Benthic community production and response to environmental forcing

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WAMSI Kimberley Marine Research Program

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Front cover images (L-R)

Image 1: Satellite image of the Kimberley coastline
Image 2: Water cascading off Tallon reef crest during ebb tide (Source: Renee Gruber).
Image 3: Humpback whale breaching, Exmouth (Source: Pam Osborn)
Image 4: Azton Howard from Bardi Jawi Rangers filtering water sample for nutrient analysis (Source: Renee Gruber).
Author Contributions: RL, RG and JF designed the field experiments, RL and RG conducted the field work and the data analysis, and RL, RG, and JF contributed to the report and wrote the related manuscripts.

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Executive Summary

Benthic primary producers such as corals, seagrasses and macroalgae, play significant roles in a variety of coastal processes. They provide habitat for numerous fauna, stabilise sediments, and form the basis of coastal food webs. Through photosynthesis, primary producers use sunlight as the energy source to transform dissolved carbon into new plant biomass. Tropical reef systems often display high rates of primary production; however, these rates are closely coupled to local physical (e.g., water motion, light, temperature) and biogeochemical (e.g., nutrient) environmental conditions. For example, previous studies have shown that high temperature and light levels can stress primary producers by negatively impacting photosynthesis. Our present understanding of environmental controls on reef productivity is based primarily on studies from the Caribbean, Hawaii, the southern Great Barrier Reef, and other Indo-Pacific regions, which contain mainly wave-dominated reefs. Tide-dominated reefs occur where the mean tidal range exceeds mean wave height, and constitute approximately 30% of tropical reefs worldwide. Tide-dominated reefs are known to experience much greater ranges in environmental conditions than wave-dominated reefs, yet the interactions between primary producers and environmental drivers that occur in these systems have been largely unstudied.

The goals of this project were to quantify the environmental variability across a macrotidal reef system in the Kimberley, and assess how benthic primary producers responded to extremes in water motion, light, and temperature. More specifically, we investigated: 1) how tides interacted with reef morphology to drive extreme daily variability in environmental conditions; 2) how this environmental variability influenced the productivity of benthic communities living in different zones of the reef platform; and 3) how water quality in coastal waters surrounding reefs varied over the wet/dry seasonal cycle.

To achieve these aims, we conducted three field experiments on an intertidal platform reef (Tallon Island) in the Sunday Island group of the west Kimberley. During each field experiment, we deployed an array of instruments measuring hydrodynamics, water temperature, light levels at the reef surface, and water column dissolved oxygen. Based on changes in water column oxygen, we were able to estimate community-scale (100s metres) rates of primary production in two distinct communities found on this reef: seagrass-dominated, and macroalgal-dominated (includes small corals and crustose coralline algae). Rates of primary production were then related to environmental data to assess whether communities were stressed during extremes in environmental conditions. In addition, we conducted intensive water quality sampling on the reef and in offshore channels to measure seasonal changes in dissolved and particulate nutrient concentrations and suspended chlorophyll.

Our findings show that macrotidal reefs in the Kimberley experience some of the most extreme conditions yet recorded for reefs worldwide. Extreme daily variability in temperature and dissolved oxygen occurs on the reef platform, and is driven by semi-diurnal tides and solar (light) cycles. The shape and friction of the platform causes water to ‘pond’ on the reef for up to 10 hours during each ebb tide (twice daily). When these extended low tide periods occur near noon, extreme warming happens on the reef, with temperatures rising by 10°C over several hours and reaching up to 38°C. This high light availability also drives high rates of benthic primary production, which releases oxygen into the water column and results in extremes in oxygen saturation (~270%). When low tide periods occur near midnight, community respiration (the consumption of oxygen and carbon to create energy) causes oxygen levels to plummet, reaching very low (hypoxic) levels; low oxygen levels are known to harm or kill organisms in other ecosystems, and are not typically recorded on reefs. We found that rates of gross primary production of the dominant communities (macroalgae and seagrass) were not adversely affected by environmental variability on this reef, emphasising the resilience of these organisms to extreme conditions. Productivity varied on a day-to-day basis, due to the timing of noon relative to low tide; this cycle lasts ~15 days, and future studies shorter than this time frame may over- or under-estimate ecological processes (such as productivity). Finally, the overall average rates of productivity we observed were similar to the global mean for tropical reefs, demonstrating that tide-dominated reefs can maintain moderate rates of production despite daily extremes in temperature.
Water quality (nutrient and chlorophyll levels) on the reef also displayed strong variability over tidal cycles, with particulate nutrients and phytoplankton (food sources for reef organisms) flooding the reef at high tide and being consumed during low tide. Towards the end of low tide, grazing depletes particles in the water column; thus, the tidal cycle is a key process that replenishes reef food sources. Our results also suggest that the seagrass zone may serve as an important nutrient subsidy to the macroalgal zone during low tides. Overall, Tallon reef was found to be oligotrophic (relatively low dissolved nutrients and chlorophyll), which is typical of tropical waters that do not receive substantial anthropogenic nutrient loads from terrestrial runoff. In waters surrounding the reef, increases in nutrient and chlorophyll concentrations occurred from the end of the dry season (Oct) through the wet season (Apr). Although significant, these differences are small, suggesting that waters at the mouth of King Sound are only mildly influenced by terrestrial inputs during low-rainfall wet seasons.

Implications for management

This study documents some of the many aspects of ‘uniqueness’ of Kimberley reefs, which experience conditions among the most extreme worldwide. Although our study was conducted on a single reef, intensive measurements of physical conditions (e.g., hydrodynamics, reef slope, reef morphology, and platform height above sea level) allow generalization of results to other Kimberley reefs. The Reef Geomorphology project (1.3.1) classified major Kimberley reefs and documented their morphology and height relative to sea level. These characteristics determine the length of low tide on a reef flat, and thus the extremes in environmental conditions such as water flow, temperature, light, dissolved oxygen, and pH that will develop there. Environmental extremes and productivity measured on other intertidal reefs, such as Montgomery Reef and Irvine-Bathurst, would likely be very similar to those measured in our study of Tallon reef. Low intertidal reefs, such as Cockatoo Reef and other reefs growing well below sea level, would experience shorter low tide periods and thus a narrower range of environmental conditions. Sloping reef platforms would experience a much more subtle version of the ‘ponding’ effect on Tallon, with the extent of ebb tide elongation depending on the slope. Thus, physical relationships described in our study can help predict tidal patterns and environmental extremes on Kimberley reefs, provided some information on reef morphology is known.

Our results also provide a foundation for developing predictive models of conditions on Kimberley reefs under future climate change scenarios. As described above, temperature variability can be predicted on these reef platforms; this can be combined with predictions of ocean warming and sea level rise to estimate future conditions on reef platforms. For Tallon (and likely for similar reefs such as Montgomery), our estimates show that sea level rise may help mitigate mean ocean warming by damping temperature extremes on the platform. Such analyses could help prioritise management action or protection status in light of climate change predictions and other anthropogenic pressures.

Our results can also be used to help inform future monitoring exercises on Kimberley reefs, such as documenting high temperature events, or planning process-based scientific field studies. For high intertidal reefs (or where platform level is above mean sea level), the spring-neap cycle does not affect temperature or other conditions on the reef. Rather, the timing of noon relative to low tide is the primary driver of reef conditions. This cycle is ~15 days in length and is an important consideration for future studies focusing on any reef process affected by temperature, light, dissolved oxygen, etc. Studies less than ~15 days may greatly over- or under-estimate the process of interest. For low intertidal reefs (or where the platform is below mean sea level), the spring-neap cycle would be an additional driver of environmental variability. A high-frequency record of water depth at the study site is a key component of any future work, as it helps interpret results and position them within tidal and light cycles. In addition, although mean ocean temperature increased through the wet season, daily ranges of environmental conditions and reef productivity did not differ between seasons. This suggests that process-based studies during the wet season do not add any significant benefit over work during the dry season, when the Kimberley is far more accessible.

Finally, while we found that overall rates of productivity in this Kimberley reef system were comparable to
other coral reef habitats worldwide, the environmental conditions under which primary producers survive and grow are extreme. This has implications for the resilience of these producers in the face of climate change across the Kimberley environment. While these organisms appear well adapted to the local environmental conditions of the Kimberley, many are still likely operating at the edge of their capacity, as evidenced by coral bleaching in the inshore Kimberley associated with the El Niño heat wave in 2016 when temperatures across northern Australia were elevated for longer periods of time. Given the extreme conditions in temperature, oxygen levels and nutrients, management should consider strategies that will ensure any additional anthropogenic pressures that could add stress to the system are minimised and also be aware of environmental conditions that may change over time.

**Key residual knowledge gaps**

The project has provided a reasonably complete assessment of how environmental variability, including temperature, light, oxygen and water motion, influence the productivity of benthic reef communities in the Kimberley. Due to the process-focused nature of the experiments, we expect the general relationships and models developed in this project to be extended to other reef systems with some confidence. In other words, if the environmental conditions and benthic primary producers of other reefs are known, or can be reasonably predicted, we believe that robust estimates of the productivity of reefs across the Kimberley can be made.

Primary producers on Tallon’s platform were highly tolerant to short-term extremes in temperature and light. This is likely due to the long time-scales (10s – 100s of years) over which communities have adapted to reef conditions. At present, it is unclear if this tolerance will convey resilience to any future ocean warming, both slow mean temperature increases and episodic warming events. Our results show that sea level rise will dampen temperature extremes on tide-dominated reefs. Will this mitigate ocean warming impacts on tide-dominated reefs in the long term? Additionally, our results suggest that low intertidal reefs are likely adapted to a more narrow range of environmental conditions. Will these reefs be more susceptible to the effects of ocean warming?

Our study was conducted during the 2013-2014 wet season, which experienced relatively low rainfall levels compared to the previous 20 years. While the results from this study indicated minor differences in reef water quality between seasons, this may not always be the case, particularly during more ‘typical’ wet years. Higher rainfall would likely result in nutrient and sediment pulses to coastal waters, driving phytoplankton production. Land-clearing and farming practices in coastal catchments increase these nutrient and sediment inputs, which has had a negative impact on reef health worldwide. At present, there is no data to assess whether changes planned for Kimberley catchments such as the Fitzroy River would substantially alter coastal water quality. Additionally, the Sunday Island group is relatively far (~100 km) from the closest major river (Fitzroy), but other reefs are much closer to river mouths, especially in the northern Kimberley. How does river discharge affect water quality surrounding those reefs? Knowledge of seasonal and spatial patterns in water quality would help better inform future management efforts of Kimberley reefs.


1 Introduction

Of all marine ecosystems, tropical reefs are among the most iconic and vibrant, and are generally considered to be among the most productive communities per area in the world (Crossland et al. 1991). Reefs provide many ecosystem services, including support of fisheries, coastal protection, and tourism (Moberg & Folke 1999). Unfortunately, many reefs have already experienced declines in health or are currently threatened by a host of anthropogenic activities (Bryant et al. 1998).

Early reef studies focused on “open ocean” systems in the Indo-Pacific region, which contained reefs that were oligotrophic, wave-dominated, and exposed to minimal terrestrial inputs. They had high coral cover, abundant light, and experienced low levels of environmental variability. Their morphology consisted of a steeply-sloping fore-reef, an energetic reef crest dominated by coralline algae, a reef flat dominated by a mosaic of coral, algae, and sand patches, and a sandy back-reef slope with scattered coral heads and outcrops (Odum & Odum 1955). Work over the past two decades has shown that productivity and nutrient uptake rates on reefs are related to mass transfer that controls how materials (e.g., nutrients, dissolved gasses, etc.) are exchanged between the water column and reef communities (a function of water velocity, bottom friction, solute concentration and diffusion characteristics).

With only a few exceptions, most reef studies have found that the circulation of reefs is dominantly forced by either surface waves or tides, or a combination of the two (Monismith 2007, Falter et al. 2013, Lowe & Falter 2015). There are strong regional differences in the global distribution of significant wave heights ($H_s$) within the ±30° latitude band where warm-water coral reefs are found (Figure 1). Reefs in the Pacific and Indian Oceans tend to be dominantly wave forced (annual mean $H_s > 1.5$ m), whereas reefs in the Caribbean, Southeast Asia, and northern parts of Australia tend to experience only weak to moderate wave forcing (typical annual mean $H_s < 1$ m). These patterns likely drive substantial regional-scale differences in the wave-driven flows and residence times within reefs, depending, of course, on the morphologies of individual reefs within each region (Lowe & Falter 2015).

Despite the dominance of wave-driven reef systems globally, there are numerous reef systems worldwide that can be considered tide-dominated, defined here simply as locations where mean tidal range ($MTR$) exceeds annual mean $H_s$ (Figure 1). Such systems have, for the most part, remained neglected in the literature; they include some extreme examples of macrotidal reef systems ($MTR > 3$ m) in regions off northern Australia, off east Africa, and off Central and South America (Figure 1). An iconic example of these tide-dominated reefs is the Kimberley region of northwestern Australia, where numerous reefs are subject to tidal ranges of ~10 m and water frequently drains off reef terraces during ebb phases of the tide (Purcell 2002). The dynamics of these tide-dominated reef systems are fundamentally different from those of traditional wave-driven coral reef flows, thus creating very different physical conditions within these reef habitats. As a consequence, the distinct environmental conditions within these tide-dominated reef habitats (such as in the Kimberley) would be expected to influence reef ecology, benthic productivity, and nutrient dynamics very differently from traditional wave-dominated tropical reefs.

The reefs of the Kimberley region of Western Australia are also one of the few remaining pristine coastal areas worldwide (Bryant et al. 1998, Halpern et al. 2008). The tidal regime of the west Kimberley causes reefs to undergo large fluctuations in environmental variables within a single tidal cycle. The purpose of this study was to estimate benthic fluxes of oxygen and nutrients and determine which environmental drivers (i.e., temperature, light) exert the strongest control on productivity. Understanding how variability in environmental conditions regulates the productivity of reef communities living in the western Kimberley provides a critical foundation for modelling and predicting both present productivity rates for reefs more broadly across the Kimberley, as well as how these rates may change into the future when environmental conditions change.
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Figure 1. Global spatial distributions and frequency distributions of (a,b) annual mean significant wave height ($H_s$), (c,d) mean tidal range (MTR), and (e,f) relative tidal range ($MTR/H_s$) experienced by warm-water coral reef systems. Adapted from Lowe and Falter (2015).

2 Materials and Methods

2.1 Site selection

Reef systems of the Kimberley region of northwest Australia experience large and rapid variations in environmental conditions. Tides on the Kimberley coast are among the largest in the world, and ranges of up to 12 m have been measured in the central Kimberley (Kowalik 2004). Due to its complex ria coastline (Brocx & Semeniuk 2011), this region contains over 2600 islands, many with associated fringing reefs (Wells et al. 1995). The climate in northwest Australia is dry monsoonal, with rain primarily concentrated between November – March during the ‘Wet’ season, but with little rain during the rest of the year (the ‘Dry’ season). Human populations are sparse; thus, the Kimberley is among <4% of marine areas worldwide with little to no human impact (Halpern et al. 2008). These reef systems represent a unique opportunity to study pristine reef communities in a strongly tide-dominated setting. Though Australia’s indigenous Traditional Owners have maintained ecological knowledge of these reefs for tens of thousands of years (Horstman & Wightman 2001), no studies have previously assessed the reef-scale productivity of these systems or the relationship between productivity and daily extremes in temperature, tide, and light.

The field study took place at Tallon Island (Figure 2), a high intertidal fringing reef located in the western Buccaneer Archipelago. This reef is characterised by its large platform (~1.5 km cross-shore), which sits slightly above mean sea level (0.25 m AHD) and experiences semi-diurnal tides of >8 m range during spring phases (Collins et al. 2015). The platform is nearly flat and its offshore boundary is sharply delineated by a steep crest (1-4 m high), over which water cascades during ebb tide (Lowe et al. 2015). This crest formation (coupled with bottom friction from benthic communities) is a hydraulic control for the draining reef and ensures that the platform remains submerged over a tidal cycle, despite the water level offshore falling up to several meters below the elevation of the reef (Lowe et al. 2015). Tidal duration is thus highly asymmetric on the reef platform, with ebb durations lasting ~10 hours regardless of phasing of the spring-neap cycle.

Although previous studies had recorded the dominant benthic species on Tallon Island reef (Wells et al. 1995), no map of benthic communities was available. The coverage of benthic community types on the reef platform
was determined by photographic transects during low tide. In the cross-shore direction, community type changed rapidly, so a 10-30 m spatial resolution was used, whereas a 100-200 m resolution was sufficient in the along-shore direction. Communities were characterised as being dominated either by seagrass or macroalgae (also containing coralline algae and small corals). This information was used to create a map of general habitat categories to guide site selection for experimental design and placement of equipment.

![Map of Tallon Island reef study site](image)

**Figure 2.** Location of Tallon Island reef study site in the Sunday Island group of the west Kimberley.

### 2.2 Field experiment #1 (2nd - 22nd Oct 2013)

Our first field experiment was conducted on Tallon reef at the end of the dry season from 2-22 Oct 2013 involving 5 UWA staff and students, as well as Bardi Jawi Rangers and Traditional Owners. As part of this, the following instrument array was deployed throughout the reef (Figure 3):

- 3 dissolved oxygen (DO) sensors;
- 1 downwelling light sensor;
- 9 temperature sensors;
- 2 acoustic Doppler current profilers (ADP); and
- 4 pressure sensors.
Water levels and mean current velocities were sampled at 1 Hz by upward-looking 2 MHz Nortek Aquadopp current profilers (Nortek AS) weighted to the bed (Figure 4). A small vertical bin size (0.1 m resolution) allowed capture of current velocities when water levels were low atop the reef. A small neutrally-buoyant drifter (mandarin) was used to estimate water velocity during slow, nearly stagnant periods of low tide on the reef platform (Figure 5). The drifter was released repeatedly at each of the DO logger sites and allowed to travel for 30 min (n = 25).
Dissolved oxygen (DO) concentration was measured optically at 5 min intervals using Troll 9000 LTS packages (InSitu Inc.), which also logged temperature. Three DO sensors were deployed to capture conditions in the major reef communities: seagrass, algal ridge, and mixed assemblage (seagrass and algae). Sensors were located along the major axis of reef draining during ebb tide. DO sensors were calibrated before deployment using a two-point method of 100% (aerated with pump) and 0% (deoxygenated with sodium sulphite) saturation. In addition to pre-deployment calibration, DO measurements were corrected for drift by placing all loggers together for 30 min at periods during and after the deployment. Drift in DO measurements was minor (≤ 0.32 mg L\(^{-1}\)) during the study. Due to sensor failure of one instrument, only estimates for the seagrass community were available during the first field experiment.

Light levels were directly measured at the Tallon Island study site. Downwelling photosynthetically active radiation (PAR) was recorded at the reef surface at 5 min intervals using a QCP-2300 cosine collector (Biospherical Instruments Inc.).

In addition to logger deployments, a suite of water quality parameters were measured at fixed stations on the reef (Figure 3). A fixed station in the adjacent channel was also sampled for comparison. Measurements of dissolved nutrients (ammonia, nitrate/nitrite, phosphate, total dissolved nitrogen and total dissolved phosphorus), particulate nutrients (particulate nitrogen and phosphorus), total suspended solids (TSS), and chlorophyll-\(a\) (chl-\(a\)) were undertaken regularly over 3 weeks.

Water samples for dissolved nutrients (25 mL) were filtered (0.45 μm Minisart syringe filter) on-site within 1 hour of being collected. Samples (2.3 L) for chl-\(a\) and TSS were shaken to homogenize and filtered using a vacuum pump onto pre-weighed and ashed (4 h at 550° C) filters (47 mm diameter 0.75 μm GF/F). Filters were rinsed with deionized water to remove salt and then placed in protective plastic containers for transport. Particulate nutrient samples were collected by passing 0.3-0.5L seawater through a filter (pre-ashed 25mm diameter 0.75 μm Whatman GF/F). Filters were retained, rinsed with ~5 mL deionized water to remove salt, and wrapped in foil for storage. All samples except TSS were immediately placed on ice after collection and then stored frozen at Cygnet Bay until transported back to Perth to be analysed within labs at UWA.
During field experiment #1, benthic chamber measurements were also obtained during different parts of the light cycle to provide additional estimates of benthic productivity within beds of different seagrass species (i.e., *Thalassia* and *Enhalus*) and densities (i.e., dense and sparse). Clear chambers were used for estimates of net community production, while darkened chambers were used for estimates of net community respiration (Figure 6).

2.3 Field experiment #2 (3rd – 10th February 2014)

A second field trip was conducted in early 2014 during the middle of the wet season involving 1 UWA student working closely with Bardi Jawi Rangers and a UWA volunteer. Road closures from heavy rain and difficulty shipping large amounts of research gear up the peninsula meant that this trip was at a much smaller scale than the others, and was mainly focused on measuring seasonal changes in water surrounding the reefs. Measurements of nutrients, chl-α, and total suspended solids were taken from the same fixed stations as Experiment 1 (Figure 3). In the last days scheduled for this trip, a large tropical low pressure system passed over Cape Leveque, requiring the research team to evacuate and ending this trip slightly early.

2.4 Field experiment #3 (22nd March – 12th April 2014)

A third field experiment was conducted on Tallon reef at the end of the wet season, involving 5 UWA staff and students. This experiment involved collaborative work with the Bardi Jawi Rangers, with Rangers participating on most days of the work. In addition, this experiment was conducted alongside an intensive hydrodynamic
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study of the reef between UWA and CSIRO as part of project 2.2.1 on reef hydrodynamics, thereby providing very comprehensive hydrodynamic dataset that provided an opportunity to quantify in further detail how the reef hydrodynamics modify reef community productivity and nutrient fluxes in this project.

During these field experiments, the following instruments were deployed (Figure 3):

- 3 dissolved oxygen sensors;
- 2 optical fluorescence (chl-a) and turbidity loggers;
- 1 downwelling light sensor;
- 2 acoustic Doppler current profilers (plus 5 additional current measurements from 2.1.1);
- 2 pressure sensors (plus 8 additional pressure sensor measurements from 2.1.1);
- 64 temperature loggers; and
- ISCO automated water sampler (Figure 7).

Water sampling was conducted similar to field experiment #1, where samples were collected for the same parameters at fixed stations both across and off the reef, and with Lagrangian measurements from drifters (n=16). In addition, an automated water sampler was mounted on top of scaffolding (deployed for support of a weather station for project 2.2.1) to take water samples overnight for improved estimates of diel nutrient and chlorophyll-a fluxes (Figure 7). Wind speed and direction were measured 10 m above the reef during this study period.

In addition, during this experiment we deployed a very high resolution array HOBO U22 Water Temp Pros (Onset) temperature loggers over the reef in a regular grid, in order to assess the drivers of spatial patterns of temperature variability and extremes, and how it relates to patterns of benthic productivity (Figure 8). Following Lentz et al. (2013), each temperature logger was calibrated in a controlled temperature water bath both before and after the deployment, such that the root-mean-squared errors were estimated to be <0.1°C in all cases. The instruments were deployed on top of small lead weights (sampling ~0.1 m above the bed), and were also wrapped in white tape to minimize any solar heating.
2.5 Laboratory analyses

Samples of water column nutrients and chl-\(a\) from every field experiment arrived in Perth still fully frozen. At the laboratory, TSS filters were re-weighed to determine suspended solid concentration. Within one month of collection, chl-\(a\) filters were thawed, extracted in the dark with 100% acetone, sonicated, filtered, and analysed on a fluorometer (Turner Designs). Nutrient concentrations from water samples were determined through automated flow-injection analysis (LACHAT Quick Chem 8000). Particulate nitrogen filters were analysed using a Shimadzu CN analyser. Particulate phosphorus samples were digested in hot persulfate and concentrations determined colorimetrically on a spectrophotometer (Shimadzu UV-Vis).

Differences in coastal water quality among seasons (Dry, Wet, late Wet) were tested using water samples taken from the channel adjacent to Tallon Island. Statistical significance (\(\alpha = 0.05\)) was determined with one-way analysis of variance (ANOVA), where data met assumptions of homoscedasticity and normality. Tukey-Kramer adjusted least-squared means were calculated and all possible pair-wise comparisons among seasons computed.

2.6 Community productivity estimates

The northern reef platform experienced flow in a consistent northeasterly direction during ebb tide (80° ± 30°, mean ± standard deviation) for ~10 hours each tidal cycle, with flow speeds becoming negligible as water depth became very low (<0.4 m) (Lowe et al. 2015). This prolonged period of unidirectional flow made the reef platform an ideal natural setting to capture community-scale (100s of meters) benthic oxygen fluxes. A one-dimensional control volume approach was taken with loggers placed along the major axis of ebb tide flow.

Depth-averaged flow speed along the major axis of the instrument transect (\(u_x\)) was available for all but ~5% of the field study, when water depth approached instrument blanking distance (\(h <0.4\) m). For periods of low \(h\), \(u_x\) was estimated from a significant linear relationship (\(p<0.001, R^2 = 0.79\)) between along-transect transport \(q_x\) and change in depth \(dh/dt\) (implied from conservation of mass, assuming horizontal dispersion was negligible) for 30-min averaged data from the Aquadopp and drifters (Figure 9). Transport was

\[
q_x = u_x h
\]
Time-series of benthic oxygen flux \( \dot{J}_{O2} \), in mmol O\(_2\) m\(^{-2}\) hr\(^{-1}\) were computed for the dominant communities (seagrass and macroalgae) from (Falter et al. 2011)

\[
\dot{J}_{O2} = \frac{d\bar{C}}{dt} + q_x \frac{dC}{dx} - k_{O2}(C_{sat} - \bar{C}). \quad (2)
\]

The first term on the right side of the equation represents the local accumulation in oxygen concentration, where \( \bar{C} \) is mean oxygen concentration along the transect (RDO1 and RDO2 for seagrass, RDO2 and RDO3 for macroalgae). The second term is the advective oxygen flux, where \( \frac{dC}{dx} \) is the gradient in oxygen concentration over the transect (\( dx \) is 270 m for both transects). The final term is the air-sea gas flux, where \( k_{O2} \) is the gas transfer velocity and \( C_{sat} \) is oxygen saturation estimated from water temperature and salinity (Weiss 1970). Previous work (Zappa et al. 2003, Borges et al. 2004) demonstrated that wind speed has a small effect on \( k_{O2} \) when speeds are low (<4 m s\(^{-1}\)), which occurred during the majority (~75%) of both field study periods. Estimates of gas transfer velocity that describe turbulent open channel flow were therefore more analogous to the reef platform than open-ocean studies. The reference gas transfer velocity \( k_{O2,ref} \) was

\[
k_{O2,ref} = k_w + k_a, \quad (3)
\]

the sum of gas transfer velocities due to water flow \( k_w \) and wind \( k_a \) (Chu & Jirka 2003). Flow-induced gas transfer was

\[
k_w = 2.15\left( u_s^3 / h \right)^{0.25}, \quad (4)
\]

where \( u_s \) is bottom shear velocity defined as

\[
u_s = u_x(C_D/2), \quad (5)
\]

and the drag coefficient \( C_D \) for the reef platform was taken as 0.02 (Lowe et al. 2015). Wind-induced gas transfer velocity \( k_a \) was

\[
k_a = 0.00183u_s^{2.25} \quad (6)
\]
where \( u_{a} \) is air surface shear velocity

\[
u_{a} = 0.01u_{10}\sqrt{(8 + 0.65 \times u_{10})}
\]

(7)

estimated from the wind speed at 10 metres \( (u_{10}) \) (Chu & Jirka 2003). Reference gas transfer coefficient \( k_{O2ref} \) was corrected to in situ conditions using estimated Schmidt numbers for oxygen under reference \( (S_{C2ref}) \) and in-situ \( (S_{C2}) \) conditions as

\[
k_{O2} = k_{O2ref}(S_{C2ref}/S_{C2})^{n}
\]

(8)

where \( n = 2/3 \) for a smooth air-water interface (Wanninkhof 1992). Schmidt numbers were estimated from a linear relationship with temperature (Ramsing & Gundersen 1994) as commonly-used estimates (Wanninkhof 1992) are only valid for temperatures <30°C. As wind speed measurements were not available for Oct 2013, the median wind speed from Apr 2014 was used to estimate \( k_{O2} \).

Logged data were averaged at 30 min intervals for calculations of local and air-sea components of \( J_{O2} \). The time-scale for averaging the advective flux term needed to reflect the transit time of a water parcel between oxygen loggers, which increased over the ebb tide period as \( u_{a} \) slowed and \( h \) decreased. For depths \( h > 0.90 \) m, the signal strength of \( J_{O2} \) was diluted by overlying water, so these periods were excluded from flux estimates. For \( 0.90 > h > 0.70 \) m, the advective flux term was estimated at 30 min intervals (Figure 10), which was roughly the transit time between loggers. As \( u_{a} \) slowed and \( 0.70 > h > 0.50 \) m, the advective flux term was estimated on a longer 80 min interval and linearly interpolated to the common 30 min interval used for all other terms. Towards the end of ebb tide, the depth became low \( (h < 0.5 \) m) and \( u_{a} \) became very slow (~1.5 cm s\(^{-1}\)); \( J_{O2} \) became dominated by the local accumulation term, though the advective term was included and estimated with 30 min averaged data. This method allowed for direct estimates of \( J_{O2} \) for ~50% of each field experiment. Due to the daily advancement of tides by 50 min, \( J_{O2} \) estimates covered a range of temperature and light conditions characteristic of this reef platform.

Daytime net production (NP in mmol O\(_2\) m\(^{-2}\) hr\(^{-1}\)) was defined as \( J_{O2} \) occurring between sunrise and sunset (PAR > 5 \( \mu \)E m\(^{-2}\) s\(^{-1}\)), while nighttime community respiration rate (CR\(_{dark}\) in mmol O\(_2\) m\(^{-2}\) hr\(^{-1}\)) was oxygen flux occurring between sunset and sunrise (PAR < 5 \( \mu \)E m\(^{-2}\) s\(^{-1}\)). Gross primary production rate (GPP in mmol O\(_2\) m\(^{-2}\) hr\(^{-1}\)) was defined as
Benthic community production and response to environmental forcing

\[ \text{GPP} = \text{NP} + \text{CR}_{\text{light}} \]  \hspace{1cm} (9)

where \( \text{CR}_{\text{light}} \) is the daytime community respiration rate (Staehr et al. 2012). Although it is common practice in metabolism studies to assume \( \text{CR}_{\text{light}} = \text{CR}_{\text{dark}} \) (Cole et al. 2000), this assumption is unlikely to be valid given the extreme diel variations in temperature, flow, and DO characteristics of this site. It was therefore important to find an appropriate way to estimate \( \text{CR}_{\text{light}} \) given the above forcing factors.

The dependency of \( \text{CR}_{\text{dark}} \) on flow was examined with the maximum uptake rate coefficient of oxygen from mass transfer limitation (\( S_{\text{MTL}} \)) as (Falter et al. 2004),

\[ S_{\text{MTL}} = \frac{\sqrt{u_x}}{2 \frac{c_f}{Sc_{O_2}} \frac{v}{D_{O_2}}} \]  \hspace{1cm} (10)

where \( c_f \) is the friction coefficient, the Schmidt number of \( O_2 \) (\( Sc_{O_2} \)) was the kinematic viscosity (\( v \)) divided by the diffusion constant of oxygen, and the Reynolds roughness number (\( Re_k \)) is defined as

\[ Re_k = \frac{u^* \kappa_k s}{v} \]  \hspace{1cm} (11)

where \( \kappa_k \) is a bottom roughness length scale, assumed to be 0.5 m for the reef platform (Lowe et al. 2015). In reefs systems where water depth approaches the maximum reef organism height, \( c_f \) increases dramatically and can be estimated from empirical relationships to the ratio of \( h \) and maximum reef height (McDonald et al. 2006). First-order oxygen uptake rate coefficients (in m day\(^{-1}\)) for measured fluxes of \( \text{CR}_{\text{dark}} \) [where \( i = \text{SG} \) (seagrass) or \( \text{AC} \) (macroalgae)] were then estimated as (Thomas & Atkinson 1997)

\[ k_i = \frac{R^2 C_{O_2}^2}{C_{\text{C}}} \times 24 \]  \hspace{1cm} (12)

and could be compared to \( S_{\text{MTL}} \).

Respiration has an Arrhenius relationship with temperature often described by \( Q_{10} \), the factor by which \( CR \) would increase if temperature were increased by 10°C (Berry & Raison 1981); in most aquatic systems (Valiela 1995) and for seagrass on Tallon reef platform (Pedersen et al. 2016), \( Q_{10} \approx 2 \). The degree to which temperature (\( T \)) changes likely affected \( CR \) was assessed by computing a temperature-adjusted version of \( \text{CR}_{\text{dark}} \) (\( CR_T \)) as (Burd & Dunton 2001)

\[ CR_T = CR_{\text{dark}} + [CR_0 - CR_0 e^{\zeta (T - T_0)}] \]  \hspace{1cm} (13)

where \( CR_0 \) and \( T_0 \) were the mean \( \text{CR}_{\text{dark}} \) and temperature, respectively, when fresh ocean water was present on the reef at nightfall and \( \zeta = \log_e(Q_{10})/10 \).

The introduction of \( CR_T \) allowed analysis of patterns in \( \text{CR}_{\text{dark}} \) with the effect of temperature removed. A linear correlation between DO and \( CR_T \) was present in both reef communities (Atkinson et al. 1994) and was used to estimate \( \text{CR}_{\text{dark}} \) for instances of nighttime flood tides and \( \text{CR}_{\text{light}} \) using daytime DO levels (Figure 16). A saturating Monod-type relationship may also be appropriate (Zimmerman et al. 1989) and was plotted for completeness (Figure 16), though not used to estimate \( CR \).

The relationship between GPP and irradiance was described by a hyperbolic tangent function (PI curves) (Jassby & Platt 1976). As discussed above, estimates of \( J_o2 \) were not available during rising tide or periods of high water depth (\( h > 0.90 \) m), so GPP was estimated from PI curves using irradiance. Daily GPP (in mmol O\(_2\) m\(^{-2}\) d\(^{-1}\)) was then calculated as the sum of all GPP (measured and estimated) in each day. Daily \( CR \) (in mmol O\(_2\) m\(^{-2}\) d\(^{-1}\)) was similarly the sum of measured and estimated \( CR \) in each day. Net community production (NCP, in mmol O\(_2\) m\(^{-2}\) d\(^{-1}\)) was daily GPP less daily \( CR \).

Uncertainty in metabolism estimates due to random sampling error and statistical modelling error was quantified using the Monte Carlo method with 10000 sets of input variables. For variables directly measured in this study (\( T, h, \text{DO}, \text{PAR}, u_n, u_{10} \)), the standard deviation was drawn from the normally-distributed measurements used to create 30 min bin averages. For variables derived from statistical relationships (\( \text{CR}_{\text{light}}, \text{GPP}/\text{CR}_{\text{dark}} \) during high water depth), standard error of regression coefficients were used.
2.7 Reef heat budget analysis for Tallon reef

To assess the physical drivers of temperature variability over the reef, a reef heat budget model was derived from the integral form of the thermal energy equation that considers changes in the water temperature \( T \) within the reef, heat exchange by the advection of water on/off the reef, and the net surface air-sea heat fluxes \( Q_{\text{net}} \) (Kundu & Cohen 1990, McCabe et al. 2010):

\[
\frac{d}{dt} \int_V T \, dV + \int_{A_{\text{open}}} T \, u \, dA = -\frac{1}{\rho_0 c_p} \int_{A_{\text{net}}} Q_{\text{net}} \, dA ,
\]

which were evaluated over the reef control volume shown in Figure 11. Here \( \rho_0 \) is the seawater density, \( c_p \) is the specific heat capacity of seawater, \( A_{\text{open}} \) is the vertical perimeter area along the reef edge, \( A_{\text{reef}} \) is the plan area of the reef platform area and \( V \) is volume of water on the reef. Following McCabe et al. (2010), we simplify the model by assuming 1) the temperature is approximately uniform throughout the reef, i.e. \( T(x,y,z) \approx T_r \) where \( T_r \) is a representative (spatially-averaged) reef temperature, and 2) that \( Q_{\text{net}} \) is likewise approximately spatially-uniform and that incoming water from of temperature \( T_{\text{in}} \) rapidly mixes with water on the reef.

Meteorological data was obtained using the weather station mounted on the scaffolding deployed on the reef (Figure 7). The weather station measured wind speed and direction, air temperature, relative humidity, barometric pressure and net solar radiation (via a radiometer) every minute. The radiometer was deployed on a horizontal pole \( \sim 1 \) m away from the scaffolding, and measured the net incoming and outgoing shortwave and longwave (infrared) radiation over the reef water surface \( Q_{\text{sw+lw}} \). Following an approach similar to Zhang et al. (2013), the meteorological and water temperature data were used to compute the individual terms that comprise net surface heat fluxes \( Q_{\text{net}} \), i.e.:

\[
Q_{\text{net}} = Q_{\text{sw+lw}} + Q_{\text{in}} + Q_{\text{sb}}
\]

where \( Q_{\text{in}} \) is the latent heat flux and \( Q_{\text{sb}} \) is the sensible heat flux. The latent and sensible heat flux terms were calculated using the COARE (Coupled Ocean Atmosphere Experiment) bulk algorithms (Fairall et al. 2003), as detailed in Zhang et al. (2013).

For a quasi-1D reef where water flows in/out of the reef approximately uniformly along the edge, the incoming/outgoing flow speed \( U_{\text{in/out}} \) can be estimated from the rate of change of the water level on the reef \( h_r \) according to mass conservation:

\[
U_{\text{in/out}} = \frac{A_{\text{reef}}}{W_r (h_r - h_{\text{MSL,L}})} \frac{dh_r}{dt}
\]

where \( W_r \) is the reef perimeter width and \( h_{\text{MSL,L}} \) is depth of the back reef (or lagoon) relative to the reef crest (see Figure 1). With Eq. (16) and these uniform-property assumptions, the evaluation of the integrals in Eq. (14) lead to the following governing reef heat balance equation

\[
\frac{d}{dt} (T_r h_r) - T_{\text{in/out}} \frac{dh_r}{dt} = \frac{Q_{\text{net}}}{\rho_0 c_p}
\]

subject to the inflow/outflow temperature conditions

\[
T_{\text{in/out}} = T_0 \quad \text{if} \quad \frac{dh_r}{dt} \geq 0
\]

\[
T_{\text{in/out}} = T_r \quad \text{if} \quad \frac{dh_r}{dt} < 0
\]
where $T_0$ is the surrounding (offshore) temperature and $T_r$ is the reef water temperature.

Equation (18) was used to assess the reef temperature variability on Tallon reef and was also used to investigate the broader role of tides on reef heat balances. For Tallon reef, $T_r$ and $h_r$ were based on the spatially-averaged temperature and water depth, measured respectively by instruments on the reef, $T_0$ was from the temperature measured by instruments on the reef slope and $Q_{net}$ was from the measured net heat fluxes. The numerical solution for $T_r$ was obtained using Euler forward differencing.

Figure 11. (a) Schematic diagram of a reef cross-section identifying the reef depth $h_r(t)$ that varies between a maximum amplitude ($\eta_{tide}$) above the depth $h_{MSL}$ at mean sea level (MSL) and the minimum depth $h_{min}$ of the lagoon/platform (refer to the text for details). The total depth $h_r$ at the back of the reef (or lagoon) is comprised of the depth over the crest ($h_{MSL,C}$), if present, and the depth of the back of the reef relative to the crest ($h_{MSL,L}$). (b) Oblique view of a reef with surface area $A_{reef}$ where water instantaneously exchanges over an area $A_{open}$ between the reef and the ocean, with flow velocity $u_{in/out}$ and temperature $T_{in/out}$. Air-sea heat exchange on the reef occurs through $Q_{net}$.

2.8 General model of temperature variability in tide-dominated reefs

To more generally investigate how properties of the tide, solar cycle and reef morphology interact to regulate temperature variability within a wide range of reef systems, we recast the governing heat budget equations (Eqs. 17 and 18) into their dimensionless forms. To further simplify our analysis and identify a set of fundamental dimensionless parameters that control temperature variability, we consider idealized tidal and solar heating cycles that nonetheless retain the periodicity and magnitude exhibited by our in situ data.

Variation in reef tidal depth ($h_r$) is thus assumed to occur at a tidal period of $\tau_{tide}$ with tidal amplitude $\eta_{tide}$ (equivalent to one-half the range) and phase difference $\phi_0$ relative to solar noon. The offshore mean sea level coincides with a reef depth $h_{MSL}$ and we assume that there is a minimum water depth $h_{min}$ where water is effectively trapped on the reef (Figure 11), i.e. this reef water depth variability is governed by

\[
h_r(t) = h_{MSL} + \eta_{tide} \cos \left( \frac{2\pi t}{\tau_{tide}} + \phi_0 \right) = h_{MSL} + \eta_{tide} \cos \left( 2\pi \frac{t}{\tau_{tide}} + \phi_0 \right)
\]

with minimum depth $h_{min}$ (such that $h_r \geq h_{min}$ always), leading to tidal truncation for reefs with $h_{MSL} \leq h_{min} + \eta_{tide}$. In this general model, we consider cases where the minimum depth $h_{min}$ may be due to either water ponding within a shallower reef crest (in which case $h_{min}$ is roughly equal to the mean lagoon
depth $h_{\text{MSL,t}}$ in Figure 11) or due to bottom friction restricting the flow of water off the reef (as in the case of Tallon reef). Likewise we assume a sinusoidal variation in the net surface heat flux term $Q_{\text{net}}$ occurring with period $\tau_{\text{solar}}$

$$Q_{\text{net}}(t) = Q_{\text{net,max}} \cos \left( \frac{2\pi t}{\tau_{\text{solar}}} \right) \quad t_{\text{sunrise}} \leq t \leq t_{\text{sunset}}$$

$$Q_{\text{net}}(t) = -Q_{\text{net,night}} \quad \text{at night}$$

(20)

where $Q_{\text{net,max}}$ and $Q_{\text{net,night}}$ are the magnitudes of the maximum net daytime heating (mostly short and longwave radiation) and constant nighttime net cooling rates (mostly latent), respectively (see below). While this is an idealized representation of the diurnal heating cycle (see Results), it nonetheless captures the dominant diurnal behavior of the physics driving net heat fluxes within reefs (including Tallon), and has also been successfully used to model heat budgets and temperature variability within other reefs, such as in the GBR (McCabe et al. 2010).

With these definitions, we can further simplify our general heat budget model by constructing dimensionless variables (denoted with the *) for time ($t^*$), reef water depth ($h^*_r$) and temperature ($T^*$) to reduce the number of free parameters needed to constrain the prevailing physics:

$$t^* = \frac{t}{\tau_{\text{solar}}}, \quad h^*_r = \frac{h_r}{\eta_{\text{ride}}}, \quad Q^*_{\text{net}} = \frac{Q_{\text{net}}}{Q_{\text{net,max}}}, \quad \tau^*_r = \frac{T_r - T_0}{Q_{\text{net,max}} \tau_{\text{solar}} \left( \rho_c c_p \eta_{\text{ride}} \right)}$$

(21)

where $c_p$ and $\rho$ are the specific heat capacity and density of seawater, respectively, and by definition $\tau^*_r$ is equivalent to one (solar) day. Substituting dimensionless variables into the governing Eqs. (17) and (18) then gives:

$$\frac{d(T^* h^*_r)}{dt^*} - T^*_{\text{in/out}} \frac{dh^*_r}{dt^*} = Q^*_{\text{net}}$$

(22)

with

$$T^*_{\text{in/out}} = 0 \quad \text{if} \quad \frac{dh^*_r}{dt^*} \geq 0$$

$$T^*_{\text{in/out}} = T^*_r \quad \text{if} \quad \frac{dh^*_r}{dt^*} < 0$$

(23)

where $T^*_r$ and $T^*_{\text{in/out}}$ are the dimensionless forms of the reef water temperature and inflow/outflow temperature, respectively. For the idealized (sinusoidally varying) tidal forcing considered in Eq. (19), the reef water depth in dimensionless form becomes

$$h^*_r(t^*) = h^*_{\text{MSL}} + \cos \left( \frac{2\pi t_{\text{solar}} + \phi_t}{\eta_{\text{ride}}} \right)$$

(24)

such that tidal truncation occurs when $h^*_r \leq h^*_{\min} + 1$. From Eq. (20), the net surface heat flux term sinusoidally varies during the day (reaching a maximum $Q_{\text{net,max}}$) and is constant at $-Q_{\text{net,night}}$ night; the constant night cooling rate was chosen to be equal to 25% of the daytime maximum warming rate ($Q_{\text{net,max}}$), which is typical of many tropical reef systems worldwide (McGowan et al. 2010, Davis et al. 2011, Zhang et al.
2013), including for Tallon reef in the present study (see Results). Therefore, the dimensionless net surface heat flux term becomes

\[ Q_{\text{net}}^* (t^*) = \cos(2\pi t^*) \]

\[ t_{\text{sunrise}}^* \leq t^* \leq t_{\text{sunset}}^* \]

\[ Q_{\text{net}}^* (t^*) = -0.25 \quad \text{at night} \]

Equations (22)-(25) can be solved using numerical integration to predict temperature variability on the reef as a function of the three dimensionless input parameters \( \tau_{\text{tide}}^*, \tau_{\text{solar}}^*, h_{\text{MSL}}^*, h_{\text{min}}^* \) defined in Eq. (21).

## 3 Results

### 3.1 Reef communities

Tallon Island reef supported two distinct communities atop the reef platform: seagrass and macroalgae. The seagrass community was larger in area, stretching from the shoreline to ~400 m from the reef crest, and was dominated inshore by *Enhalus acoroides* (L.f.Royle) and offshore by *Thalassia hemprichii* (Ehrenb.) Aschers (Wells et al. 1995). The macroalgal community covered a band ~200 m wide at the reef crest and was dominated by brown macroalgae (*Sargassum* spp.) but also contained coralline algae, many macroalgal species, and small corals. A mixed assemblage of sparse *Thalassia* and *Sargassum* was found in the sandy ~200 m band separating seagrass and macroalgal communities (Figure 3).

### 3.2 Hydrodynamic context (current and water level variability)

A detailed study of tide-dominated reef hydrodynamics, including the hydrodynamics of Tallon reef, was the focus on Project 2.1.1. Therefore, only a brief summary of the main features of the current and water level variability on the reef are described here.

Field measurements captured a full spring-neap tidal cycle in both seasons, with tidal ranges of ~7 m during springs and ~2.5 m during neaps (Table 1). Water depth on the reef platform \( (h) \) ranged from 3.4 - 0.3 m during spring tides and from 0.3 – 1.3 m during neap tides (Figure 13a). Due to reef platform elevation 0.3 m above mean sea level, there were large asymmetries in duration of tide phases (Lowe et al. 2015). Ebb tide was elongated to ~10 h (Figure 12), with flow speed tapering from ~0.4 m s\(^{-1}\) to 0.015 m s\(^{-1}\) over the ebb period (Figure 10). Wind speed \( (u_{10}) \) was generally low (Figure 13b) with a mean of 3.3 m s\(^{-1}\) (Table 1) and wind-driven waves did not occur due to the sheltering of Tallon Island by surrounding islands.

<table>
<thead>
<tr>
<th>Date</th>
<th>( n )</th>
<th>Tidal range (m)</th>
<th>Wind speed ( u_{10} ) (m s(^{-1}))</th>
<th>Temperature (°C)</th>
<th>DO (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Spring</td>
<td>Neap</td>
<td>Mean ± SD</td>
<td>Mean ± SD</td>
</tr>
<tr>
<td>Oct 2013</td>
<td>14</td>
<td>6.7</td>
<td>2.6</td>
<td>nd*</td>
<td>28.5 (± 2.00)</td>
</tr>
<tr>
<td>Apr 2014</td>
<td>12</td>
<td>7.3</td>
<td>2.7</td>
<td>3.3 (± 1.56)</td>
<td>30.8 (± 1.47)</td>
</tr>
</tbody>
</table>

Table 1. Summary of environmental conditions on Tallon reef platform during both field experiments. *no data available
Environmental conditions on the reef platform were characterised by large diel changes, the magnitude of which was determined by the position of the solar cycle relative to the tidal cycle. Light levels were generally high, reaching 1800 €\mu$E m$^{-2}$ s$^{-1}$ when low water depth ($h < 0.4$ m) coincided with approximately mid-day. Water temperature on the reef platform was on average ~2° C warmer in the wet season than during the dry, though a wide range of temperatures were recorded during both seasons (Table 1). Temperature increases of >10° C within a single tidal cycle occurred when low tide coincided with high light levels (Table 1). Diel oxygen concentrations varied up to 440 €\mu$M (Figure 13c) and became supersaturated (> 100%) when low tide coincided with high light levels, reaching up to 280% saturation. Water column hypoxia (DO < 63 €\mu$M) developed on the reef platform each night for 1 - 7.5 h (Figure 13c), depending on the overlap of low tide and nighttime.
Figure 13. Time-series of conditions on Tallon Island reef during the wet season field experiment including: a) water depth on the reef platform (h, in black) and current speed in the along-transect direction (\(u_x\), in blue); b) wind speed and direction at 10 m (\(u_{10}\)); c) dissolved oxygen concentration (blue) at RDO1 (seagrass site) with reef water depth (black); and d) water temperature (red) at RDO1 (seagrass site) with reef water depth (black). Red dashed line indicates hypoxia (DO < 63 µM).

3.3 Benthic community productivity

Environmental conditions and responses in benthic community metabolism were similar between seasons. In order to focus on metabolic response to environmental forcing, most results presented are from the wet season (Field Experiment #3) with implications of seasonality discussed towards the end. The relative importance of the three benthic oxygen flux components (local, advective, and air-sea exchange) to \(J_{O2}\) differed between benthic community zones (seagrass versus algae). In the seagrass community, overall \(J_{O2}\) estimates were dominated by the local accumulation term, with advective and air-sea components each contributing <10% (Figure 14A). Local and advective terms were both important to \(J_{O2}\) estimates in the macroalgal community (~55% and 35%, respectively), while air-sea exchange again contributed <10% (Figure 14B). Despite playing a minor role in overall \(J_{O2}\) estimates, air-sea exchange became important for short periods in the diel cycle, (e.g., when water column DO became supersaturated and flow was > 2 cm s\(^{-1}\)) (Figure 14B), and was therefore retained in the calculations.
Hourly rates of reef community daytime net production and nighttime respiration showed distinct diel patterns (Figure 14C). The daily advancement of the tidal cycle by 50 min each day and length of both field studies resulted in metabolism estimates from a wide range of flow, temperature, DO, and light conditions characteristic of this reef platform. Comparison of first-order oxygen uptake rate coefficients derived from CRdark ($k_{SG}$ and $k_{AC}$) and those based on the limits of mass transfer ($S_{MTL}$) showed that CRdark approached mass transfer limitation when flow speeds were low (Figure 15), but was not strongly related to flow speed in general.
Rates of CRdark tended to decline through the night, and approached zero when nighttime overlapped fully with ebb tide (Figure 14C). Water temperatures on the reef platform cooled by up to 4°C during these periods due to air-sea heat exchange. However, estimates of temperature-adjusted CRdark (CRt based on a $Q_{10} = 2$) demonstrated that only <8% of the pattern of decline observed in CRdark could be attributed to falling temperatures. A strong linear trend was observed between CRdark and DO in seagrass and macroalgal communities ($p < 0.0001$, $R^2 = 0.73$ and 0.61, respectively) over the range of DO (8 – 300 µM O₂, 10 – 150% saturation) measured (Figure 16). To assess whether depletion of labile photosynthates over the night could have contributed falling CRdark (Figure 14C), this DO/CR relationship was also limited to only estimates from the first 2 hours after sunset, yet the linear trend remained similar.
estimates of net daytime production (NP) from field measurements (as opposed to statistical relationships) revealed responses related to DO (Figure 17). Hourly net production rates were greater overall in the macroalgal community than the seagrass community (Figure 17). Both communities showed declining rates of NP when DO was high (>350 µM), which typically occurred in the afternoon and indicated some hysteresis in NP.

Daytime community respiration rates (CR_{light}) were estimated based on linear relationships with DO (Figure 16) and used to calculate rates of hourly gross primary production (Figure 18). Like NP estimates, the macroalgal community showed greater rates of GPP than seagrass at all light levels. Photosynthesis-irradiance relationships using GPP exhibited a classic hyperbolic tangent form, with no hysteresis evident (Figure 18). This indicated that the pattern of hysteresis observed in NP at high light levels (Figure 17) was driven by increases in daytime CR relative to photosynthesis.
Benthic community production and response to environmental forcing

Mean daily rates of metabolism (the combination of field measurements and estimates from statistical models) showed similar net community production between seagrass and macroalgae (Table 2). The macroalgal community had higher overall rates of both GPP and CR than seagrass, yet both communities had balanced P:R ratios over the whole experiment (Table 2).

<table>
<thead>
<tr>
<th>Community</th>
<th>n</th>
<th>NCP (±)</th>
<th>CR (±)</th>
<th>GPP (±)</th>
<th>P:R</th>
</tr>
</thead>
<tbody>
<tr>
<td>Seagrass</td>
<td>12</td>
<td>50</td>
<td>440</td>
<td>490</td>
<td>1.1</td>
</tr>
<tr>
<td>Macroalgae</td>
<td>12</td>
<td>30</td>
<td>670</td>
<td>700</td>
<td>1.0</td>
</tr>
</tbody>
</table>

Table 2. Summary of daily net community production (NCP), community respiration (CR), and gross primary production (GPP) during the wet season. Daily fluxes (mean of n days) in units of mmol O₂ m⁻² d⁻¹ (± standard error). Ratio between daily GPP and CR (P:R) also shown.

NCP also varied on a daily basis due to changes in environmental forcing (Figure 13) driven by the mismatch between the length of tide and light cycles (50 min daily). This is best visualized by plotting daily NCP as a function of the phase difference between light and tide on that day (Figure 19). A phase difference of 0° represents a day when high tide was aligned with solar noon/midnight, and thus ebb tide occurred from ~12:00 – 22:00 and 0:00 – 10:00. A phase difference of 180° represents a day when high tide was aligned with ~18:00/6:00, and thus ebb tide occurred from ~18:00 – 04:00 and ~6:00 – 16:00. Maximum daily NCP was observed near “0” and decreased as tide-light phase difference advanced, reaching a minimum near “180°” (Figure 19). A phase difference of 180° means that the water depth was low when the highest light levels occurred, resulting in large increases in DO and temperature on the reef platform. This strong cosinusoidal pattern in NCP was seen in both communities over wet and dry seasons (Figure 19).
3.4 Nutrient and chlorophyll-α dynamics

Changes in dissolved nutrient concentration due to uptake and release by benthic communities were most readily measureable during slow flow speeds ($u_e$) occurring in low tide periods. In the macroalgal-dominated zone, strong signals of dissolved inorganic nitrogen release as nitrate/nitrite (NO$_x$), and ammonia (NH$_3$) were measured, whereas seagrass communities showed slight nitrogen uptake (decrease in concentration) (Figure 20). In both community zones, phosphate (PO$_4^{3-}$) concentrations remained stable during low tide periods and were similar on and off the reef platform (Figure 20C). Offshore samples were taken from all phases of the tidal cycle, and there was no difference in nutrient concentration with phase of tide.

By utilizing the one-dimensional control volume approach described previously to measure community productivity, we were able to estimate nutrient uptake or release rates during ebb tide periods. Release rates of NO$_x$ increased through the ebb period, reaching a maximum in the late afternoon (Figure 21). Ammonia uptake and release rates were lower overall than NO$_x$ by an order of magnitude. There was net community uptake of NH$_4^+$ early in the ebb tide period, which switched to net release around midday. Phosphate uptake / release rates were negligible.
Figure 20. Representative time-series of water column dissolved a) ammonia, b) nitrate/nitrite, and c) phosphate concentration over an ebb tide period during the dry season. Samples taken from offshore reference (blue), macroalgae-dominated (orange), and seagrass-dominated (green) communities. The depth of water on Tallon reef is shown (black line).
Benthic community production and response to environmental forcing

Using the automated water sampler during Field Experiment #3 in the late-wet season, we were able to measure water quality changes over full diel and tidal cycles. These measurements were used to verify logged data from optical chl-\(\alpha\) sensors deployed on the reef platform. Suspended chlorophyll-\(\alpha\) (mainly from phytoplankton, but also including small plant-derived detritus) is a primary food source of reef filter-feeding organisms. Chlorophyll-\(\alpha\) concentration showed substantial variability over each tidal cycle, with peaks in concentration occurring near peak flood and ebb velocities, presumably due to resuspension and mixing on the reef platform (Figure 22). Flood tides also flushed fresh suspended labile material onto the reef flat, which was then grazed down almost to zero during ebb tide (Figure 22).

Seasonal differences in water quality in the vicinity of the reef platform were evident despite strong mixing in channels surrounding Tallon Island. Concentration of dissolved inorganic nitrogen species (nitrate/nitrite and ammonia) increased significantly through the wet season, while phosphate concentrations did not change significantly (Figure 23A). Chlorophyll-\(\alpha\) and phaeophytin (suspended dead photosynthetic material, mainly comprised of dead phytoplankton) levels also increased significantly through the wet season (Figure 23B). Ratios of chl-\(\alpha\) to phaeophytin indicated that the majority of suspended photosynthetic material was detrital in
3.5 Role of interacting tidal and solar heating cycles on temperature variability

3.5.1 Tallon reef temperature and heat flux observations

During Field Experiment #3 when a detailed reef heat budget for Tallon was quantified, the offshore tidal range reached a maximum of ~7.5 m near the middle of the study during spring (~1 Apr) and a minimum of ~4 m during neap towards the beginning and end (Figure 24a). Water levels over the reef platform were highly asymmetric, with water draining very slowly off the reef during ebb tide, as a result of the large ‘truncation’ of tidal variability over the reef (i.e. the duration of the ebb was ~10 hours while the flood was only ~2 hours) (Figure 24a). Due to the 12.4 hour period of the dominant M2 tide, the time of the offshore low tide minima
gradually shifted forward over the study period. Initially, the first low tide of the day occurred prior to sunrise and gradually shifted to just before noon by the end of the experiment (Figure 24d).

Diurnal variations in the net surface heat fluxes $Q_{net}$ did not vary substantially over the course of the experiment, typically ranging from -200 W m$^{-2}$ at night to +600 W m$^{-2}$ during the day (Figure 24b). This consistency was due to the cloud-free skies that persisted over most of the experiment; conditions that are typical during the ‘dry season’ in the Kimberley (April through October) due to its tropical monsoon climate. Variations in $Q_{net}$ were dominated by net heating from the combined short and long wave radiation ($Q_{sw+lw}$) during the day and net cooling from latent heat fluxes ($Q_{le}$) during the night (Eq. (15)).

Daily temperature fluctuations in reef water temperature $T_r$ (spatially-averaged over the reef platform) ranged from as small as $\sim 2^{\circ}$ C during the initial period of the study, to as large as $\sim 8^{\circ}$ C during the latter period and exhibited fairly complex but periodic behavior (Figure 24c). Nonetheless, our simple reef heat budget model (Eqs. (17)-(18)) was able to accurately reproduce both the shape and magnitude of the complex diurnal temperature reversals over the course of the study. The maximum diurnal temperature fluctuations were greatest towards the end of the study when low tide occurred slightly before solar noon despite the tidal range being lower at this time (Figure 24d). Therefore, the maximum observed temperature fluctuations were not in phase with the maximum daily tidal amplitude (Figure 24a,c); i.e., diurnal temperature fluctuations were not maximal during the largest spring tides. This is because once the offshore sea level drops below the reef crest, the depth and duration of the remaining ebb tide period on the reef is determined solely by the morphology of the reef regardless of how low offshore sea levels reach (i.e., -2 m during neap vs. -4 m during spring). More generally, we find that the phase difference between the semi-diurnal (M2) tide and solar cycles generally has the most substantial influence on the reef heat balance, such that the magnitude of daily temperature variation ($\Delta T = T_{max} - T_{min}$, where $T_{max}$ and $T_{min}$ are the daily temperature maxima and minima) is mostly independent of the spring-neap phasing at Tallon Island.
3.5.2 Model predictions of Tallon reef temperature variability

To further demonstrate the general responses of the model and assess how tides and solar heating cycles drive temperature variability on these macrotidal reefs, we first consider the application to a simplified Tallon reef where \( h_{\text{min}} = 0.5 \text{ m} \) and \( h_{\text{MSL}} = 0 \text{ m} \), the tide consists solely of an M2 mode (the dominant mode at Tallon) such that \( \tau_{\text{tide}} = 12.42 \text{ hr} \), and a tidal amplitude \( h_{\text{tide}} = 3 \text{ m} \) that is halfway between the spring maxima (4 m) and neap minima (2 m); the results of which yield \( \tau_{\text{solar}} / \tau_{\text{tide}} = 0.52, \ h_{\text{MSL}} = 0 \) and \( h_{\text{min}} = 0.17 \) (Table 3). To
assess the variability in $T_{r}^{*}$ over multiple tidal cycles (days), we initialize each simulation with $\phi = 0^\circ$ and advance forward in time, such that there is an additional 0.84 hr (~50 min) lag in the solar cycle relative to the tidal cycle each day. The results reveal the presence of a low frequency modulation of the diurnal fluctuations in $T_{r}^{*}$ with a beat period of $\tau_{\text{low}} = \tau_{\text{solar}} \tau_{\text{tide}} / (2\tau_{\text{tide}} - \tau_{\text{solar}}) \approx 14.8$ days, which are generated by the daily phase drift between the M2 tide and the diurnal solar cycle (Figure 25a). Importantly, while this low frequency modulation of temperature occurs at the same frequency of spring-neap tidal cycles, which are governed by analogous interactions between solar and lunar tidal constituents (Kvale 2006), the temperature variability here is generated solely by interactions between solar heat fluxes and the dominant lunar tides (since the daily tidal amplitude here is constant).

<table>
<thead>
<tr>
<th>Label</th>
<th>Site</th>
<th>$\eta_{\text{tide}}$</th>
<th>$h_{\text{min}}^{*}$</th>
<th>$h_{\text{MSL}}^{*} = h_{\text{MSL}} / \eta_{\text{tide}}$ mean</th>
<th>present</th>
<th>MSL +0.7 m</th>
<th>MSL +1.5 m</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Tallon Island, Kimberley, northwestern Australia</td>
<td>3.0 m</td>
<td>0.17</td>
<td>0.00 0.23 0.50</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>B</td>
<td>Warraber Island, Torres Straight</td>
<td>1.2 m</td>
<td>0.33</td>
<td>0.33 0.92 1.58</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>Cocos (Keeling) Islands, eastern Indian Ocean</td>
<td>0.6 m</td>
<td>0.42</td>
<td>0.42 1.58 2.92</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>Lady Elliot, Great Barrier Reef</td>
<td>0.8 m</td>
<td>0.50</td>
<td>0.50 1.38 2.38</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>Ofu, American Samoa</td>
<td>0.5 m</td>
<td>2.00</td>
<td>2.00 3.40 5.00</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>Rarotonga, Cook Islands</td>
<td>0.4 m</td>
<td>2.50</td>
<td>2.50 4.25 6.25</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

Table 3. Tidal amplitudes and reef morphology parameters for Tallon reef and other example tidally-forced reefs globally. Values of dimensionless reef depth relative to mean sea level ($h_{\text{MSL}}^{*}$) are included for present conditions (0 m MSL) and two sea level rise scenarios (+0.7 m MSL, +1.5 m MSL).

We can use the Tallon reef example ($h_{\text{MSL}}^{*} = 0$ and $h_{\text{min}}^{*} = 0.17$) to further investigate how the magnitude of the maximum diurnal temperature fluctuations $\Delta T_{r}^{*}$ (defined as the difference in the daily maximum and minimum of $T_{r}^{*}$) plotted for different instantaneous phase differences $\Delta \phi_{i}$ between the solar and tidal cycles.
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(Figure 25b). Here $\Delta\phi_i$ is defined as the instantaneous phase mismatch between the maximum tide and solar noon on a given day. The maximum dimensionless temperature fluctuation ($\Delta T_r^* \approx 1.8$) occurs when $\Delta\phi_i$ is slightly less than 180° (roughly equivalent to when low tide occurs just before solar noon), which defines a general condition of optimal warming (or cooling) when tidal truncation leads to extended periods of low water depth ($h_t = h_{min}$). Alternatively, we can consider the hypothetical case where offshore mean sea level is sufficiently high enough that tidal truncation does not occur (i.e., the reef platform becomes too deep at low tide); for this example, $h_{MSL}^*$ is increased to $h_{min}^* + 1 = 1.17$ so that the low tide depth is exactly equal to the minimum depth of $h_{min} = 0.5$ m (or $h_{min}^* = 0.17$) for Tallon (Figure 25b). For this alternate case, the maximum dimensionless temperature fluctuations are substantially reduced ($\Delta T_r^* \approx 0.5$), with the maximum warming / cooling for this case instead occurring earlier in the tidal cycle, or when $\Delta\phi_i \approx 120^\circ$. In other words, the combination of a symmetric tidal variation (with reduced low tide duration) and deeper water column leads to much lower temperature variations for a given maximum net heat flux $Q_{net,max}$ and tidal amplitude $\eta_{tide}$.

Figure 25. Response of reef temperature to interacting light and tidal cycles. (a) Reef temperature variability (in dimensionless form $T_r^*$ per Eq. (21)) over 30 days, for an idealized Tallon reef ($h_{MSL}^* = 0$ and $h_{min}^* = 0.17$), illustrating the 14.8 day modulation of the temperature fluctuations caused by the phase drift between the maximum tidal elevation and solar irradiance. (b) The maximum diurnal temperature variation $\Delta T_r^*$ (defined as the difference between the daily maximum and minimum value of $T_r^*$), as a function of the instantaneous phase difference between the tidal and solar cycle ($\Delta\phi_i$). Results shown are for an idealized Tallon reef under present-day sea level ($h_{MSL}^* = 0$) shown in (a), as well as a hypothetical scenario where mean sea level is increased so that the depth at low tide is equal to $h_{min}$ (equivalent to $h_{MSL}^* = 1.17$) and tidal truncation no longer occurs.
Table 4. Tidal amplitudes ($\eta_{tide}$) represent average values (i.e., intermediate between spring and neap). $F_{tide}$ denotes the tidal form factor (see text for details). Diurnal temperature range changes (% $\Delta T_r$ change) are relative to present conditions with 0 m MSL. Present temperature ranges ($\Delta T_r$) are drawn from literature values and projected for different mean sea level rise scenarios (+0.7 m and +1.5 m) using the model.

### 3.5.3 General influence of sea level rise on reef temperature extremes

We further apply the model over a realistic range of reef depths and tidal amplitudes to more generally assess how tides, surface heat fluxes and reef morphology interact to drive reef temperature variability across a broader range of reef morphologies and tidal amplitudes. Using this more global model, we then assess the sensitivity of the diurnal temperature extremes to sea level rise for a number of other reef systems from across the Indo-Pacific region. We consider six example reefs that have both detailed tidal records and accurately-surveyed reef profile bathymetry relative to mean sea level; thus providing us with robust estimates of both $h^*_{\text{MSL}}$ and $h^*_\text{min}$ (Tables 3 and 4). Fortunately, four of these six reefs also have detailed temperature data showing diurnal temperature extremes ranging from 1.5-6.5°C depending on the reef morphology, local tidal amplitude, time of year and weather conditions (Table 4). In particular, we investigate how maximum diurnal temperature fluctuations (in dimensionless form $\Delta T^*_r$) over the 14.8 day cycle respond to variations in the
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Dimensionless reef depth parameters $h_{\text{MSL}}^*$ and $h_{\text{min}}^*$ (Eq. (21)). Prior analyses have shown that tidal truncation occurs only when the depth of the reef falls below a common physical boundary of $h_{\text{MSL}}^* \leq h_{\text{min}}^* + 1$ (Lowe et al. 2015), or in dimensional terms, when $h_{\text{MSL}} \leq h_{\text{min}} + \eta_{\text{tide}}$. The tidal-thermodynamic model reveals that $\Delta T_r^*$ increases when either $h_{\text{MSL}}^*$ and/or $h_{\text{min}}^*$ is decreased. This is mainly due to the net incident heat flux being distributed over a shallower water column (Figure 26). However, we further note that there is a very abrupt increase in $\Delta T_r^*$ when $h_{\text{MSL}}^* < h_{\text{min}}^* + 1$ (i.e., when tidal truncation occurs). This is due to tidal truncation extending the duration of low water depths on the reef, thus elongating periods of enhanced heating/cooling of the shallow water column, and as a consequence, making the response of diurnal temperature variability to a changing sea level highly nonlinear.

To assess how sea level rise will specifically affect temperature extremes within each of these reef environments, we assume that both future tidal conditions and $Q_{\text{net, max}}$ remain unchanged. The former is determined solely by celestial mechanics while the latter is reasonable given that, with the exception of potential regional changes in cloud cover and decadal changes in solar radiation, $Q_{\text{net, max}}$ should remain relatively constant. Two sea level rise scenarios are considered based on predicted end of the century scenarios (see Methods): a 0.7 m rise based on IPCC projections from CMIP5 (Church et al. 2013) and a 1.5 m rise that captures the upper end of semi-empirical projections (see summary by Nicholls et al. 2011). In both cases, these rates of sea level rise far exceed typical reef accretion rates (Quataert et al. 2015). One very important aspect of our simple tidally-driven heat exchange model is that if $Q_{\text{net, max}}$ is kept constant (albeit approximately so), then the ratio of a given temperature extreme ($\Delta T_r$) following sea level rise to present day values is independent of the particular local value of $Q_{\text{net, max}}$, due to the linear form of governing differential equation (Eq. (17)). Thus, although detailed in situ measurements of $Q_{\text{net, max}}$ may not be available for most reef sites, this data is not actually necessary for predicting how much temperature extremes at a given reef site will change as a consequence of sea level rise on a percentage basis.

The results indicate that 0.7 m of sea level rise would reduce present maximum diurnal temperature ranges ($\Delta T_r$) by between 7-65% across the six reef sites while 1.5 m of sea level rise would reduce $\Delta T_r$ by 18-86% (Table 4). Reefs that already experience moderate tidal truncation now (i.e., where the offshore low tidal elevation falls just below the minimum depth) will experience the most substantial reductions in temperatures under future sea level rise. Such examples of reef systems near the ‘tipping point’ include Lady Elliot in the Great Barrier Reef and the Cocos Islands in the eastern Indian Ocean (Figure 26). Thus for reefs such as Lady Elliot where $\Delta T_r$ typically ranges from 2.5-5.5 °C at present, these extremes are predicted to be reduced to 1.5-3.3 °C under 0.7 m of sea level rise and to just 0.5-1.2 °C under 1.5 m of sea level rise.

A more general consequence of these results is how diurnal temperature extremes within different types of reefs worldwide will be affected by a rise in mean sea level depending on their mean depth and regionally-specific tidal amplitudes. From Figure 26a it is clear that those reefs presently located in a regime near the cusp of tidal truncation (i.e. those reefs with $h_{\text{MSL}}^* \approx h_{\text{min}}^* + 1$) will experience the most substantial reductions in temperature extremes under future sea level rise due to the abrupt transition in $\Delta T_r^*$ that occurs when $h_{\text{MSL}}$ becomes deeper than $h_{\text{min}} + \eta_{\text{tide}}$ and tidal truncation ceases, as illustrated in Figure 26b. Given that many coral reef crests are presently located near mean sea level (Falter et al. 2013), it is very common for the offshore sea level to fall below the elevation of many reefs at low tide, and hence experience some form of
tidal truncation regardless of the regional tidal amplitude.

Figure 26. Diurnal temperature extremes for various reefs worldwide and their response to sea level rise. (a) The magnitude of the maximum diurnal temperature fluctuation (in dimensionless form), as a function of the normalized minimum reef water depth ($h_{\text{min}}^*$) and normalized reef depth relative to mean sea level ($h_{\text{MSL}}^*$). Tidal truncation occurs below the black dotted line ($h_{\text{MSL}}^* \approx h_{\text{min}}^* + 1$). The location of four reefs within this parameter space (A-D, see Tables 3 and 4) are shown for three mean sea level scenarios (0 m relative to present; a future +0.7 m rise; a future +1.5 m rise). Ⓐ = Tallon island, Ⓑ = Warraber island, Ⓒ = Cocos Islands, Ⓓ = Lady Elliot island. (b) The response of $\Delta T_r^*$ for the four reefs with fixed $h_{\text{min}}^*$ but varying $h_{\text{MSL}}^*$.

4 Discussion and Conclusions

4.1 Observed water quality patterns and variability

4.1.1 Temperature and dissolved oxygen

Environmental conditions measured in this study easily rank among the most extreme recorded for reef systems worldwide (Kleypas et al. 1999). When low water depths occurred near midday, temperatures on the reef platform reached in excess of 36°C and showed high variability (in some cases >10°C) over a single day (Figure 13d). Extreme reef temperatures (>36°C) have been previously documented in the Red Sea and Arabian Gulf (Coles & Fadlallah 1991, Ateweberhan et al. 2006), and diel fluctuations of ~8°C can occur in the Gulf of Oman (Coles 1997) and shallow backreef environments such as Ofu, American Samoa (Craig et al. 2001) or One Tree Island, Australia (Silverman et al. 2012). Similar patterns have also been recorded in some seagrass studies, as the species present on Tallon reef (E. acoroides and T. hemprichii) often occur in intertidal environments that can experience elevated temperatures (>35°C) on sub-daily timescales (Bridges & McMillan 1986, Collier & Waycott 2014).

This study documented extremes in DO saturation on sub-daily timescales, where supersaturation (up to 280%) occurred with midday low water levels and up to 7.5 hours of hypoxia occurred at night (Figure 13c). Similar ranges in DO have been recorded in shallow backreef areas (Piniak & Brown 2009, Koweek et al. 2015b) and
lagoon waters isolated during semidiurnal low tides (Silverman et al. 2012) or spring low tides (Ohde & van Woesik 1999). Hypoxia is known to occur in the diffusive boundary layer around reef organisms and in organism tissue (Kühl et al. 1995, Ulstrup et al. 2005, Pedersen et al. 2016), but is not typically reported in reef water columns, with very few exceptions (Koweek et al. 2015b) and not to the extent of Tallon reef. The length of both field studies presented here allowed capture of water quality over nearly the full set of tide/light interactions (~14.8 days) (Lowe et al. 2016). Although temperature and DO extremes over the year undoubtedly exceed what is presented here, these water quality patterns can nonetheless be considered representative for this reef.

### 4.1.1 Dissolved nutrients and chlorophyll

Nutrient and chlorophyll-a concentrations showed strong patterns associated with tidal flooding on the reef platform. The chlorophyll-a signal seen in daily time-series indicated the importance of twice-daily tidal inundation for replenishing labile algal and detrital material for reef filter-feeders.

Overall, Tallon reef was found to be oligotrophic (low dissolved nutrients and chl-a), which is normal for tropical waters that do not receive substantial anthropogenic nutrient loads from terrestrial runoff. Nutrient loading negatively impacts coral health in several ways (Szmant 2002, Fabricius 2005), and is one of many stressors causing coral decline in reefs worldwide. Large rainfall events, such as the monsoon or tropical cyclones, cause pulses of nutrients to enter coastal waters; these nutrient delivery events have received much attention in the Great Barrier Reef Marine Park (Devlin & Brodie 2005, Devlin & Schaffelke 2009, Kroon et al. 2012), and hundreds of millions of dollars have already been spent in attempts to ameliorate water quality.

The statistically significant increase in DIN (more than doubled) and chl-a measured through the wet season period (Figure 23) indicated that wet season terrestrial discharge did reach the Sunday Island reef group, although it may have been substantially transformed and remineralised along the way. The fact that a measureable change in nutrients occurred was surprising given the strong mixing, both vertically and between King Sound and offshore waters. Additionally, the 2013/2014 Wet season had low rainfall levels relative to wet seasons from the previous 18 years (Figure 27). Higher rainfall would very likely result in larger pulses of labile terrestrial material to King Sound than what was measured here. A great deal more research is necessary before we can understand how coastal water quality is affected by the wet season in the Kimberley region.

![Figure 27](image)

Figure 27. Fitzroy River discharge measured at Willare (site 802008 Dept. of Water) from 1998 – 2014. Upper right inset shows years ranked by annual discharge, with 2013/2014 Wet season highlighted in red.
4.2 Productivity and respiration

4.2.1 Components of benthic oxygen flux

The use of a one-dimensional control volume approach in this study allowed estimation of rates of community production at high frequency (half-hourly) during ebb tides by accounting for changes in oxygen due to local accumulation, cross-reef advection, and air-sea gas transfer. The local component dominated in both communities (Figure 14), likely due to the dense biomass of primary producers, the shallow water depth, and slow flow (<5 cm s\(^{-1}\)) during large portions of ebb tide.

The advective component was relatively larger in the macroalgal community compared to the seagrass community. This was likely partially due to the location of loggers relative to community boundaries; water that entered the macroalgal control volume (advected past RDO2) had previously overlain a different and less productive community (seagrass), which led to an oxygen concentration gradient between RDO2 and RDO3. Whereas, water that entered the seagrass control volume (advected past RDO1) had previously overlain similar seagrass community, and thus a strong oxygen concentration gradient did not develop between RDO1 and RDO2. This result suggests that reef flat metabolism on similar high intertidal reefs (Solihuddin et al. 2016) may be estimated for portions of the tidal cycle using Eulerian methods alone provided flow patterns are well-established, the community of interest is of sufficient size, and sensor positions relative to community boundaries are carefully considered. For other tide-dominated reefs where flow speeds remain moderate throughout the tidal cycle (e.g., ≥ 5 cm s\(^{-1}\)) and transit times are short, advective components may comprise a large proportion of \(J_{O2}\) (Falter et al. 2012).

The air-sea gas flux was a relatively minor component (<10%) of diel oxygen patterns in both communities (Figure 14), which is typical of reef systems (Frankignoule et al. 1996). As this reef flat was characterised by shallow water depths (during \(J_{O2}\) estimation), negligible wave action, low wind speeds (~3 m s\(^{-1}\), Table 1), and variable flow speeds, a parameterization of air-sea gas flux for turbulent open-channel flow (Chu & Jirka 2003) was deemed more appropriate than those from high wave and wind conditions commonly-used in reef studies (Wanninkhof 1992, Ho et al. 2006). The minor contribution of air-sea gas flux to \(J_{O2}\) estimates despite diel extremes in oxygen saturation was surprising, but was likely due to the coincidence of low flow (Figure 13), and thus small \(k_{O2}\), with these saturation extremes.

4.2.1 Patterns in hourly community respiration

Estimates of nighttime community respiration in this study showed large variability expressed in a regular pattern with diel and tidal cycles. When nighttime coincided fully with ebb tide, rates of \(CR_{dark}\) were greatest directly after sunset and declined steadily through the night (Figure 14C). Similar patterns have been documented in previous studies and are generally attributed to declines in internal dissolved organic carbon stores (photosynthates) of benthic organisms (Falter et al. 2011, Silverman et al. 2012, Long et al. 2013). In our study, these changes could be due to: 1) declines in nighttime temperatures from air-sea heat exchange, 2) changes in flow speed over ebb tide, 3) changes in oxygen concentration, or 4) DOC (photosynthate) limitation in benthic organisms. Each explanation is discussed in detail below.

The wide range of environmental conditions on Tallon Island reef and large number of observations resulted in \(CR_{dark}\) estimates from a broad range of temperature (28.5 – 32.5°C), DO (7 – 300 µM), and flow speed (0.01 – 0.20 m s\(^{-1}\)). Falling nighttime temperatures due to air-sea heat exchange explained merely ~8% of the diel pattern in \(CR_{dark}\). Uptake rate coefficients for \(CR_{dark}\) did not increase with \(u_x\), indicating that \(CR_{dark}\) was not controlled by flow speed with the exception of low flows where it approached mass transfer limitation (Figure 15). We found strong trends between \(CR_{dark}\) and DO concentration (Figure 16), which potentially indicate that \(CR_{dark}\) was DO-limited by low DO and stimulated by high DO. These relationships did not change when only the first two hours after sunset were used, which suggests that internal DOC limitation was not involved in \(CR_{dark}\) patterns. The dependence of \(CR_{dark}\) on DO concentration has been shown in a few previous studies, either as a Monod-type saturating (Zimmerman et al. 1989) or linear (Atkinson et al. 1994) relationship. This is an important finding.
because it helps us understand controls on daytime community respiration (which cannot be estimated directly) and allows us to more rigorously estimate gross primary production. It also helps us to better predict the effects of climate change and other future stressors of Kimberley reefs based on an understanding of which environmental conditions most affect metabolism.

Daytime community respiration is commonly estimated in metabolism studies as the constant mean of $CR_{\text{dark}}$ (Long et al. 2013). We believe that for this reef (and other Kimberley reefs experiencing large diel swings in oxygen), $CR_{\text{light}}$ can be represented as a function of DO and is therefore generally high during the daytime. A small number of studies show that daytime respiration of reef organisms exceeds that of nighttime (Kühl et al. 1995), but much more research is necessary. This finding is important in the context of the following subsection.

### 4.2.2 Diurnal variations in hourly production

Hysteresis in photosynthesis-irradiance curves (declines in NP in the afternoon relative to NP in the morning, Figure 17) is often seen in seagrass and reef metabolism studies (Long et al. 2013, Koweek et al. 2015a, Adams et al. 2016) and is attributed to many different things, including high light/temperature inhibition of photosynthesis, or light history-dependent respiration. In this study, we have shown that elevation of $CR_{\text{light}}$ due to high afternoon DO levels most likely caused hysteresis in NP at upper irradiances.

When gross primary production of reef platform communities was estimated accounting for increases in $CR_{\text{light}}$, hysteresis in production rates was not evident in either community (Figure 18). Thus, seagrass and macroalgal communities were able to maintain high rates of photosynthetic carbon fixation across a wide range of temperatures and high irradiance. *Thalassia hemprichii* is an intertidal species that has previously been documented in environments with short term temperature increases to ~35°C (McMillan 1984, Campbell et al. 2006, Collier & Waycott 2014, Pedersen et al. 2016). Despite periods of high temperature, small coral thrived on the reef platform; research suggests that water flow and thermal acclimation (Nakamura & Van Woesik 2001, Nakamura et al. 2003) may mitigate coral bleaching when temperatures are elevated. Regardless, this finding illustrates that Kimberley reef platform communities are well-adapted to short-term exposures to high temperature/light and expands our understanding of reef community tolerance to stressors.

### 4.2.3 Metabolism over daily to weekly time-scales

Rates of daily GPP and CR from the macroalgal community (700 and 670 mmol O$_2$ m$^{-2}$ d$^{-1}$, respectively) were within the range of estimates from most reef systems from across the Indo-Pacific (300-1300 mmol O$_2$ m$^{-2}$ d$^{-1}$) (Falter et al. 2013), and similar to mean values for reef flats (740 and 640 mmol O$_2$ m$^{-2}$ d$^{-1}$, respectively). Rates of GPP and CR for the seagrass community (490 and 440 mmol O$_2$ m$^{-2}$ d$^{-1}$, respectively) were well above the global mean for tropical seagrass systems (252 and 217 mmol O$_2$ m$^{-2}$ d$^{-1}$, respectively) (Duarte et al. 2010). At the reef scale, our study estimated daily metabolism similar to the mean of previous studies, despite the extreme conditions present (Figure 28). The production to respiration ratio estimated in our study (1.0) was slightly less than the mean of Indo-Pacific reefs (1.24). The P:R ratio estimated for seagrasses (1.1) was slightly lower than the mean of 1.61 from tropical seagrass studies (Duarte et al. 2010). The balanced P:R ratios of Tallon Island reef communities indicate that the reef is neither a net source nor sink of carbon to adjacent systems.
One of the most interesting findings from this study was the relationship between daily net community production and the light-tide phase difference (Figure 19), which has cycles of ~15 day length (the same as for temperature in section 3.5.2). This shows that the reef platform varies between net autotrophic and net heterotrophic depending on location within this ~15 day envelope. The oscillation between positive and negative NCP is likely driven by the time of day when low water depths occur. For example, low water depth during sunrise/sunset (~0°), would result in high NCP due to low rates of CR (low temperatures and DO). Conversely, low water depth during ~noon/midnight (~180°) would result in low NCP due to high rates of CR.

Many productivity studies last only a few days, which could lead to large overestimates in net production if the study occurred ~0° phase difference or large underestimates if the study occurred ~180° phase difference. This finding suggests that metabolism or other biogeochemical studies should be conducted over a ~2 week period to gain a representative estimate of the biogeochemical process of interest. Additionally, point measurements of temperature, oxygen, nutrients, and other water quality variables on Kimberley reefs will have little ecological relevance without a physical context.

In summary, this study demonstrated that Tallon reef is a moderately productive and metabolically balanced system thriving in some of the most extreme environmental conditions measured worldwide. Through use of the one-dimensional control volume method, we were able to estimate metabolism over a natural range of conditions that would be very difficult to replicate in the laboratory. Relationships between reef metabolism and environmental variables determined in this study can most likely be extended to other Kimberley reefs that experience periods of low water depth. We found that reef gross primary productivity was not affected by high temperature in the short-term, most likely because communities are well-adapted to extreme diel variations. The reef communities varied between net autotrophic and net heterotrophic depending on the interaction between tidal and solar cycles. Our study shows that a strong understanding of reef hydrodynamics is necessary to interpret measurements of temperature, oxygen, and nutrient levels on Kimberley reefs. The estimates of metabolism and key environmental drivers presented here give us a basis to understand reef response to future stressors.
4.3 Physical drivers of extreme temperature and water quality variability in tide-dominated reefs

The results demonstrate how the phase differences between solar heating and tidal cycles can drive variability in temperature extremes within the Kimberley’s reefs, whose amplitude oscillates over a ~14.8 day period as a result of differences between Earth’s rotational and moon’s orbital period. The present study focused specifically on the interaction associated with the principal lunar semi-diurnal (M2) tidal constituent that tends to be the dominant constituent in most locations globally, particularly in the tropics. While other tidal constituents with different frequencies can similarly induce different frequency interactions, the general framework and model described here can be readily extended to investigate other tidal scenarios, including under mixed (semi-diurnal + diurnal) conditions. For the generally dominant M2 constituent considered here, the interaction between the different tidal and solar forcing periods leads to a ~14.8 day modulation of the diurnal temperature fluctuations. We note that our results are similar to the ~14.8 day thermal cycle theorized and observed by Vugts (1975) within periodically drying tidal flats in estuaries; however, the physical mechanisms driving temperature variations in these estuarine systems are not the same as in the reef systems considered here. In Vugts (1975), the tidal modulation in the heat balances occurs via large tidal changes in the wetted surface area of shallow tidal flat systems (which alters air-sea heat exchange), with the response to heat advection being neglected over the large-scales considered in these studies (i.e., at scales larger than a tidal excursion length). However, most reef systems have well-defined shorelines bounded by sloping beaches, or in the case of Tallon Island rocky coastal topography, that results in only minimal changes in reef surface area over a tidal cycle. Instead, tidally-driven advection drives substantial heat exchange between a reef and the ocean, which plays a dominant role in the tidal modulation of these reef heat balances by modifying reef water residence times.

The particular morphology of a reef also plays a very important role in regulating temperature variability. As a consequence, tidal variability over tide-dominated reefs is frequently asymmetric, with longer fall durations than rise durations (Ludington 1979, Wilson 1985, Callaghan et al. 2006, McCabe et al. 2010, Lowe et al. 2015). This effect of tidal truncation arises from the restriction of water draining out of a reef, either by bottom friction or topographic constraints such as a shallower crest (Lowe et al. 2015), which can have a very important influence on the magnitude of diurnal reef temperature variations. Such tidal asymmetry acts to extend the low tide period when shallow water persists on a reef, thereby enhancing daytime warming (or night time cooling). This not only substantially increases the magnitude of diurnal temperature extremes, but also the phase difference between the solar and tidal cycle when these maximum temperatures occur.

An important implication of these results is how sea level rise could alter reef heat balances in the future, and hence, may partially contribute to limiting local reef temperature extremes in the presence of a globally warming ocean. It may seem intuitive, at first, that reefs with large tidal exposure would be less sensitive to sea level changes, given that the degree of mean sea level change will be comparatively smaller than natural sea level variations these reefs already experience on a daily basis. However, in terms of diurnal temperature variability, our results indicate that reefs subject to large tidal amplitudes can be affected just as much by a sea level rise as those subject to small tidal amplitudes (Figure 26a). Reefs that are already close to the critical depth at which tidal truncation ceases (i.e., near \( h_0^* = h_{\text{min}}^* + 1 \)) are particularly close to a ‘tipping point’ where even a relatively small increase in mean sea level can lead to a substantial reduction in the magnitude of diurnal temperature fluctuations. In some cases, reefs that presently experience +5°C diurnal temperature extremes could see a >2 °C reduction in these temperatures from a 0.7 m sea level rise, or a >4 °C reduction by a 1.5 m rise (Table 4); amounts that are comparable in magnitude to projected warming of the tropical oceans by the end of the century under a business-as-usual scenario (2-4°C, Bopp et al. (2013)). It is not yet known, however, how much future reductions in local temperature variability will ameliorate the more chronic thermal stresses imposed by long-term increases in regional ocean temperature.

Finally, while the focus of this study has been on the temperature extremes of tide-dominated reefs, the results are also relevant to understanding the role of tides on many other reef water quality parameters. For example,
changes in reef water column oxygen, $pCO_2$ and pH are also governed just as much by the morphology of the reef and the hydrodynamic forces driving circulation as they are by the fluxes driving their non-conservative behavior (Falter et al. 2013). Furthermore, the specific processes responsible for driving changes in oxygen, $pCO_2$, and pH (i.e., production, respiration, and net calcification) also exhibit light-dependent diurnal periodicities similar to the net atmospheric heat fluxes used in the present model, albeit with much greater spatial variation in the biologically-driven bottom boundary fluxes (Lowe & Falter 2015). Thus, we expect sea level rise to also cause reductions in daily extremes of oxygen, $pCO_2$, and pH in similar proportion to the reductions in daily temperature extremes that we report here. Although there is little doubt that rising levels of atmospheric $pCO_2$ will cause substantial changes to ocean temperature and chemistry over the following century, we conclude that rising sea levels will likely help moderate the extreme conditions presently found in many shallow, tide-dominated reef habitats.

### 4.4 Recommendations for future work

The project has provided a reasonably complete assessment of how environmental variability, including temperature, light, oxygen and water motion, influence the productivity of benthic communities in the Kimberley. Due to the process-focused nature of the experiments, we expect the general relationships and models developed in this project to be extended to other reef systems with some confidence. In other words, if the environmental conditions and habitat characteristics of other reefs in the Kimberley are known, or can be reasonably predicted, we believe the robust estimates of the productivity of other reefs in the Kimberley can be made.

Therefore, a natural future step will be to integrate future information about reef characteristics across the broader Kimberley region, including properties of reef morphology, habitats and benthic community structure. While beyond the scope of the present project, this will enable our results to be extended more broadly across the Kimberley to, for example, obtain quantitative estimates and maps of reef productivity across the region.

Further work is also needed to develop a more comprehensive view of nutrient dynamics for reefs in the Kimberley. Given the intensive nature of the two year field program, which happened to focus on the 2013-14 period that was unusually dry, future work should compare these observations to more typical wet seasons. While the results from this study did not indicate any substantial differences in reef nutrient dynamics between seasons, this may not always be the case, particularly during anomalously wet years. To understand this inter-annual variability, future research would benefit from regular water quality monitoring across the coastal Kimberley.
5 References


Bryant D, Burke L, McManus J, Spalding M (1998) Reefs at risk: a map-based indicator of threats to the world’s coral reefs. World Resources Institute, Washington, DC


Koweek DA, Dunbar RB, Monismith SG, Mucciarone DA, Woodson CB, Samuel L (2015b) High-resolution physical and biogeochemical variability from a shallow back reef on Ofu, American Samoa: An end-member perspective. Coral Reefs 34:979-991


Ramsing N, Gundersen J (1994) Seawater and gases. Tabulated physical parameters of interest to people working with microsensors in marine systems.
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7 Communication

7.1 Students supported

- Renee Gruber, The University of Western Australia, PhD, expected completion December 2016, “Productivity and biogeochemical processes on a macrotidal tropical reef platform”.

7.2 Journal publications

7.3 Submitted manuscripts


7.4 Presentations

- Lowe, R.J. Benthic community productivity of the Kimberley’s macrotidal reefs”, DPaW Lunch and Learn seminar, April 2016.
- Lowe, R.J. Exploring the circulation and heat balances of northwestern Australia’s remote macrotidal reefs, ARC Centre of Excellence for Coral Reef Studies Symposium, Hobart, October 2015.
- Lowe, R.J. Environmental forcing of benthic community productivity within the Kimberley’s macrotidal reefs, WAMSI Research Conference, March 2015.
- Lowe, R.J. Oceanic drivers of coral reef systems along Western Australia, ARC Centre of Excellence for Coral Reef Studies Symposium, Canberra, October 2014.

7.5 Other communications achievements

- Lowe, R.J. “Kimberley tides and ocean circulation”, Science on the Broome Coast – Public Talk, August 2016