Coral community composition and reef development at the Similan Islands, Andaman Sea, in response to strong environmental variations

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ABSTRACT: The Similan Islands, a Thai archipelago in the Andaman Sea located near the shelf break, are subjected to frequent (up to several events per hour) and abrupt changes in physico-chemical conditions, particularly during the dry season (NE monsoon, January through April) and to an intense monsoon season with strong surface wave action (May to October). The exposed west slopes of the islands feature more coral species, but lack a carbonate reef framework. By contrast, the sheltered east sides show a complex reef framework dominated by massive Porites. Our results suggest that the sudden changes in temperature, pH and nutrients (drops of up to 10°C and 0.6 U and increases of up to 9.4 µmol NOx l⁻¹, respectively) due to pulsed upwelling events may rival the importance of surface waves and storms in shaping coral distribution and reef development.

KEY WORDS: Benthic−pelagic coupling · Coral community · Similan Islands · Andaman Sea · Upwelling · Internal waves

INTRODUCTION

Coral reefs are highly diverse and productive benthic ecosystems thriving in shallow, clear, warm, nutrient-poor, tropical waters (Veron 2000). Many reefs, however, are subjected to natural disturbances that affect hydrodynamics, light, temperature, nutrients and pH on various scales, e.g. monsoon-driven rainfall, mixing by storms and upwelling of cold, deep waters (Leichter et al. 1996). Surface gravity waves can strongly affect coral morphologies and reef development in exposed areas (Storlazzi et al. 2001). Internal waves have been shown to play an important role in the cross-shore exchange of nutrients, larvae and particulate food (Pineda 1991) and can be a source of nutrition for coral reefs (Leichter et al. 2003, Wolanski et al. 2004). However, when upwelling is strong, positive fertilizing effects may be counteracted by the negative effects of low temperature, low pH and high nutrient loads favouring algal rather than coral growth (McCook et al. 2001).

Reef growth is commonly most vigorous on the exposed offshore (or windward) side of an island (Veron 2000). Along the offshore islands in the Andaman Sea, however, satellite images (www.reefgis.reefbase.org) and monitoring studies (Phongsuwan et al. 2008) suggest that reef development is restricted to the sheltered east sides, while the west sides appear to be devoid of reef formations (Fig. 1).

The Andaman Sea features 2 major climatic and oceanographic phenomena: a strong SW monsoon from May to October, with strong swell and heavy rain (Wu & Zhang 1998), and the occurrence of large-amplitude internal waves (LAIW) of >60 m amplitude.
(Osborne & Burch 1980, Jackson 2004) generated by tidal currents across the shallow ridges of the Andaman–Nicobar island arc and the reefs NW of Sumatra (Jackson 2004). The LAIWs travel as waves of depression in packets of 5 to 8 waves eastward across the deep Andaman basin at speeds of 2 m s⁻¹ (Osborne & Burch 1980) and are observable as rip bands on the ocean surface (Jackson 2004). Near the continental shelf break, the rip bands align with the isobaths and proceed in water shallower than 90 m (Jackson 2004), suggesting transformation into waves of elevation, breaking and generation of turbulent bores spilling onto the shelf (Vlasenko & Hutter 2002). The resulting diapycnal mixing was shown to enhance pelagic production near the shelf break (Jackson 2004, Nielsen et al. 2004) in the swash zone of breaking solitons (Vlasenko & Hutter 2002, Jackson 2004). The further transport of cold, deep waters due to LAIW up to shallow shelf areas, however, has not been proven yet. But 2 recent investigations at the Similan Islands, an offshore archipelago in the Andaman Sea located near the shelf break, found strong-pulsed upwelling at the exposed western side of the islands (Roder et al. 2010, 2011), suggesting that those LAIW may reach the island chain. These studies showed differences in the nutritional status of corals growing on the exposed west and sheltered east sides during a period of intense, pulsed upwelling.

This pulsed upwelling might be an important factor affecting not only the physiology of the corals but also the ecology of the coral communities. Low temperatures have been shown to limit reef development (Kleypas et al. 1999) and may select for species tolerant to low temperatures in areas subjected to upwelling of cold water (Brown 1997). Because upwelled waters can be low in oxygen (Levin et al. 1991) and pH (Levitus & O’Brian 1998), they may select for corals adapted to cope with large fluctuations in these parameters. Further corals may benefit from enhanced nutrient supply due to increasing zooxanthellae densities (Muscatine et al. 1989, Ferrier-Pagès et al. 2001) and photosynthetic efficiency (Muscatine et al. 1989). But higher nutrient concentrations also favour the growth of competitively superior macroalgae (McCook et al. 2001) and enhance bioerosion (Chazottes et al. 2002). Tropical shallow platforms subjected to intense (wind-driven) upwelling have thus been shown to be devoid of coral reefs (Hallock & Schlager 1986), but moderate and/or intermittent upwelling may allow for moderate to extensive reef development (Andrews & Gentien 1982).

Nevertheless, it cannot be ruled out that physical forcing due to storms, mainly during the monsoon season, or caused by tsunamis, impacts reef development on the exposed western sides of these islands as well (Allen & Stone 2003); although the recent tsunami in 2004 showed only moderate impact on Andaman Sea offshore coral reefs (Allen & Stone 2005, Plathong 2005), with <13% of the reefs significantly damaged, and this damage affecting mostly the narrow north and south passages between islands and not the broad west island faces.

The present study explores the coral abundance and diversity in coral communities of the Similan Islands subjected to marked differences in temperature fluctuations and monsoonal storm exposure. We propose that the pulsed fluctuations due to upwelled waters and the impact of surface waves differentially affect shallow (~7 m) and deep (~20 m), west and east coral communities and carbonate framework development. This was tested by comparing coral community composition, cover and framework development...
on the west, east, shallow and deep faces of the islands and relating the biological findings to the physico-chemical characteristics of these habitats.

MATERIALS AND METHODS

Study area

The Similan Islands in the Andaman Sea, Thailand, are located about 60 km west of the Thai coast and 400 km east of the Andaman–Nicobar Islands. The 9 rocky islands composing the archipelago are aligned from north to south over a distance of about 24 km, almost perfectly perpendicular to the direction of the LAIW (coming mainly from the west; Jackson 2004), and at an oblique angle (45°) to the SW monsoon (Fig. 2). Their topography and surrounding bathymetry are characterized by a generally steep slope (>40°) down to 20 m depth along the western sides and a slightly gentler slope (<35°) in the east, with broad sandy beaches and shallow reefs. The upper western slope features large boulders until about 15 m depth before giving away to rubble and coarse sand. The average depth around and between the islands is 70 m (Fig. 2). In the east, the upper slope is shallow (5 to 10 m), with reefs and fine carbonate sands extending about 30 m from the island shores. The deeper slope is steep and covered with corals to a depth of about 30 m. Annual rainfall is 3560 mm and mostly restricted to the SW monsoon between May and October.

Benthic sampling

For the coral community study, 16 localities were selected: 9 were distributed along the eastern sides of the Similan Islands (E 1.1 to E 9.1; Fig. 2) and sheltered from direct upwelling and monsoon impact; 7 were distributed along the western sides (W 2.1 to W 9.1; Fig. 2) and exposed to the full impact of deep waters and monsoon. Although precaution was taken to obtain a representative dataset, logistical constraints caused some sites to be more clustered (W 8.2 and W 8.3, E 8.1 and E 8.2) than preferable on theoretical grounds. To explore the west–east differences in the benthic composition, one 100 m line transect was established at 14 m depth at each of the 16 localities (marked in each case with W 14 m and E 14 m, respectively) and marked with steel stakes, one at either end. The line-point-intercept method (LPI; Loya 1972) was adopted with measuring points every 50 cm (n = 200 data points transect−1). At each point the benthic component (i.e. scleractinian corals, soft corals, macroalgae, rock, sand and rubble) under the transect line was recorded, and the number of individuals was counted. Scleractinian corals were identified to species level. To establish vertical differences in the composition of the main macrobenthic groups at low taxonomic resolution (no species identification), additional triplicate transects of 50 m length were conducted at Ko Miang on both west (W 4.1) and east (E 4.1) sides at 20 m and 7 m depth (W 20 m and E 20 m, W 7 m and E 7 m,
respectively). These transect lines were positioned one after another along the respective isobath with a distance of 10 m between the end of one and the beginning of the next. Again, the LPI method was adopted as described above (n = 100 data points transect⁻¹, analysis of similarity [ANOSIM] between triplicate transects to exclude auto-correlation; see Table S5 in the supplement at www.int-res.com/articles/suppl/m456p113.pdf).

Carbonate framework development was assessed on both west and east sides of islands 4.1 and 8.1 (at 4.1 at 7 and 20 m depths and at 8.1 at 14 m depth) from scaled photographs, showing the elevation of the coral framework above its basement (sediment or rock) and a measuring stick positioned perpendicular to the isobaths in front of the framework. Images were taken every 5 m along the transects (where applicable, i.e. corals were present). The height of the framework was calculated by image analysis with the software ImageJ as the closest distance (in cm) between the top 10 tips of corals and the basement. A total of 207 framework measurements was carried out — 21, 67 and 59 in the west at 7, 14, and 20 m depth, respectively, and 16, 28 and 16 each in the east at 7, 14 and 20 m depth, respectively.

Physico-chemical sampling

Temperature measurements

A total of 20 temperature recorders (TidbiT v2, Onset computers; resolution 0.2°C within a range of 0 to 50°C) recording temperature at 6 min intervals were deployed at 2 depths (20 and 7 m) and exposures (west and east) at 5 study sites (2.1, 4.1, 7.1, 8.1 and 8.3; Fig. 2). Loggers were attached about 20 cm above the substrate and kept for 21 mo from February 2007 to November 2008. Additional temperature loggers were deployed at the island Ko Miang (W 4.1 and E 4.1) at 7, 14 and 20 m depths logging for 4 mo from December 2007 through March 2008. Intercalibration of the TidbiT loggers after the long-term deployment showed small temperature differences between loggers (max. 0.8°C) contrasted by up to 10°C temperature variations in the course of the pulsed upwelling events.

CTD and light measurements

Temperature, salinity, pressure, oxygen, pH, chlorophyll a fluorescence and optical backscatter were measured in 1 min intervals with a CTD (Seacat SBE 19plus, Sea-Bird Electronics) deployed at 20 m depth on the west side (W 4.1) of Ko Miang (Fig. 2) in March 2007 for 4 wk. Light loggers (Onset pendant light logger: 0 to 320 000 lux [lm m⁻²]) were deployed on both sides of Ko Miang (W 4.1, E 4.1) at 2 depths (20 and 7 m) about 20 cm above the substrate recording light values every minute from December 2007 until April 2008. The loggers were cleaned daily to keep fouling organisms from interfering with light measurements. Additionally, a photosynthetic active radiation (PAR, µmol quanta m⁻² s⁻¹) sensor (Bio-photometric Instruments) was mounted on a CTD (SEACAT, SBE 19plus, Sea-Bird Electronics) in January 2008 over a period of 20 d on the west side of Ko Miang (W 4.1) at 20 m depth.

Chemical measurements

A total of 89 water samples was collected by SCUBA during daily dives at Ko Miang in February and March of 2007 and 2008 (Fig. 2; at W 4.1: 21 samples at 20 m, 18 samples at 7 m depth; at E 4.1: 25 samples each at 20 and 7 m depths). Samples were taken with 1 l PE-bottles about 1 m above the reef substrate in the vicinity of the temperature loggers. The sampling time was recorded on each occasion for later correlation with water temperature. Immediately after collection, water samples were filtered through pre-combusted and pre-weighed glass-fibre filters (Whatman GF/F, 45 µm) for determination of total suspended matter (TSM). The filters were kept frozen before drying at 50°C and determining dry weight of TSM gravimetrically (Mettler, AT21 Comparator, 1 µg accuracy). Aliquots of the filtrate were stored in sterile polypropylene bottles after poisoning them with mercuric chloride (Kattner 1999) until inorganic nutrients (nitrate, nitrite, ammonium, phosphate and silicate) were analyzed according to Grasshoff et al. (1999) and Parsons et al. (1989) in a spectrophotometer (GBC Model UV/VIS981 with autosampler Model FS3000 in 2007, and Evolution III autoanalyzer, Alliance Instruments, in 2008). Further filtrate aliquots were transferred to pre-combusted glass ampoules spiked with phosphoric acid and flame sealed for determination of dissolved organic carbon (DOC). DOC samples were analyzed by means of high temperature catalytic oxidation using a Dohrmann DC-190 Total Organic Carbon Analyzer equipped with a platinum catalyst. Before injection into the furnace, the acidified samples were decarbonated by purging with oxygen. The evolving CO₂
was purified, dried and detected by a non-dispersive infrared detection system.

Data processing and statistical analysis

Biological and environmental data were tested for the assumption of normality and homogeneity of variances using Kolmogorov-Smirnov and Levene’s tests, respectively, and subjected to parametric and non-parametric analyses, as required.

Transect data were processed as percentage abundances per site and analyzed with the software PRIMER v6 for non-parametric multivariate datasets (Clarke & Gorley 2006). Analysis of similarities (ANOSIM) permutation tests based on Bray-Curtis similarities were used to detect spatial differences in substrate cover and species compositions between island sides (1-way analysis with 25 iteration steps). It was further used to clarify depth-dependent benthic cover and distribution of substrate types on the west and east sides of Ko Miang (2-way-crossed design implicating the differing exposure depending on orientation: W versus E; depth: 20 versus 7 m). ANOSIM calculates a global R statistic that reflects the differences in variability between groups as compared to within groups (so R values are proportional to differences between the groups) and checks for the significance of R using permutation tests (Clarke & Gorley 2006). Non-metric multidimensional scaling (NMDS) was used to further describe the benthic communities. Based on a similarity matrix, NMDS generates plots in which the distance between points is proportional to their degree of similarity (Clarke & Gorley 2006). Similarity percentage (SIMPER) analyses were consulted to assess the respective contributions of substrate types and coral species to the similarities and dissimilarities within and among the west and east sites studied. Descriptive coral community factors such as the Shannon index, evenness and species richness (Rogers 1993) were calculated for every site and tested for exposure (west vs. east) using Student’s t-tests. The effect of side (west and east) and depth (7, 14 and 20 m) on coral framework development was analyzed using ANOVA, with side and depth as treatment factors; post hoc, pairwise comparisons were performed via Tukey’s HSD tests.

The calculation of degree days cooling (DDC; °C d) as a site-specific indicator of the upwelling involved the following steps: decomposing the temperature data into moving average and residuals, splitting the residuals into positive (warm) and negative (cold) values, and summing up the values for the cold residuals. Given the highly skewed distribution of the temperature data and the sensitivity of the arithmetic mean to outliers, it was appropriate to use the most common (or mode) temperature to represent the average. The moving average was calculated using a slide function (by Jos van der Geest; www.mathworks.com/matlabcentral/fileexchange/12550) with a 1 d time window for the mode value. The daily sum of cold residuals was divided by the number of samples per day. As not all loggers were recording without failure at all sites, the calculation of DDC was done for the period when data were available from all sites and depths (20 wk, February 2007 to July 2007, covering the SE and SW monsoon periods in roughly equal proportion). The relationship between physical (DDC), chemical (nutrients) and biological (coral community composition and cover) descriptors was assessed using linear-regression analyses. Possible temperature differences (monthly minima, maxima and mode values) between sites corresponding to the same upwelling exposure (e.g. W 20 m) were tested with Student’s t-tests (Table S1 in the supplement at www.int-res.com/articles/suppl/m456p113.pdf), and data for all sites that were not statistically different (at p = 0.01) were pooled for further analyses. Temperature data, nutrient concentrations and light recordings were statistically tested for differences between west and east using non-parametric Kruskal-Wallis ANOVA by ranks. Linear-regression analyses were undertaken to test the relationships between both chemical and physical parameters in the water as the dependent variables.

Fig. 3. NMDS ordination of coral communities at the Similan Islands based on species abundance data (%) and Bray-Curtis similarities from 1 × 100 m transect at 14 m depth at each study site. West (▼) and east (▲) sites (see Fig. 2) grouped in separate clusters.
with temperature as the independent variable, followed by a Student’s $t$-test to examine their statistical significance. If not stated otherwise data are always displayed as means (±SE).

Table 1. Coral diversity indices (Shannon index [$H_k$], evenness [$E_k$], species richness [$S$]), and coral cover (as a fraction of hard substrate) determined from 1 × 100 m transect at 14 m depth at each study site (all sites). At island Ko Miang (Sites W 4.1 and E 4.1, see Fig. 2) additional determination of coral cover at 20 m and 7 m depth from in each case 3 × 50 m transects (mean ± SE). Linear regression model with $H_k$, $E_k$, $S$ and coral cover as dependent and degree days cooling (°C d) as independent variables ($n$ = number of samples; significance levels are *0.05 > $p$ ≥ 0.01, **0.01 > $p$ ≥ 0.001)

<table>
<thead>
<tr>
<th>Depth</th>
<th>Index</th>
<th>Site location</th>
<th>n</th>
<th>$R^2$</th>
<th>$p$</th>
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<td>West</td>
<td></td>
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<tr>
<td>All sites</td>
<td>$H_k$</td>
<td>2.69 (0.15)</td>
<td>10</td>
<td>0.4</td>
<td>*</td>
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<tr>
<td></td>
<td>$E_k$</td>
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<td>0.07</td>
<td>0.46</td>
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<tr>
<td></td>
<td>$S$</td>
<td>38.4 (2.4)</td>
<td>10</td>
<td>0.63</td>
<td>**</td>
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<tr>
<td></td>
<td>Coral cover</td>
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<td>10</td>
<td>0.13</td>
<td>0.29</td>
</tr>
<tr>
<td>Ko Miang</td>
<td>Coral cover</td>
<td>0.32 (0.09)</td>
<td>6</td>
<td>0.42</td>
<td>0.08</td>
</tr>
<tr>
<td></td>
<td>14 Coral cover</td>
<td>0.35</td>
<td>30</td>
<td>0.38</td>
<td></td>
</tr>
<tr>
<td></td>
<td>7 Coral cover</td>
<td>0.54 (0.04)</td>
<td>54</td>
<td>0.03</td>
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</table>

Table 1. Coral diversity indices (Shannon index [$H_k$], evenness [$E_k$], species richness [$S$]), and coral cover (as a fraction of hard substrate) determined from 1 × 100 m transect at 14 m depth at each study site (all sites). At island Ko Miang (Sites W 4.1 and E 4.1, see Fig. 2) additional determination of coral cover at 20 m and 7 m depth from in each case 3 × 50 m transects (mean ± SE). Linear regression model with $H_k$, $E_k$, $S$ and coral cover as dependent and degree days cooling (°C d) as independent variables ($n$ = number of samples; significance levels are *0.05 > $p$ ≥ 0.01, **0.01 > $p$ ≥ 0.001)

A total of 144 hermatypic coral species was recorded, belonging to 40 genera and 17 families (Tables S2 & S3 in the supplement). Coral species composition differed between island sides (ANOSIM, 1-way analysis: west versus east: global $R = 0.585$, $p < 0.001$, dissimilarity 83.2%; Table S4) as seen also in the distinct clustering in the NMDS plot between west and east sites (Fig. 3). In the west, a higher species diversity ($t$-test, $p < 0.01$), richness ($t$-test, $p < 0.02$) and evenness ($t$-test, $p < 0.04$; Table 1) was found compared to in the east. In the east all sites were characterized by a dominance of *Porites* species (64.2% cover as a fraction of total coral cover compared to 15.7% in the west; Table S2 & S3). The within-west and within-east similarities in species composition were low (24.3 and 31.9%, respectively; Table S4), due to a variable number of rarer species mainly within the genera *Acropora* (28.6% cover as a fraction of total coral cover in the west and 11.5% in the east) and *Porites* distributed

![Fig. 4. Benthic composition at Ko Miang on exposed west (W 4.1) and sheltered east (E 4.1) sites (see Fig. 2), grouped by depth: 7, 14 and 20 m: (A) general benthic composition and (B) coral morphologies as fractions of total number of living colonies.](image-url)
unevenly between the sites, particularly in the west (Tables S2 & S3).

Benthic substrate composition

From the 2 sides (west, east) and 3 depths investigated at Ko Miang (7, 14 and 20 m), the lowest living coral cover was found at W 20 m (ANOSIM, 2-way-crossed analysis: W 4.1 versus E 4.1: global R = 0.639, p < 0.02; Fig. 4A). Significantly higher living coral cover was found at W 7 m (35.7 ± 5.6% versus 12 ± 4.4% at 20 m depth), E 20 m and E14 m (36.7 ± 4.6% at 20 m depth and 33.6% at 14 m depth versus 18.7 ± 5.5% at 7 m depth; 20 m versus 7 m: global R = 0.37, p < 0.05; Fig. 4A). Living coral cover was the second most powerful contributor to the dissimilarity of 72.5% between west and east, with a contribution of 25.1%; the most important contribution to this dissimilarity between west and east was achieved by sand and rubble with 35.9%, followed by rock (17.9%) in third position. At 14 m depth, differences in the benthic composition failed to be significant between west and east sides (ANOSIM, 1-way analysis: global R = 0.079, p = 0.22; Table S5 in the supplement at www.int-res.com/articles/ suppl/m456p113.pdf) due to similar distributions of total hard substrate consisting of dead coral, living coral and rock (with 67.7 ± 6.6% in the west and 72.8 ± 3.3% in the east), and sand including rubble (with 29.8 ± 6.9% in the west and 25.9 ± 3.3% in the east). Living coral cover exhibited higher differences between the west (25.6 ± 4.2%) and east (35.9 ± 4.6%) sides. Rock structures were only found in the west, while in the east hard substrate consisted exclusively of dead coral (Fig. 4A, Table S6).

Coral framework and morphologies

A dense 3-dimensional coral framework characterized the east, whereas in the west hard corals were distributed as solitary colonies without developing an actual carbonate framework (ANOVA, Tukey HSD-tests: W versus E, p < 0.001; Fig. 5, Table S7). This was especially distinct at W 20 m and W 7 m, whereas at W 14 m depth the framework was higher (ANOVA, Tukey HSD-test: p < 0.001; Fig. 5, Table S7). Coral morphologies differed significantly between sides (ANOSIM 2-way-crossed analysis: W versus E, global R = 0.311, p < 0.05) and depths (7 versus 20 m, side data pooled, global R = 0.253, p < 0.05) (Fig. 4B). The dominance of massive and encrusting colonies in the west and of branching species in the east of Ko Miang fell short of being significant (t-test, p = 0.064 and p = 0.1, respectively). Large massive and encrusting hard coral species sparsely covered areas at W 20 m, and small colonies of all morphological types grew very close to each other on the rocks in shallower west waters (Fig. 4B). Here, branching hard corals, especially within the genera *Millepora* and *Acropora*, often displayed flattened morphologies with pancake-like broadened bases and strongly reduced ramification (Fig. 1A).

Physico-chemical environment

The pattern of high- and low-variability periods and order of magnitude in amplitude in the temperature time series was consistent among all sites of the same depth and orientation (Fig. 6, cf. synchronous monthly mode, minimum and maximum temperature values, Student’s t-tests: Table S1). This is also reflected by the differences between mean values and ranges (maximum to minimum temperature) of monthly temperature between west and east sites (Table 2; Kruskal-Wallis W 20 m versus E both depths: p < 0.00; Table S8). Mode values differed only between W 20 m and E 7 m sites (Table 2; Kruskal-Wallis, p < 0.01; Table S8) and showed small (<3.5°C) seasonal differences, irrespective of side or depth. The low seasonal variability contrasts with the high variability found at higher frequencies, as reflected by the large differences between mode and minima values showing declining temperature fluctuations: W 20 m > W 7 m > E 20 m > E 7 m (Fig. 6,
Fig. 6. Temperature record covering 21 mo along the Similan Islands (March 2007 to November 2008): monthly minimum (○), maximum (●) and mode (○) values at 5 west (W 2.1, W 4.1, W 7.1, W 8.1, W 8.3) and east (E 2.1, E 4.1, E 7.1, E 8.1, E 8.3) sites (see Fig. 2), respectively, in 20 and 7 m depth (orientation south to north). Blank periods are missing values due to lost or broken loggers. Vertical line in each panel indicates change of year 2007 to 2008.
Table 2). Corresponding DDCs showed 5-fold higher values at W 20 m compared to E 7 m (Kruskal-Wallis test: p < 0.001; Table S8) and about 2- and 3-fold higher values than at W 7 m and E 20 m, respectively (Table 2).

Fig. 7 shows a time series section of raw data at W 20 m at Ko Miang during strong temperature fluctuations. Synchronization of abrupt temperature decreases of up to 5−9°C occurred with drops in oxygen concentration of down to 12% (down to 21.8 µmol l−1), in pH of up to 0.6 U (down to 7.75) and increases in salinity of up to 1.5 to values of 34.4. Statistics did not support temperature-dependent variations in optical backscatter and chl a fluorescence (Fig. S1 in the supplement).

Highest light intensities (lux, 1 m m−2), were found at E 7 m and lowest at W 20 m. Light conditions were significantly different in the west and east at both depths (7 and 20 m) (Kruskal-Wallis test, p < 0.001; Table S9A), with over 3-fold higher light values in the east during midday and a longer lasting light environment at W 7 m at the end of the day (Fig. 8). This pattern can be explained by the topographical characteristics of all the islands, with high, steep slope-shaped west sides shading the west for longer than the east and flattened island topography in the east. During a period of high temperature variations in January 2007, PAR never exceeded 141.5 µmol quanta m−2 s−1 at W 20 m, with a mean of 102.5 ± 3.4 µmol quanta m−2 s−1.

Nutrient concentrations revealed significant differences between the west and east for nitrate and nitrite, as well as for silicate (Kruskal-Wallis test, p < 0.001 and p < 0.04, respectively). Mean concentrations of silicate were >60% higher at W 20 m than at E 20 m, supported by an increase of nitrate and nitrite (40%) and of phosphate (50% higher at W 20 m than at E 7 m) concentrations (Table 3). In

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<th>West</th>
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<tr>
<td></td>
<td>20 m</td>
<td>14 m</td>
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<tr>
<td>All sites</td>
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<tr>
<td>Mean</td>
<td>28.5 (0.7)</td>
<td>28.8 (0.6)</td>
</tr>
<tr>
<td>Mode</td>
<td>28.7 (0.6)</td>
<td>28.9 (0.1)</td>
</tr>
<tr>
<td>Range</td>
<td>5.2 (1.7)</td>
<td>4.3 (1.7)</td>
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<tr>
<td>DDC</td>
<td>−58.3 (1.9)</td>
<td>−31.6 (3.8)</td>
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<tr>
<td>Ko Miang</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>28.0 (0.8)</td>
<td>28.3 (0.7)</td>
</tr>
<tr>
<td>Mode</td>
<td>28.2 (0.4)</td>
<td>28.1 (0.5)</td>
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<tr>
<td>Range</td>
<td>5.6 (1.1)</td>
<td>4.8 (1.1)</td>
</tr>
<tr>
<td>DDC</td>
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shallow waters the differences may be still noticeable but are far less pronounced and statistically not significant. Nutrient concentrations exhibited a negative relationship with temperature when correlated to the in situ temperature during sampling (Table 3). No significant differences were found between the different sides of the island in ammonium concentration, dissolved organic carbon or suspended particulate matter which varied independently of temperature variations (Table 3, Table S9B in the supplement).

DISCUSSION

The Similan Islands reveal a lack of coral reef framework development along their western sides, in contrast to the complex 3-dimensional coral framework in the east. The development of coral reefs in general has been reported to be undermined by cold water influence (Burns 1985, Kleypas et al. 1999). Although several tropical corals are known to occur over a wide geographical range of temperatures (e.g. up to 12°C differences in maximum summer temperatures for species co-occurring in the Arabian Sea and Lord Howe Island, Australia; Hughes et al. 2003), and can tolerate large annual ranges of temperature (up to 25°C) and survive severe cold periods (13°C for several days; Coles & Fadlallah 1991), cold water stress (6 to 10°C under normal conditions) may disrupt the sensitive association between corals and their endosymbiotic zooxanthellae and result in a decreased fitness and growth performance of the coral host (Coles & Fadlallah 1991, Gates et al. 1992, Saxby et al. 2003). The reduced coral cover in the west at 20 m depth and the absence of reef framework in the west may therefore be partly due to the observed temperature fluctuations. The cold temperatures in the present study, however, were only one stressor out of several co-occurring environmental factors. Temperature was negatively correlated with nutrient input (Table 3) and salinity, and positively with pH and oxygen concentration (Fig. 7). Thus, the cold water masses reaching the west of the Similan Islands may be derived from sub-pycnocline sources containing re-
mineralized nutrients (Jackson & Williams 1985) due to respiration and decomposition processes in areas isolated from additional oxygen supply (Levin et al. 1991). The covariance of temperature with these environmental parameters (Fig. 7, Table 3) allowed an assessment of the physico-chemical environment from temperature alone for periods where no other data were available. Exceptions were the dissolved organic matter and suspended particulate matter concentrations (see also optical backscatter; Fig. S1), which were independent of temperature and similar between west and east. Reduced oxygen and high nutrient concentrations in the sub-thermocline waters transported into shallow areas may possibly have led to a mutual compensation of depletion (low oxygen) and production processes (plankton production due to higher nutrient concentrations; Eppley et al. 1979) in the west. This could have been a reason for the lack of difference in dissolved organic matter concentrations between west and east. The similar particulate matter concentrations appeared inconsistent with the differences in the physical oceanographic variables and the varying light conditions between west and east sides. This could be explained by higher suspended particulate loads in the west than in the east caused by higher currents (Roder et al. 2011) and due to more intense resuspension. Mean nutrient concentrations for nitrate, nitrite and phosphate in the west were above the average concentrations of most tropical reefs (Table 3, Kleypas et al. 1999). During periods of high temperature fluctuations (February, March) they were close to or even above the extreme values assessed for coral reef communities (Kleypas et al. 1999). Increased nutrient concentrations entail neutral or positive effects on coral nutrition, physiology and zooxanthellae concentrations (Muscatine et al. 1989, Ferrier-Pagès et al. 2001). The combination of higher current velocities and nutrient concentrations can enhance coral nutrient uptake (Hearn et al. 2001) and photosynthesis (Szmant 2002). Accordingly, higher pigment concentrations in western corals were indicative of their similar net photosynthetic rates compared to sheltered eastern corals (Jantzen 2010). Several coral species of different morphology and growth rates along the Similan Islands displayed higher pigment concentrations, protein content and biomass under the influence of increased nutrient concentrations and higher planktonic food supply in the west (Roder et al. 2010, 2011). However, these positive effects on coral nutrition and photosynthetic performance may be small compared to the negative direct effects of nutrients and low pH on coral calcification and growth (Kinsey & Davies 1979, Ferrier-Pagès et al. 2000). Values of pH decreased during cold-water intrusions in the west (up to 0.6 U; Fig. 7). This effect in combination with reduced light intensities (Fig. 8) may be the main reason for the significantly reduced coral growth at W 20 m depth (Schmidt et al. unpubl. data), as both processes are tightly coupled in zooxanthellate corals and their efficiency suffers from lowered pH and light levels (Marubini et al. 2001). This might also be a major reason for the inability of the corals in the west to develop a reef framework (Manzello et al. 2008).

The pulsed extreme conditions in the chemical and physical environment are replaced each time within 15 to 30 min, again by warm surface water (see Fig. 7; nearly identical mode temperature values in the west and east; Table 2). The intensity of these fluctuations increases with depth (7 m < 14 m < 20 m), but does not seem to cause severe physiological damage or mortality as observed for long-term disturbances following cold spells (Coles & Fadlallah 1991) or eutrophication (Szmant 2002). Monsoonal storms, on the other hand, occur at monthly or seasonal frequencies (Wu & Zhang 1998) and last for days. Their impact diminishes with depth. When looking at the intermediate depth of 14 m in the west, the coral framework was higher than in deep (20 m) and shallow (7 m) waters, and high coral diversity was found in contrast to the sheltered eastern reefs (Table 1). In the east species diversity may be lower in the absence of disturbance due to competitive exclusion in an equilibrated system (Connell 1978), with high coral cover but less species dominating the communities and no available free substrate for new settlers (no bare rock in the east; Table S6). In the west, however, the seasonal alternation of temperature fluctuations (January through April) and monsoonal surface waves (May through October) may be a disturbance which represents a selective regime (Karlson 1980) to which organisms adapt and generate a variety of competitive mechanisms (Karlson & Hurd 1993). This non-equilibrium state may reset succession, with regularly new settling corals on the available free substrate (bare rock in the west; Table S6), and thus enhance coral development and coral species diversity (Connell 1997).

Going deeper to 20 m depth along the western Similan Islands, the reef assemblage is impoverished, with single corals growing apart from each other (Figs. 4B & 5). This is reminiscent of coral communities described for upwelling-influenced sites dominated by only one or a few species, such as mainly Porites, Acropora, and Pocillopora (Benzoni et al. 2003). Slow-growing, massive and sub-massive species dominated the community here (Fig. 4, Tables S2
& 3). Corals of this type are often described as stress-tolerators (Riegl & Piller 2000, Veron 2000, Benzoni et al. 2003), with high adaptive tolerance to different types of environmental stress (Marshall & Baird 2000, Alutoin et al. 2001, Fabrizius et al. 2011). This was also shown by Roder et al. (2011) who found high metabolic plasticity in the species *Porites lutea* and *Diploastrea heliopora* to the fluctuating chemical conditions in the west of the Similan Islands. The general abundance of branching species was reduced in the west compared to in the east (Fig. 4B); however, *Acropora* species showed comparatively higher abundances in the west. Possibly *Acropora* corals were able to benefit more than other coral species from the changes in environmental conditions between upwelling and monsoon-impact periods. Most of them are fast-growing species (Veron 2000) and able to recover rapidly from disturbances and spread by using broken coral fragments as new settling colonies (Highsmith 1982). Light-catching table-like morphotypes of branching colonies which would have been due to the reduced light conditions with increasing depth (Macintyre & Smith 1974) were rare in the west. This indicates that hydrodynamics in the west, due to the high current speeds correlated with cold water (Roder et al. 2010) and due to surface gravity waves, may have reduced the number of sheltered sites for these exposed growth forms. Especially the shallow (7 m), western corals showed growth patterns as described for high-wave-energy areas (Storlazzi et al. 2001, 2003) nestled and disjointed directly to the granite basement with robust or fast-growing colonizing species. However, the impact of surface gravity waves can only be estimated as we lack knowledge of their actual intensity and maximum depth range for our study area. We suppose that, in contrast to most other reef sites, coral morphologies, species distribution and reef development in the west are affected by both hydrodynamics in shallow areas causing coral breakage and leading to hampered 3-dimensional reef development (Fig. 5, Grigg 1998) and the highly fluctuating physico-chemical conditions with increasing depth.

In summary, the composition of a coral community reflects the interaction of the governing physical, chemical and biological drivers (De Vantier et al. 1998). The coral communities growing along the western Similan Islands are exposed to a stressful physico-chemical environment. They appear to be adapted to mechanical disturbances, such as strong surface waves, and they deal successfully with the pulsed frequent supply of sub-ypnocline waters into their low-light environment. They appear to exist in species-rich communities where regular disturbance produces a highly successional community structure.

### Caveats to the conclusions

The western Similan Islands were found to be exposed to temperature fluctuations (see also Roder et al. 2010, 2011) which are more frequent and severe than reported earlier elsewhere (Leichter et al. 1996, 2003), and rank among the largest ones so far reported in tropical shallow reef areas above 30 m depth (Wolanski et al. 2004, Sheppard 2009). Seasonal variations, revealing the strongest temperature fluctuations during the late NE monsoon (March; Fig. 6) may be related to variations in the depth and strength of the pycnocline: the shallow pycnocline during the NE monsoon corresponds to the strongest LAIW activity (Nielsen et al. 2004). The pulsed, short-term fluctuations of temperature to the order of minutes, periodic occurrences in several events per day (Fig. 7) and seasonal variations (Fig. 6; see also Wolanski et al. 2004 and Storlazzi et al. 2003) indicate that they may be caused by LAIW reaching these shallow-water areas. The especially high temperature variability in 2007 compared to 2008 (Fig. 6) may be related to the untypical positive Indian Ocean Dipole event in 2007 (Bhera et al. 2008) with stronger upwelling events (Webster et al. 1999) and likely internal waves.

Even though LAIW seem to be the likely source of the observed temperature fluctuations, other possible factors, such as current-driven advection of cold waters (Andrews & Gentien 1982) or seasonal wind-driven upwelling (D’Croz & O’Dea 2007), cannot be excluded. Research in this area has just started and needs to be expanded further to include data not only on the NE monsoon period but also the SW monsoon, to complement our knowledge on surface wave impacts and to disentangle the differential influences of upwelling and surface waves on coral reef development. In addition, verification of the primary drivers of the temperature fluctuations — these may be currents, tides or internal waves — would complement the present approach and set it in its oceanographic context.

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