Plasticity in the Sclerites of a Gorgonian Coral: Tests of Water Motion, Light Level, and Damage Cues

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Abstract. The gorgonian coral *Briareum asbestinum* contains skeletal elements (sclerites) that vary in length and density within and among local populations. Data from previous work suggested that the sclerite compositions of colonies may be altered in response to environmental cues such as predator damage, water motion, and light level. To test these hypotheses, colonies from shallow reefs were transplanted to racks at a single location where the three environmental factors of interest were artificially manipulated. After 9–14 weeks of growth, sclerite morphologies and densities had not changed in response to shading or to water-motion reductions that mimicked deep-water conditions. However, colonies did respond significantly to two types of simulated predator damage. Following tip amputation, sclerites in the regenerated tips of damaged colonies were shorter and more dense than in the controls. In contrast, mid-branch scarring caused colonies to produce longer sclerites at lower densities. Since long sclerites deter feeding by predatory snails, the increase in sclerite length in response to scarring of mid-branch regions may function as an inducible defense.

Introduction

Phenotypic plasticity—or differential phenotypic expression of a genotype under varying environmental conditions—has been a subject of increasing interest to evolutionary ecologists since Bradshaw’s (1965) extensive review (e.g., Bradshaw, 1974; Via and Lande, 1985; Schlichting, 1986; Stearns, 1989; Scheiner, 1993). Although plasticity can be nonadaptive or maladaptive, researchers have been primarily interested in adaptive plasticity as a mechanism by which organisms cope with changing environments. Adaptive plasticity is especially well documented in plants, for which the literature contains numerous examples of induced responses to physical factors such as shade (Turesson, 1920), desiccation (Harlan, 1945), soil fertility (Sorensen, 1954), temperature (Mooney and West, 1964), and water stress (Roy and Mooney, 1982). Bradshaw (1965) reasoned that plasticity should often be favored in plants because they are sessile organisms whose autotrophic lifestyle requires that they inhabit relatively open spaces where they may be exposed to environmental extremes. When the scale of temporal variability is shorter than the lifespan of the organism, or where spatial variability is at a scale smaller than the dispersal range of the organism, adaptive plasticity is a viable strategy.

In marine habitats, sessile colonial invertebrates, such as scleractinian and gorgonian corals, share many characteristics with plants. Colonies are permanently sessile, often with upright branched growth forms (Barnes, 1987); and many corals depend on symbiotic algae (zooxanthellae) for a large proportion of their energy (Muscatine, 1974; Muscatine et al., 1975; Svoboda, 1978; Sebens, 1987) and are thus largely autotrophic. In addition, populations with wide distributions are routinely exposed to variation in abiotic factors such as light level (McCloskey and Muscatine, 1984; Miles, 1991), wave exposure (de Weerdt, 1981; Sebens, 1984), and sedimentation rate (Foster, 1979). Indeed, variation in colony morphology or physiology in response to temperature, depth, or water movement has been documented in sponges (Bavestrello et al., 1993), scleractinian corals (Wijsman-Best, 1974; Foster, 1979; Lesser et al., 1994), and gorgonian corals (Grigg, 1972; West et al., 1993).

In both plants and corals, inducible defenses represent a special category of adaptive plasticity that involves biotic rather than abiotic cues. Defined as environmentally triggered phenotypic responses that defend against spe-
cific biotic selective agents (Adler and Harvell, 1990), inducible defenses have emerged as prominent phenomena only in the past 20 years. A variety of plant responses to real or simulated herbivore damage include increased morphological defenses such as spines or resistant growth forms (Young, 1987; Lewis et al., 1987) and production of physicochemical deterrents such as polyphenolic compounds, alkaloids, or silica (McNaughton and Tarrants, 1985; Schultz, 1988; Van Alstyne, 1988; Baldwin and Ohnmeiss, 1993). Among colonial invertebrates, various species respond to the presence of competitors by producing agonistic weaponry such as digestive filaments, stolons, or sweeper tentacles (Ivker, 1972; Francis, 1973; Lang, 1973; Wellington, 1980; Sebens and Miles, 1988; Harvell and Padilla, 1990; Miles, 1991). However, although predator-induced defenses, such as spines and helmets, have been documented in various clonal freshwater invertebrates (e.g., Gilbert and Stemberger, 1984; Havel, 1986), examples of induced defenses against predators in colonial marine invertebrates (Harvell, 1984) are surprisingly rare.

In the Caribbean coral *Briareum asbestinum* (Gorgonacea), both biotic and abiotic environmental agents may contribute to plastic variation in skeletal features. Unlike the hard corals with their massive calcium carbonate skeletons, gorgonian corals consist of a central axis surrounded by an outer cortex of a soft, polyp-bearing matrix, the structural integrity of which is maintained by small (0.2–1.2 mm), calcitic skeletal elements (sclerites) (Fig. 1). Within and among local populations, *B. asbestinum* colonies vary significantly in the mean lengths and densities of their sclerites. Furthermore, a reciprocal transplant of colonies between shallow and deep sites indicated that sclerite morphology is phenotypically plastic in response to one or more environmental cues (West et al., 1993; West, 1996).

One such cue may be damage from predators. In addition to skeletal support, sclerites fulfill a second role as structural defenses against predators. *B. asbestinum* contains ichthyodeterrent, diterpene secondary compounds (Pawlik et al., 1987), and predation by fishes appears to be minimal (C. D. Harvell, unpubl. data). However, the gastropod *Cyphoma gibbosum* is a major predator of gorgonian corals (Kinzie, 1970; Lasker and Coffroth, 1988; Lasker et al., 1988) and is an important source of damage to *B. asbestinum* (Hazellet and Bach, 1982). *Cyphoma gibbosum* contains biotransformation enzymes capable of detoxifying *B. asbestinum* allelochemicals (Vrolijk and Targett, 1992) and consumes artificial foods containing *B. asbestinum* extracts and pure compounds just as readily as it consumes control foods (C. D. Harvell, unpubl. data). Therefore, the sclerites of *B. asbestinum* may be its only defense against this predator, such that differences in skeletal composition will correlate with the ability of colonies to deter snail feeding (West et al., 1993; West, 1996). As sclerites increase in size, they render artificial foods less palatable to *C. gibbosum* (Van Alstyne and Paul, 1992; West et al., 1993; West, 1996).

Hence, the inducibility of long, defensive sclerites as a reaction to damage would be a potentially advantageous plastic response.

In addition to biotic induction, sclerite plasticity among *B. asbestinum* populations may also be cued by changing abiotic conditions. At two islands in the Caribbean, colonies of *B. asbestinum* are distributed along a depth gradient of 1–30 m. At the shallow end of the gradient, sclerites are short and of high density, whereas deep colonies contain sclerites that are long and of low density. Sclerite length and density within colonies are negatively correlated, such that colonies with both long and densely packed sclerites do not occur (West et al., 1993; West, 1996). Although biotic induction by a patchily distributed predator could account for variability within sites, the pattern of increasing sclerite length with depth cannot be explained solely by the presence of predators. Snails are indeed present at all of my study sites, but their densities and damage actually decrease with depth, whereas sclerite length increases with depth (West, 1996; C. D. Harvell, unpubl. data). Hence, largescale patterns of sclerite variability with depth may be generated by abiotic environmental factors.

Along the depth cline, declining light penetration and
water motion both show some degree of correlation with sclerite variation (West, 1996). Both factors might affect skeletal composition in different ways and might function as cues that induce a plastic response in sclerite length and density. Observed correlations between sclerite variation and percent penetration of photosynthetically active radiation (West, 1996) may be related to effects of light on colony growth rates (C. D. Harvell, unpubl. data), because gorgonians depend on their symbiotic zooxanthellae for most of their energy (Sebens, 1987). In addition, strong correlations between sclerite variation and water motion (West, 1996) may relate to the function of sclerites as skeletal support structures. The sclerites of B. asbestinum act as rigid reinforcing points of attachment within the soft matrix, providing resistance to deformation; both smaller sclerites and greater densities of sclerites confer greater stiffness (Wainwright et al., 1976; Koehl, 1982; Palumbi, 1986). Therefore, colonies may display depth-related shifts in skeletal composition according to water motion and light level cues.

In this study, I tested the ability of one biotic agent (predator damage) and two abiotic factors (water motion and light) to induce skeletal modifications in the soft coral B. asbestinum. Because the large collections needed for this work would have denuded the sparsely populated deep reefs at my sites, I focused on shallow-water B. asbestinum. To examine separately the effect of each type of cue, two large transplant experiments were conducted in which colonies from a shallow population were grown on racks at a single site where water motion, light, and damage were manipulated under controlled conditions. I hypothesized that shallow-water colonies subjected to reduced water motion and reduced light (simulating deep-water conditions) and colonies subjected to mechanical scarring (simulating predator damage) should respond in each case by producing longer sclerites at lower densities compared to controls.

Materials and Methods

Experiment I: water motion and damage

To assess whether differences in water motion and simulated predator damage would induce plastic changes in sclerite composition, I conducted a transplant experiment at San Salvador, Bahamas, from 1 June to 3 August, 1991. Branches from a shallow (1–3 m) population of B. asbestinum were transplanted to racks on which they reattached themselves and grew as independent new colonies. The racks were arrayed at a single location where water motion and damage were varied. The 12 racks were constructed according to a design modified from West et al. (1993). Each rack consisted of an acrylic plate, dimensions 30.0 × 23.0 × 2.5 cm, that snapped into an aluminum angle frame. Each plate accommodated 12 (2 rows of 6) colonies, each inserted into a recessed well and secured with cushioned cable ties to an acrylic post.

The racks were rigidly affixed to cement blocks at a shallow (3 m) site that exposed the colonies to high-energy waves and surge. Water motion was reduced inside half of the racks by clear, small-mesh (0.20 mm) nylon screening that was attached to form walls 15 cm tall (Fig. 2). Water motion inside control and walled racks was quantified by recording the percent dissolution of plaster of Paris (a water-motion index) using methods adapted from Muus (1968), Doty (1971), Day (1977), Bushek (1988), and Jokiel and Morrissey (1993). For this integrated relative measure of water movement due to current velocities and turbulence (Doty, 1971), I measured the weight loss of five replicate plaster domes (24 g) for each rack type over a 24-h period (see West, 1996, for a more detailed description). When compared with measurements made at various other B. asbestinum habitats, the reduction in water motion between walled and control racks was found to be similar to differences in water motion between colonies growing deep within crevices and colonies out on the open reef flat. Naturally occurring B. asbestinum colonies from these habitat types differ significantly in sclerite composition (West, 1996).

All racks were fitted with clear, large-mesh (2.5 cm) roofs that afforded protection from disturbance by fishes without affecting light penetration. Furthermore, placement of the racks in a sand patch several meters from the reef prevented discovery of the colonies by benthic predators (confirmed through weekly monitoring). Light measurements with a quantum/radiometer/photometer (LI-185B with LI-192SB underwater sensor; LI-COR, Inc., Lincoln, Nebraska) showed that penetration of photosynthetically active radiation (PAR) was the same inside and outside of walled and control racks. During the experiment, the walls and roofs were scrubbed regularly with a stiff brush to remove fouling organisms.

The effects of damage due to simulated predation or breakage were tested by assigning each colony to one of three damage treatments: (1) control; (2) scar; and (3) tip amputation (Fig. 2). The controls were left undamaged. The scar treatment simulated damage by Cyphoma gibbosum (Gastropoda), a predator that is typically found on the middle region of colony branches (Harvell and Suchanek, 1987; Gerhart, 1990), where it can rasp the cortex as deep as the axis (Harvell and Suchanek, 1987; West, 1996). Hence, starting about 3 cm from the tip and working downward, I damaged the mid-regions of colonies by gouging them to the axis to create a 1- × 3-cm scar. Finally, tip amputation involved severing the top 1 cm of the colony to simulate predation or breakage. B. asbestinum is a preferred prey species of Her-
modice carunculata (Polychaeta), which commonly feeds by a characteristic removal of branch tips (Vree-land and Lasker, 1989). Thus, tip removal may be similar to the damage inflicted on colonies by worms. Yet, tip removal also lays bare a cross-sectional area of the colony, so amputation also mimics the end product of breakage resulting from heavy wave action. The original cortex material from each scar or tip amputation was preserved in 70% ethanol for later analysis.

Because the number of samples was large, the experiment was set up over 2 days. Each day, 72 undamaged branches, 10 cm long, were collected from a shallow (1–3 m) reef of high wave energy. To maximize the genetic diversity of the collection, divers swam linearly along the reef and sampled single branches from colonies that were separated from any others by at least 3 m. After collection, the branches were suspended in flow-through mesh bags from the side of the boat at the nearby transplant site. They were then assembled onto racks (equal numbers of damage types, randomized with respect to rack position) inside large containers of seawater in the boat, and each completed rack was immediately conveyed to the transplant site below. Virtually all branches began extending their polyps within 2 h of transplantation. Within a few weeks, the branches had attached themselves to the racks and were growing as independent colonies. The colonies were allowed to heal and grow for 9 weeks, and survival exceeded 95%. During the experiment, more than 50% of the colonies increased in height by 0.1–1.5 cm; and all survivors showed positive growth in the sense that they produced new cortex material that encrusted their supportive posts and ties, filled in their scars, and covered over their tip wounds. At the end of 9 weeks, the colonies were collected and preserved in 70% ethanol until they could be processed in the laboratory.

Experiment II: light and damage

To test the effects of reduced light and colony damage, another transplant experiment was conducted from 7 February to 20 April, 1991. The methods were the same as those described for experiment I with the following modifications. Branches were collected from throughout a shallow, relatively calm bay and transplanted to a 12-m location nearby. The colonies were subjected to the same three types of damage and were distributed similarly within the racks, but instead of walls, half of the racks were fitted with roofs made of a double layer of dark window screening (Fig. 2). This screening (1-mm
Sclerite measures

At the end of each experiment, subsets of 1–3 colonies per damage type per rack were measured. Sclerite lengths were recorded in the laboratory with the MORPHOSYS image analysis program (Meacham and Duncan, 1990; version 1.26). For colonies subjected to scarring or tip amputation, both the original cortex material collected when the colonies were damaged (“before” sample) and newly regenerated material from the healed wound (“after” sample) were analyzed. In experiment I, a mid-branch cortex sample from the side opposite the scar (“opposite” sample) was also analyzed to see whether the response to scarring was regionwide. Hence, cortex material was sampled from the following regions of the colonies: (1) tip region of damaged colonies, before and after amputation; (2) mid-branch region of damaged colonies, before, after, and opposite the scars; and (3) mid-branch and tip regions of control colonies at the end of the experiment (after).

For sclerite length measurements, cortex subsamples were taken from the edges of scars before and after healing, from original and healed tips at a distance of 1 cm down from the apex, and from the same locations (by distance from the apex) in the controls. In each case, two small (3–4 mm³) cortex samples were excised, and the organic matter was dissolved away with a solution of 2.6% sodium hypochlorite. The isolated sclerites were rinsed and distributed in their entirety across six slides (three slides per tissue sample). Video images of the first 4 intact sclerites encountered per slide were measured, for a mean of 24 sclerites per region per colony.

From the same regions of the colonies, the proportion of cortex weight consisting of sclerites (the sclerite weight fraction, a measure of density) was estimated according to Harvell and Suchanek’s (1987) protocol. From mid-branch regions, a 1- × 3-cm cortex scraping at the edge of the original scar (or from the middle region of the control) was taken, and from tip regions, the distal 1 cm of cortex material was used in all cases. The samples were separately dried for 24 h at 60°C and weighed; a drying test on a subset of samples showed that they did not continue to lose weight after 24 h. Each sample was then ashed in a muffle furnace at 450°C for 1 h. This process burned the organic matter away, but left the sclerites intact (Harvell and Suchanek, 1987). The sclerite weight fraction was calculated as the proportion of the total cortex weight consisting of sclerites (ash weight/dry weight).

Statistical analyses

All length and weight fraction distributions were normal and homoscedastic, so groups were compared using parametric tests. The “after” data from experiments I and II were subjected to split-unit (or split-plot) analysis of variance (Neter et al., 1990). Blocks, which tested for microenvironmental effects within the experimental site, consisted of pairs of adjacent racks (units), one unit being a control rack while the other was a reduced exposure (experiment I) or reduced light (experiment II) rack. Within racks, individual colonies (subunits) received the different damage treatments. Because previous work had indicated that different regions within colonies may differ in sclerite composition (West, 1996), mid-branch and tip regions were examined separately. Hence, for each experiment, four split-unit analyses were performed: (1) mid-branch sclerite length; (2) mid-branch sclerite weight fraction; (3) tip sclerite length; and (4) tip sclerite weight fraction. Because the Unit*Damage term was not significant in any of the analyses (P > 0.29 in all cases), it was removed from the model and its sum of squares was pooled with the error sum of squares for final calculation of F-statistics (Neter et al., 1990).

Mid-branch and tip-damage responses detected in the split-unit ANOVAs were confirmed through comparison of “before” and “after” material from within damaged colonies. Both before and after measurements were taken for each mid-damaged or tip-damaged colony, and my a priori expectation was that after and before samples would differ in the same direction as would the after and control samples; hence, I analyzed the before and after data using paired one-tailed t-tests. Data from opposite the scars (experiment I only) were also compared to before data using a paired one-tailed t-test.

In both experiments, the different contrasts of interest involved multiple comparisons and contained variables that lacked independence from variables in other tests (e.g., sclerite length and sclerite weight fraction are from the same colony; control mid-branch and control tip are from the same colony). Hence, a sequential Bonferroni correction for multiple tests was performed (Holm,
The sequential Bonferroni method is less conservative and more powerful than the standard Bonferroni method yet still restricts the probability of a type-I error for the entire test as well as each step of the test to $\alpha = 0.05$ (Holm, 1979; Rice, 1989).

Results

Damage

In both experiments I and II, mean sclerite length increased 23% in regenerated mid-branch scars (see Fig. 1), whereas sclerite weight fraction decreased 2.3% (Fig. 3). The increase in sclerite length was significant for the after and control comparisons of the split-unit ANOVAs (Tables I, II) as well as for the before and after paired comparisons (Fig. 3A). Mean sclerite weight fraction decreased in cortex material after scar damage, but the effect was statistically significant only for the before and after comparison of experiment II (Tables I, II; Fig. 3B). Thus, simulated predator damage did alter the average lengths and (in one case) the average weight fractions of *B. asbestinum* sclerites. Moreover, the mid-branch increase in sclerite length was not restricted to newly formed cortex material within the healed scars. Sclerites

Table I

<table>
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<tr>
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<th>P value</th>
<th>Length</th>
<th>Wt. fraction</th>
</tr>
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<td></td>
<td>Damage</td>
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Note: Tip region: comparison of protected to exposed colony tips and comparison of damaged colony tips to undamaged control tips; mid-branch region: comparison of protected to exposed mid-regions and comparison of damaged colony mid-regions to undamaged control mid-regions. Blocks (pairs of adjacent racks) test for microenvironmental effects. *denotes significance under sequential Bonferroni adjustment.

Table II

<table>
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<tr>
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<td>Damage</td>
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Note: Tip region: comparison of shaded to unshaded colony tips and comparison of damaged colony tips to undamaged control tips; mid-branch region: comparison of shaded to unshaded mid-regions and comparison of damaged colony mid-regions to undamaged control mid-regions. Blocks (pairs of adjacent racks) test for microenvironmental effects. *denotes significance under sequential Bonferroni adjustment.
from material opposite the scar were also significantly longer than the sclerites before damage (Fig. 3A), indicating that this length response was regionwide.

The response to tip removal was opposite to that of mid-branch scarring. Colonies subjected to tip amputation produced new tips that contained sclerites that were up to 16% shorter and made up 2.5% more of the total cortex weight (Fig. 4). The significant difference between tips after damage and control tips is reflected in the damage term of the split-unit ANOVA for experiment II (Table II). The difference between damaged tips and controls was not statistically significant in experiment I (Table I; Fig. 4A). However, within colonies, both experiments showed a significant decrease in sclerite length from before to after damage (Fig. 4A). Increases in mean weight fraction in healed tips were statistically significant only for the before and after comparison of experiment I (Tables I, II; Fig. 4B).

**Water motion**

Dissolution of plaster of Paris domes over 24 h showed that water motion inside the walled racks was significantly reduced compared to water motion within control racks ($P = 0.01$, unpaired $t$-test; Fig. 5). The magnitude of the reduction was similar to water-motion differences measured for colonies growing within crevices versus on the open reef flat at a variety of sites around San Salvador (Fig. 5). Such microhabitat differences have been correlated with significant sclerite variation, with microprotected colonies containing longer sclerites at lower densities than microexposed colonies (West, 1996).

When colonies from a shallow site of high-energy waves and surge were shielded from water motion within the walled racks, their sclerites did not differ from the sclerites of control colonies after 9 weeks. This result was consistent for both mid-branch and tip regions of colonies (Table I). In summary, the skeletal composition of *Briareum asbestinum* colonies was not modified in response to the degree and duration of water-motion reductions tested here.

**Light**

In experiment II, shallow-water colonies were subjected to greatly reduced light levels by shading them with dark screens that had been previously determined to reduce the light levels typical of 12 m to those typical of 30 m (J. Miles, pers. comm.). After 14 weeks, shaded colonies did not differ significantly from unshaded controls in either sclerite length or sclerite weight fraction. This result was consistent for both mid-branch and tip regions of colonies (Table I). Thus, as with water motion, there was no indication that the light reductions tested in this experiment triggered a plastic response in sclerite composition.

**Discussion**

**Environmental cues**

*Briareum asbestinum* colonies responded to one of the three types of environmental cues tested in this study. Simulated predator damage caused clear and significant changes in sclerite morphology, and these changes actually reversed patterns of sclerite length that were observed in colonies without mid-branch or tip damage. Samples collected from naturally growing colonies in the field display a consistent pattern in which the sclerites are longer at the tips of colonies than at the basal regions; similarly, the experimental controls contained longer sclerites at the tips than at the mid-branch regions (Fig. 285).
In contrast to the damage responses, there was no indication that the sclerites of shallow-water colonies were altered in response to the abiotic cues of reduced water motion or reduced light. Depth-related sclerite variation may therefore include a component of genetic differentiation. With its ability to reproduce asexually through fragmentation (Brazeau, 1989) and its brooded, negatively buoyant larvae (Brazeau and Lasker, 1990), *B. asbestinum* is probably relatively philopatric. Indeed, an electrophoretic study performed concurrently with this work indicates significant genetic differentiation between shallow and deep populations (Brazeau and Harvey, 1994). Yet there is also some indication that sclerites are altered in response to wounds that mimic breakage (see discussion below), and reciprocal transplantation revealed a plastic change in sclerites with very extreme differences in depth (West et al., 1993). Here, I tested the more usual variation in light and water motion that colonies in highly populated shallower environments encounter. As such, the exposure and light treatments applied to *B. asbestinum* over 9–14 weeks did not induce significant changes in sclerite length or sclerite weight fraction. Further testing is warranted to determine whether a longer duration of acclimation could eventually result in plastic shifts in sclerite composition.

**Ecological and evolutionary implications**

When colony tips regenerate, the new cortex contains sclerites of decreased length and increased weight frac-

### Figure 6. *Briareum asbestinum*. Comparison of mean sclerite length (mm) in different regions of naturally growing colonies versus experimental colonies. Source groups: Bonefish Bay = samples from tip and base regions of colonies \( (n = 16) \) growing naturally near the transplant sites, Control Colonies = samples from tip \( (n = 41) \) and mid-branch \( (n = 42) \) regions of pooled experimental controls, Damaged Colonies = samples from regenerated tips \( (n = 39) \) and healed mid-branch scars \( (n = 43) \) of pooled damage treatments. Error bars are 1 standard error.
tion. Since tip amputation is a characteristic feeding mode of the worm *Hermodice carunculata*, then perhaps increased sclerite densities are an induced defense that makes feeding difficult for worms, or renders the food less palatable (Vreeland and Lasker, 1989). This hypothesis seems unlikely for the following reasons. First, *H. carunculata* does not feed exclusively on colony tips, but also feeds on middle regions of *B. asbestinum* branches (Vreeland and Lasker, 1989) where the response to damage is a *decrease* in sclerite density. Second, Vreeland and Lasker (1989) found no relationship in this predator between preferences for particular gorgonian species and the ash content (sclerite density) of those species. Finally, *H. carunculata* routinely feeds on milleporid fire corals (Witman, 1988), and one hungry worm in the laboratory attempted to engulf the tip of a plastic pen cap (pers. obs.). Hence, a 2.5% increase in sclerite weight fraction at colony tips would probably be a poor defense against this predator.

Instead, the plasticity of sclerites in colony tips after amputation may be a response that decreases further breakage. Survival is lower for small fragments than for larger fragments in other gorgonians (Lasker, 1990), thus breakage of *B. asbestinum* colonies may result in partial mortality as water motion abrades loose branch fragments against the reef. In addition, Gerhart (1990) found that fouling by epibionts such as algae is a serious problem for gorgonian colonies that have suffered major damage such as the baring of large wounded areas at scars or break points. For these reasons, it may be advantageous for colonies to cover such wounds with new cortex material that is fortified against further breakage through increased packing of smaller sclerites.

Biomechanical assays using both artificial and real tissues have indicated that small sclerites at high densities confer greater stiffness than longer and more sparsely packed sclerites (Koehl, 1982). This may be because small sclerites at high densities provide more surface area for tissue attachment and also leave smaller spaces consisting solely of deformable soft matrix. In more recent work, Koehl (1996) has also shown that the strength and toughness of sclerite-reinforced materials increases—and then decreases—with increasing sclerite density. Biomechanical testing of actual *B. asbestinum* branches is needed to determine whether the tip response renders the surrounding matrix stronger and tougher, or more brittle and breakable. A finding of increased strength and toughness in regenerated tips would support the hypothesis that the observed sclerite modifications function in colony reinforcement. This hypothesis could be further tested by breaking additional colonies at the mid-branch region instead of the tip region and observing their response. To adopt a fortification response of shorter, denser sclerites at mid-branch, colonies would have to be capable of reversing the scar-induced sclerite shifts detected in this study (see below).

At the mid-branch region, *B. asbestinum* colonies respond dramatically to mechanical scarring, producing sclerites that are 23% longer. The middle region of colonies is the typical feeding location of the snail *Cyphoma gibbosum* (Harvell and Suchanek, 1987; Gerhart, 1990), and longer sclerites have been shown to reduce snail feeding significantly (West et al., 1993; West, 1996). At St. Croix (U.S. Virgin Islands), *B. asbestinum* colonies from two habitats contain sclerites that differ in length by about 22% (see Fig. 1), and individuals of *C. gibbosum* that are given a choice feed at a higher rate and spend more time on the colonies with shorter sclerites than on those with longer sclerites (West, 1996). It is unlikely that this effect is due solely to chemistry because *C. gibbosum* has gorgonian-detoxifying enzymes (Vrolijk and Targett, 1992) and appears indifferent to *B. asbestinum* extracts in sclerite-free artificial foods (C. D. Harvell, unpubl. data). Conversely, artificial food assays that employed isolated sclerites in the absence of chemistry have indicated that this predator is significantly deterred by long sclerites (West et al., 1993; West, 1996). Hence, the production of 23% longer sclerites in the mid-branch regions of scarred San Salvador colonies may represent an induced defense.

Harvell (1984) hypothesized that predator-induced defenses should be favored in clonal taxa that suffer intermittent, unpredictable nonfatal encounters with predators. *Cyphoma gibbosum* frequently causes significant damage to colonies of *B. asbestinum*, but it seldom completely kills them (Kinzie, 1970; Birkeland and Gregory, 1975). At shallow sites of greatest snail activity, the average amount of surface area scarred per *B. asbestinum* colony can approach 30% (West, 1996). Although the tenure time of individual *C. gibbosum* on particular gorgonian colonies averages only about 10 days (Harvell and Suchanek, 1987), snails tend to form aggregations that have been observed to stay together in a localized area (and even on a single colony) for up to 4 months (Kinzie, 1970; Birkeland and Gregory, 1975; Hazlett and Bach, 1982; Gerhart, 1986). Therefore, colonies within such a feeding area may be grazed upon multiple times, by multiple snails, over a period of weeks to months. Meanwhile, *B. asbestinum* in areas adjacent to large snail aggregations may be virtually free of predator damage (pers. obs.). Experiments on feeding behavior have shown that when given a choice, *C. gibbosum* tends to move off of colonies containing long sclerites and onto colonies containing short sclerites (West, 1996). Therefore, production of snail-deterrent sclerites by *B. asbestinum*, occurring over a period of months, would be an appropriate and advantageous defensive response.

A potential cost to this induced defense may relate
once again to the effects of changes in skeletal composition on the biomechanical behavior of colonies. Tissues that contain long sclerites at low densities will be less stiff and more elastic than tissues with short sclerites at high densities. In the region of a shift to long, sparse sclerites, a branch will indeed be in little danger of breaking due to brittleness; yet too great a propensity for pliability might itself be disadvantageous. A branch that is too soft may not be able to support the optimal orientation for light or prey capture, and excessive bending and swaying of the colony in waves and surge could lead to damage as tissue is abraded against adjacent coral heads. If so, this may explain why long sclerites are an induced rather than a constitutive defense.

The evolution of adaptive plasticity is favored in many plants and corals because of their distributions across changing habitats and their inability to escape extreme environmental conditions through movement (Bradshaw, 1965). In plants, inducible chemical and morphological defenses against predators have been widely reported (Schultz, 1988). In colonial invertebrates such as bryozoans, gorgonians, and scleractinians, it is morphological defenses against competitors rather than against predators that have been most widely documented (Adler and Harvell, 1990). Examples of predator-induced defenses have been uncommon in colonial invertebrates and have been cited only for temperate bryozoans (Harvell, 1984). Now, *B. asbestinum* may represent a new example of a tropical colonial invertebrate with an inducible defense against a predator. Another gorgonian (* Plexaura dichotoma*) also responds to snail damage by increasing both the size and density of its sclerites (Nowlis, West, and May, unpubl. data). These results for two different genera of gorgonians suggest that the functioning of sclerites as an inducible defense may be widespread in this order of corals.

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