

Effects of experimental CO2 enrichment on the PSII photochemical

efficiency of Symbiodinium sp. in Acropora millepora

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Abstract

Enrichment of seawater with CO₂ decreases the concentration of the carbonate ion while increasing that of hydrogen and bicarbonate ions. We use pulse-amplitude-modulation (PAM) fluorometry to investigate whether, in the absence of warming, and in sub-saturating light, these changes affect the PSII photochemical efficiency of Symbiodinium sp. in the reefbuilding coral Acropora millepora. We assessed this experimentally with 30-min-interval saturation pulse analyses at 25 °C, a daily peak in the intensity of the photosynthetically active radiation (PAR) at ~65 μ mol quanta m⁻² s⁻¹, and a seawater pCO₂ that we gradually increased over nine days from ~496 to ~1290 µatm by injection of CO₂-enriched air. Nine 14day time series, which, except one, were recorded at the growing apices of a coral branch, revealed diel oscillations in the PSII photochemical efficiency characterized by a steep nocturnal decrease followed by a steep increase and peak in the morning, a daily minimum at midday ($\Delta F/F_{m}$ ',midday), and a daily maximum at the onset of darkness at 19:00 h ($F_v/F_{m,19:00 h}$). An inadvertent shift in the position of one of the PAM fluorometer measuring heads revealed differences between the basal part and the growing coral apices of a coral branch in $\Delta F/F_m$ 'midday and Q_m. In ambient seawater (Control) Symbiodinium sp. exhibited a gradual decrease, over the course of the experiment, in $\Delta F/F_m$, midday, $F_v/F_{m,19:00 h}$, and the slope of the linear regression between the relative electron transport rate and the intensity of PAR (rETR/PAR). Although two of three successive experiments indicated that CO₂ enrichment counteracted these trends, statistical analyses failed to confirm an influence of pCO_2 on $\Delta F/F_m$ ',midday, $F_v/F_{m,19:00 h}$, and Q_m , rendering this experiment inconclusive.

Keywords

CO2 enrichment; PAM fluorometry; photophysiology; reef-building coral; ocean acidification

Introduction

Enrichment of seawater with CO₂ decreases the concentration of the carbonate ion while increasing that of hydrogen and bicarbonate ions [1]. This shift in the seawater carbonate system occurs in conjunction with ocean warming and can affect coral calcification and photosynthesis, which are intimately coupled [2]. Past experiments have revealed that the coral symbiosis is more susceptible to thermal stress than CO₂ enrichment, and that the physiological plasticity which influences its resilience is species-specific [3–7]. One physiological process of particular interest is the upregulation of the calcifying fluid pH [8,9]. A CO₂ induced increase in seawater [H⁺] may increase the energy required for this upregulation [10] and if so, then such additional energy demand must be compensated by photosynthesis of the symbiotic dinoflagellates, which provide most of the coral's energy by transferring photosynthetic products to their hosts [11].

Coral species apparently differ in their photosynthetic response to CO₂ enrichment due to a host-specific regulation of the symbionts' carbon concentrating mechanism (CCM) [12,13]. The CCM uses active bicarbonate transport and carbonic anhydrase to increase the concentration of CO₂ at the site of type II RuBisCO—an enzyme with a low affinity for CO₂ (14–18]. A high carbonic anhydrase activity may indicate that the coral symbiont lives in a carbon-scarce environment and therefore invests energy in concentrating carbon [13]. In *Porites porites* (Pallas, 1766) and *Acropora* sp., enrichment of their environment with CO₂ may then increase the gain from photosynthesis for the benefit of the holobiont [19]. In *Acropora muricata* (Linnaeus, 1758), for example, under conditions of sub-saturating light, CO₂ enrichment can increase chlorophyll pigments and the de-epoxidation of xanthophylls, thus increasing the capacity of the symbiont to photoacclimate to low irradiance [20]. The species *Stylophora pistillata* Esper, 1797 responded similarly with an increase in photosynthetic

efficiency [21], and the symbionts in *A. millepora* (Ehrenberg, 1834) and *Seriatopora hystrix* Dana, 1846 apparently increase their maximum PSII quantum yields and light-limited electron transport rates in response to CO₂ enrichment [22].

Other studies support the view that corals do not respond to CO_2 enrichment [23–26], and yet others have demonstrated negative effects. Kaniewska et al. [27], for example, suggested that in *A. millepora*, CO_2 enrichment caused widespread changes in gene expression consistent with metabolic suppression, an increase in oxidative stress, apoptosis and symbiont loss, and a decrease in respiration and photosynthesis. Furthermore, Edmunds [28] reported negative effects of CO_2 enrichment—in this study, CO_2 enrichment decreased both the symbiont's maximum and effective photochemical efficiencies.

Here, we follow the studies by Edmunds [28], Hoadley et al. [25], and Noonan and Fabricus [22], asking if, in the absence of warming, CO₂ enrichment affects the PSII photochemical efficiency of *Symbiodinium* sp. in the reef-building coral *A. millepora*. To investigate this experimentally, we conducted time series of saturation pulse analyses (pulse-amplitude-modulation fluorometry) monitoring the symbiont's maximum photochemical efficiency, F_v/F_m , midday effective photochemical efficiency, $\Delta F/F_m$ ', and the relationship between the relative electron transport rate and the intensity of the photosynthetically active radiation (rETR/PAR) while gradually increasing the seawater *p*CO₂.

Material and methods

Experimental design

We conducted three consecutive laboratory experiments in each of which we acclimated one fragment (~7 cm tall, 4 cm wide) of the coral *A. millepora* for seven days in each of three seawater recirculation tanks to a simulated daily light cycle that peaked midday at a photosynthetically active radiation (hereafter, PAR) of ~65 μ mol quanta m⁻² s⁻¹ (Fig 1, S1 Fig

1). The nine coral fragments originated from one large *A. millepora* specimen collected off the East Coast of Australia, and therefore, we assume that they hosted the same *Symbiodinium* clade.

After their collection, and until acclimation in the laboratory, the coral fragments were kept under a constant 12/12 h dark/light regime (PAR ~ 90 μ mol quanta m⁻² s⁻¹, Kessil A160WE Tuna Blue LED). In the laboratory, they were placed on a gridded plate under the measuring head of a pulse-amplitude-modulation (PAM) fluorometer (Monitoring-PAM aquatic version, Walz GmbH, Germany) and a Kessil A80 Tuna Blue controllable LED (S1 Figs 1, 2). Because only three PAM fluorometer measuring heads were available for this study, three sets of three time-series measurements were conducted consecutively. For a recent review of the strengths and limitations of chlorophyll fluorescence measurement, see Bhagooli et al. [29].



Fig 1. Experimental design and timeline. In each of three consecutive 16-day experiments, one coral fragment was tested in each of three tanks. Transitions from white to blue shades indicate an increase in seawater *p*CO₂. Numbers, seawater [CO₂] at 25 °C (μmol kg⁻¹); numbers in parentheses, seawater [H⁺] at 25 °C (nmol L⁻¹). N-dashes indicate missing data.

Following 7 days of acclimation, the seawater pCO_2 was gradually raised and then maintained in two of the three tanks from ~500 to ~1200 µatm by computer-controlled injection of CO₂- enriched air over the next nine days in three steps. Computer feedback control adjusted CO_2 additions to target pH values, and the three steps resulted in pH = 7.8 on day 10, pH = 7.7 on day 13, and pH = 7.6 on day 16. In each experiment, one of the three tanks remained at ambient pH = 8.0 (Control). Once the first experiment was completed, the CO_2 injection was shut off, allowing the *p*CO₂ in seawater and atmosphere to equilibrate before starting the next experiment on day 19 with three new coral fragments. These steps were repeated before starting Experiment 3 on day 38 (Fig 1).

Laboratory setup

Each experimental tank (S1 Fig 1) contained ~450 L seawater collected from Okahu Bay, New Zealand. A submerged pump (1260, Eheim) moved ~9 L min⁻¹ from the main tank (112 \times 72 \times 60 cm) through a water cooler (HC-300A, Hailea) and UV sterilizer (Pond One UV-C 9W, ClearTec) into an elevated mixing barrel (210 L) from which the seawater returned to the main tank by gravity. The sizes of the tank and mixing barrel, and the flow rate of the pump, were chosen to ensure that short-term fluctuations in the *p*CO₂ of the seawater in the mixing barrel (due to CO₂ enriched air injection, as explained below) did not affect the tank.

A heater (500 W GH Quartz Glass heater, Aqua One) was placed at the floor of the main tank. The chiller and heater maintained the seawater temperature at 25 ± 0.5 °C. The seawater was also pumped from the main tank through an external particle filter (Professional 4+ 350 Cannister filter, Eheim) into a small, elevated plastic container ($30 \times 20 \times 10$ cm) that contained the coral fragment. The tube returning seawater from the particle filter was aimed towards the coral fragment to ensure rapid flow across the coral surface. The overflow from this container returned the seawater into the main tank.

The distance between the measuring head of the PAM fluorometer and the coral surface was \sim 40 mm. The PAM fluorometer measures ambient light immediately adjacent to the area under test using a small sheet of Teflon placed flush with the measured tissue and reflecting

light to an internal PAR sensor. This sensor was calibrated against a Li-Cor Li-192 underwater quantum sensor. The PAM recordings show that the LED mounted above each coral fragment gradually increased the intensity of PAR from 5 am to a midday maximum of ~65 μ mol quanta m⁻² s⁻¹ and then gradually decreased this intensity until 7 pm, when the LED was turned off (S1 Fig 2). Note that the lowest irradiance emitted from the LED was ~20 μ mol quanta m⁻² s⁻¹. The resulting daily flux, as measured by the internal PAR sensor of the PAM fluorometer, ranged between 1.7 and 2.6 mol quanta m⁻² day⁻¹.

Saturation pulse and induction/recovery analyses

The Walz software WinControl-3.0 ran a batch routine to automatically perform one saturation pulse analysis every 30 minutes between 03:00 and 24:00 h and one daily induction/recovery analyses between 02:00 and 02:30 h. The following settings were applied: saturation pulse intensity = 12, saturation pulse width = 0.6 s, gain = 1, measuring light intensity = 6, measuring light frequency = 3. The measuring light was turned off between saturation pulses and initial measurements of the baseline florescence confirmed that the intensity of the measuring light did not cause an actinic light effect. The daily induction/recovery analyses started with an F₀ determination, followed by a series of 12 saturation pulses (delay = 40 s) 20 seconds apart with the actinic light on (65 μ mol quanta m⁻² s⁻¹). After ~4 minutes, the actinic light was turned off, and a series of 8 saturation pulses occurred at increasing intervals between 0.5 and 9 minutes. Prior to deployment, each PAM sensing head was zeroed in the experimental setup.

We derived the maximum PSII photochemical efficiency, F_v/F_m , from measurements of F_0 and F_m in darkness: $F_v/F_m = (F_m - F_0)/Fm$ (Table 1) [30]. The effective photochemical efficiency, $\Delta F/F_m$ ', was derived from the maximum (F_m ') and minimum (F') fluorescence yields at ambient light intensity: $\Delta F/F_m' = (F_m' - F') / F_m'$ [30]. F_v/F_m measured at 19:00 h, and $\Delta F/F_m'$ measured at midday gave the midday excitation pressure, Q_m : $Q_m = 1 - [(\Delta F/F_m', midday) / (\Delta F/F_m', midday)]$

 $(F_v/F_{m,19:00 h})$] [31]. The PAR recorded by the internal sensor of the PAM and the effective photochemical efficiency, $\Delta F/Fm'$, were used to derive the relative electron transport rate: rETR = $\Delta F/F_m' \times PAR \times 0.5$ [32].

Table 1. Summary of fluorescence parameters measured or derived in conditions of darkness or actinic light. AL, actinic light; SP, saturation pulse.

Symbol	Fluorescence parameter	Equation/comments	Reference
Darkness, measured variables			
F_0	Minimum fluorescence		
F_{m}	Maximum fluorescence	Saturation pulse	
Darkness, derived variables			
$F_{\rm v}$	Variable fluorescence	$\mathbf{F}_{\mathbf{v}} = (\mathbf{F}_{\mathbf{m}} - \mathbf{F}_{0})$	
F_v/F_m	Maximum photochemical efficiency	$F_v/F_m = (F_m - F_0)/F_m$	[30]
Actinic light, measured variables			
F _m '	Maximum fluorescence yield	Saturation pulse	
F0'	Minimum fluorescence yield		
Actinic light, derived variables			
Fv'	Variable fluorescence	$F_{v}' = (F_{m}' - F_{0}')$	
$\Delta F/F_m$ '	Effective photochemical efficiency	$\Delta F/F_{m}' = (F_{m}'-F')/F_{m}'$	[30]
Q_m	Excitation pressure	$Q_m = 1 - [(\Delta F/F_{m,'midday})/(F_v/F_{m, 19:00 h})]$	[31]
		$(F_v/F_m \text{ measured at } 19:00 \text{ h},$	
		F _m ' measured midday)	
rETR	Relative electron transport rate	$rETR = \Delta F/F_m' \times PAR \times 0.5$	[54]

Seawater carbonate system

The pH of the seawater in the mixing barrel of each circulation unit was continuously measured with a SenTix HWD electrode connected to a pH 3310 meter (WTW). These measurements were sent to a computer with CapCtr software (Loligo® Systems ApS) controlling the opening and closing of a solenoid valve when the seawater pH increased above or decreased below the daily set point. The solenoid valve released CO₂-enriched air (5% CO₂, 21% O₂ in nitrogen) from a gas cylinder to a perforated tube in the mixing barrel. The pH electrodes were calibrated using NIST/DIN pH buffers to test for theoretical Nernstian electrode behavior and then conditioned in seawater before determining the electrode-specific offset between the potential measured in NIST/DIN pH buffer and that measured in certified seawater reference material (TRIS in synthetic seawater). The electrodes were recalibrated at the start of each experiment.

Determination of seawater DIC, TA and salinity

To determine the seawater carbonate system, we collected a one-liter sample from each circulation unit at the start of each experiment, each night before CO₂-enriched air injection was increased, and at the end of the experiment. These samples were preserved with mercuric chloride and later analyzed for dissolved inorganic carbon (DIC) with a SOMMA (Single Operator Multiparameter Metabolic Analyzer) coulometer system and for total alkalinity (TA) with a closed-cell potentiometric titration system following the SOP's 2 and 3a procedures [33]. We used these DIC and TA measurements and D. Pierrot's adaptation of the CO2Sys.BAS program [34] to compute the seawater pCO_2 and pH (total scale, mol kg-SW⁻¹). The dissociation constant for HSO4⁻ was taken from Dickson [35]; the values of K1 and K2 of carbonic acid were from Mehrbach et al. [36] refitted by Dickson and Millero [37]. Note that water samples were not collected on the final day of Experiment 3 due to a COVID-19 pandemic lockdown, and four seawater samples were destroyed during transport to the analytical lab (missing data in Fig 1 and S1 Table 2). The seawater salinity was measured with a handheld conductivity meter (Knick, Germany) and maintained at 34.5 ± 0.5 by daily addition of ultrapure water.

Statistical analysis

Although saturation pulse analyses were conducted every 30 minutes, we only used the data collected during the last day of each step- pCO_2 increase (days 7, 10, 13, 16), considering that during this day, conditions in the experimental tanks had been fully established as per the pH set point.

The daily induction/recovery routines were used to assess the effects of CO_2 enrichment on the PSII efficiency using two variables: the variable fluorescence, F_v , determined by analysis of the first saturation pulse of the induction, and the average of the plateaued F_v measured during recovery.

All statistical analyses were performed with R statistical software (version 1.3.959). The Shapiro-Wilk test was used to assess if the data were normally distributed, and homogeneity of variance was tested with the Levenes test. The $F_v/F_{m,19:00 h}$, $\Delta F/F_m$ ',midday, Qm, and the slope of the linear regression of rETR versus incident PAR were analyzed with a four-factor, nested ANOVA in which individual coral fragments were the random factor nested in tank, treatment, and day of measurement, which were fixed factors. The same ANOVAs were then used to analyze the variables derived from the induction and recovery analyses. We used an Akaike information criterion model selection to determine the best model possible to describe the relationship between the fluorescence parameters, the three tanks, the treatment, and the individual coral fragment. The tank effect was not significant and was removed from the model.

Results

Seawater CO₂ enrichment

The measured and derived seawater carbonate chemistry parameters at the start of each of three consecutive CO₂ enrichment experiments (days 7, 26, and 45; Fig 1) and 3, 6, and 9 days later are summarized in Figure 1 and S1 Table 2. The CO2Sys.BAS computations confirmed that the stepwise increase in the injection of CO₂-enriched air increased the *p*CO₂ of the ambient seawater in the tanks of the Treatments from 493 ± 44 µatm (n = 6) to 799 ± 101 µatm (n = 5, days 10, 29, and 48), 1109 ± 89 µatm (n = 6, days 13, 32, and 51), and 1290 ± 69 µatm (n = 4, days 16, 35, and 54; S1 Table 2).

Saturation pulse analyses

Time series of saturation pulse analyses revealed diel oscillations in the PSII photochemical efficiency of *Symbiodinium* sp. characterized by a steep nocturnal decrease followed by a steep increase and peak in the morning, a daily minimum at midday, and a daily maximum at the onset of darkness at 19:00 h (Fig 2, S1 Fig 3). We note that the F_v/F_m times series in Fig 2 and S1 Fig 3 were interrupted at 02:00 h by photosynthesis induction routines.



Fig 2. *Symbiodinium* sp. in *Acropora millepora*. Diel variations in the PSII photochemical efficiency (darkness: F_v/F_m ; light: $\Delta F/F_m$ ', see Table 1) of nine coral fragments, one

placed in each of three tanks (tank 1, blue; tank 2, black; tank 3, orange) for each of three consecutive experiments. Daily induction and recovery analyses caused gaps in time series from midnight to 03:00 h.

Both variables, the maximum PSII photochemical efficiency ($F_v/F_{m,19:00 h}$ recorded at 19:00 h, Table 1) and the midday effective PSII photochemical efficiency, $\Delta F/F_m$ ',midday, gradually decreased over the course of the experiment in both Controls and Treatments (Fig 3). This decrease, which produced a significant effect on days 13 and 16 (S1 Table 1), was independent of CO₂ enrichment. Similarly, the midday excitation pressure, Q_m, which was derived from $F_v/F_{m,19:00 h}$ and $\Delta F/F_m$ ',midday (Table 1), was not affected by CO₂ enrichment (Fig 3, S1 Table 1).



Fig 3. *Symbiodinium* sp. in *Acropora millepora*. Time series of the maximum PSII photochemical efficiency measured at 19:00 h, $F_v/F_{m,19:00 h}$, the midday PSII effective photochemical efficiency, $\Delta F/F_m$ ',midday, and the midday excitation pressure, Q_m (see Table 1), of nine coral fragments, one placed in each of three seawater circulation units (circles, Tank 1; squares, Tank 2; triangles, Tank 3) in each of three consecutive experiments. In each experiment, the seawater in two tanks (filled symbols) was

gradually enriched with CO₂ (Treatment). The pCO₂ in the third unit (open symbols) remained at ~506 µatm over the duration of the experiment (Control).

Our measurements on the coral fragment in Tank 1 of Experiment 2 (Treatment, Fig 1) differed from all other measurements in that the PAM measuring head accidently changed its position so that it was not directed toward the distal but a more basal part of a coral branch. This apparently resulted in much lower PSII photochemical efficiencies (Fig 3). Because the difference in $\Delta F/F_m$ ',midday was greater than that in $F_v/F_{m,19:00 \text{ h}}$, the derived Q_m exceeded that of the other two coral fragments tested in Experiment 2 (Fig 3).

The slope of the linear regression between rETR and PAR ($R^2 > 0.997$, S1 Fig 4) decreased over the course of the experiment under ambient *p*CO₂ conditions (Fig 4, open symbols). Such decrease was also observed in CO₂-enriched seawater (Fig 4, closed symbols), but in Experiments 1 and 3, this decrease was less steep so that the difference in slope between the Control and Treatments increased over the course of the experiment.



Fig 4. *Symbiodinium* sp. in *Acropora millepora*. Time-series of the slope of the linear regression of relative electron transport rate (rETR) versus incident photosynthetically active radiation (PAR, μ mol quanta m⁻² s⁻¹) shown in S1 Fig 4. The rETR was derived

from measurements immediately after (day 0) and three, six and nine days after acclimation of nine coral fragments, one placed in each of three seawater tanks (circles, tank 1; squares, tank 2; triangles, tank 3) for each of three consecutive experiments. In each experiment, the seawater in two tanks (filled symbols) was gradually enriched with CO₂ (Treatment, see text and S1 Table 2).

Induction-recovery dynamics

Like $F_v/F_{m,19:00 h}$ and $\Delta F/F_m$ ',midday, the variable fluorescence, F_v , recorded during the induction–recovery routine decreased over the course of the experiment in both the Control and Treatment groups (S1 Table 3). F_v measured at the beginning of each nocturnal photosynthesis induction was not affected by CO₂ enrichment (S1 Table 3, S1 Fig 5). Once the actinic light was switched off, F_v gradually recovered, approaching pre-light exposure values (F_v between 600 and 800) within 40 min (S1 Fig 5). Again, CO₂ enrichment did not affect the average postinduction recovery F_v (S1 Table 3, S1 Fig 5).

Discussion

We observed that in ambient seawater (Control) under conditions of a sub-saturating diel light cycle and ~25 °C, *Symbiodinium* sp. exhibited a gradual decrease, over the course of the experiment, in the midday and maximum PSII photochemical efficiency ($\Delta F/F_m$ ',midday and $F_v/F_{m,19:00 h}$), and the slope of the linear regression between the relative electron transport rate and the intensity of PAR (rETR/PAR). Although two of three successive experiments indicated that CO₂ enrichment counteracted these trends, statistical analyses failed to confirm an influence of *p*CO₂ on $\Delta F/F_m$ ',midday, $F_v/F_{m,19:00 h}$, and the midday excitation pressure, Q_m , rendering this experiment inconclusive.

The midday excitation pressure, Q_m, is an indicator of symbiont performance at maximal irradiance [31]. The near-zero values measured in this study suggest a high proportion of open

PSII reaction centers and possible light limitation. The plots in Fig 3 demonstrate that one of the three coral fragments tested in Tank 1 of Experiment 2 exhibited relatively low $\Delta F/F_{m}$, midday and a higher Q_{m} (Fig 3; filled circles). In this case, the measuring head of the PAM fluorometer had inadvertently changed its position at the onset of the time series, so that it pointed towards the basal part instead of the growing coral apices of a coral branch. This basal part may have exhibited greater light scattering than the distal parts, which would explain the low $\Delta F/F_m'$, midday and higher Q_m [38]. This accidental observation emphasizes the importance of accurate placement of the measuring heads for measurement replication. If we exclude these data for that reason, and consider each experiment separately, then it appears that $\Delta F/F_m$ ', midday measured in ambient pCO₂ seawater (Control) was lower than that measured in CO₂-enriched seawater, in each of the three experiments. On the other hand, the midday excitation pressure, Q_m, was higher than that measured in CO₂-enriched seawater. Similar effects have been reported for A. muricata by Crawley et al. [20] and for P. *damicornis* by Jiang et al. [39], showing that CO₂ enrichment can decrease the PSII excitation pressure. In the former study, this decrease was caused by a reduction in F_v/F_m, while in the latter $\Delta F/F_m$ ' increased, as observed in our study.

We also note that in Experiments 1 and 3, the slope of the $\Delta F/F_m$ ',midday and $F_v/F_{m,19:00 h}$ time series measured in increasingly CO₂ enriched seawater was smaller than that of the time series measured in ambient *p*CO₂ seawater (Fig 3). Similarly, the difference in the slope of the rETR/PAR relationship between Treatments and Control increased as the *p*CO₂ increased (Fig 4). This points to a possible positive effect of CO₂ enrichment; the increasing seawater *p*CO₂ may have counteracted the gradual decrease in $\Delta F/F_m$ ' and rETR/PAR slope that was observed under ambient *p*CO₂ conditions. However, it remains unclear why such a trend was not observed in Experiment 2. If the coral fragments were to host different clades of Symbiodinium, then this may explain a difference in response. However, since the fragments used in our experiment came from the same parental coral, this possibility seems unlikely. We observed that under conditions of ambient pCO_2 F_v/F_{m,19:00 h} and Δ F/F_m',midday gradually decreased over time. Possible causes for this include incomplete acclimation to laboratory conditions. Before our experiment, the coral fragments lived in a constant 12/12 h dark/light regime with a PAR of approximately 90 µmol quanta m⁻² s⁻¹, providing a daily photon flux of around 3.5 mol quanta m⁻² day⁻¹. In our experiment, the PAR intensity was modulated around a midday peak, resulting in a smaller flux of 1.7–2.6 mol quanta m⁻² day⁻¹. Although the coral *A. millepora* appears to tolerate low-light conditions [40,41], it seems that acclimation may take up to 20 days [40], which exceeds the acclimation period in our experiment. Seawater CO₂ enrichment may have supported such acclimation in Experiments 1 and 3 [20], preventing Δ F/F_m',midday of the coral fragments from decreasing as steeply as in the Control under conditions of ambient *p*CO₂.

The observed diurnal decline in $\Delta F/F_m$ ' (Figs 2, S3) correlated with the daily peak in radiation exposure and possibly the development of reversible and photoprotective non-photochemical quenching [42,43]. On the other hand, the sharp decline in F_v/F_m during the night points to chlororespiration, which can create a trans-thylakoid [H⁺] gradient in the dark through cyclic electron transport around PSI, thereby promoting ATP production [44–46]. Chlororespiration requires oxygen and darkness or at least very low light [47]. In our experiment, these conditions were met at 18:30 h when PAR decreased below ~20 µmol quanta m⁻² s⁻¹ and was shut down at 19:00 h (S1 Fig 2).

Chlororespiration can deplete the accessible oxygen in the coral tissue [48,49]. Without an electron acceptor, electrons may have accumulated in the PSII–PSI electron transport chain, reducing the pool of plastoquinones. This will have initiated the transition of light harvesting complexes from PSII to PSI [44], decreasing the absorption cross section available for PSII

[50]. At dawn, F_v/F_m increased rapidly (Figs 2, S1 Fig 3) perhaps following the stimulation of PSI, which oxidized the plastoquinone pool and reversed the transition of light harvesting complexes [45]. Although the function of chlororespiration is still debated [44,51,52), its associated reduction of O₂ accumulated during the day may have lowered the risk of reactive oxygen damage to PSII, and the induced state transition may have supported an efficient onset of photosynthesis and O₂ production at the onset of light (44,53].

Conclusion

Our time series of saturation pulse analyses revealed evidence for chlororespiration of *Symbiodinium* sp. in the reef-building coral *A. millepora*. An inadvertent shift in the position of one of the PAM fluorometer measuring heads revealed differences between the basal part and the growing coral apices of a coral branch in $\Delta F/F_m$ ',midday and Q_m—an accidental observation that emphasizes the importance of accurate sensor placement for measurement replication. Although two of three successive experiments indicated that CO₂ enrichment counteracted the gradual decrease in $\Delta F/F_m$ ',midday, $F_v/F_{m,19:00 \text{ h}}$, and the slope of the linear rETR/PAR regression, observed in the Control over the course of the experiment, statistical analyses failed to confirm such effect, rendering this experiment inconclusive. We believe that the possibility of such an effect warrants further experimentation.

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Author contributions

A.M. and K.V. conceived and performed the experiment and analyzed the data. K.V. wrote the paper with assistance from A.M. and D.B.

Data availability

The datasets are available from the corresponding author on reasonable request.

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