The effect of temperature compensation on the circadian rhythmicity of photosynthesis in \textit{Symbiodinium}, coral-symbiotic alga

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Circadian rhythms, which are found in most eukaryotes, are defined as rhythms that persist under constant conditions with a periodicity close to 24 h. One central key characteristic of all circadian rhythms is "temperature compensation", which allows organisms to maintain robust rhythms with a period close to a diel cycle over a broad range of physiological temperatures. To better understand the response of the circadian clock in corals to temperature elevation, photosynthesis as an output process of the circadian clock was studied both in \textit{Stylophora pistillata} corals and in cultured \textit{Symbiodinium} algae. The time period of photosynthesis was not affected by temperature elevation in the cultured algae or in the corals harbouring \textit{Symbiodinium}. However, the photosynthetic system responded to temperature elevations by adjusting the photosynthetic apparatus. These findings suggest that the endogenous algal circadian clock regulates the photosynthetic rhythm and compensates for temperature elevations that occur in the natural environment.
A common feature of the circadian clock under temperature fluctuations is that high temperatures are interpreted as daylight and low temperatures are interpreted as darkness. These responses suggest that light and temperature signals are interpreted through common entrainment pathways. The molecular component of the input pathway entrained by temperature has not been defined because changes in temperature affect nearly every chemical and physical process in the cell. Therefore, it may be impossible to define a single "temperature receptor" or a single input pathway. Hastings and Sweeny described a similar effect of temperature on two reactions that have antagonistic effects on the time period of the clock; these antagonistic effects result in temperature compensation. This idea was the basis for the antagonist balance model later proposed by Ruoff and Ruoff. This model is consistent with the findings of Gould et al. and provides an explanation for the temperature compensation phenomenon.

Coral reefs that harbour endosymbiotic dinoflagellate algae of the genus *Symbiodinium* are more susceptible than land plants to temperature elevations. Symbiotic corals evolved and live in tropical marine environments that are characterised by temperature stability. Strong temperature anomalies can therefore lead to drastic responses, including a breakdown of the coral (host)/algae symbiosis, known as bleaching. Coral bleaching, in which corals lose their colouration because of the expulsion of symbiotic algae, results from many physical and biological factors, including high or low sea surface temperatures (SSTs), UV irradiation, bacterial infection, lowered salinity and pollution. Mass bleaching events resulting from SST changes are thought to originate from acute photosynthetic damage to *Symbiodinium*. Since photosynthesis in dinoflagellates is a clock-regulated process, the efficiency of temperature compensation (an important property of all circadian rhythms) may allow *Symbiodinium* to acclimate to temperature elevations in a manner similar to that of photoacclimation. Because the circadian clock machinery controls numerous biochemical and physiological pathways, the present study was designed to elucidate the basic mechanism of the
Symbiodinium clock machinery as manifested in Figure 1 (A–D), under the growth temperature of 24°C and the ways in which SST fluctuations (above 24°C) affect the clock apparatus, which eventually can lead to a mismatch in photosynthetic efficiency or, in extreme cases, to the breakdown of the coral/algae symbiosis. The mechanisms of the circadian clocks in Symbiodinium have not been systematically investigated, and the level of circadian control in this genus is still unknown. Temperature can entrain, reset and phase-shift the circadian rhythm, indicating that the clock recognises and responds to changes in temperature at several levels².

The specific aim of this study was to characterise the effects of elevated temperature on the circadian rhythm of photosynthesis under constant LL. We monitored oxygen evolution during photosynthesis in cultured Symbiodinium clade A and the clade D that is considered as temperature-tolerant²⁶–³⁰ and in the symbiotic Stylophora pistillata coral possess clades A and C³¹. In addition to the measurement of oxygen oscillation, an array of photosynthetic fluorescence parameters (Y(II), NPQ and ETR) were examined during a diel cycle under various physiological temperatures with constant light conditions.
**Results**

Temperature compensation effects were investigated in *Symbiodinium* cultures and in *S. pistillata* corals. Cultures were grown under LD cycles with a light intensity of 100 µmol quanta m⁻² s⁻¹ at 24°C and then transferred to LL conditions with elevated temperatures. The Q₁₀ value was calculated with the formula $Q_{10} = R_2/R_1^{10(T_2-T_1)}$, where $R_2$ is the rate of the process under the elevated temperature ($T_2$) and $R_1$ is the rate under the reference temperature ($T_1$) under LD cycles. $Q_{10}$ was found to be 1 for the ambient temperature (24°C) under constant light, 1.11 ± 0.08 for 27°C and 1.05 ± 0.02 for 30°C (Figure 1A). When tested at a temperature of 6°C above the ambient temperature, clades A and D, exhibited the same periodicity behaviour when the $Q_{10}$ values were calculated (paired t-test, $P > 0.05$). Both clades compensated for the temperature elevation and maintained a photosynthetic rhythm. However, oxygen production was lower under conditions of elevated temperature than at the ambient temperature, more dramatically so in clade A than in clade D (Figure 1). The decline in oxygen production was 23.5% in clade A and 16.9% in clade D, corresponding to the average of the maximal value. Corals were tested at the ambient temperature with a light intensity of 100 µmol quanta m⁻² s⁻¹, and the fragments were subjected to an elevated temperature of 30°C. The calculated $Q_{10}$ values were 1.02 ± 0.01 for the corals grown at ambient temperature and 1.03 ± 0.005 for those grown at 30°C (Figure 2). Increasing the temperature by 6°C did not cause any significant difference in the diel pattern (paired t-test, $P > 0.05$), and the circadian clock system maintained a circadian rhythm under this constant condition of 24.3 ± 0.02, compared with a rhythm of 24.3 ± 0.15 at ambient temperature. Fluorescence measurements were performed on *S. pistillata* fragments to test their ability to conserve a diel rhythm under an LD cycle (12:12). These fragments showed a clear diel pattern in the effective PSII quantum yield Y(II), peaking between 08:00–16:00, then decreasing until 04:00, when an increase in the yield was observed before the light came on (paired t-test, $P = 0.014$, $N = 5$) (Figure 3). Several photosynthetic parameters of the corals were measured under LL conditions: the effective PSII quantum yield Y(II), non-photochemical quenching (NPQ) and the electron transport rate (ETR). In all of the experiments, the coral fragments were subjected to an LD cycle of 12:12 with an ambient sea-water temperature of 22°C and a light intensity of 100 µmol quanta m⁻² s⁻¹. The fragments were subsequently transferred to LL conditions and subjected to a temperature of 22°C (control) or elevated temperatures of 26°C, 29°C and 31°C. The *S. pistillata* fragments were analysed with the Imaging-PAM device under constant light conditions in the absence of dark adaptation to avoid the influence of a darkening period that could potentially entrain the circadian rhythm. Similar patterns were expected for each of the temperatures tested. The temporal diel measurements of effective quantum yield Y(II) showed increasing values toward midnight (12:00) and late afternoon (16:00). The quantum yield subsequently decreased during the dark period. Three hours before the onset of the light period (at 04:00), the Y(II) value began to increase. The maximal Y(II) value (at 16:00) but not the minimal value at 04:00 under the highest temperature was significantly different only from the value at 24°C (one-way ANOVA followed by Tukey’s HSD test, $P < 0.05$, $N = 4$), whereas there was a significant difference between the 16:00 to 04:00 Y(II) values at all temperatures tested (paired t-test, $P < 0.01$, $N = 4$ per temperature) (Figure 4A; Figure 5). The estimated non-photochemical quenching (NPQ) measured at the light level (501, photosynthetic active radiation (PAR)) at which the curve reached a plateau revealed a diel cycle for each temperature tested (Figure 4B; Figure 5). The NPQ values at the lowest temperature were significantly different both at 16:00 and at 04:00 when compared with the respective values at higher temperatures (one-way ANOVA followed by Tukey’s HSD test, $P < 0.01$, $N = 4$). At the highest temperature (31°C), no reduction was observed at 16:00 (8 h after the light was turned on), and the system maintained a high value for this time point (Figure 4B). The final fluorescence parameter measured was the electron transport rate (ETR) at 16:00 and 04:00 (Figure 6A, B). The rate was faster at lower temperatures (22°C, 26°C, 29°C, 31°C) than at higher temperatures (29°C, 31°C). A significant difference was observed between the rates at 16:00 and 04:00 (paired t-test, $P < 0.01$, $N = 4$ per temperature). The ETR decreased dramatically (by 20–30%) between these times. There is a correlation between the ETR and oxygen production in the photosynthetic process. The maximal value of oxygen evolution occurred at noon, and the minimal value occurred at midnight, when the electron transport rate was lowest.

**Discussion**

The photosynthetic apparatus is very sensitive to thermal stress, and even moderate changes in temperature disrupt the system32. Many studies have addressed the effects of thermal stress on coral photosynthesis and the associations between the host and the symbiotic
algae\textsuperscript{18,33–35}. Other studies have focussed on photosynthetic photochemical efficiency measurements with chlorophyll fluorescence techniques and have identified diurnal changes in the efficiency of dinoflagellates in the host coral tissues\textsuperscript{26–38}. The goal of the above studies was mainly to understand the impacts of temperature and light synergism on the bleaching phenomenon, which is lethal for corals. We did not find any previous study that characterised the relationships among the circadian clock, photosynthesis and thermal stress by testing the temperature compensation potential in symbiotic corals.

Compensation for temperature fluctuations (the $Q_{10}$ factor) is the backbone of the circadian clock system machinery. Photosynthesis is a clock-regulated process in \textit{Symbiodinium} (as shown in Figure 1 (A–D), under the growth temperature of 24°C) and can compensate for temperature changes. This ability was confirmed in our temperature experiments, which demonstrated oxygen production with a cycle close to 24 h (24°C = 24.3°C, 27°C = 23.5°C and 30°C = 23.5°C) under constant light (Figure 1A–D) and a cycle of 24.3 h under LD. The calculated $Q_{10}$ values for 27°C and 30°C were close to 1 (1.1 and 1.05, respectively). Oxygen evolution decreased when temperature increased in both clades. The decline in oxygen evolution as the temperature increased from 24°C and 30°C was 16.9% for clade D and 23.5% for clade A. Thus, the decline was much more significant for clade A. Many researchers\textsuperscript{26–30} have claimed that clade D is more thermal-resistant than other clades, while other researchers have argued that the thermal sensitivity of \textit{Symbiodinium} is not clade-specific but rather is dependent on physiological-biochemical features such as the thylakoid lipid composition, which determines the sensitivity of photoinhibition to heat stress\textsuperscript{26}. Differences in the sensitivity of photoinhibition to thermal stress were reported between two cultured species of \textit{Symbiodinium} clade A\textsuperscript{26}. The effects of short-term exposure of \textit{Symbiodinium} to elevated temperatures included an increase in oxygen evolution\textsuperscript{26}; however, our longer exposure experiment revealed a reduction in photosynthetic oxygen evolution, as was also found in freshly isolated \textit{Symbiodinium} from Caribbean corals\textsuperscript{26}. Our findings revealed a clear difference between the photosynthetic performances (based on oxygen production) of clades A and D and demonstrated the robustness of clade D under temperature elevation. No significant difference was observed in the ability of the two clades to compensate for temperature changes. An increase in temperature from 24°C to 30°C revealed a $Q_{10}$ value of 1.05, indicating the presence of a robust compensation system for the \textit{Symbiodinium} endogenous clock under constant conditions in the absence of external stimuli (in the dark), (Figure 1), even when the organisms are part of the coral host tissue (Figure 2). A similar reduction in oxygen evolution was found for cultured algae as was noted in “hospite” \textit{Symbiodinium} between the two temperatures. The circadian clock system successfully compensated for temperature changes in both \textit{Symbiodinium} systems (as free-living organisms and as part of the coral tissue), even though the second system included corals, which have an endogenous clock in addition to the algal clock.

The sensitivity of the photosynthetic machinery to elevated temperatures is primarily attributed to photoinhibition of PSII\textsuperscript{25}. Under elevated temperatures, \textit{Symbiodinium} in its symbiotic form with corals displayed photoinhibition through inhibition of the PSII repair process; therefore bleaching tolerance is a consequence of high rates of repair relative to the photodamage, rather than differential degrees of photodamage process\textsuperscript{25} as suggested by earlier works\textsuperscript{35,45}.

Diurnal patterns in steady-state quantum yield have been documented in corals that harbour endosymbiotic dinoflagellate algae of the genus \textit{Symbiodinium}, suggesting a photoprotective function pattern that is linked to the ambient light course in shallow water environments\textsuperscript{26–39}. Diel cycles were observed under LD conditions of artificial light (100 $\mu$mol quanta m$^{-2}$ s$^{-1}$); $Y(II)$ values during the day revealed an increase towards noon, followed by a decline until 4:00, when $Y(II)$ began to increase (Figure 3). The diel fluorescence measurements under constant dim light demonstrated that $Y(II)$ values increased from the time when the lights came on until noon and then dropped during the afternoon. The increasing values of $Y(II)$ at 4:00 before the appearance of the first light indicate anticipation by the photosynthetic apparatus, which prepares the system for the coming “dramatic event” of a change from dark to light (Figure 3, 4A).

The cycles under both LD and LL reveal a similar pattern and demonstrate the endogenous circadian rhythm of the photosynthetic apparatus. In this experiment, we used dim light to avoid the synergistic effect of high light levels and temperature elevation, which can mask the circadian rhythm. Because of the low-light conditions and constant temperature, the temporal patterns of $Y(II)$ were different from those of other studies that used higher or ambient light intensities\textsuperscript{18,40}. Nevertheless, the steady state of the quantum yield in \textit{S. pistillata} was circadian under LL conditions at all four
Thermal stress usually causes a decrease in the electron transport rate, as shown by Jones et al.\textsuperscript{45}. Our light-curve analysis revealed clear oscillations in the ETR parameter values. Differences between the rates at the elevated temperatures were demonstrated at two time points (16:00 and 04:00) under LL conditions (Figure 6A,B), and the electron transport rates were lower at lower temperatures (22°C, 26°C) than at higher temperatures (29°C, 31°C). The electron transport rate also decreased between 16:00 and 04:00 under constant light, similar to the patterns of the ETR curve under the natural daily cycle of corals\textsuperscript{36}, in which the capacity for electron transport increases at noon and then decreases during sunset. The temporal circadian behaviour of the ETR reflects the photosynthetic machinery, which is governed by the \textit{Symbiodinium} endogenous core clock oscillator. These findings are consistent with the oxygen production behaviour, \textit{i.e.}, higher photosynthetic production at noon and lower production during the subjective night under constant light (Figure 6).

Under elevated temperature and constant light, the photosynthetic parameters of the photosynthetic system exhibited a strongly diurnal behaviour. Similar results have been obtained in photosynthetic organisms such as macroalgae and, specifically, in two species of symbiotic corals\textsuperscript{46,47,48}.

An asymmetric temporal response of photosynthesis was observed at similar light intensities between morning and afternoon; this response has been termed the “hysteresis effect”, the “afternoon nap” or the “afternoon depression” of photosynthesis\textsuperscript{49}. The present results indicate that the hysteresis effect can be attributed to the endogenous circadian rhythm. In our experimental conditions, both temperature and light were constant. Therefore, a desynchronisation between those two physical factors was suggested by Vonshak et al.,\textsuperscript{43} but the photosynthetic system did respond to elevated temperature by altering the photosynthetic quantum yield and oxygen production. The circadian clock system thus successfully compensated for a temperature elevation of 6°C in free-living \textit{Symbiodinium} as well as in corals (temperature elevation of 9°C) harbouring \textit{Symbiodinium}. We did not attempt to determine the “breakpoint” temperature at which the circadian machinery no longer controls the rhythmicity of photosynthesis, leading to a cascade of biochemical and transcriptional changes and, finally, to bleaching. We have successfully described the circadian clock capabilities for temperature compensation displayed by free-living \textit{Symbiodinium} and the symbiotic coral/algae association. Future studies should consider integrating the relationships among thermal stress, temperature compensation, the circadian clock of \textit{Symbiodinium} photosynthesis and the potential breakdown of the coral/algae symbiosis.

**Methods**

**Collection and maintenance of corals and \textit{Symbiodinium} cultures.** The \textit{Symbiodinium} cultures used in this study were from clade A (CCMP 2467) and clade D (CCMP 2556). All cultures were grown in 2 L Fernbach flasks containing 1 L of half-strength medium (I2) without silica. Illumination was provided by lateral fluorescent lamps under a 12:12 h LD cycle. The cultures were maintained in a culture room with a controlled temperature set to 24°C for all indoor experiments. The irradiance was measured by a quantum sensor (LI-COR, Lincoln, NE, USA) as 100 \mu mol quanta m\textsuperscript{-2} s\textsuperscript{-1}. The \textit{S. pistillata} corals tested were collected by scuba divers from a depth of 5 m in front of the Interuniversity Institute (IUI) for Marine Sciences, Eilat, Gulf of Elat, Red Sea, Israel. After collection, the corals were fragmented, placed in running seawater and then placed in 900 L of aquaria for a period of one week for acclimation.

**Monitoring of oxygen evolution.** For the cultured \textit{Symbiodinium}, oxygen evolution was monitored with an OXY-4 4-channel oxygen meter coupled to 4 optods (optical oxygen sensors; PreSens, Germany) and connected to 4 dipping probes (DP-PSIs). The optods continuously monitored the oxygen evolution every 5 min. The effects of temperature and compensation ability were tested in clades A and D. For each of the temperatures tested, \(N = 4\) cultures were exposed to two LD cycles under an ambient
temperature of 24°C followed by two cycles of LL with the temperature set at 24°C, 27°C or 30°C. Similar experiments were performed with corals at temperatures of 24°C and 30°C (For each temperature N=4 corals were tested).

**Imaging-PAM measurements.** Photosynthetic efficiencies were measured in S. pistillata corals during an LD period (12 : 12) with culture under a light intensity of 100 μmol quanta m−2 s−1. The corals were analysed every 4 h with Imaging-PAM (pulse amplitude modulation; Maxi-PAM, Walz Gmbh, Effeltrich, Germany). The resulting images were analysed with the Imaging-Win software program (v.2.00 m, Walz Gmbh, Effeltrich, Germany) and recorded for each of the branches (N=5).

Light-response curves were measured with increasing illuminations of 20 s intervals (0, 1, 16, 41, 81, 141, 221, 276, 351, 426, 501, 601, 701, 801, and 901 μmol quanta m−2 s−1) under an ambient temperature of 22°C and analysed for the effective PSII quantum yield (Y(II)) (with no dark adaptation). Under an LL cycle, four ambient temperatures of 22°C, 26°C, 29°C and 31°C were examined (For each tested temperature N=4). After irradiation, the following parameters of the coral fragments were analysed: effective PSII quantum yield (Y(II)) (with no dark adaptation), non-photochemical quenching (NPQ) and electron transport rate (ETR).

**Data analysis.** Oxygen data sampled in the time domain were transformed into the frequency domain via Fourier transform. For this purpose, we coded an application in the Python environment with the ‘scipy’ library function ‘fft’. From the output of the application, we extracted the dominant frequencies that characterised the examined rhythms. The frequency of each of the experiments was used in the formula for calculating Qm = R/RI = (RI/R1)1/2 (T/2π). For statistical analysis, paired t-tests and a one-way ANOVA followed by Tukey’s HSD test were used to assess the differences among the resulting images were analysed with the Imaging-Win software program (v2.00 m, Walz Gmbh, Effeltrich, Germany).

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M.S. and O.L. designed the research; M.S. performed the research; M.S. analysed the data; M.S. and O.L wrote the paper.

Additional information
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