

## Temporal patterns in coral reef, seagrass and mangrove communities from Chengue bay CARICOMP site (Colombia): 1993-2008

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**Abstract:** Few monitoring programs have simultaneously assessed the dynamics of linked marine ecosystems (coral reefs, seagrass beds and mangroves) to document their temporal and spatial variability. Based on CARICOMP protocol we evaluated permanent stations in coral reefs, seagrass beds and mangroves from 1993 to 2008 in Chengue Bay at the Tayrona Natural Park, Colombian Caribbean. Overall, the studied ecosystems showed a remarkable stability pattern over the monitoring period. While there were annual variations in coral reefs (coral cover) and mangroves (litterfall) caused by hurricane Lenny in 1999, particular trends in seagrass (leaf area index and leaf productivity) appear to reflect the natural variability in this ecosystem. We suggest that monitoring sites at the three marine ecosystems had in general a healthy development in the last 16 years. Our results are critical to locally improve the management strategies (Tayrona Natural Park) and to understand the long-term dynamics of closely associated marine ecosystems in the Caribbean. *Rev. Biol. Trop.* 58 (Suppl. 3): 45-62. Epub 2010 October 01.

**Key words:** monitoring, coral reef, seagrass, mangrove, Chengue, CARICOMP, Colombian Caribbean.

Coral reefs, seagrass beds and mangroves are undoubtedly the most important ecosystems on the coastal areas of tropical seas. They contribute diverse benefits to the coastal zone of many regions including protection of shores from erosion, supporting high biodiversity and sustaining fisheries and tourism (Ogden & Gladfelter 1983, Kjerfve *et al.* 1998). Although each ecosystem can exist independently and has specific characteristics that make it different from the others, they frequently occur together in the same area and the resulting interactions could favor their own development and balance (Mumby *et al.* 2004, Harborne *et al.* 2006). For example, coral reefs

create barriers that reduce the impact of wave action and currents and therefore facilitate the development of sedimentary substrates where seagrasses and mangroves can flourish (Ogden 1988). Mangroves and seagrass beds function as sedimentary traps that retain materials from coastal runoff, improving water transparency and favoring coral reef growth (Granek *et al.* 2009). Furthermore, it has been proposed that transfer of materials, nutrients and energy that occur among these three ecosystems are very important to sustain the high productivity and biodiversity in the coastal zone (Granek *et al.* 2009). On the other hand, Halpern *et al.* (2007) explain that interactions

among these ecosystems will affect the impact of a threat in a particular area, both positive (dispersal-mediated recovery) and negative (threat transfer). Since mangroves and seagrass are capable of providing high-quality nursery habitats for juvenile reef fishes, their loss could have a large impact on coral reefs community fishes that depend upon these nursery habitats (Nagelkerken *et al.* 2000, Dorenbosch *et al.* 2004, Mumby *et al.* 2004).

Despite their importance to human welfare, these ecosystems have been strongly impacted by anthropogenic activities. Important portions of mangroves around the world are threatened, and present-day mangrove forest areas have declined substantially, with an average loss of 35% (Valiela *et al.* 2001). Destruction or loss of seagrass meadows have also been reported for most parts of the world; even if natural causes are often related with this condition (e.g. “wasting disease” or high energy storms), the destruction has been commonly associated with human activities due to coastal development and growing populations (Short & Coles 2001, Duarte *et al.* 2006). Ecological degradation has led to a decrease between 30 and 60% of seagrass beds around the world (Lotze *et al.* 2006, Waycott *et al.* 2009), and locally, some areas of the Colombian Caribbean have lost even more than 90% of seagrasses in the last 70 years (Díaz & Gómez 2003). During the last three decades of the past century, coral reefs have been facing unprecedented changes and mortality events when compared with geological timescales (Aronson *et al.* 2004, Pandolfi & Jackson 2006, Pandolfi *et al.* 2006). An unequivocal sign of these changes is the loss of living coral cover which was estimated around 5% per year until 2001 in the Caribbean (Gardner *et al.* 2003, Côté *et al.* 2005), 2% per year until 2004 in the Indo-Pacific (Bruno & Selig 2007) and has been significant over the last 40 years in the GBR (Bellwood *et al.* 2004). Although Colombian coral reefs have not been exempt from this worldwide process, no clear patterns of coral decline have been observed in recent years by monitoring programs (Rodríguez-Ramírez *et al.* 2010, Zapata *et al.* 2010).

An international cooperative research network to advance understanding of the interactions and importance of the three major coastal ecosystems of the Caribbean was formulated at the beginning of the 1990s. The Caribbean Coastal Marine Productivity Program (CARICOMP) was created by this initiative and monitoring activities began in many countries of the region in 1992 (CARICOMP 1997a, Kjerfve *et al.* 1998). The Institute of Marine and Coastal Research (INVEMAR) has represented Colombia and contributed to the CARICOMP program through provision of long term monitoring at Chengue Bay (Tayrona Natural Park) since 1992 (Garzón-Ferreira 1998). This site includes sampling stations in coral reef, seagrass and mangrove ecosystems. A second CARICOMP site was established in 1998 in Colombia at San Andrés Island under the responsibility of the Corporation for the Sustainable Development of the San Andrés and Providencia Archipelago (CORALINA). The experience obtained by INVEMAR while participating for several years in CARICOMP was essential for the creation and launch of the National Reef Monitoring System of Colombia (SIMAC). This program has been operating since 1998 without interruption and includes monitoring stations in the Caribbean and the Pacific (Garzón-Ferreira & Rodríguez-Ramírez 2010).

This paper presents a synthesis of the results obtained during 16 years of environmental monitoring in Chengue Bay. Temporal patterns displayed by each studied variable are analyzed and discussed for each of the three ecosystems: coral reef, seagrass beds and mangrove forest.

## MATERIALS AND METHODS

**Study site:** Chengue is a small bay (surface area 3.3km<sup>2</sup>) located in the Parque Nacional Natural Tayrona, 14km northeast of the city of Santa Marta on the Caribbean coast of Colombia (11°20' N, 74°08' W) (Fig. 1). The coastal topography is heterogeneous, with steep relief due to the closeness of the Sierra

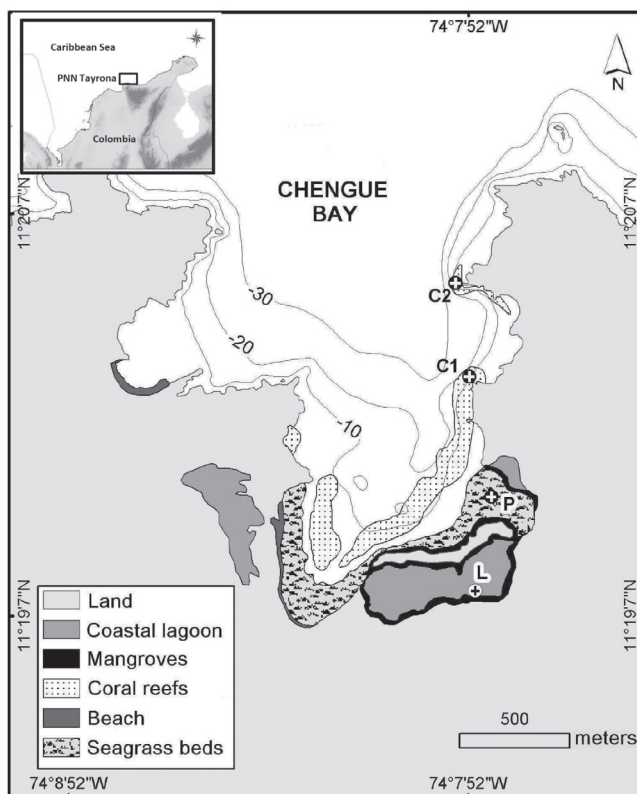


Fig. 1. Chengue Bay indicating bathymetry (m), distribution of principal coastal marine ecosystems, location of CARICOMP monitoring sites (plus signs): mangrove wetlands on the lagoon (L), seagrass beds (P) and coral reefs (C1 and C2).

Nevada de Santa Marta, the highest mountain system in Colombia (5 800m above sea level). Consequently, the landscape is shaped by numerous rocky headlands, islets, inlets, and bays with sedimentary beaches, small lagoons and small river discharges (Garzón-Ferreira & Cano 1991). Regional climate and oceanography are determined by coastal topography and trade wind incidence, with a dry season from December to April, when strong NE trade winds ( $3.5\text{ms}^{-1}$ ) reduce precipitation and generate an upwelling that transports cold water (mean  $25.5^{\circ}\text{C}$ ) from 100-200m depth to the surface (Bula-Meyer 1977, Ramírez 1983); and a rainy season from May to November, with lower-velocity trade winds (mean  $1.5\text{m s}^{-1}$ ), higher precipitation (>80% of the annual total)

and sea surface temperature (mean  $28.0^{\circ}\text{C}$ ), and frequently turbid currents (continental run-off) entering the bay (Garzón-Ferreira 1998). A narrow continental shelf, with depths of 200m only 2km from shore, supports small but highly diverse biological communities and ecosystems. Chengue has low human intervention, although some activities have an impact on marine ecosystems.

CARICOMP monitoring in Chengue Bay is conducted in three ecosystems: mangroves, seagrass beds, and coral reefs, which have developed at the eastern and southern shores (Fig. 1), where trade wind-induced wave action is low; instead, the bay's NW shore is influenced by high wave energy and clearly dominated by rocky bottoms. Coral reef

communities in Chengue have been reviewed by several authors (Werding & Erhardt 1976, Solano 1987, Garzón-Ferreira & Cano 1991), concluding in a general description of thirty-one species of hermatypic corals and three hydrocorals. Two main types of reefs are identified in Chengue (Garzón-Ferreira 1998): 1) narrow, short coastal fringing reefs growing over the belt of metamorphic rocks in the northern deep half of the bay, dominated by massive and encrusting corals (mainly *Diploria*, *Montastraea*, and *Colpophyllia*), which form a reef slope extending to a depth of 15-25m and ; (2) extensive fringing reefs growing away from the shore on sedimentary flats of the bay, dominated by foliaceous and branching corals (*Acropora palmata*, *Agaricia tenuifolia* and *Millepora*), which become reef slopes covered with massive corals at depths of 6-8m.

All five species of seagrass beds occurring in Colombia are found in Chengue. *Thalassia testudinum* beds (where CARICOMP monitoring is conducted) are the most extensive, although restricted to shallow waters in the southern portion of the bay (<3m depth) (Rodríguez-Ramírez & Garzón-Ferreira 2003). Within these beds are found patches of *Syringodium filiforme* and, less frequently, *Halodule wrightii* and *Halophila baillonis*. In deeper waters of the eastern bay wide stands of *Halophila decipiens* also occur. *Thalassia* beds grow over calcareous sediments, mainly coarse sand and coral rubble. The calcareous algae *Halimeda opuntia* is particularly abundant, and in many places, is the dominant live component covering the bottom of *Thalassia* beds (Garzón-Ferreira & Cano 1991). Chengue exhibits the most important mangrove formation in the national park, which is associated with a coastal lagoon (Fig. 1) surrounded mostly by Red mangrove *Rhizophora mangle* reaching heights of 13m and densities of 3800 trees/ha (Garzón-Ferreira & Cano 1991, Garzón-Ferreira 1998). Nearby, in less swampy terrain, three additional species are found: *Avicennia germinans*, *Conocarpus erectus*, and *Laguncularia racemosa*. More detailed information about coastal ecosystems in Chengue is provided by

Rodríguez-Ramírez & Garzón-Ferreira (2003) and Rodríguez-Ramírez *et al.* (2004).

#### **Coral reef sampling and data analysis:**

The cover of benthic components was estimated annually along five 10 m permanent marked transects on two reef sites (Fig. 1). Transects were sampled using a light chain; components beneath each link were recorded following categories used by CARICOMP (2001). Hard corals were identified to the lowest possible taxonomic level (genus or species). Incidence of coral diseases and bleaching were estimated by examining every hard coral colony (>5cm) within a 2m wide band along each 10m transect. A PVC 1-m pipe marked every 5cm was used as a reference to estimate colony size and band width while swimming at each side of the transect line. Each colony was identified to species level and its surface examined carefully to record the presence of any disease or bleaching. The abundance of important mobile invertebrates (lobsters, octopuses, crabs and sea urchins) was also recorded along each belt transect (10x2m).

Prior to analyses, cover estimates were arcsine transformed. Overall means were calculated by averaging annual means values (averaging all transects per year) by each variable. Temporal trends of coral, algae (pooling turf, fleshy, encrusting and calcareous algae) and abiotic substrate were evaluated by repeated measures ANOVA. We used the univariate approach and the Greenhouse-Geisser corrected probability to test F (adjustment for multiple comparisons: Bonferroni).

#### **Seagrass sampling and data analysis:**

Seagrass community was monitored in two sampling stations to the SE in Chengue Bay which had luxuriant *Thalassia testudinum* beds (Fig. 1). In each station, net above-ground productivity and shoot densities of *T. testudinum* were determined annually (each September) using the leaf marking technique from CARICOMP (2001). Shoots within six tagged wire quadrats (20x10cm) were marked a short distance above the green/white interface at the leaf base with a

single punch of a needle. After a growth period of 8 days, the shoots were counted and extracted in order to estimate shoot densities. Each leaf was cut at the marked point and sorted into three fractions, -new leaves (unmarked leaves that had emerged after marking), old growth (the part of the leaves between the needle mark and meristem (=green-white interface) and old standing crop (the leaf part above the needle mark)-. Leaf fractions were decalcified in 10% hydrochloric acid, rinsed and dried at 75°C to constant weight for dry weight determination. Additionally, five shoots were collected from each sampling station to estimate Leaf Area Index (LAI). Leaf blades were counted, and total length and width of each leaf measured. Leaf area per shoot and LAI, which is the product of leaf area per shoot and shoot density, were calculated. Environmental data at the sampling stations were registered; temperature and salinity were measured with data loggers and a hand refractometer, respectively. Light attenuation (water transparency) was measured horizontally with a Secchi disk and expressed as the length at which the disk no longer could be seen.

Mean values of productivity (g/m<sup>2</sup>/day), shoot density (shoots/200cm<sup>2</sup>) and LAI were used to estimate temporal trends. A polynomial model was used to fit the data if R<sup>2</sup>≥0.70. Pearson's correlations were run to identify relationships between environmental variables and biological data.

**Mangrove sampling and data analysis:** Mangrove assessment was conducted following the CARICOMP monitoring methods (CARICOMP 2001). The site was established along the southeastern shore of Chengue Bay, where *Rhizophora mangle* appeared to be dominant in a fringing forest (Fig. 1). To determine the structural attributes of the red mangrove, three plots of 10x10m (0.01ha) were established randomly inside the forest and monitored at one year intervals. Inside each plot, all trees with trunk diameter greater than 2.5cm inside a square 10m side were marked with consecutive numbers, and mapped to facilitate their

relocation the following year. Height above sediment of the highest prop root (m), length of trunk (m), diameter at breast height or DBH (cm), basal area (ground space covered by trees in m<sup>2</sup>/ha), density (trees/100m<sup>2</sup>) and biomass (kg/m<sup>2</sup>) were all recorded annually. The latter variable was calculated using the DBH conversion factor of Golley *et al.* (1962). The complexity of the forest in terms of the structure of trees was estimated with the Holdridge index (Holdridge *et al.* 1971). The mean annual structural variables were calculated averaging the three plots. Within each same plot, primary productivity was estimated based on litterfall (g/m<sup>2</sup>/day). Ten fall traps of 50x50cm (0.25m<sup>2</sup>) were deployed at regular intervals and collected one month later (October through November). Additionally, the interstitial salinity was measured. In the laboratory, the samples were classified (leaves, bracts, fruits, flowers, wood/twigs, frass and branches) and oven dried to 70°C for 48 hours, before being weighed. Subsequently, means data for the traps in the three plots were calculated to obtain overall means for October and November.

## RESULTS

**Coral reefs:** Overall means of coral and algal cover at Chengue were 33% (SE±0.42) and 48.8% (SE±0.78) respectively. Algal turf was the most abundant biotic and algal component (mean=30.6% (SE±1.41) and the dominant coral species belonged to the *Montastraea annularis* complex (18.3% SE±0.47). Live coral cover has remained stable during the study period (no significant differences  $p>0.05$ , Fig. 2A). Only a slight decrease (~4%) in coral cover was registered between 1999 and 2000 (Fig. 2A). In contrast, algal cover and abiotic substrate have showed high variability (Fig. 2B,C). Algae ranged from 43.8% in 1998 to 53.5% in 2002 (Fig. 2B) and abiotic substrate from 11% in 1993 to 20.7% in 2008 (Fig 2C). These components displayed several significant differences among years but no

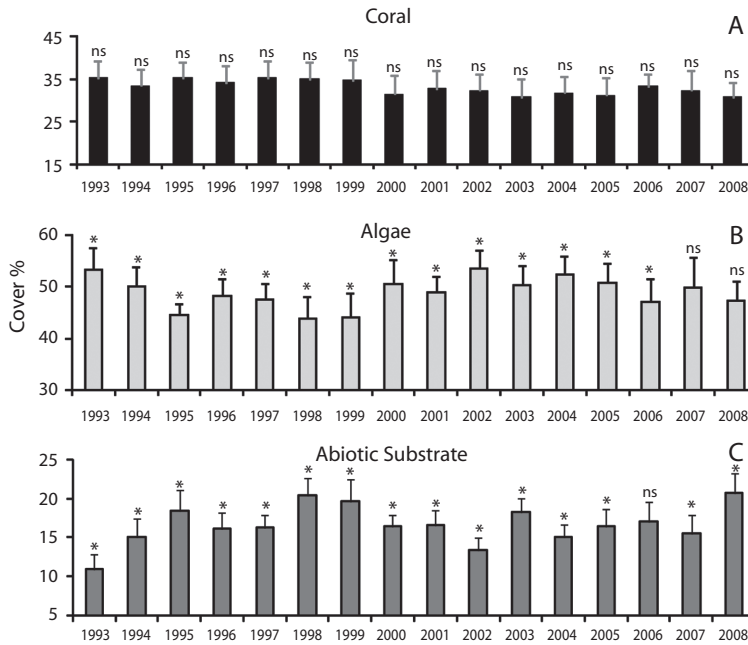


Fig. 2. Mean percentage cover (+ 1 SE) for major benthic components at Chengue Bay during the monitoring period (1993-2008). (A) Coral, (B) Algae, and (C) Abiotic Substrate. ns= year similar to others years,  $p > 0.05$ . \*= year significantly different to one or more years,  $p < 0.05$ .

clear trends (increase or decrease) were evident (Fig. 2B,C).

Few changes in disease prevalence, coral bleaching and paled corals were observed between 1998 and 2008 (Fig. 3). The major bleaching event occurred during 1998, the first year of monitoring; with 10% of corals being affected. In 1999, this percentage decreased to about 4%, and continued to decrease in successive years. Thereafter, bleaching rates were under 4%, until 2008 when bleached corals reached 5% of the total of colonies in the transects. Paled corals registered similar tendencies to bleached corals, with values over 12% in 1998, increasing to close to 15% in 1999. After 1999, the registered percentage of paled corals stayed below 2%, except in 2005 and 2008, when the percentage of paled corals increased to over 3%. Coral diseases such as Black Band Disease, White Band Disease, Yellow Band Disease and White Plague have

been observed in Chengue. The most prevalent coral diseases during the 10 years of monitoring were Dark Spots Disease (0.0%-2.5%) and White Plague (0.2%- 2.3%).

Lobsters (*Panulirus* spp.) and octopuses (*Octopus* spp.) were sporadically observed in transects. Sea urchins (*Echinometra* spp. and *Diadema antillarum*) were scarce (maximum density 0.025 individuals/m<sup>2</sup> in 2007).

**Seagrass beds:** Structural parameters of *T. testudinum* beds on Chengue Bay showed different trends over the monitoring time (Fig. 4); thus: 1) LAI had periodic maximum peaks every 3-4 years, during 1996, 1999, 2003 and 2007; however, analysis showed no time trend (polynomial regression;  $R^2 < 0.70$ ). Overall mean value for LAI was  $4.0 \pm 0.5$ , with fluctuations from  $2.4 \pm 0.3$  (in 2002) to  $6.7 \pm 0.5$  (in 2003). 2) Density of shoots showed several fluctuations, but not as marked as LAI; maximum peaks were



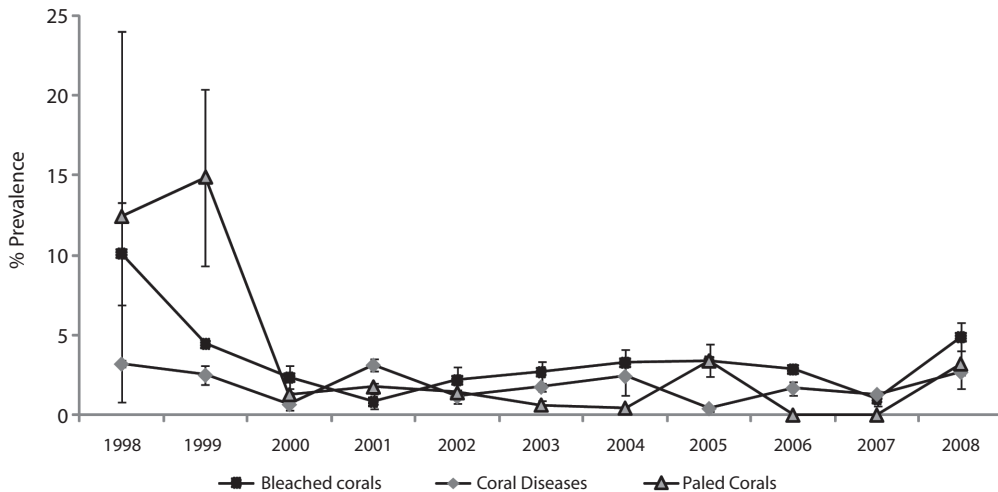


Fig. 3. Mean ( $\pm 1$  SE) of bleached, paled and disease coral prevalence at Chengue Bay during the monitoring period (1998-2008).

found in 1996, 1999, 2003, 2004 and 2005, and no time trend was evident (polynomial regression;  $R^2 < 0.70$ ). Overall mean value for shoot density was  $13.5 \pm 0.5$  shoots/200cm<sup>2</sup>; with fluctuations from  $11.3 \pm 0.5$  (in 2002) to  $16.1 \pm 1.1$  (in 2003) shoots/200cm<sup>2</sup>. 3) Productivity was constant during most of the monitoring period (near 2.5 g/m<sup>2</sup>/day); however, during 1999 an extremely high value was evident, and since 2005 mean values showed a significant progressive increase ( $R^2 > 0.70$ ). No correlations were identified between environmental variables and biological data (Pearson, all  $r$  values between  $-0.390$  and  $0.135$  and  $p$  values  $> 0.05$ ). Means of environmental data during the sampling periods are shown in Table 1.

**Mangrove:** Red mangrove (*R. mangle*) formations on the southeastern coastal lagoon, at Chengue, have been assessed annually since 1995 (13 years). The structural complexity according to the Holdridge index had a mean value of 1.07 (Table 2). Key average characteristics of the Red mangrove included: 1) 1.57m highest prop root, 2) 3.58m trunk length, 3)

12.3cm of DBH, 4) 43.8m<sup>2</sup>/ha basal area, 5) 13.1kg/m<sup>2</sup> biomass and 6) a density of 32.1 trees/100m<sup>2</sup>. The mean value of interstitial salinity was 39.3. No major structural changes in the mangrove stand have been recorded over the monitoring period. Detailed information on the values of each structural component is shown in Table 3. The mean litter production remained constant between 4.0 and 6.0g/m<sup>2</sup>/day; however, a minimum value of 3.0 g/m<sup>2</sup>/day was registered during 1997 and a remarkable maximum value of 13.7g/m<sup>2</sup>/day during 1999 (Fig. 5).

## DISCUSSION

The results from this monitoring program are significant because they document the temporal variation of key structural and functional variables in local coral reefs, seagrass beds and mangroves over a 16 year monitoring period. Furthermore, they constitute a valuable baseline to assess the long-term dynamics of the most important marine ecosystems in the Caribbean.

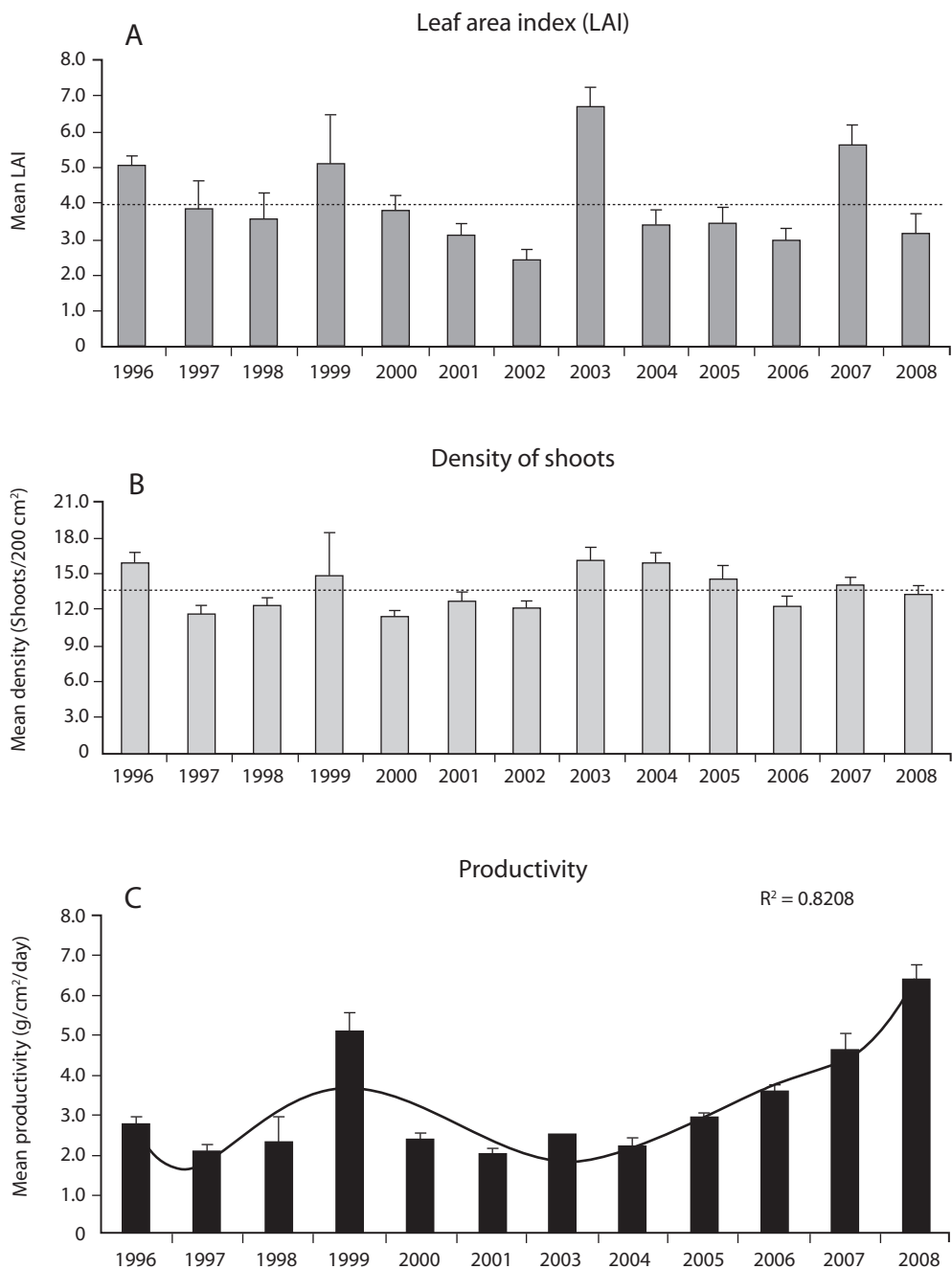


Fig. 4. Mean (+ 1 SE) for structural attributes of *Thalassia testudinum* at Chengue Bay during the monitoring period (1996-2008). (A) Mean leaf area index, overall mean indicated by dashed line; (B) Density of shoots, overall mean indicated by dashed line; and (C) Leaf productivity, fluctuation is visualized as trend line with its respective R-squared value.



TABLE 1  
*Mean values ( $\pm$  S.E.) of environmental variables assessed on Chengue Bay each September during the monitoring period (\*= no data collected)*

Year	Temperature (°C)	Salinity (PSU)	Transparency (m)
1996	29.3 (0.02)	35.5 (0.50)	7.0 (1.60)
1997	28.7 (0.02)	36.3 (0.48)	7.3 (1.40)
1998	30.5 (0.03)	32.9 (0.90)	4.9 (0.50)
1999	*	33.8 (2.00)	4.0 (0.50)
2000	29.4 (0.03)	36.2 (0.40)	5.1 (1.00)
2001	29.1 (0.03)	37.0 (0.90)	7.1 (0.90)
2002	29.7 (0.03)	36.5 (0.30)	7.6 (1.30)
2003	29.4 (0.02)	37.6 (0.90)	5.8 (1.90)
2004	*	36.5 (0.40)	6.2 (1.20)
2005	29.3 (0.03)	35.0 (0.70)	4.9 (1.10)
2006	29.1 (0.02)	37.8 (0.30)	5.0 (0.00)
2007	28.9 (0.03)	35.5 (0.50)	4.0 (1.00)
2008	29.3 (0.04)	34.5 (0.50)	9.6 (2.40)

TABLE 2  
*Comparison of the Holdridge complexity index of the mangrove forest from Chengue Bay, CARICOMP site, with respect to other forest sampled in the Colombian Caribbean*

Area	Station	HCI	Physiographic type	Source
Alta Guajira	Bahía Portete	5.20	Riverine	Sánchez-Paez & Álvarez-Leon (1997)
Alta Guajira	Bahía Tukakas	2.58	Fringe	Sánchez-Paez & Álvarez-Leon (1997)
Baja Guajira	La Enea	7.17	Basin	Sánchez-Paez & Álvarez-Leon (1997)
Baja Guajira	Playa de Holandeses	2.34	Fringe	Sánchez-Paez & Álvarez-Leon (1997)
CGSM	Rinconada	6.50	Basin	CGSM 2009 (unpublished data)
CGSM	Caño Clarín 1	6.70	Riverine-Basin	Sánchez-Paez & Álvarez-Leon (1997)
CGSM	Los Micos	2.50	Fringe-Basin	Sánchez-Paez & Álvarez-Leon (1997)
CGSM	Punta Blanca	2.58	Riverine	Sánchez-Paez & Álvarez-Leon (1997)
CGSM	Mengajito	2.47	Fringe	Sánchez-Paez & Álvarez-Leon (1997)
PNNT	Chengue	2.95	Fringe	Sánchez-Paez & Álvarez-Leon (1997)
PNNT	Cinto	5.21	Riverine	Sánchez-Paez & Álvarez-Leon (1997)
PNN I. Salamanca	Caño Loro	3.76	Riverine-Basin	Sánchez-Paez & Álvarez-Leon (1997)
Ciénaga La Virgen	Marlinda	0.08	Fringe	Sánchez-Paez & Álvarez-Leon (1997)
PNNCR y SB	Isla Rosario	3.03	Fringe-Basin	Sánchez-Paez & Álvarez-Leon (1997)
Bolívar	Bahía de Barbacoas	4.32	Fringe-Basin	Sánchez-Paez & Álvarez-Leon (1997)
Golfo Morrosquillo	Ciénaga La Caimanera	6.46	Fringe-Basin	Sánchez-Paez & Álvarez-Leon (1997)
Golfo Morrosquillo	Ciénaga de Pablo	5.00	Fringe-Basin	Sánchez-Paez & Álvarez-Leon (1997)
Golfo Morrosquillo	Caño Lobo	6.07	Riverine-Basin	Sánchez-Paez & Álvarez-Leon (1997)
Golfo de Urabá	Bocas de Atrato	1.96	Fringe	Sánchez-Paez & Álvarez-Leon (1997)
PNNT	Chengue	1.07	Fringe	This study

HCI=Holdridge complexity index.

CGSM=Ciénaga Grande de Santa Marta.

PNNT=Parque Nacional Natural Tayrona.

PNN I. Salamanca=Parque Nacional Natural Isla de Salamanca.

PNNCR y SB=Parque Nacional Natural Corales de Rosario y San Bernardo.

TABLE 3

Mean and standard error (SE) of each of the structural attributes evaluated and interstitial salinity in the CARICOMP *Rhizophora mangle* site for each year of monitoring at Chengue Bay

Year	Prop root		Trunk		DBH (cm)		Basal		Biomass <sup>1</sup>		Density		Interstitial	
	Lenght (m)		Lenght (m)		DBH (cm)		Area (m <sup>2</sup> /ha)		(kg/m <sup>2</sup> )		(Ind/100m <sup>2</sup> )		Salinity (PSU)	
	Mean	SE	Mean	SE	Mean	SE	Mean	SE	Mean	SE	Mean	SE	Mean	SE
1995	1.59	0.11	3.82	0.41	11.6	0.44	42.6	3.6	13.5	0.48	34.7	2.4	40.2	2.7
1996	1.60	0.11	3.87	0.44	11.7	0.43	43.4	3.8	13.5	0.52	34.0	2.1	40.3	1.6
1997	1.61	0.11	3.90	0.44	11.8	0.43	42.3	3.0	13.2	0.68	33.3	2.7	43.9	0.9
1998	1.61	0.11	3.56	0.09	12.1	0.42	43.8	2.7	13.4	0.61	33.3	2.7	38.5	3.2
1999	1.56	0.10	3.56	0.09	12.0	0.44	43.9	3.2	13.4	0.68	33.3	2.7	35.0	3.2
2000	1.56	0.17	3.50	0.51	12.1	0.79	44.1	3.3	13.4	0.68	33.3	2.7	39.9	2.1
2001	1.55	0.17	3.40	0.24	13.5	1.16	44.1	3.6	13.2	0.81	33.3	2.7	35.3	3.9
2002	1.57	0.17	3.44	0.47	12.3	0.81	44.7	3.4	13.3	0.80	31.7	2.4	38.3	3.8
2003	1.60	0.18	3.50	0.45	12.4	0.80	44.8	3.4	13.3	0.80	33.7	3.0	41.8	0.3
2004	1.60	0.17	3.50	0.44	12.4	0.74	42.1	3.1	12.6	0.86	30.3	1.7	40.7	1.2
2005	1.55	0.18	3.57	0.47	12.5	0.88	45.1	3.6	13.2	0.83	30.6	2.4	39.0	2.5
2006	1.54	0.20	3.52	0.49	12.9	1.05	43.7	2.9	12.6	0.57	29.5	2.5	37.1	3.1
2007	1.54	0.19	3.53	0.47	13.1	0.96	44.0	3.0	12.7	0.57	29.0	2.5	39.9	0.7
2008	1.51	0.20	3.44	0.51	12.2	0.80	45.1	3.2	12.9	0.55	29.0	2.5	39.7	2.3
OVERALL														
MEAN	1.57	0.01	3.58	0.04	12.3	0.14	43.8	0.3	13.1	0.09	32.1	0.5	39.3	0.63
95-08														

DBH=diameter at breast height.

1=according to Golley *et al.* (1962).

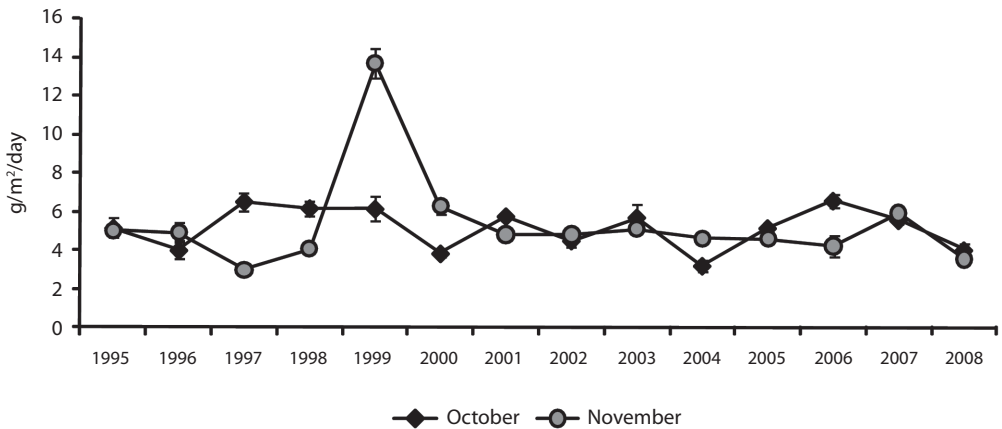


Fig. 5. Mean ( $\pm$  1 SE) of mangrove *R. mangle* productivity in terms of rate of litterfall during October and November from 1995 to 2008 Chengue Bay.

**Coral reefs:** Monitored sites in Chengue resemble many reefs across the Western Atlantic, Wider Caribbean and the Gulf of Mexico in terms of major benthic components. Overall, means of coral and algae cover are within maximum and minimum values registered on a regional scale by CARICOMP and AGRRA programs. Algal cover may range from less than 5% in Cayo Sombrero, Venezuela up to 90% in Puerto Morelos, México (CARICOMP *et al.* 2002, Linton & Fisher 2004) and coral cover may vary from 2.5% in Costa Rica to 57.8% in Flower Garden Banks, USA (Kramer 2003). Both programs have found a dominance of *Montastraea annularis* complex and algal turfs (CARICOMP *et al.* 2002, Kramer 2003).

Our findings showed that there is a pattern of stability for coral cover but there is no clear trend in algal cover at monitoring sites between 1993 and 2008. Thus coral cover has remained fairly unchanged during the last 16 years. Previous temporal analyses of coral cover at Chengue found a similar trend between 1993 and 2000 (CARICOMP *et al.* 2002, Rodríguez-Ramírez & Garzón Ferreira 2003). Although this appears to conflict with two meta-analyses pointing out significant trends of decline in the Caribbean region, in particular the loss of coral cover and/or increase in algae abundance (Gardner *et al.* 2003, Côté *et al.* 2005), recent studies on coral reef dynamics in the Caribbean and Florida Keys have documented similar patterns of stability in coral cover at local scales (CARICOMP *et al.* 2002, Roggers & Miller 2006, Edmunds & Elahi, 2007, Somerfield *et al.* 2008, Rodríguez-Ramírez *et al.* 2010). The minor reduction registered in coral cover was due to Hurricane Lenny, which touched marginally the north coast of South America by November 14-16, 1999 (Rodríguez-Ramírez & Garzón-Ferreira 2003). However, the lack of changes in coral cover at the monitoring sites does not mean that these reefs may not have been under a process of decline. Temporal analyses of coral species show that significant losses have occurred in other reef assemblages in Chengue (Rodríguez-Ramírez *et al.* 2010).

Although previous analysis had revealed a decreasing trend in algal cover between 1993 and 1999 for monitoring sites in Chengue (CARICOMP *et al.* 2002, Rodríguez-Ramírez & Garzón-Ferreira 2003), our results showed that there is no overall pattern of decrease or increase from 1993 to 2008 (Fig. 2B). Clearly, algal variations were not related to changes in coral cover. Instead they reflected transitions with the abiotic substrate. According to Rodríguez-Ramírez & Garzón-Ferreira (2003) this concomitant variation between algae and abiotic substrate may represent some sampling bias laying out the chain on the substratum. On the other hand, algal variations could be associated with seasonal changes because surveys have been done during the rainy season (October-December) where algal communities experience noticeable changes in composition and abundance (Díaz-Pulido & Garzón-Ferreira 2002). However further analyses are required to explain fully the relationship between algal and abiotic substrate variations.

During the study period, Chengue reefs suffered from bleaching (in 1995, 1998, 1999, 2005 and 2008) and have been affected by coral diseases. Earlier studies recognized these stress factors caused minor coral mortality (CARICOMP 1997b, Rodríguez-Ramírez & Garzón-Ferreira 2003, Gil-Agudelo *et al.* 2006, Rodríguez-Ramírez *et al.* 2008). In general, the prevalence of bleached corals remained below 4% and paled corals below 2%, which is considered normal for the Caribbean (Weil *et al.* 2006). In 2005, a major bleaching event occurred in the Caribbean region (Wilkinson & Sauter 2008), but corals in Chengue Bay showed only a slight increase in the average of bleaching and paled corals (both close to 3.5%). This event did not have an effect in this area as large as in other places, probably due to local seasonal upwelling (Rodríguez-Ramírez *et al.* 2008). Regarding coral diseases, all the most important coral diseases reported for the Caribbean (Black Band Disease, White Band Disease, Yellow Band Disease; see Garzón-Ferreira *et al.* 2001, Weil *et al.* 2006) were observed in Chengue, but the most prevalent

coral diseases were Dark Spots Disease and White Plague. The prevalence of coral diseases remained relatively low and stable during these 10 years of monitoring, usually staying below 2% and increasing to over 3% (in 1998 and 2001). Thus, the bleaching and diseases data suggest that, in comparison with other Caribbean coral reefs, Chengue reefs have exhibited an important degree of resistance to mass bleaching events and coral diseases.

Because there are no historical records about the abundance of mobile invertebrates of commercial value before the beginning of CARICOMP activities, it is not possible to assess if these organisms have suffered from overfishing or if their low density is a natural condition in Chengue. However, during the years of the monitoring program it has been possible to observe that Chengue is frequently visited by fishermen divers who capture lobsters, crabs, octopuses and other large invertebrates.

Sea urchins results are consistent with composition and density patterns found elsewhere in the Caribbean by CARICOMP (CARICOMP *et al.* 2002, Linton & Fisher 2004) and support previous findings for these invertebrates at Chengue (Rodríguez-Ramírez & Garzón-Ferreira 2003, Rodríguez-Ramírez *et al.* 2006). Although few urchins were recorded in the monitoring transects, important populations of these organisms (including *D. antillarum*) have been observed in shallow reefs of Chengue. This suggests a depth-related spatial distribution of urchins that is probably controlled by the competitive interactions with the damselfish *Stegastes planifrons*. The territories of this fish species are abundant at the depth interval where monitoring transects are located in Chengue (Santodomingo *et al.* 2002, Rodríguez-Ramírez *et al.* 2006). Herbivorous sea urchins, such as *D. antillarum* and *Echinometra viridis*, are aggressively excluded from damselfish territories (see Ceccarelli *et al.* 2001 for review).

**Seagrass beds:** Structural parameters found in this study reveal a healthy and productive

seagrass bed on the Chengue Bay monitoring site. LAI historical data showed periods of relative stability followed by peaks (Fig. 4A); this pattern could be a sign of a cyclical renewal tendency in the ecosystem, in which the lower values for some years (*e.g.* 1998, 2002 and 2006) indicate a greater number of new leaves (low leaf height and width) in a developing community. Height and width may also be indicators of stress in seagrass communities, both decreasing when plants are exposed to unfavourable environmental factors; however, considering the relative homogeneous fluctuation of LAI values during the monitoring period, stress may not be the cause of lower values in this case. Additionally, overall mean of leaf area index was high compared to other CARICOMP sites (CARICOMP 1997c, Linton & Fisher 2004). Overall mean density of shoots in Chengue was higher compared to other CARICOMP sites (Linton & Fisher 2004), and fluctuation among years remained near the mean value. These conditions could also be a sign of relative stability and healthy stands of *T. testudinum* on the Bay. Progressive increase of productivity of *T. testudinum* during the last years of monitoring reflects a healthy community developing under favourable environmental conditions. It is clear that production of seagrass beds is usually limited by resource availability (mainly nutrient concentration and light quality) and/or heavy losses caused by physical disturbances (Powell *et al.* 1989, Duarte & Chiscano 1999); according to this, it could be assumed that conditions within the Bay such as nutrient inputs from adjacent mangroves and from seasonal upwelling waters, low exposure to river discharges, and low human intervention, favour the productivity potential of *T. testudinum* beds. Our observations of productivity are limited to the above ground vegetative expansion, but wider quantitative measures including growth of the underground rhizomes and roots should be carried out to reflect other aspects of plant expansion and maturation (Short & Coles 2001).

Since no significant correlation was found between environmental data (temperature,

water transparency and salinity) and biological parameters of *T. testudinum* beds (density of shoots, net above-ground productivity and LAI), it is assumed that fluctuations detected for some structural parameters are better explained by cyclical renewal tendencies more than a response of the community to inter-annual environmental changes. However samplings along the year are required to establish potential relationships between environmental variables and biological data because our results are constrained to a single period during the year.

Monitoring results may characterize the seagrass beds on Chengue Bay as a lush, healthy and productive ecosystem on the Colombian Caribbean. Its inclusion in a legally protected area with low human intervention, moderate continental influx, seasonal upwelling and its proximity to productive mangroves and coral reefs, are special characteristics that must favour the current condition of these seagrass beds.

**Mangrove:** The Red mangrove forest in Chengue Bay has a low structural complexity compared with other mangrove forests along

the Colombian Caribbean (Table 2). Relatively small tree sizes, poorly defined wood, very intricate and branched forest (Garzón-Ferreira 1998) are essential characteristics of the site. These features are related to deficiencies in water flow, high salt content and soil conditions (Table 3). In contrast with the mean structural attributes of some of these variables in other mangrove forests in the Caribbean (Table 4), most of the data in this study are close to the highest values reported for other CARICOMP network locations. The results showed that no major structural changes have occurred since 1995 (Table 3). However, variations in biomass, basal area and density, were due mainly to the death of 18% of trees assessed inside the plots. This may have been caused by insect damage (Rodríguez-Ramírez & Garzon-Ferreira 2003). Attacks from xylophagous insects and other pests of some trees are part of the natural mangroves cycles that do not have a major influence on mangrove populations (Villalobos *et al.* 1996). Human intervention could not have affected directly the structural development of the forest, considering that no evidence of human disturbance was observed on the sample plots. At global level, the mangrove ecosystem

TABLE 4  
Comparison of the red mangrove (*R. mangle*) forest structure and productivity (litter fall) at different CARICOMP sites in the Caribbean

Country	Basal Area (m <sup>2</sup> /ha)	Biomass <sup>1</sup> (kg/m <sup>2</sup> )	Density (Ind/100m <sup>2</sup> )	Litter fall rate (g/m <sup>2</sup> /day)	Source
	Mean	Mean	Mean	Mean	
Barbados	33.5	10.0	23.0		CARICOMP (1997d), Linton & Fisher (2004)
Bermuda	28.4	10.6	40.0	1.6	CARICOMP (1997d), Linton & Fisher (2004)
Belize	36.8	15.7	56.3	2.5	CARICOMP (1997d), Linton & Fisher (2004)
Costa Rica	22.9	4.6	9.3	4.0	Fonseca <i>et al.</i> (2007)
Jamaica	13.2		49.5	1.3	Juman (2005), Linton & Fisher (2004)
Mexico	18.6	8.1	28.2		CARICOMP (1997d)
Panamá	12.3	4.6	59.3	2.7	Guzmán <i>et al.</i> (2005)
Puerto Rico	20.4	8.3	23.7	4.1	Juman (2005), Linton & Fisher (2004)
Tobago	24.5	14.1	26.3	3.6	Juman (2005), Linton & Fisher (2004)
Venezuela	24.2	12.6	46.2	4.3	Juman (2005), Linton & Fisher (2004)
Colombia (Chengue)	43.8	13.1	32.1	5.2	This study

1=according to Golley *et al.* (1962).

is severely threatened and has lost at least 35% of the area of mangrove forests in the past two decades. This is due partly to global changes, but mainly to human activities such as construction, mariculture and agriculture (Valiela *et al.* 2001, Halpern *et al.* 2007). In this context, it is suggested that the mangrove in Chengue differs from this scenario because it has not undergone major disturbances over the period of the study, with some alterations primarily influenced by natural degradation factors.

The cause of the decline in productivity (litterfall) recorded in November 1997 is unknown, but the high values obtained during November 1999 were related to the rains and strong winds associated with hurricane Lenny (Rodríguez-Ramírez & Garzon-Ferreira 2003). Such factors may lead to defoliation, fragmentation and the downfall of the wood of the mangrove trees (Cintrón & Schaeffer-Novelli 1983, Salazar-Vallejo 2002, Rodríguez-Ramírez & Reyes-Nivia 2008). Compared to other CARI-COMP areas in the Caribbean, Chengue is one of the most productive mangroves in terms of litterfall (Table 4). This condition is related to the interstitial salinity (overall mean 39.3, Table 3), which is higher than the optimum range mentioned by Prah (1990) (15 and 25) and a consequence of the fresh water deficit that causes defoliation. Moreover, October and November are rainy months, which promote litterfall and their transport out of the system. The high rate of litter production indicates that the mangrove is providing a significant input of energy and nutrients to adjacent ecosystems (seagrass beds and coral reef) within the bay.

The results of this study are meaningful because they illustrate a remarkable stability of the study sites at Chengue Bay contrasting with the recent context of decline at regional and global scales for marine ecosystems (Valiela *et al.* 2001, Gardner *et al.* 2003, Lotze *et al.* 2006, Waycott *et al.* 2009). Although an analysis of the global network of Marine Protected Areas (MPAs) showed that management is very limited for the Tayrona Park -including Chengue bay- (Mora *et al.* 2006), our outcomes suggest this park can preserve

coastal marine communities as long as they are subjected to moderate levels of natural and anthropogenic disturbances. Given that human populations are largely dependent on marine ecosystems, the loss of these habitats and their associated biodiversity may have enormous implications for human societies around the world. Therefore, Chengue bay appears to be an essential refuge of biodiversity in the Colombian Caribbean and may have an important role in terms of resistance and resilience to the effects of climate change.

In summary, we did not find evidence of decline trends in the coral reef, seagrass and mangrove sites at Chengue over the monitoring period. Annual variations in coral reefs and mangroves were related to the effects of an isolated event, namely Hurricane Lenny. The observed patterns in seagrasses appear to characterize the natural variability of cyclical renewal processes. Therefore, our results suggest that monitoring sites at the three ecosystems have had a healthy development in the last 16 years. This information is a unique support for management and conservation purposes at Tayrona Natural Park. As Chengue ecosystems can be critical for ecological processes and also tourism benefits in the region, an effective enforcement on the marine ecosystems inside the Tayrona Park is required.

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## RESUMEN

Pocos programas de monitoreo han estudiado simultáneamente la dinámica de ecosistemas marinos estrechamente relacionados (arrecifes coralinos, pastos marinos y manglares) para documentar su variabilidad espacial y temporal. Siguiendo el protocolo de monitoreo del programa CARICOMP, estaciones permanentes de monitoreo en estos ecosistemas fueron evaluadas entre 1993 y 2008 en la Bahía de Chengue del Parque Nacional Natural Tayrona (Caribe Colombiano). En general los ecosistemas monitoreados han presentado un patrón de estabilidad durante los años de estudio. Mientras los arrecifes coralinos (cobertura de coral) y manglares tuvieron algunas variaciones anuales debidas al paso del huracán Lenny en 1999, los pastos marinos registraron tendencias particulares de cambio (índice de área foliar y productividad de hojas) que podrían estar reflejando la variabilidad natural de la pradera estudiada. Por lo tanto se sugiere que los sitios monitoreados en cada ecosistema han tenido un desarrollo saludable en los últimos 16 años. Estos resultados son importantes para mejorar localmente las estrategias de manejo (Parque Nacional Natural Tayrona) y para evaluar la dinámica a largo plazo en los ecosistemas marinos del Caribe.

**Palabras clave:** Monitoreo, arrecifes coralinos, pastos marinos, manglares, Chengue, CARICOMP, Caribe colombiano.

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