Anthropogenic stressors and eutrophication processes as recorded by stable isotopes compositions in coral skeletons

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Abstract

The northern Gulf of Aqaba, in the northeastern branch of the Red Sea, is a clear example of humans effecting on the health of fringing reefs. Our results demonstrate the effect of an anthropogenic stressor on the carbon and oxygen stable isotopes compositions, namely net pen fish farming with annual fish production of $2.4 \times 10^6$ kg yr$^{-1}$. We sampled and studied long coral cores from stressed, remote and intermediate localities and short-term transplanted Porites sp. colonies from the west side of the Gulf of Aqaba to a remote and a polluted sites, respectively. The data shows that mariculture and other human related stressors did not influence the oxygen isotopic signature over a period of two decades. However, the carbon fractionation changed along a geographical gradient and depended on proximity to the source of contamination. We suggest that $\delta^{13}$C of coral skeleton is a promising proxy for identifying long term processes of coral growth under high nutrient loads and potential disturbances to the coral reef ecology.

1 Introduction

Coral reefs dominated tropical oceans over 200 million years and flourished especially during interglacial periods. Scleractinian (reef building), coral reefs can flourish in the nutrient-poor (Darwin, 1842; Odum and Odum, 1955) environments due to their mutualistic symbiosis with dinoflagellates (Symbiodinium spp.), known as zooxanthellae. Their carbon and energy requirements are met by different species-specific combinations of algal photosynthetic products, as well as by predation on zooplankton, supplemented in some cases by minor contributions of dissolved organic carbon compounds and bacteria (Muscatine et al., 1989). Scleractinia, a major member of phylum Cnidaria, forms the framework of coral reef communities, in which they from massive structures that monopolize coastal zones in tropical seas (Allemand et al., 2004). During the last two decades coral reefs have undergone an increase in various
anthropogenic environmental disturbances (Goreau et al., 2000). These have placed unprecedented stresses on coral reefs, primarily through declining water quality, pollution, overexploitation, and more recently by changes to the atmospheric concentration of greenhouse gases through ocean warming and acidification (Hoegh-Guldberg et al., 2007). On the local scale, pollution by human activities leads mainly to eutrophication and siltation, exposing many fringing and offshore reefs to severe stress (Brown, 1997; Loya, 2007). Corals provide record of the chemical and physical conditions that existed in the local seawater at the time of accretion of their calcium carbonate skeletons (Mcconnaughey, 2003; Allemand et al., 2004; Meibom et al., 2007). Due to their high rate of growth and accretion of solid calcium carbonate, corals serve as an oceanic recorder with enhanced time resolution (monthly to seasonal) (Mcculloch et al., 1999; Cohen et al., 2001; Felis et al., 2003). Coral skeletal variations in stable isotopes, such as oxygen and carbon, reflect surrounding seawater temperatures, evaporation, precipitation, light intensity, and coral metabolic activity but in a complex and non-linear way (Levy et al., 2006; Rosenfeld et al., 2006).

A particularly relevant site for investigating reef health is the rich coral reefs in the Israel port city of Eilat, situated at the northern edge of the Gulf of Aqaba, Red Sea. Although coral ecosystems are usually affected by one or another stress-producing element, the Eilat reefs provide a convenient laboratory for studying damage, for they have been subjected to multiple natural and man-made disturbances.

Although the coral reefs at Eilat (Red Sea, 29° N) are situated near the northern limit of reef building by stony (scleractinian) corals, they are highly diverse (Loya, 1972) and characterized by a high degree of endemism (Fishelson, 1995). They have been exposed to various natural and anthropogenic disturbances (Loya, 2007). The shallow reefs are affected by occasional winter storms that cause large surge (Friedman, 1968), which can dislodge a considerable number of corals (Wielgus and Levy, 2006). Anthropogenic stresses from intensive tourism activity and poor water quality from sewage pollution, flood waters, ballast and bilge water from boat activity and discharges of fuel, oil, detergents, and phosphates have all damaged the reefs. In the last several
years, water clarity has deteriorated, along with the symbiont photosynthesis and coral calcification (Atkinson et al., 2001). Also, in the last 10 years, mariculture effluents of $2.4 \times 10^6$ kg yr$^{-1}$ of in situ net-pen fish cages (mainly *Sparus aurata*, a non indigenous species in the Gulf of Aqaba (Eilat) were deployed in the northeast part of the Gulf. These facilities discharge organic matter ($4 \times 10^4$ kg of phosphorous and $2.4 \times 10^5$ kg of nitrogen) and particulate matter into the water column (Loya et al., 2004). The fish farms have “replaced” the urban sewage, which stopped in 1995 because land facilities for urban sewage treatment were developed (Loya et al., 2004). There is a general consensus that Eilat's scleractinian corals have suffered in recent years, characterized by 70% loss of abundance, over 50% decrease in coral cover, low rates of larval settlement and recruitment (Ben-Tzvi et al., 2004), a decrease in calcification (Silverman et al., 2004), an increase in coral mortality, macro-algal blooms smothering the corals, and the increasing percentages of corals affected by pathogenic diseases related to continuous eutrophication originating from the fish farming (Loya et al., 2004).

We undertook to measure and compare the isotopic signatures and the growth parameters formed under relatively disturbed and remote conditions of *Porites* sp. colonies which were transplanted in 2001 in the vicinity of the fish farm cages in the northern tip of the Gulf (high nutrient load, low clarity, high phytoplankton productivity (Genin and Silverman, 2005; Genin and Shaked, 2006, 2007) and on the reef across from the Interuniversity Marine Institute (IUI) “reference site” (relatively remote and undisturbed conditions). Furthermore we also undertook a historic study of gradients in the $\delta^{18}O$ and $\delta^{13}C$ isotopic records of corals of the northern part of the Gulf of Aqaba, going from north to south. Long-drilled records from three colonies were examined in order to identify and trace possible environmental stress in the region. This allowed us to obtain, for the first time, the isotopic signature of coral growth under high nutrient loads. The outcome of these environmental and “human affect” influences is of concern to the future of coral reefs found in the north tip of the Red Sea and they may be indicative of future anthropogenic patterns associated with regions which are considered to be under high human impact.
2 Materials and methods

In 2001, several single *Porites* sp. colonies were stained with alizarin red, acclimated and later transplanted from the Navy harbor (see Fig. 1) to the vicinity of the fish farm area and in front of the IUI research station (Rosenfeld et al., 2003). The corals were transplanted 20 m below the surface in order to preserve their original depth. The *Porites* corals were monitored every month by scuba diving to determine how well they survived the transplantation procedure. During 2005 five of the transplanted colonies were harvested, two from the IUI and three from the fish farms. The severe flood of winter 2006 killed the remaining corals, which were intended to serve as a long-term monitor of eutrophication. All five corals were analyzed for their isotopic compositions during the years before and after transplantation. Addition set of *Porites* sp. cores were sampled in order to obtain the modern north–south gradients from the fish farms to Taba, along with corals that were transplanted near the fish farms cages and in IUI (“reference site”). The three large colonies that have been drilled provided 20–25 years of history pattern records. All coral cores were washed; air dried, and sectioned into 7 mm thick slabs. The cores were x-rayed, with positive prints of the slabs revealing the annual density-banding patterns characteristic of massive corals, such as *Porites*. The prints were used as a guide for microdrilling along the major axis of growth and later for the chronological ordering, age modeling, of the coral bands (Linsley et al., 1999).

For high-resolution reconstructions, samples of 100–150 µg CaCO$_3$ were extracted every 0.6 mm with a dental drill (Swart, 1983). Because corals grow 8–12 mm per year, this sampling method provides bimonthly–monthly resolution. CaCO$_3$ samples for δ$^{18}$O and δ$^{13}$C analyses were reacted with 100% orthophosphoric acid and CO$_2$ was measured using a Finnigan GasBench II gas preparation unit connected on line to a Finnigan MAT 252 isotope ratio mass spectrometer. Calibration was maintained by routine analyses of internal and international standards. The long-term precision of our internal laboratory standard is 0.06‰ and 0.10‰ for carbon and oxygen, respectively. Even-spaced time series (monthly) were obtained by interpolating the raw data using
the Analyseries time series analysis tool (Paillard et al., 1996) from isotopic minima and maxima. Results are reported in per-mill units relative to the international Vienna–Peedee Belemnite Limestone Standard (V-PDB). Interpretation and correlation of the data were related to temperature, light, water currents, salinity and nutrient data which are available at high temporal resolution from the long term monitoring program in Eilat, running by the Interuniversity Institute (IUI) which was also acting in this project as the “reference site”.

3 Results

We have obtained 6–7 years of high-resolution records from each colony. The transplanted colonies at the IUI reference site exhibited no change in their $\delta^{13}$C pattern, neither in amplitude nor absolute values (Figs. 2a, b and Fig. 3a, t-test $p > 0.05$); the annual average growth rate was $9.86 \pm 1.72$ mm yr$^{-1}$. In contrast, the carbon isotopic measurements of the corals transplanted to the fish cages (FCI, FCII, and FCIII) clearly demonstrated a “high nutrient load” growth phase in the skeletons. There is a marked alteration in the pattern and amplitude of the annual $\delta^{13}$C relative to “normal” growth period prior to transplantation (t-test $p < 0.05$), (Fig. 2c–e, Fig. 3a) although average growth rates did not decline and amounted to $9.94 \pm 3.32$ mm yr$^{-1}$. Hence, we are able to identify the $\delta^{13}$C isotopic signature of a period of high nutrient load. There was no significant alteration of $\delta^{18}$O signals of both transplanted groups, IUI and FC, due to transplantation (Fig. 2f, t-test $p > 0.05$). Overall, all corals isotopic composition show a relatively constant annual water temperature throughout the past 6–7 years, and they capture the normal annual cycle of sea surface temperature (Fig. 3b). Our measurements are in good agreement with the long term in situ SST measurements carried out at the IUI peer.

North–south gradients of three cores drilled from large Porites sp. colonies at a water depth of 5–7 m were used to document the isotopic records characteristic of the last 20–25 years in the Gulf Aqaba. The first core, point A (Fig. 1), is the northernmost site.
where large *Porites* were identified in the Gulf; the second core was sampled to the south at point A, some 700 m north of the Navy harbor, near point B (Fig. 1). The third core was taken at the southernmost part of the study site, point C, (Fig. 1). Positions of the three cores allowed us to detect the possible influence of the high nutrient load plume that could have originated from the fish cages to the north. We have obtained an average resolution of 15 samples per year after assigning an absolute age to each sample, which allows an accurate reconstruction, with monthly details, of the annual cycles of oxygen and carbon isotopes. The average results of $\delta^{18}O$ from the three cores does not exhibit significant long-term annual trend, implying that temperature, evaporation and salinity remained constant during the past two decades (Fig. 4a, one-way ANOVA, $p > 0.05$). The $\delta^{13}C$ values (Fig. 4b) show significant differences between the south cores, B and C, and the north core A (one-way ANOVA, $p < 0.05$). B and C core samples do not exhibit a long-term trend and the annual variability is very similar over the past 25 years, with annual growth ranging 12.2 and 10.28 mm yr$^{-1}$, respectively. This suggests that the physiological parameters of the corals including skeletal growth did not significantly change during this period and that they were probably exposed to the same light levels, nutrient loads, and trophic activity throughout the entire period. In contrast, carbon isotopic record from core A, which is the closest site to the fish cages and the northern hotel line, exhibited an annual growth of 9.28 mm yr$^{-1}$ (Fig. 4b). Before 1994, the signal is similar to the patterns presented by all other corals. However, after 1994 the signal is characterized by large anomalous annual amplitudes, as well as an overall shift towards heavy $\delta^{13}C$ values (one-way ANOVA, followed by Tukey HSD test, $p < 0.05$).

4 Discussion

The results presented here imply that the carbon signature is more sensitive to environmental stress than the annual changes in the skeletal growth. In fact, this change resembles the pattern in the $\delta^{13}C$ record of the corals transplanted to the fish-cage
area and is similar to the pattern of high heterotrophy caused by increased levels of zooplankton (Grottoli, 2002). Thus, the corals in the vicinity of the northern part of the Gulf themselves are witnesses to increasingly higher nutrient load and discharge, which is known to have risen rapidly since 1994 (2 × 10⁵ kg yr⁻¹ of fish in 1991 to annual 4.5 × 10⁵ kg yr⁻¹ in 1994, Fig. 5) with maximum production of 2400 tons in 2001 (Atkinson et al., 2001). However this influ- ence was not observed in the carbon isotope measurements of the southern cores (B and C), ~2.5, 8 km south-west to the fish cages. There are two ways of explaining this observation: (a) the excess nutrients from the north are diluted over 2.5 km to a level that does not affect coral growth at depth of about 7 m or (b) the excess nutrients move eastward or to the center of the Gulf without reaching coastal areas to the southwest. However, the prevailing winds and the surface current regime do not support eastward diversion of the surface water. Furthermore our study shows unequivocally that Porites sp. colonies were affected at the site close to the fish farming compared to those growing to the southwest, such as sites B, IUI, and C. The impact of Gulf eutrophication seems to be local (scale of 2 km), but with time chronic nutrient enrichment may affect a larger reef area south of point A. Several years of monthly monitoring light intensity did not reveal any significant change among 8 coastal sampling sites along the Israeli coast, including the FC (Genin and Shaked, 2008, Fig. C11). Hence, we rule out the possibility that light may have affected the carbon signals (Grottoli, 2002). Global climatic factors may interact with local stresses to hasten the decline of marine communities and the degradation of coral reefs around the planet; coastal marine eutrophication has drastic consequences on corals reefs, in particular. Therefore using carbon isotopes in coral skeleton will be important in assessing anthropogenic stressors in coral reefs and associated ecosystems, and can be used to distinguish between local and global stresses. In addition, δ¹³C of long coral skeletal records can be used to identify past periods of high nutrient loads.

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References


Grottoli, A. G.: Effect of light and brine shrimp on skeletal delta C-13 in the Hawaiian coral...


Fig. 1. The study site at the northern tip of Eilat, Gulf of Aqaba. In red: long cores, drilled from un-manipulated coral colonies at the northern and southern part of the study area (5–7 m depth). In blue: coral colonies transplanted from the navy base to under the fish cages and to the reference site in front of the IUI laboratory (20 m depth). Core A is located close to the north beach in front of the hotels line (29°32’ 54.88” N, 34°57’ 25.54” E), core B is close to the navy harbor (29°32’ 50.20” N, 34°57’ 16.45” E), core C is located, southwest of the fish farms (29°29’ 48.62” N, 34°54’ 41.81” E).
Fig. 2. $\delta^{13}$C and $\delta^{18}$O compositions of the transplanted coral colonies. The gray line represents the alizarin stain line imprinted in the skeleton in January 2001 during the transplantation procedure. (a, b) Corals transplanted to the IUI reference site; (c–e) corals transplanted near the fish farm area; (f) average skeletal $\delta^{18}$O since 1994 (black line) of transplanted corals (FC and IUI) compared with monthly SSTs (top black line), ($n = 5$ colonies), (black line represents the mean value).
Fig. 3. Centered values (mean ± SD) of oxygen and carbon isotopes from IUI and FC colonies before and after transplantation in 2001, alizarin stain line is marked in gray. (a, b) Top graphs represents the FC (c, d) represents IUI.
Fig. 4. Monthly variability in skeletal δ¹⁸O and δ¹³C averaged over three long *Porites* sp. coral cores retrieved from north to south in the Gulf of Aqaba, Eilat. (a) δ¹⁸O (mean ± SD) since 1985 compared with monthly SSTs (gray line). The correlation ($r^2 = 0.74$) between SST and δ¹⁸O after averaging the three corals since 1985. The slope is $-0.116\%$ per °C. (b) Monthly anomalies of δ¹³C from core C: core B; core A, coral cores (black line representing moving average 5 points).
Fig. 5. Net feeding and nutrient discharge by the fish farming over the years calculating from the feeding versus the yield growth of the fishes in kg. Open triangles represent feed in kg, open cycles represent yield in kg, and open squares represent discharge in kg.