NOTE

Fluctuations in algal chlorophyll and carotenoid pigments during solar bleaching in the coral *Goniastrea aspera* at Phuket, Thailand

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ABSTRACT: HPLC analysis of pigments of the symbiotic algae of the coral *Goniastrea aspera* (Verrill) showed the presence of the chlorophylls a and c₂ (chl a and c₂) and the carotenoids peridinin, diadinoxanthin, diatoxanthin and β-carotene. Increased production of diatoxanthin was seen in algae extracted from the western surfaces of corals, exposed to high levels of solar radiation, during a solar bleaching event. In addition, an increase in the ratio of diatoxanthin to the total xanthophyll pool was observed in bleached tissues, together with a progressive increase in the total xanthophyll pool (relative to total chlorophyll levels) over the bleaching period. These results support a potential photoprotective function for xanthophylls in the coral-algal symbiosis. Chlorophyll breakdown products, phaeophytin a and pyrophaeophytin a, were recognised in considerable quantities only in bleached tissues. Computation of a simple chlorophyll budget showed that 45% of chl a was lost in partially bleached tissues and approximately 62% in fully bleached tissues, indicating a very rapid photodegradation of chlorophyll 24 h after exposure to high irradiance.

KEY WORDS: Symbiotic algae · Pigments · Xanthophylls · Coral · Bleaching

Coral bleaching is characterised by a loss of endosymbiotic algae, loss of algal pigmentation or a combination of the 2 processes. Pigment changes have frequently been monitored in bleached corals and are usually quantified as alterations in algal chlorophyll content (Hoegh-Guldberg & Smith 1989, Szmant & Gassman 1990, Fitt et al. 1993). Fluctuations in pigments other than chlorophylls have rarely been analysed in bleached corals. One exception is the work of Kleppel et al. (1989) who showed significant reductions in algal chlorophyll c₂ (chl c₂) content and in concentrations of the carotenoids peridinin and diadinoxanthin in algae extracted from bleached *Montastrea annularis*. A decrease in algal chlorophyll content was also reported by Fang et al. (1995) during bleaching induced by exposure of corals to reduced salinity in laboratory studies. No significant changes in algal carotenoid concentrations were noted in these experiments.

In the field, predictable solar bleaching events, which result in the formation of localised bleaching areas on coral colonies subject to elevated solar irradiance (Brown et al. 1994), have allowed changes in algal chlorophyll to be monitored by spectrophotometry before, during, and after the bleaching event (Le Tissier & Brown 1996). In the present paper, changes in algal chlorophyll and carotenoid composition, and measurement of chlorophyll breakdown products have been examined, using HPLC, in the coral *Goniastrea aspera* during a period of solar bleaching. This type of stress results in bleached areas (solar lesions) on the western surfaces of corals as a result of afternoon subaerial exposure to high solar irradiance (Brown et al. 1994).

Materials and methods. Study site and field sampling: The study site at Phuket, Thailand (7° 50'N, 98° 25.5'E), factors responsible for solar bleaching and the temporal progression of bleaching in corals at the site have been described in detail by Brown et al. (1994) and Le Tissier & Brown (1996). In 1996 a major solar bleaching event was predicted to occur on the spring tide of 17 to 19 February. Coral cores were sampled on 17 February and on 2 consecutive days following initial bleaching. No bleaching was evident.
on 17 February but partially bleached (pale colour) areas were obvious on the western sides of colonies on 18 February and fully bleached (white) solar lesions on 19 February. Cores were extracted by methods described in Le Tissier & Brown (1996), from normally coloured tissues on the east and west sides of 5 coral colonies on 17 and 19 February and from west sides only on 18 February. Additional cores were collected from partially bleached areas and normally coloured tissues on the west sides of 7 colonies on 18 February, and from fully bleached areas and normally coloured tissues on the west sides of 5 coral colonies on 19 February. Coral cores were transported back to the laboratory on ice in the dark and were stored in a freezer (−20°C) before processing.

Preparation of samples for HPLC analysis: Cores were air dried and the tissues collected and homogenised in 10 mM Tris-HCl and 1% NaCl (Sharp et al. 1994) by means of a tissue tearer (Ultra-Turrax T25) at 13 500 rpm. The skeleton was allowed to settle and the supernatant centrifuged at 600 × g for 10 min. The resulting algal pellets were collected and maintained in liquid nitrogen prior to transportation to the UK. Before HPLC analysis the pellets were homogenised in buffer and subsamples extracted for algal counts; the remaining cells were centrifuged for 5 min at 1800 × g and the resulting pellets extracted in 90% acetone.

HPLC analysis: Frozen samples were extracted in 90% acetone and analysed for pigments by reversed phase HPLC on a Pecosphere C-18 column according to Barlow et al. (1993b, 1995). Aliquots of extracts were mixed 1:1 with ammonium acetate buffer and injected into a Shimadzu HPLC system. Two solvents, A and B, were used in the separation. Solvent A consisted of 80% methanol and 20% 1 M ammonium acetate, and solvent B contained 60% methanol and 40% acetone. Pigments were separated by a linear gradient from 0% B (100% A) to 100% B (0% A) over 10 min followed by an isocratic hold at 100% B for 7.5 min at a flow rate of 1 ml min⁻¹. Chlorophylls and carotenoids were monitored by absorbance detection at 440 nm. while phaeopigments were detected by fluorescence at 405 nm excitation and 670 nm emission. Pigment identification and calibration were performed with standards obtained from the Water Quality Institute, Denmark, and pigments isolated from microalgal species in the Plymouth Culture Collection (Barlow et al. 1993b, 1995). Peak identity was further confirmed on selected samples by on-line diode-array spectroscopy (Waters 990). Chl a and b standards (Sigma Chemical Co.) were quantified spectrophotometrically using the extinction coefficients of Jeffrey & Humphrey (1975).

Statistical analysis: Paired t-tests were used for comparing the amount of algal pigments in east and west paired samples and between normally coloured and bleached pairs of cores collected from the same colonies on 18 and 19 February, following the procedure of Zar (1984). On 19 February, the α-value was adjusted for the number of multiple comparisons made. The differences computed between paired samples were derived from a normally distributed population of differences and so the assumption for this statistical test was satisfied.

Results. Algal pigment concentrations identified by HPLC are summarised in Table 1. Chl a and c₂, the carotenoids peridinin, diadinoxanthin, diatoxanthin and β-carotene and the chlorophyll breakdown products phaeophytin a and pyropheophytin a were recognised in all samples. There were no significant differences in the pigment concentrations of algae from normally coloured tissues extracted from east and west sides of colonies on 17 and 19 February, apart from diatoxanthin which showed significant differences in concentration between east and west sides on 17 February (p < 0.05). The mean values of diatoxanthin (Table 1) indicated an increase on the west sides in comparison

Table 1. Concentrations of pigments in pg cell⁻¹ (standard deviation in parentheses) in samples collected during the bleaching event (17 to 19 February 1996) from east (E) and west (W) sides for normally coloured samples, and from west sides for partially bleached (W-P.bl.) and bleached samples (W-Bl.).

<table>
<thead>
<tr>
<th>Pigment</th>
<th>17 Feb W (n = 5)</th>
<th>18 Feb W (n = 7)</th>
<th>18 Feb W-P.bl. (n = 7)</th>
<th>19 Feb W (n = 5)</th>
<th>19 Feb W (n = 5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chlorophyll a</td>
<td>5.46 (1.42)</td>
<td>4.78 (2.15)</td>
<td>4.87 (1.24)</td>
<td>5.37 (1.39)</td>
<td>4.23 (1.84)</td>
</tr>
<tr>
<td>Chlorophyll c₂</td>
<td>1.17 (0.29)</td>
<td>1.02 (0.44)</td>
<td>0.90 (0.32)</td>
<td>1.43 (0.44)</td>
<td>0.92 (0.39)</td>
</tr>
<tr>
<td>Peridinin</td>
<td>3.06 (0.84)</td>
<td>2.64 (1.12)</td>
<td>1.56 (0.92)</td>
<td>2.50 (0.93)</td>
<td>0.62 (0.49)</td>
</tr>
<tr>
<td>Diadinoxanthin (Dd)</td>
<td>1.15 (0.32)</td>
<td>0.94 (0.40)</td>
<td>1.25 (0.29)</td>
<td>0.65 (0.38)</td>
<td>0.31 (0.22)</td>
</tr>
<tr>
<td>Diatoxanthin (Dt)</td>
<td>0.17 (0.04)</td>
<td>0.28 (0.13)</td>
<td>0.15 (0.04)</td>
<td>0.26 (0.04)</td>
<td>0.18 (0.12)</td>
</tr>
<tr>
<td>β-Carotene</td>
<td>0.12 (0.03)</td>
<td>0.12 (0.05)</td>
<td>0.15 (0.04)</td>
<td>0.12 (0.05)</td>
<td>0.08 (0.04)</td>
</tr>
<tr>
<td>Dt/(Dd+Dt)</td>
<td>0.13 (0.03)</td>
<td>0.23 (0.05)</td>
<td>0.37 (0.08)</td>
<td>0.11 (0.00)</td>
<td>0.25 (0.08)</td>
</tr>
<tr>
<td>Phaeophytin a</td>
<td>0.03 (0.02)</td>
<td>0.09 (0.07)</td>
<td>0.09 (0.06)</td>
<td>0.05 (0.03)</td>
<td>0.13 (0.13)</td>
</tr>
<tr>
<td>Pyropheophytin a</td>
<td>0.01 (0.01)</td>
<td>0.02 (0.02)</td>
<td>0.02 (0.01)</td>
<td>0.04 (0.04)</td>
<td>0.15 (0.05)</td>
</tr>
</tbody>
</table>
to the east sides. The ratio of diatoxanthin to the total xanthophyll pool (i.e. diatoxanthin plus diadinoxanthin) was also significantly different ($p < 0.005$) between east and west sides of colonies sampled on both 17 and 19 February, with west sides showing higher ratios compared to east sides by approximately 2-fold.

On subsequent days (18 and 19 February), significant differences in chlorophyll pigments between normally coloured tissues and partially bleached ($p < 0.05$) and fully bleached ($p < 0.01$) tissues on the west sides of colonies were noted. Both partially bleached and fully bleached tissues contained algae with decreased chlorophyll pigment values compared to normally coloured tissues. In parallel with these alterations, an increase in the concentration of chlorophyll breakdown products in partially bleached and bleached areas (compared to normally coloured tissues on the western sides of colonies) was evident. A simple chlorophyll budget was calculated (after Barlow et al. 1993a) by using estimates of the molar ratio of chlorophyll and its breakdown products for partially bleached (18 February) and fully bleached samples (19 February) compared with normally coloured tissues extracted from western surfaces on these dates (Fig. 1). Approximately 45% of chl a was lost, presumably as a colourless residue, in partially bleached tissues with approximately 62% lost in fully bleached tissues. In these latter samples, 10% chl a was converted into phaeophytin a and 4% into pyrophaeophytin a.

Comparison of algal carotenoid concentrations between bleached tissues and normally coloured tissues revealed decreased mean values for peridinin ($p < 0.01$) in bleached samples. Increases in the ratio of diatoxanthin to the total xanthophyll pool were apparent in partially bleached ($p < 0.001$), but not in fully bleached, tissues. No significant differences in the concentration of $\beta$-carotene in different tissues were observed. The carotenoid composition of partially bleached and fully bleached tissues from the west sides of colonies is compared with that of normally coloured coral samples in Fig. 2. The major differences between the samples relate to the size of the xanthophyll pool components, diadinoxanthin and diatoxanthin, which are greatest in the fully bleached tissues sampled on 19 February. $\beta$-Carotene is also present in greater proportions in fully bleached tissues than in normally coloured and partially bleached tissues.

**Discussion.** Significant reductions in algal chl a and c$_2$ and the light-harvesting accessory pigment peridinin were observed in partially and fully bleached tissues following solar bleaching, when compared with normally coloured tissues from the western surfaces of colonies. Previous studies of bleached corals from the Caribbean, by HPLC, detected reduced chl c$_2$ and peridinin levels in algae but no attempts were made to analyse chlorophyll breakdown products (Kleppel et al. 1989). The present study is the first to identify chlorophyll breakdown products in symbiotic algae of bleached coral tissues. Phaeophytin a is produced *in vitro* by acidification (Lorenzen 1967) while enzymatically mediated pyrolysis of phaeophytin a may be responsible for the production of pyrophaeophytin a (Barlow et al. 1993a). The mechanism for specific production of phaeopigments *in vivo* remains unknown.
The degradation of chl a and the resulting formation of breakdown products under conditions of high irradiance has been well documented in higher plants (Brown et al. 1991) but the mechanism of chlorophyll breakdown in bleached corals is unknown. Although the majority of coral bleaching responses has been correlated with elevated temperatures, increasing physiological and field evidence points to an interaction between elevated temperature and irradiance as a major factor in the extensive bleaching of corals worldwide over the past 15 yr (Glynn 1993, Brown et al. 1995, Brown 1997, Iglesias Prieto & Trench 1997). For corals subject to solar bleaching in the present study, the generation of colourless residues within 24 h of exposure to high irradiance suggests a very rapid photodegradation of chlorophyll (Jen & Mackinney 1970). Such colourless residues are produced in higher plants in the presence of active oxygen radicals (Brown et al. 1991) which have already been invoked as mediators of the bleaching process in symbiotic anthozoans, subject to elevated temperatures and irradiance (Lesser et al. 1990).

Also of interest in the present study is the possible photoprotective role of the algal xanthophylls diadinoxanthin and diatoxanthin. The conversion of diadinoxanthin to diatoxanthin in the light and the resulting dissipation of excess absorbed light energy as heat has been cited as having a photoprotective role in algae (Demers et al. 1991, Arsalane et al. 1994, Young & Frank 1996) similar to that of the well established xanthophyll cycle in higher plants (Demmig-Adams & Adams 1993). Increased production of diatoxanthin in algae extracted from the western surfaces of corals subjected to high levels of solar radiation, the increase in the ratio of diatoxanthin to the total xanthophyll pool in bleached tissues and the progressive increase in the total xanthophyll pool over the bleaching period (relative to total chlorophyll levels) all support a potential photoprotective function for diadinoxanthin and diatoxanthin in the coral/algal symbiosis. A progressive increase in β-carotene proportions in bleached tissues over the bleaching period bears similarities with higher plants where the xanthophyll cycle pool and β-carotene have been reported to vary concomitantly (Demmig-Adams et al. 1989, Thayer & Bjorkman 1990). The fact that corals were bleached on the western surfaces of colonies suggests that any photoprotective function was stretched to its limits and, in the case of the bleached tissues, defense mechanisms failed ultimately to offer algal cells adequate protection from the damaging effects of solar radiation.

Carotenoids in symbiotic algae clearly have the potential of playing an important role in the coral bleaching process. Already Warner et al. (1996) have shown that symbiotic algae from different coral species...
vary in their photoprotective ability and that this characteristic is correlated with their susceptibility to bleaching at elevated temperatures. Such an ability is most likely influenced by the efficiency of the xanthophyll cycle and the size of the xanthophyll pool, but more detailed study of carotenoids is needed before the significance of the photoprotective function of the xanthophylls can be established in the coral/algal symbiosis. It should be noted that while diadinoxanthin and diatoxanthin have been identified in coral symbiotic algae, the presence of an active xanthophyll cycle can only be demonstrated by complementary studies of algal photosynthetic efficiency under high irradiance stress and recovery. Furthermore, differences in photoprotective ability are most likely correlated not only with the presence of the xanthophyll cycle but also with a suite of important mechanisms that include morphological changes in the chloroplasts, presence and activity of antioxidant enzymes and ability to repair photochemical damage.

Acknowledgements. We thank the Director and staff of the Phuket Marine Biological Centre for their support, particularly Dr Hansa Chansang, Mr Niphon Phongsuwan, and Mr Ukrit Satapoomin.

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