites (2-way ANOVA: F = 24.5, p = 0.001). Although these values declined by June 2001, mean percentage colony mortality remained >50% for this species at 2 ridge sites. Partial colony mortality was much lower for *A. agaricites* at ridge sites.

The size distribution of *Agaricia tenuifolia* colonies shifted toward smaller colonies from 1999 to 2001 at reef ridge sites ($\chi^2 = 30.2$, p = 0.0001, all sites combined) (Fig. 6). For example, all *A. tenuifolia* colonies at Tunicate ridge were in the smallest size class (<20 cm) in 2001. At Wee Wee and Peter Douglas ridges, larger colonies were still in evidence during both 1999 and 2001, suggesting higher colony survival at these sites following the 1998 disturbances.

### Heat shock proteins

*Agaricia agaricites* showed higher levels of HSC 70 than *A. tenuifolia* under ambient (28°C) and heat-treated (+6°C) treatment conditions (2 h heat shock, 2 h ambient) (Figs. 7 & 8). Ambient *A. agaricites* showed significantly ($t = 2.6$, p = 0.04) higher amounts of HSC 70 than *A. tenuifolia* and heat-treated *A. agaricites* showed induced HSC 70 accumulation ($t = 4.29$, p = 0.005).

HSP 90 was detected in *Agaricia agaricites* (August initial samples and heat shock experiment) but not in *A. tenuifolia* (Fig. 7). Due to the low sample size (n = 1 to 3), no pattern of expression related to treatment was evident; however, HSP 90 concentrations in *A. agaricites* were ~5 to 10 µg cm⁻² (Robbant 2002).
Table 1. Mean (±SE) percentage partial colony mortality for *Agaricia agaricites* (Aa) and *A. tenuifolia* (At) from fore reef and coral reef ridge sites off Belize during June 1999 and 2001. The number of coral colonies sampled is indicated. ND: no data.

<table>
<thead>
<tr>
<th>Sites</th>
<th>1999 Aa</th>
<th>1999 At</th>
<th>2001 Aa</th>
<th>2001 At</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. of colonies</td>
<td>No. of colonies</td>
<td>No. of colonies</td>
<td>No. of colonies</td>
</tr>
<tr>
<td>Fore reef</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tobacco</td>
<td>11 ± 5</td>
<td>24</td>
<td>20 ± 7</td>
<td>16</td>
</tr>
<tr>
<td>South Water</td>
<td>41 ± 10</td>
<td>16</td>
<td>24 ± 6</td>
<td>32</td>
</tr>
<tr>
<td>Curlew</td>
<td>18 ± 5</td>
<td>21</td>
<td>47 ± 6</td>
<td>36</td>
</tr>
<tr>
<td>Ridges</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wee Wee</td>
<td>18 ± 12</td>
<td>9</td>
<td>75 ± 5</td>
<td>69</td>
</tr>
<tr>
<td>Peter Douglas</td>
<td>ND</td>
<td>0</td>
<td>90 ± 3</td>
<td>70</td>
</tr>
<tr>
<td>Tunicate</td>
<td>ND</td>
<td>0</td>
<td>95 ± 2</td>
<td>89</td>
</tr>
</tbody>
</table>

HSP 70, HSP 25, HSP 27 and plant HSC 70 (SPA-812, SPA-801, SPA-800 and SPA-817, respectively, StressGen Biotech) were not detected in samples of either *Agaricia agaricites* or *A. tenuifolia* from the August heat shock experiment (Robbart 2002). The absence of plant HSC 70 suggests that zooxanthellae HSC 70 was not present, occurred at undetectable levels in samples or was not possible to detect using the plant HSC antibody.

**DISCUSSION**

Immuneblotting results reported here indicate that variable HSP expression in *Agaricia agaricites* and *A. tenuifolia* may have been important in their differential response and recovery to the ENSO warming event and the subsequent stressor of Hurricane Mitch on the Belize barrier reef system. Notably, *A. agaricites* had twice as much HSC 70 as *A. tenuifolia* at ambient temperatures (28°C) and 6 times as much HSC 70 after heat shock at +6°C. Further, no accumulation of HSC 70 was detected in *A. tenuifolia* during 2 h of heat shock. HSP 90 was only detected in *A. agaricites*. Aronson et al. (2000) documented nearly 100% cover of *A. tenuifolia* prior to the thermal anomaly, and subsequently (Aronson et al. 2002a) reported both a higher maximum SST and longer periods of time above the HotSpot threshold for an area near the ridge sites (Channel Caye) compared with a fore reef site (Carrie Bow Caye) during the thermal anomaly. Populations of *A. agaricites* and *A. tenuifolia* as well as total coral cover, surveyed in the present study at well-flushed fore reef sites, showed no significant change in percent live cover between June 1999 and 2001. In addition, a January 1999 survey at South Water Caye soon after the 1998 autumn disturbances suggested only a minor decline in live cover. In contrast, *A. tenuifolia*, formerly the space dominant coral at ridge sites (Aronson et al. 2002b), suffered major mortality associated with these disturbances (Aron-
son et al. 2000, 2002a [see their Fig. 3], Peckol et al. 2003). *A. agaricites* showed ~18% partial colony mortality (surveyed at 1 ridge site only), while *A. tenuifolia* exhibited ~75 to 95% colony mortality (June 1999 survey). Both *Agaricia* species showed some population recovery at Peter Douglas and Wee Wee ridge sites but percentage live cover at Tunicate remained extremely low during the June 2002 survey. Population and immunoblotting data together suggest that *A. tenuifolia* was poorly equipped to respond to the intermittent environmental stress characteristic of ENSO events.

Of the 3 ridge sites, Tunicate ridge off Cat Caye had the highest mortality of *Agaricia tenuifolia* (Aronson et al. 2000, 2002a, Peckol et al. 2003) and negligible recovery. This site is part of a reef system located in the Pelican Cayes: a series of rhomboid shoals, forming enclosed and protected ‘ponds’ (Macintyre et al. 2000). Shyka & Sebens (2000) found that reef ridge sites in the Pelican Cayes were low-flow environments (1 to 3 cm s⁻¹) compared with nearby reefs. Flow rates have major effects on several aspects of coral physiology (Sebens 1997), and reduced rates of water exchange would intensify the impact of a thermal stress and increased rainfall from Hurricane Mitch. The continued decline in live *A. tenuifolia* cover at Tunicate ridge following the 1998 ENSO episode was also related to decreased surface area available for coral colonization due to high cover (>75%) of the encrusting sponge, *Chondrella ct. nucula Schmidtk* (Aronson et al. 2002a, Peckol et al. 2003). This shift from coral cover to non-coral cover represents another major transition at this site, similar in scope to the change from the space-dominant *Acropora cervicornis* to *A. tenuifolia* following decimation of *A. cervicornis* to White-band Disease (Aronson & Precht 1997). Sucha shift will likely have significant implications for future recovery (Curran et al. 2004).

The size-distribution data revealed distinctive recovery patterns at the 3 reef ridge sites. At Tunicate ridge, all live *Agaricia tenuifolia* colonies surveyed during 2002 were <10 cm in diameter (with 67% of the colonies <5 cm) and occurred below 7 m, suggesting that recovery was limited to deeper (cooler) waters and was largely from coral recruits (see also Aronson et al. 2002a). Wee Wee and Peter Douglas ridges had greater numbers of residual colonies that survived the thermal stress as well as new recruits, contributing to the more rapid recovery at these sites. The agaricids are early colonizers of open substrata in reef environments, due to asexual fragmentation and a brooding mode of reproduction that promotes high settlement rates (Hughes & Jackson 1985, Richmond & Hunter 1990). Lower encrusting sponge cover on dead *A. tenuifolia* substrata at Wee Wee and Peter Douglas ridges compared with Tunicate (Curran et al. 2004) likely further enhanced coral recovery at these sites.

*Agaricia agaricites* and *A. tenuifolia* exposed briefly (2 h) to heat shock (+6°C above ambient) showed differential expression of HSC 70. The experiment was terminated at 4 h, so it is possible that *A. tenuifolia* has a delayed response not evidenced in this study. This is the first HSP research conducted on *Agaricia* spp.; however, other corals, such as *Goniopora djboutiensis*, *Montastraea annularis* and *Acropora grandis* showed elevated expression of HSP 70/HSC 70 within 1 to 4 h of heat shock (Hayes & King 1995, Fang et al. 1997, Sharp et al. 1997). HSC 70 is known as a constitutive heat shock protein, expressed during normal cellular functioning of cells. Additionally, HSC 70 has been shown to move from its origin in the cytoplasm to the nucleolus (Lindquist & Craig 1988) and in this location, may play a protective role in stabilizing the proteins necessary to assemble ribosomal subunits. Therefore, HSC 70 may act as a 'first defender' as it is already being produced constitutively.

In the present study, the antibody used was specific for HSC 70 (SPA-615, StressGen Biotech) and no HSP
Acknowledgements. We thank C. Fraço, A. Jugovic, R. Lukšic, M. Nestelerođe and S. Ristau for field and data-processing assistance; M. Carpenter for boat operation; and H. Thompson for help in developing HSP protocols and test statistic analysis. We gratefully acknowledge facilities support at Carrie Bow Caye Marine Field Station (Smithsonian Institution, K. Ritzler, Director) and the Wee Wee Caye Marine Laboratory (P. and M. Sleave, Directors). This research was supported by Blakeslee grants (M.L.R., P.P., S.F.S.), a Culppea Foundation grant to Smith College (P.P. & H.A.C.), the Smith College Summer Science Program (J.B.-S.), and the Elizabeth Horner Fund (M.L.R.). Wells PG (1999) identified #654 to the Caribbean Coral Reef Ecosystem Program at the National Museum of Natural History, Smithsonian Institution.

LITERATURE CITED


Bosch TCG, Krylov SM, Bode HR, Steele RE (1988) Thermal tolerance and synthesis of heat shock proteins: these responses are present in Hydro attenuta but absent in Hydro oligactis. Proc Natl Acad Sci (USA) 85:7927–7931


Enkin CM (1996) Where have all the carbonates gone? A model comparison of carbonate budgets before
Thompson HS, Scordilis S (2001) Downhill running elicits distinct but similar protein and mRNA expression patterns of HSP 25, HSP 70 and HSP 90. Fish Physiol Biochem 23:15–41

Submitted: July 8, 2003; Accepted: August 10, 2004
Proofs received from author(s): November 5, 2004

Editorial responsibility: Otto Kinne (Editor), Oldendorf/Luhe, Germany