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***Diadema antillarum* 17 years after mass mortality: is recovery beginning on St. Croix?**

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Abstract *Diadema antillarum* was once ubiquitous in the Caribbean, but mass mortality in 1983–84 reduced its numbers by >97%. We measured *Diadema* abundance on back reefs and patch reefs that have been well studied for >25 years. From June 2000 to June 2001, populations on back reefs have increased >100% (June 2001 mean densities 0.004–0.368/m²), while patch reef populations increased >350% (June 2001 densities 0.236–0.516/m²). Populations are dominated by small urchins, suggesting high recent recruitment. Increased *Diadema* densities appear to be affecting macroalgae abundance. The general spatio-temporal pattern of recovery around St. Croix seems to be following that of the die-off, suggesting that the same oceanographic features that spread *Diadema*'s pathogen are now carrying urchin larvae.

Keywords *Diadema antillarum* · Recovery · Herbivory · Caribbean

Introduction

In 1983–84, *Diadema antillarum*, the Caribbean long-spined sea urchin, experienced one of the most devastating mortalities ever recorded in a marine animal.

Almost certainly caused by an unknown pathogen, the mass mortality reaped >97% of the total Caribbean population, and was not followed by rapid recovery (Lessios 1995).

Its prolonged absence has affirmed that *Diadema* was a major controller of macroalgae on Caribbean reefs, as experiments had suggested earlier [reviewed in Lessios (1988)]. Intensive ecological studies of *Diadema* on St. Croix reefs began in the 1970s, and showed that *Diadema* grazing caused the mysterious halos of bare sand found around Caribbean patch reefs (Ogden et al. 1973). Removal of *Diadema* on reefs at St. Croix and Jamaica resulted in shifts in dominance from algal turfs to filamentous or foliose macroalgae such as *Dictyota*, *Padina*, and *Turbinaria* (Sammarco et al. 1974; Sammarco 1982; Carpenter 1986). Since *Diadema* is also a reef bioeroder (e.g. Ogden 1977), and can influence coral settlement (Sammarco 1980), it shapes the entire reef community. Recent and apparently ongoing recovery of *Diadema* populations in Jamaica has been accompanied by reductions in macroalgal cover and increased coral abundance (Aronson and Precht 2000; Edmunds and Carpenter 2001).

Diadema mass mortality began in St. Croix in mid-January 1984, generally spreading west to east and offshore to onshore. In about 3 weeks the disease traveled approximately 18 km along the north (leeward) shore from Salt River to Tague Bay, (N.B.O., unpublished data; Fig. 1)—opposite to the direction of prevailing currents (Lugo-Fernández et al. 1998). A similar pattern was documented for the spread of the disease in Jamaica, where the mortality took about 2 weeks to travel from Negril to Discovery Bay, moving from west to east, about 25 km a week, against prevailing currents (Hughes et al. 1985). On a larger inter-island scale, the mortality did follow prevailing surface currents, as documented by Lessios et al. (1984). With the exception of Barbados, where the population recovered to 57% of pre-dieoff numbers by 1985 (Hunte and Younglao 1988), recovery has not occurred [reviewed in Lessios (1995)]. Stricken *Diadema* populations have experienced many

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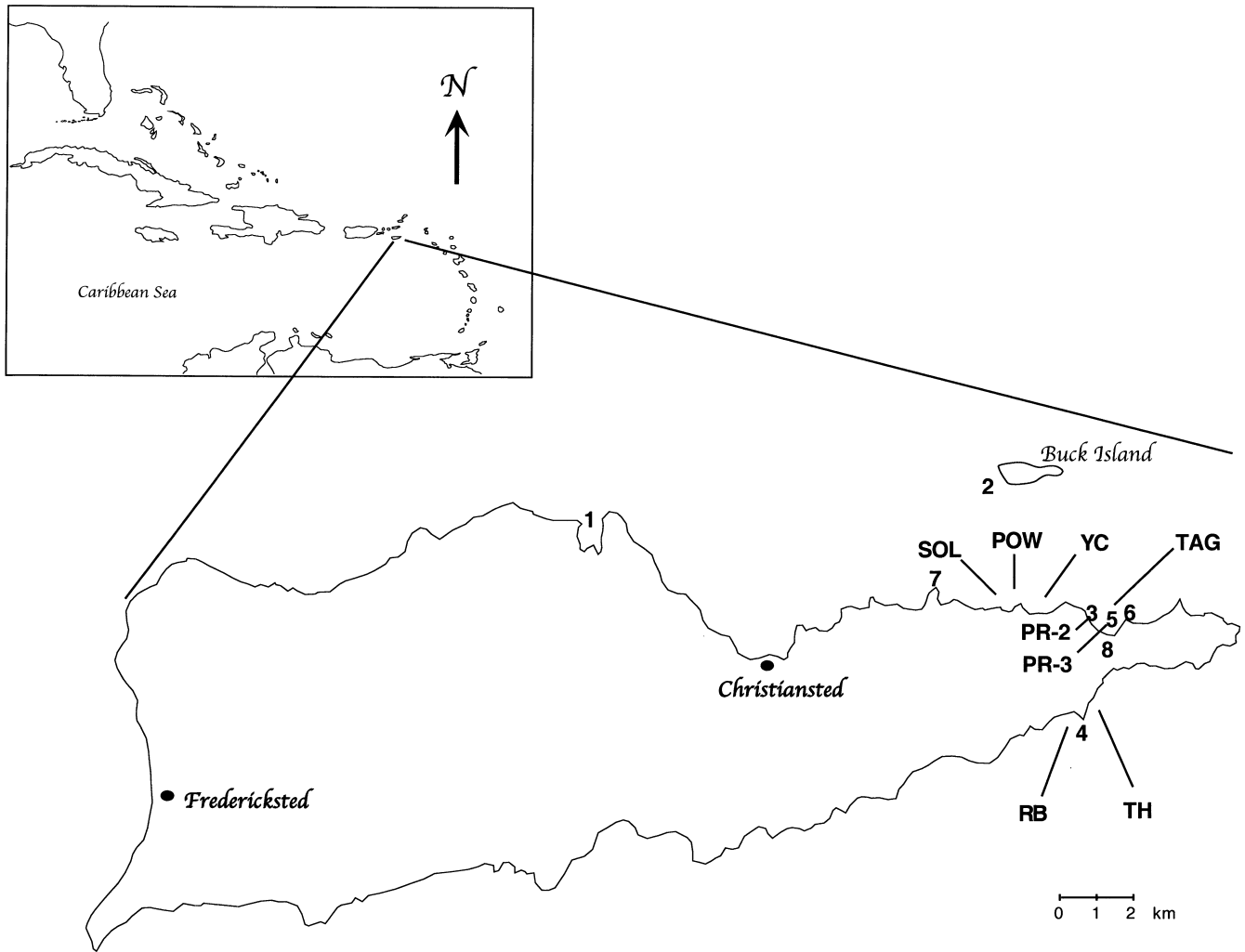


Fig. 1 Map of St. Croix, showing study sites and progression of *Diadema* die-off in 1984. Back-reef sites are: SOL Solitude; POW Pow Point; YC Yellowcliff; TAG Tague Bay; TH Turner Hole; RB Rod Bay. The two patch reef sites, PR-2 and PR-3, are labeled with their ranks in progression of the die-off, as follows. Approximate dates of beginning of die-off: 1) Salt River, 1/18/84; 2) Buck Island, 1/30/84; 3) PR-2, 2/2/84; 4) Grassy Pt., 2/3/84; 5) PR-3, 2/6/84; 6) Romney Pt., 2/13/84; 7) Pull Pt., 2/25/84; 8) West Indies Laboratory flow-through aquaria, 3/3/84

years of low recruitment, failing to produce enough larvae to overcome planktonic mortality (Lessios 1995). Limited fertilization success of widely spaced individuals [Allee effect, e.g. Levitan (1991)] or lack of grazed substrate for settlement (Bak 1985) may have acted to keep populations low (Lessios 1995).

This report documents the possible beginnings of recovery of *Diadema* populations in St. Croix, comparing spatial patterns of recovery with that of the mass mortality, evaluating effects on macroalgae, and comparing recent data (collected by R.J.M., A.J.A., and J.P.E.) with historical data (collected by N.B.O., J.C.O., and R.C. Carpenter). If such a recovery is persistent and widespread in the Caribbean, it may induce a phase

shift on coral reefs, back to coral dominance from the present macroalgal dominance (Edmunds and Carpenter 2001).

Methods

Diadema antillarum was counted four times (in June 2000, October 2000, February 2001, and June 2001) at each of eight locations on St. Croix, USVI: two south-shore back-reef sites, four north-shore back-reef sites, and two north-shore patch reefs (Fig. 1). At each back-reef site at each sampling time, SCUBA divers counted *Diadema* in 14 randomly located 50×2-m transects. Densities from Tague Bay's back-reef from 1983–86 were obtained from Carpenter (1990); these data were collected in March 1986, and in December of all other years. *Diadema* were also counted on two patch reefs in Tague Bay, PR-2 (area ~800 m²) and PR-3 (area ~1,500 m²). On the two patch reefs, SCUBA divers counted all visible *Diadema* in 2000/2001, and complete counts of *Diadema* were made on a yearly basis during the years 1973 and 1982–84 (Ogden et al. 1973; N.B.O., unpublished data).

Diadema test diameters were measured in October 2000 at the same six back-reef sites as above (Fig. 1). All *Diadema* in seven haphazardly located 30×2-m transects at each site were counted and measured to the nearest millimeter with long-jawed calipers. *Triptenustes ventricosus* and *Echinometra* spp. (*lucunter* and *viridis*) were also counted. Pre-mortality (1983) and 1984–86 size frequencies from Tague Bay were obtained from Carpenter (1990).

Test size frequencies for 1983, 1984–86, and 2000 were divided into three classes: small [< 40 mm; mean size of 1-year-old *Diadema* = 48.6 mm (Karlson and Levitan 1990)], medium (41–60 mm), and large (> 60 mm) and compared between years with chi-square tests (Sokal and Rohlf 1995).

Algal percent cover was estimated from the same transects as above at five sites: Pow Point, Yellowcliff, Tague Bay, Turner Hole, and Rod Bay. A video camera held ~ 1 m above bottom was moved along the transect line at a rate of ~ 3 m/min, filming a strip ~ 45 cm wide. Each video frame ($\sim 50 \times 45$ cm) was overlain by an acetate sheet printed with 10 randomly located dots, and the substrate under each dot was categorized as macroalgae (foliose or filamentous algae, not turfs), live coral, sand, or other. Macroalgal percent cover was calculated as the percent of total dots per transect on hard substrate (not including sand) that fell on algae.

Macroalgal percent covers were arcsine transformed ($\text{Sin}^{-1} \sqrt{x}$, Sokal and Rohlf 1995) and regressed against *Diadema* densities. Data were analyzed by transect ($n = 32$).

Results

Abundance

Diadema density at five of six back-reef sites increased by an average 100% from June 2000 to February 2001, then leveled off or decreased slightly from February to June 2001 (Table 1, Fig. 2). The exception was Rod Bay, where densities remained very low. Although Tague Bay density increased 114% during this time, density there also was still quite low at $0.004/\text{m}^2$. *Diadema* density generally increased from west to east among the north-shore back-reef sites, following the pattern of spread of the mortality (Fig. 3). In 1983, before the die-off, Carpenter (1990) recorded mean *Diadema* densities at Tague Bay back-reef of $6.4/\text{m}^2$ (± 81 SE); by 1985, Tague Bay densities were at 0. Though back-reef densities at Tague Bay, the easternmost site, are still low ($0.004/\text{m}^2$ in June 2001) densities at other sites are higher (0.2 – $0.8/\text{m}^2$), though still an order of magnitude lower than pre-mortality densities (Fig. 2).

Diadema densities on the patch reefs grew steadily during this study, increasing an average 350% from June 2000 to June 2001 (Fig. 4). Density of *Diadema* on PR-3, the more eastern patch reef, remained lower than that of PR-2 throughout the sampling. Pre-mortality densities

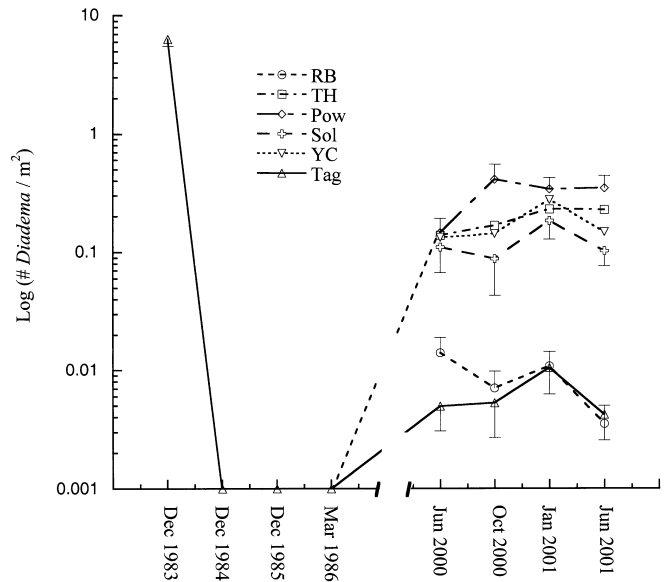


Fig. 2 Log *Diadema* density at six sites on St. Croix from June 2000 to June 2001, and at Tague Bay from 1983 (pre-mortality) to 1986 [data from Carpenter (1990)]. Error bars are 1 standard error. For clarity, most lines between 1986 and 2000 are omitted, error bars are shown for only one direction, and error bars for TH and YC, which are comparable with other sites, are not shown. Note difference in x-axis scaling across break. See Fig. 1 for site abbreviations

of PR-2 and PR-3 from 1973–83 were > 10 *Diadema*/ m^2 ; by 1985 densities had declined to near zero. June 2001 densities are $0.52/\text{m}^2$ for PR-2 and $0.24/\text{m}^2$ for PR-3 (Fig. 4).

Size distributions

Mean test diameter of *D. antillarum* on St. Croix back-reef habitats in October 2000 was 48.6 mm (SD 20.8, $n = 362$). The two southern sites (RB, TH) had larger urchins than the northern sites (Table 2). Mean test diameter was significantly different between sites (ANOVA, $p < 0.05$) but test diameter was not correlated

Table 1 Mean urchin densities per square meter ± 1 SD during this study. *Tripneustes ventricosus* and *Echinometra* sp. densities are from 30×2 -m transects ($n = 7/\text{site}$) measured in October 2000; *Diadema* densities are from 50×2 -m transects ($n = 14/\text{site}$) measured in June/October 2000 and February/June 2001. n/a Data not available

Site	<i>D. antillarum</i>				<i>T. ventricosus</i> Oct 2000	<i>Echinometra</i> sp. Oct 2000
	June 2000	Oct 2000	Feb 2001	June 2001		
Solitude	0.114 ± 0.23	0.091 ± 0.25	0.193 ± 0.30	0.107 ± 0.15	0.007 ± 0.002	0.357 ± 0.06
Pow Point	0.151 ± 0.26	0.431 ± 0.79	0.359 ± 0.31	0.368 ± 0.54	0	0.012 ± 0.01
Yellowcliff	0.138 ± 0.17	0.149 ± 0.20	0.291 ± 0.02	0.156 ± 0.17	0.007 ± 0.002	0.171 ± 0.03
Tague Bay	0.005 ± 0.01	0.005 ± 0.01	0.011 ± 0.47	0.004 ± 0.01	0	0.100 ± 0.01
Turner Hole	0.143 ± 0.24	0.175 ± 0.26	0.241 ± 0.40	0.243 ± 0.40	0	0.143 ± 0.02
Rod Bay	0.014 ± 0.03	0.007 ± 0.01	0.011 ± 0.02	0.004 ± 0.01	0	1.943 ± 0.20
PR-2	0.085	0.261	0.278	0.516	n/a	n/a
PR-3	0.082	0.185	0.198	0.239	n/a	n/a

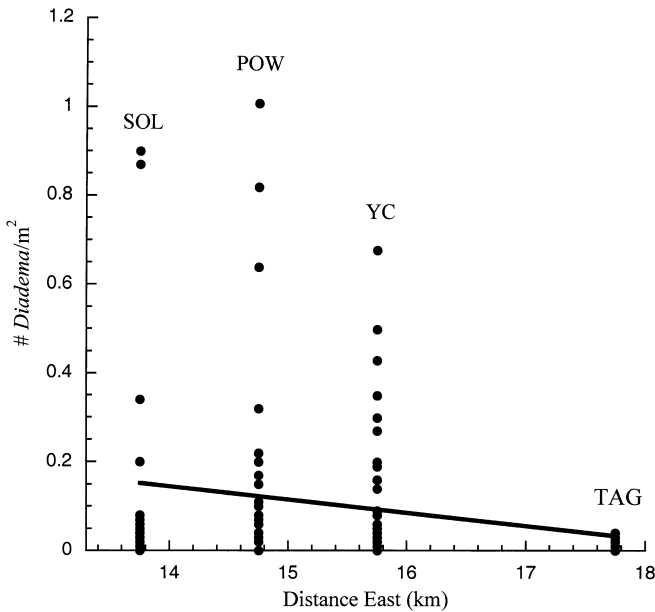


Fig. 3 *Diadema* density across the four north-shore sites from west to east. Distances are in kilometers from Salt River to the west (Fig. 1). The negative slope is significant ($F=15.74$, $p=0.007$). $Y=63.105-3.431X$, $R^2=0.08$. Data points represent individual 50×2 m transects from June 2000 ($n=14/\text{site}$)

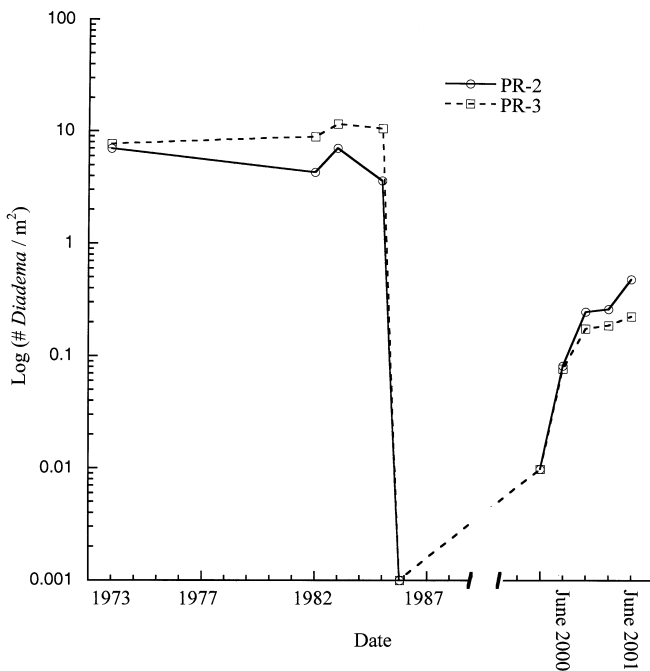


Fig. 4 *Diadema* density at patch reefs PR-2 and PR-3. [1973–84 data from Ogden (Ogden et al. 1973, N.B.O. and J.C.O., unpublished data).] Despite higher historical densities at PR-3, PR-2, whose population crashed ~1 week prior to that of PR-3, now has higher densities

with population density ($p>0.05$). Size frequency of *Diadema* pooled across all sites is shown in Fig. 5, along with historical data. The October 2000 distribution is

dominated by 31- to 40-mm urchins (31.6%). The pre-mortality size distribution [from Carpenter (1990)] is dominated by small to mid-sizes, particularly the 21- to 30-mm size class (25.5%), whereas the 1984–86 distribution (shortly after mass mortality) is dominated by very large sizes, with 32.8% >80 mm. The distribution of *Diadema*'s small (<40 mm), medium (41–60 mm), and large (>60 mm) size classes in 2000 is not significantly different from the 1983 pre-mortality distribution (chi-square, $p=0.07$). The 1984–86 distribution, however, is very different from both the 2000 distribution (chi-square, $p<0.0001$) and the 1983 distribution (chi-square, $p<0.0001$).

Grazing

Macroalgal cover showed a significant negative relationship with *Diadema* density (ANOVA, $p=0.03$, Fig. 6). Algal cover appears to be dramatically higher when urchin abundance is below about $0.1/\text{m}^2$. No significant trend was seen in coral cover ($p>0.05$), although our sampling method did not have the resolution to detect coral recruits, as Edmunds and Carpenter (2001) did.

Tripneustes abundances were very low, and *Echinometra* abundances were not correlated with either *Diadema* abundance or macroalgal cover ($p>0.05$, Table 1).

Discussion

Diadema densities on St. Croix are still far below pre-mortality numbers, but substantially higher than the near-zero densities that immediately followed the mass mortality. Recent rapid population growth ($>100\%$ increase on back-reefs, $>350\%$ on patch reefs) contrasts sharply with the very slow to nonexistent increases of the past 17 years. Recent size distributions indicate successful recruitment, suggesting that this could be the beginning of a full population recovery, potentially returning *Diadema* to its role as an important grazer on Caribbean reefs.

Small *Diadema* are now abundant in St. Croix, and the population is approaching the pre-dieoff size

Table 2 Mean *D. antillarum* test diameter (mm) for six St. Croix sites, measured in October 2000. n Number of urchins measured in seven transects; SD standard deviation

Site	Mean test diameter (mm)	SD	n
Solitude	42.1	17.8	82
Pow Point	43.0	20.3	95
Yellowcliff	53.8	20.6	89
Tague Bay	40.7	17.8	20
Turner Hole	57.1	19.6	69
Rod Bay	67.9	13.0	9

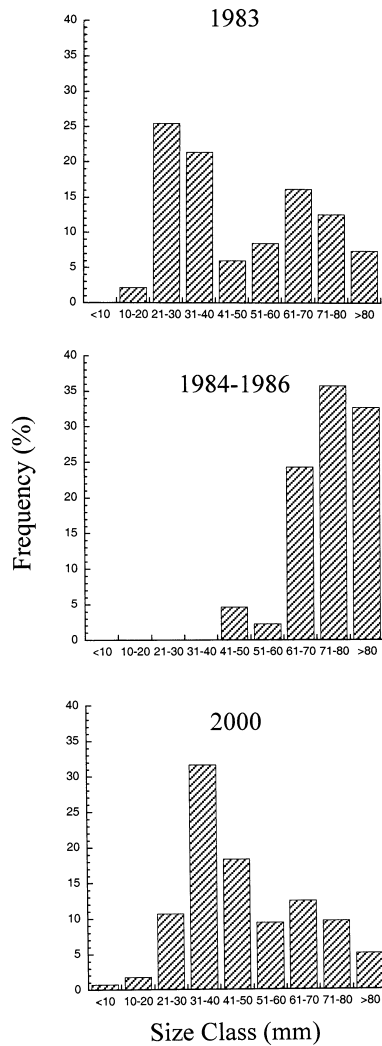


Fig. 5 Frequencies of *Diadema antillarum* test diameters for 1983 (pre-mortality, Tague Bay, $n=400$), 1984–86 (immediately post-mortality, Tague Bay, $n=149$), and October 2000 (Solitude, Pow Pt., Yellowcliff, Tague Bay, Rod Bay, and Turner Hole, $n=364$). [1983–86 data from Carpenter (1990)]

frequency distribution (Fig. 5). Urchins from 20–40 mm have become particularly common; this size corresponds with urchins less than 1 year old (Karlson and Levitan 1990). Mean test diameter is not correlated with population density across sites; this suggests that population densities are well below trophic carrying capacity, with no intraspecific competition (Levitan 1988).

The recently elevated *Diadema* densities in St. Croix are evidently affecting macroalgal cover (Fig. 6), as they did before the mass mortality (Ogden et al. 1973). We noticed that the rugose micro-sites that tend to harbor *Diadema* were often visible from a considerable distance due to the bright bare patches resulting from urchin grazing (R.J.M., A.J.A., J.P.E., J.C.O., personal observations). Of course, anecdotes and correlations do not prove causality, but decades of experimental and correlative studies on the effect of

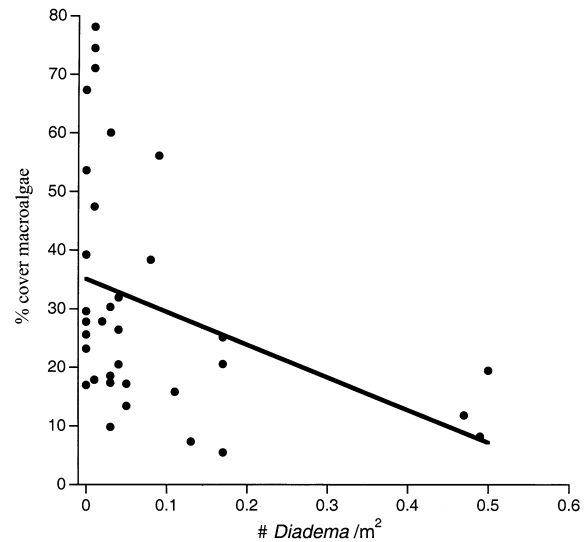


Fig. 6 Percent cover macroalgae regressed against *Diadema* density. Negative slope is significant ($F=4.9323$, $p=0.03$). $Y=0.317735-0.00612X$, $R^2=0.13$. Regression analysis was done on arcsine-transformed data

Diadema grazing [reviewed in Lessios (1988)] support this interpretation.

The 1984 mortality spread from west to east and offshore to onshore at St. Croix, as was particularly noted on the north shore (Fig. 1). Bank-barrier reef populations died off first, followed by populations on lagoonal patch reefs, and, finally, by urchins in the flow-through aquaria at the West Indies Laboratory, where the seawater intakes were located close to shore (N.B.O., unpublished data). This pattern recurred on an even smaller scale on patch reefs PR-2 and PR-3, which are less than 100 m apart: the more western PR-2 experienced the population crash ~ 1 week prior to PR-3, and urchins began dying on the seaward edge of both reefs, reflecting the offshore to onshore pattern. The beginnings of recovery documented here (June 2000) show a similar pattern to that of the mortality, with highest north-shore densities in the west on back reefs, and higher densities on PR-2, the more western patch reef, compared to PR-3 (Figs. 3 and 4).

Larval filtering—higher settlement at upstream versus downstream sites, as larvae settle on suitable substrates and are depleted en route (Gaines et al. 1985)—might explain these spatio-temporal patterns of recovery. Though net flow along the north shore of St. Croix is westward, tide-driven current reversals occur frequently (Lugo-Fernández et al. 1998), and it is also conceivable that larval behavior could result in eastward transport (e.g. Pineda 1999). Caselle and Warner (1996) noted a similar pattern of highest recruitment on western upcurrent reefs of the north shore of St. Croix in the reef fish *Thalassoma bifasciatum*, and Swearer et al. (1999) inferred from trace elements in juvenile otoliths that larval retention on western reefs caused the high recruitment. A reversal of prevailing currents off

St. Croix's north shore in 1997 was caused by a meso-scale anti-cyclonic eddy, which may be a regular event in the area (Harlan et al. 2002). This may facilitate larval retention on western reefs, or contribute larvae of distant origin to the reefs, and help explain the spatio-temporal pattern in our data.

Interestingly, the spread of mass mortality on Jamaica's north coast was also west to east in direction, and it moved with similar speed (Hughes et al. 1985). Unlike St. Croix, present densities at some of the same sites cited in Hughes et al. (1985) do not reflect a similar pattern of recovery (Edmunds and Carpenter 2001). However, the recovery in Jamaica is much more advanced than that in St. Croix [densities of $\sim 5/m^2$ in 'urchin zones'; Edmunds and Carpenter (2001)], and early patterns may no longer be visible. In addition, facilitation by *Tripneustes* or *Echinometra* grazing does not seem to have played a role in the recovery in St. Croix, as it may have in Jamaica (Aronson and Precht 2000).

What caused the sudden increase in *Diadema* densities in St. Croix, after 15 years of negligible recovery? The population may have been seeded by larvae from some upstream source—a likely candidate is Barbados, where *Diadema* populations recovered relatively quickly after the die-off (Hunte and Younglao 1988), and reefs still enjoy relatively high densities of *Diadema* ($1.94/m^2$ at Bellairs Reef in 2000; CARICOMP data). However, we do not have data from sites between St. Croix and Barbados to support this conjecture, and this recolonization route does not by itself explain why it would take so long for significant numbers of larvae to reach St. Croix. Moreover, Hunte and Younglao (1988) concluded that Barbados is not a regional larval source, since they saw no downcurrent recruitment of *Diadema* at St. Lucia or Dominica coinciding with recruitment pulses at Barbados. Seeding from distant larval sources may be episodic, with recruitment in most years largely from resident populations, as has been suggested for some reef fish populations on St. Croix (Swearer et al. 1999). This would account for a very slow initial recovery, followed by more rapid population increases when local populations achieve critical densities. The high proportions of juvenile *Diadema* appearing now in St. Croix may indicate that these critical densities have been achieved.

This research, coupled with recent reports from Jamaica, and unpublished CARICOMP data, suggest that a large-scale recovery of *Diadema* populations may be underway. Immediate and continued monitoring of *Diadema* populations throughout the Caribbean is essential if we are to understand how this recovery proceeds.

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References

- Aronson RB, Precht WF (2000) Herbivory and algal dynamics on the coral reef at Discovery Bay, Jamaica. *Limnol Oceanogr* 45(1):251–255
- Bak RPM (1985) Recruitment patterns and mass mortalities in the sea urchin *Diadema antillarum*. *Proc 5th Int Coral Reef Congr* 5:267–272
- Carpenter RC (1986) Partitioning herbivory and its effects on coral reef algal communities. *Ecol Monogr* 56:345–63
- Carpenter RC (1990) Mass mortality of *Diadema antillarum* I. Long-term effects on sea urchin population-dynamics and coral reef algal communities. *Mar Biol* 104:67–77
- Caselle JE, Warner RR (1996) Variability in recruitment of coral reef fishes: the importance of habitat at two spatial scales. *Ecology* 77:2488–2504
- Edmunds PJ, Carpenter RC (2001) Recovery of *Diadema antillarum* reduces macroalgal cover and increases abundance of juvenile corals on a Caribbean reef. *Proc Natl Acad Sci* 98:5067
- Gaines SD, Brown S, Roughgarden J (1985) Spatial variation in larval concentrations as a cause of spatial variation in settlement for the barnacle, *Balanus glandula*. *Oecologia* 67:267–272
- Harlan JA, Swearer SE, Leben RR, Fox CA (2002) Surface circulation in a Caribbean island wake. *Cont Shelf Res* 22:417–434
- Hughes TP, Keller BD, Jackson JBC, Boyle MJ (1985) Mass mortality of the echinoid *Diadema antillarum* Phillipi in Jamaica. *Bull Mar Sci* 36:377–384
- Hunte W, Younglao D (1988) Recruitment and population recovery of *Diadema antillarum* (Echinodermata; Echinoidea) in Barbados. *Mar Ecol Prog Ser* 45:109–119
- Karlson RH, Levitan DR (1990) Recruitment-limitation in open populations of *Diadema antillarum*: an evaluation. *Oecologia* 82:40–44
- Lessios HA (1988) Mass mortality of *Diadema antillarum* in the Caribbean: what have we learned? *Annu Rev Ecol Syst* 19:371–393
- Lessios HA (1995) *Diadema antillarum* 10 years after mass mortality: still rare, despite help from a competitor. *Proc R Soc Lond B* 259:331–337
- Lessios HA, Robertson DR, Cubit JD (1984) Spread of *Diadema* mass mortality through the Caribbean. *Science* 226:335–337
- Levitan DR (1988) Algal-urchin biomass responses following mass mortality of *Diadema antillarum* Phillipi at St. John, U.S. Virgin Islands. *J Exp Mar Biol Ecol* 119:167–178
- Levitan DR (1991) Skeletal changes in the test and jaws of the sea urchin *Diadema antillarum* in response to food limitation. *Mar Biol* 111:431–435
- Lugo-Fernández A, Roberts HH, Wiseman WJ Jr, Carter BL (1998) Water level and currents of tidal and infragravity periods at Tague Reef, St. Croix (USVI). *Coral Reefs* 17:343–349
- Ogden JC (1977) Carbonate-sediment production by parrot fish and sea urchins on Caribbean reefs. In: Frost SH, Weiss MP, Saunders JB (eds) *Reefs and related carbonates—ecology and sedimentology*. American Association of Petroleum Geologists Studies in Geology no 4, Tulsa, Oklahoma, pp 281–288
- Ogden JC, Brown RA, Salesky N (1973) Grazing by the echinoid *Diadema antillarum*. Formation of halos around West-Indian patch reefs. *Science* 182:715–717
- Pineda J (1999) Circulation and larval distribution in internal tidal bore warm fronts. *Limnol Oceanogr* 44:1400–1414
- Sammarco PW (1980) *Diadema* and its relationship to coral spat mortality: grazing, competition, and biological disturbance. *J Exp Mar Biol Ecol* 45:245–272

- Sammarco PW (1982) Effects of grazing by *Diadema antillarum* Philippi (Echinodermata: Echinoidea) on algal diversity and community structure. *J Exp Mar Biol Ecol* 65:83–105
- Sammarco PW, Levinton JS, Ogden JC (1974) Grazing and control of coral reef community structure by *Diadema antillarum* (Echinodermata:Echinoidea): a preliminary study. *J Mar Res* 32:47–53
- Sokal RR, Rohlf FJ (1995) *Biometry*. WH Freeman, New York
- Swearer SE, Caselle JE, Lea DW, Warner RR (1999) Larval retention dynamics drive recruitment patterns in an island population of a coral reef fish. *Nature* 402:799–802