



Photosynthetic and Respiratory Acclimation of Understory Shrubs in Response to *in situ* Experimental Warming of a Wet Tropical Forest

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Despite the importance of tropical forests to global carbon balance, our understanding of how tropical plant physiology will respond to climate warming is limited. In addition, the contribution of tropical forest understories to global carbon cycling is predicted to increase with rising temperatures, however, *in situ* warming studies of tropical forest plants to date focus only on upper canopies. We present results of an *in situ* field-scale +4°C understory infrared warming experiment in Puerto Rico (Tropical Responses to Altered Climate Experiment; TRACE). We investigated gas exchange responses of two common understory shrubs, *Psychotria brachiata* and *Piper glabrescens*, after exposure to 4 and 8 months warming. We assessed physiological acclimation in two ways: (1) by comparing plot-level physiological responses in heated versus control treatments before and after warming, and (2) by examining physiological responses of individual plants to variation in environmental drivers across all plots, seasons, and treatments. *P. brachiata* has the capacity to up-regulate (i.e., acclimate) photosynthesis through broadened thermal niche and up-regulation of photosynthetic temperature optimum (T_{opt}) with warmer temperatures. *P. glabrescens*, however, did not upregulate any photosynthetic parameter, but rather experienced declines in the rate of photosynthesis at the optimum temperature (A_{opt}), corresponding with lower stomatal conductance under warmer daily temperatures. Contrary to expectation, neither species showed strong evidence for respiratory acclimation. *P. brachiata* down-regulated basal respiration with warmer daily temperatures during the drier winter months only. *P. glabrescens* showed no evidence of respiratory acclimation. Unexpectedly, soil moisture, was the strongest environmental driver of daily physiological temperature responses, not vegetation temperature. T_{opt} increased, while photosynthesis and basal respiration declined as soils dried, suggesting that drier conditions negatively affected carbon uptake for both species. Overall, *P. brachiata*, an early successional shrub, showed higher acclimation potential to daily temperature variations, potentially mitigating negative effects of

chronic warming. The negative photosynthetic response to warming experienced by *P. glabrescens*, a mid-successional shrub, suggests that this species may not be able to as successfully tolerate future, warmer temperatures. These results highlight the importance of considering species when assessing climate change and relay the importance of soil moisture on plant function in large-scale warming experiments.

Keywords: experimental warming, photosynthesis, respiration, stomatal traits, thermal acclimation, TRACE, tropical forests

INTRODUCTION

Tropical forests cycle a disproportionate amount of Earth's carbon dioxide (CO₂) relative to their total land area and have the highest photosynthetic rates and aboveground carbon density of all terrestrial ecosystems (Beer et al., 2010; Pan et al., 2013; Schimel et al., 2015). These critical biomes are expected to approach temperatures outside their historical climate boundaries within the next decade (Diffenbaugh and Scherer, 2011; Mora et al., 2013). However, the magnitude and direction of the effects of climate warming on tropical forest carbon balance are not well constrained (Korner, 2004; Lloyd and Farquhar, 2008; Booth et al., 2012; Wood et al., 2012; Cavaleri et al., 2015). Plant carbon balance is determined by the uptake of CO₂ into the system through photosynthesis, and the release of CO₂ through respiration, however, these two processes respond differently to changes in temperature. Instantaneous photosynthetic rates increase with increasing temperatures until an optimum is reached, after which net photosynthesis rates decline (Berry and Bjorkman, 1980), whereas respiration rates rise with temperature in an exponential fashion and eventually decline at very high temperatures that cause membrane dysfunction (reviewed in Atkin et al., 2005; O'Sullivan et al., 2013; Heskell et al., 2016). Reduced photosynthesis and increased respiration with warming above photosynthetic optimum temperatures could result in CO₂ release exceeding uptake, possibly inducing a positive feedback that would exacerbate climate warming (Cox et al., 2000; Zhang et al., 2014; Drake et al., 2016; Hubau et al., 2020).

Forest upper canopies receive a higher proportion of direct solar radiation; such that, more carbon is cycled in sun leaves than shade leaves (Ellsworth and Reich, 1993). However, even though forest understory layers show lower CO₂ uptake rates per unit leaf area compared to upper canopy leaves, shaded leaves absorb the light that is scattered and refracted past the upper leaves, significantly contributing to forest productivity. In fact, due to their high leaf area index, the shaded canopy and understory leaves can contribute more than 50% of gross primary productivity of tropical forests (Chen et al., 2012; He et al., 2018). The role that shaded leaves play in tropical forest carbon cycling may further increase as temperatures rise if upper canopies surpass their thermal thresholds (He et al., 2018; Mau et al., 2018; Pau et al., 2018). Shaded leaves could continue to play an even larger role in forest carbon cycling if elevated CO₂ lowers transpiration (Kirschbaum and McMillan, 2018) and, therefore, evaporative cooling in the upper canopy leaves, as this could potentially further increase upper canopy leaf temperatures and

reduce photosynthesis (Doughty and Goulden, 2008; Fauset et al., 2019). While some *in situ* tropical warming studies have focused on upper canopy leaves (Doughty, 2011; Slot et al., 2014), no study - to date - has investigated the physiological responses of tropical understory plants to experimental warming *in situ*, and rarely do warming experiments in any ecosystem investigate acclimation responses of *both* photosynthesis and respiration within the same study.

Both photosynthesis and respiration have the capacity to thermally acclimate (i.e., make adjustments to conserve carbon balance), which could mitigate the negative effects of increasing temperatures. Over time periods ranging from days to weeks, photosynthetic thermal acclimation can be measured as an *up-regulation* of either the optimum temperature of photosynthesis (T_{opt}), the rate of photosynthesis at that optimum temperature (A_{opt}) (Berry and Bjorkman, 1980; Way and Yamori, 2014), or through a broadened width of the photosynthetic temperature response curve (Ω) (Slot and Winter, 2017; **Table 1**). Respiratory acclimation to warming, on the other hand, manifests as a *down-regulation* of either the temperature sensitivity or the basal rate of respiration (Atkin and Tjoelker, 2003). Tropical forests experience more narrow variations in temperature than other latitudinal zones and plant function is determined by their growth environment and evolutionary history. Therefore, tropical species may not have the plasticity necessary to acclimate to climate warming to the same degree as organisms in systems that experience wider diurnal, seasonal, and inter-annual temperature ranges (Janzen, 1967; Cunningham and Read, 2003; Drake et al., 2015). Global meta-analyses (Way and Oren, 2010; Slot and Kitajima, 2015) and an *in situ* tropical canopy warming experiment (Slot et al., 2014) have shown that tropical plant respiration will likely acclimate to warmer temperatures, however, there is still large uncertainty and conflicting evidence surrounding photosynthetic acclimation (Cunningham and Read, 2003; Slot and Winter, 2017; Smith and Dukes, 2017; Crous et al., 2018). Recent studies on tropical seedlings suggest that some tropical species can photosynthetically up-regulate with increased temperatures (Slot and Winter, 2017; Smith and Dukes, 2017), however, an *in situ* warming study on tropical canopy leaves resulted in lower photosynthetic rates (Doughty, 2011). Respiration often acclimates even when photosynthesis does not, because a lack of photosynthetic acclimation can lead to less photosynthate available for respiration (Dewar et al., 1999; Atkin and Tjoelker, 2003; Aspinwall et al., 2016), resulting in lower respiration rates due to substrate limitation. There are limited examples of how tropical species will respond to experimental warming *in situ* (Doughty, 2011; Slot et al., 2014), and to date,

TABLE 1 | Abbreviations and descriptions.

Variable	Description	Units
A_{net}	Net photosynthesis	$\mu\text{mol m}^{-2}\text{s}^{-1}$
A_{opt}	Rate of photosynthesis at T_{opt}	$\mu\text{mol m}^{-2}\text{s}^{-1}$
A_{25}	Rate of photosynthesis at 25°C	$\mu\text{mol m}^{-2}\text{s}^{-1}$
g_{s_opt}	Rate of stomatal conductance at T_{opt}	$\text{mol m}^{-2}\text{s}^{-1}$
J_{max}	The maximum rate of photosynthetic electron transport	$\mu\text{mol m}^{-2}\text{s}^{-1}$
J_{opt}	The rate of J_{max} at $T_{opt,I}$	$\mu\text{mol m}^{-2}\text{s}^{-1}$
Q_{10}	Factor that describes the rate respiration increases for every 10°C increase in temperature	unitless
$R:A$	Ratio of respiration at 25°C to photosynthesis at 25°C	unitless
R_d	Dark respiration	$\mu\text{mol m}^{-2}\text{s}^{-1}$
R_{25}	Rate of respiration at 25°C	$\mu\text{mol m}^{-2}\text{s}^{-1}$
T_{leaf}	Leaf temperature	(°C)
T_{opt}	The optimum temperature for net photosynthesis	(°C)
$T_{opt,I}$	Optimum temperature of maximum photosynthetic electron transport	(°C)
$T_{opt,V}$	Optimum temperature of maximum rate of Rubisco carboxylation	(°C)
T_{veg}	Surface vegetation temperature of experimental plots	(°C)
T_{vegMAX}	Mean maximum daily surface vegetation temperature	(°C)
$T_{vegMEAN}$	Mean daily surface vegetation temperature	(°C)
T_{vegMIN}	Mean minimum daily surface vegetation temperature	(°C)
V_{cmax}	Maximum rate of Rubisco carboxylation	$\mu\text{mol m}^{-2}\text{s}^{-1}$
V_{opt}	The rate of V_{cmax} at $T_{opt,V}$	$\mu\text{mol m}^{-2}\text{s}^{-1}$
VPD	Vapor pressure deficit	kPa
WWC_{10}	Soil volumetric water content from 0 to 10 cm depth	$\text{m}^3 \text{m}^{-3}$
WWC_{20}	Soil volumetric water content from 20 to 30 cm depth	$\text{m}^3 \text{m}^{-3}$
Ω	The difference in T_{opt} and the temperature where the rate of photosynthesis is 37% of T_{opt}	(°C)

no studies have investigated how tropical plants respond to larger-scale, *in situ* whole-plant warming.

A major consequence of the scarcity of data in tropical forests, particularly for *in situ* studies, is uncertainty in our capacity to accurately model tropical ecosystem carbon exchange (Cavaleri et al., 2015; Lombardozzi et al., 2015; Mercado et al., 2018). In particular, data that inform models on how vegetation will respond to climate warming are severely lacking for tropical systems (Arneth et al., 2012; Booth et al., 2012; Cernusak et al., 2013; Huntingford et al., 2013; Wood et al., 2019). Although these models are based on the same empirical acclimation functions which were derived mostly from temperate species (Kattge and Knorr, 2007), the degree of this physiological acclimation varies (Lombardozzi et al., 2015; Smith et al., 2016; Mercado et al., 2018). Using a land-surface model, Mercado et al. (2018) recently predicted that photosynthetic capacity will be up-regulated as tropical regions acclimate to warmer temperatures, positively stimulating carbon storage in tropical regions, however, emerging empirical data do not support the idea of increased carbon uptake in African and Amazonian tropical forests (Hubau et al., 2020). To more accurately model future carbon cycling of these key ecosystems, we need to understand if these systems are able to adjust, or acclimate, to warmer temperatures (Huntingford et al., 2013).

To improve the understanding of how tropical understory plant carbon cycling will respond to climate warming, we investigated plant physiological responses of two understory shrubs to elevated temperature using a field-scale warming

experiment in a wet tropical forest that heated the understory vegetation and soil using infrared heating panels (Kimball et al., 2018). While we controlled for as much variability as possible, *in situ* manipulative experiments are inherently variable both spatially and temporally. As a result, treatment differences may be obscured by high environmental variability, especially if sample size is relatively low, as is often the case with time-intensive gas exchange response curve measurements. To address this challenge, in addition to analyzing a binary treatment effect (control vs. warmed), we also assessed plant acclimation capacity by investigating photosynthetic and respiratory responses to daily environmental fluctuations using a more granular statistical approach. Physiological plasticity is directly related to thermal acclimation, as plasticity determines whether a plant has the potential to acclimate (Piersma and Drent, 2003; Atkin et al., 2006; Gunderson et al., 2010). Species that show an ability to adjust in response to environmental variation are likely to perform better under multiple aspects of climate change compared to species with limited physiological plasticity (Nicotra et al., 2010); therefore, assessing plastic responses to environmental variations provide an additional approach for examining plants' capacity for thermal acclimation.

Based on previous *in situ* studies showing a lack of up-regulation of net photosynthesis with warming (Doughty, 2011), evidence that tropical plants are operating near physiological thresholds (Doughty and Goulden, 2008; Mau et al., 2018), and the fact that tropical forests have evolved within a very narrow climatic envelope (Janzen, 1967), we expected these tropical

shrubs to have limited photosynthetic acclimation potential. We further hypothesized that respiration would acclimate to experimental warming, as respiratory acclimation has been shown in tropical plants developed *in situ* (Slot et al., 2014).

MATERIALS AND METHODS

Study Site and Meteorological Variables

This study was conducted at the Tropical Responses to Altered Climate Experiment (TRACE) site located at the USDA Forest Service Sabana Field Research Station, within the Luquillo Experimental Forest (18°18'N, 65°50'W) in northeastern Puerto Rico. This site is located at 100 m elevation in a tropical wet forest, with Ultisol soil classification (Scatena, 1989). Mean annual precipitation during 2014–2016 was 2271 mm, with each month averaging 112–324 mm rainfall. Mean annual temperature is 24°C (Harris et al., 2012). The wet season is May through November, and January through April is drier on average, however, this tropical forest experiences relatively low temperature and precipitation seasonality. In 2016, the site's basal area of trees >1 cm was 38.76 m² ha⁻¹ and stand density was 3100 trees ha⁻¹. Canopy height averages ca. 20 m, and light level at mid canopy is approximately 600 μmol m⁻² s⁻¹. The site is a secondary forest that, at the time of this study, had been regenerating from abandoned pasture land for approximately 70 years.

Tropical Responses to Altered Climate Experiment is comprised of three heated and three control 4 m-diameter plots located in the forest understory. The heated plots (initiated September 2016) were warmed +4°C using six infrared (IR) heating panels, each positioned in a hexagonal ring and raised approximately 2.6 m above the ground (**Figure 1**). Control plots had identical infrastructure, but with no electrical power cabling and non-heated black metal panels instead of IR panels. See Kimball et al. (2018) for more detail of experimental design and infrastructure. Plots experienced less than 20% canopy openness (Reed et al., 2020). In August 2017, canopy cover was similar between treatments, where leaf area index was 6.60 ± 0.27 (mean ± sd) in control and 6.34 ± 0.22 in heated plots.

Above-canopy daily rainfall and air temperatures used to display conditions throughout this study were collected from a 25 m tower weather station located approximately 2 km from the TRACE site (**Figure 2A**). Daily rainfall (mm) at the tower station was collected using a 10 cm plastic funnel draining into a 180 ml plastic bottle. Surface vegetation temperatures (T_{veg}) of each plot were monitored using infrared thermometers (SI-121, Apogee Instruments, Logan, UT), which measured vegetation (across multiple plants) and surface soil temperature at the center of each plot with an 18° half angle field of view. Below-canopy air temperature at the study site (CS215, Campbell Scientific, Logan, UT, United States; **Supplementary Table S1**) and T_{veg} were recorded using a multiplexer and datalogger (AM16/32, CR1000, Campbell Scientific; **Figure 2B**). Soil moisture and temperature were measured at the edge, center, and midway between center and edge of each plot at 0–10 cm depth, and additional



FIGURE 1 | Photograph of one of the experimental warming plots. Photo credit: Aura M. Alonso-Rodríguez.

probes were installed at 20–30 cm depth at the plot center (CS655, Campbell Scientific; **Figures 2C,D**). Air temperature (T_{air}) and relative humidity (RH) were measured inside each plot (~1 m from the ground) using dataloggers (MX2301, Onset Computer Corp., Bourne, MA, United States) from June 20 to August 31, 2017, after the majority of plant gas exchange sampling was completed. Vapor pressure deficit (VPD) was calculated from T_{air} and RH using the Tetons equation from Monteith and Unsworth (2008).

Sampling Design

We measured net photosynthesis, leaf respiration, and stomatal traits during four measurement campaigns: two before and two after the initiation of warming. Pre-warming measurements were taken in January (winter) and August 2016 (summer). Warming was initiated on September 28, 2016, and post-warming measurements were taken in January (winter), after 4 months of warming, and May–June 2017 (summer), after 8 months of warming (**Figure 2**).

Measurements were conducted on the first fully expanded leaf of the two most common species within the plots: *Psychotria brachiata* Sw., an early successional shrub that can be prevalent in the shaded understory but performs well in an open canopy environment (Devoe, 1989; Valladares et al., 2000; Pearcy et al., 2004), and *Piper glabrescens* (Miq.) C. DC., a mid-successional shrub species (Myer and Walker, 1997). Including seedlings under 30 cm height, there were 14–58 individual *P. brachiata*

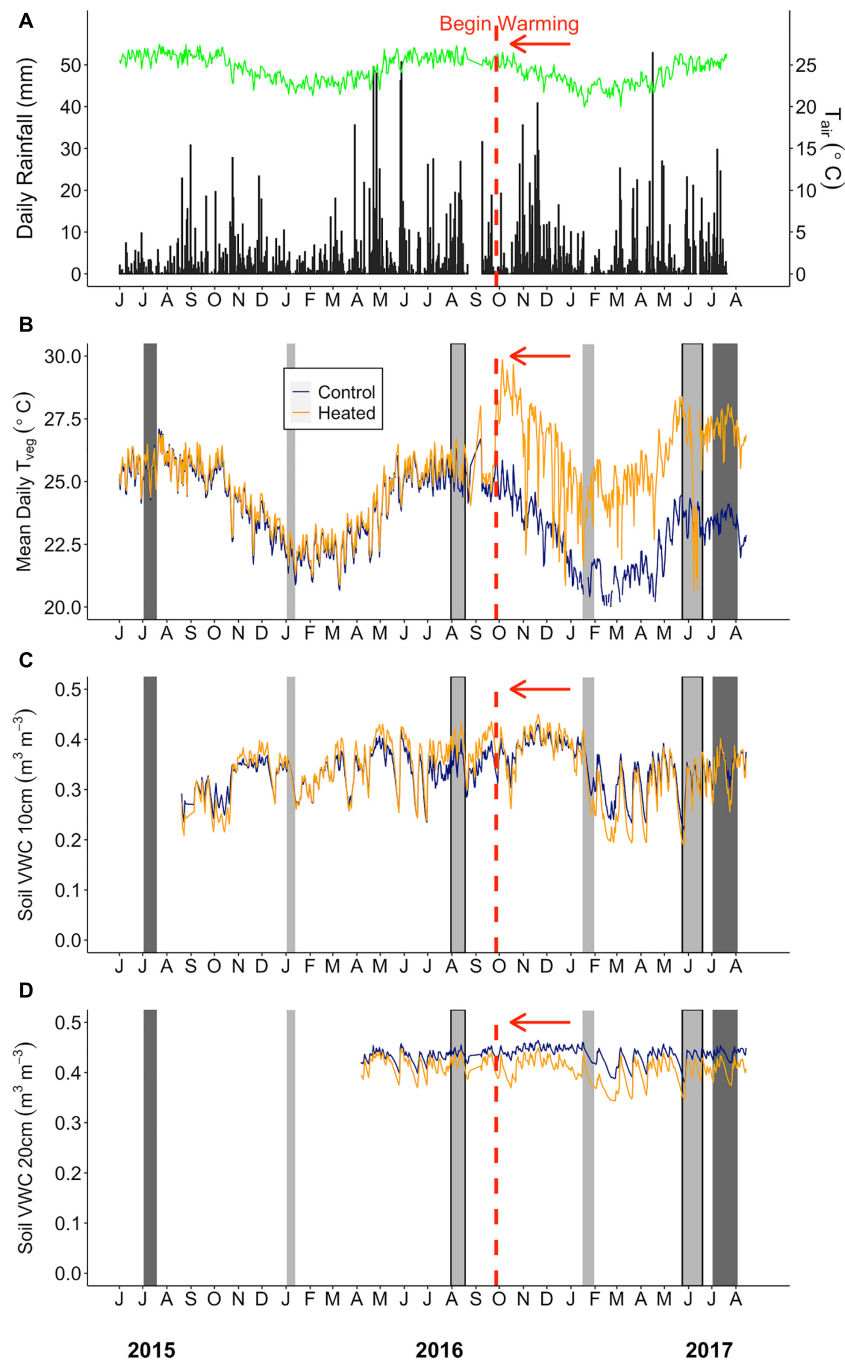


FIGURE 2 | Environmental summaries throughout the duration of the pre-warming and post-warming campaigns. **(A)** Daily rainfall (black bars) and average daily air temperature (T_{air} ; green line). **(B)** Mean daily surface vegetation temperature of the heated (orange) and control (dark blue) plots. Mean daily soil moisture at **(C)** 0–10 cm and **(D)** 20–30 cm depth for the heated and control plots. The dates shown range from July 1st 2015–August 15th 2017. The vertical red dashed line depicts the beginning of the warming treatment in the heated plots. The light gray bars depict A_{net} and R_{dark} sampling campaigns. The sampling campaigns that are light gray bars outlined in black (August 2016 and June 2017) depict campaigns where stomatal size and density was measured. The dark gray bars depict the V_{cmax} and J_{max} sampling campaigns. Rainfall and air temperature were collected from an above canopy weather station. Air temperature (°C) (HMP50-L, Campbell Scientific) was logged using a datalogger (CR1000, Campbell Scientific).

and 2–84 individual *P. glabrescens* individuals per plot in 2017. *P. brachiata* comprised 30% and *P. glabrescens* comprised 5% of the number of woody plants greater than 30 cm height

within the plots. The heights of individuals measured within each plot ranged from 25.5 to 209.0 cm for *P. brachiata* and 12.0 to 110.0 cm for *P. glabrescens*. Average stem diameter of

individuals measured 2 cm above the soil surface ranged from 3.6 to 37.6 mm for *P. brachiata* and 3.6 to 10.4 mm for *P. glabrescens*. Two to four leaves per species per plot were sampled during each campaign from separate individual plants whenever possible (Supplementary Table S2). In the cases where three leaves for a species were not available within a plot, extra leaves were measured from a separate plot. In some instances, there were not enough individual plants throughout the plots to get an adequate samples size; in this case, two leaves from the same individual plant were measured (Supplementary Table S2).

Net Photosynthesis and Stomatal Conductance Response to Temperature

We measured photosynthetic temperature response at 20, 25, 27, 30, 33, 35, 37, and 40°C on attached leaves using an LI6400XT infrared gas analyzer fitted with the 6 cm² leaf chamber (6400-02B, Li-COR Inc., Lincoln, NE, United States). Temperature was controlled by cycling hot or cold water through the Expanded Temperature Control Kit (6400-88, Li-COR Inc.) using gravity (Mau et al., 2018). Photosynthetic photon flux density was controlled at saturating irradiance (800 μmol m⁻² s⁻¹) based on photosynthetic light response curves performed on these understory plants (data not shown), CO₂ concentration at 400 ppm, and flow rate between 150 and 500 μmol m⁻² s⁻¹ to keep chamber VPD between 1 and 2 kPa. It was sometimes difficult to keep VPD below 2 kPa at temperatures above 35°C, however. Each leaf was allowed approximately 5 min to equilibrate to new chamber conditions. Measurement duration for a single temperature response curve ranged between 40 and 75 min, and measurements were conducted between 8 am and 4 pm. Each measurement campaign lasted 21–35 days. Environmental conditions were variable throughout the day and throughout campaigns.

Net photosynthetic temperature response parameters were extracted using the following equation from June et al. (2004):

$$A_{net} = A_{opt} \times e^{-\left(\frac{T_{leaf} - T_{opt}}{\Omega}\right)} \quad (1)$$

where A_{net} is net assimilation at the instantaneous leaf temperature (T_{leaf}), and Ω is the difference in T_{opt} and the temperature where photosynthesis is reduced to 37% of A_{opt} . The parameter Ω is a measure of the width of the temperature response curve, where a larger value of Ω indicates a wider curve, or broader photosynthetic thermal niche. In eight of the 124 curves, A_{net} peaked outside the range of measured temperatures, and in these instances, T_{opt} and A_{opt} were determined as the temperature at the maximum rate of photosynthesis, and Ω was not extracted. Therefore, Ω statistical analyses were based on 116 of the 124 temperature response curves. For each curve, we also extracted stomatal conductance at T_{opt} (g_{s_opt}). The parameter g_{s_opt} was extracted by fitting linear regressions to each g_s - temperature responses and extracting the rate of g_s at the photosynthetic optimum temperatures. Before g_{s_opt} was extracted, $g_s > 3$ standard deviations away from the mean were

determined to be outliers outside the range of instrumental error and were removed.

A-C_i Curves

To further investigate mechanisms underlying photosynthetic acclimation, we performed photosynthetic CO₂ response curves (A-C_i curves), at multiple temperatures to measure temperature responses of the maximum rates of Rubisco carboxylation (V_{cmax}) and the maximum rates of electron transport (J_{max}). Due to the time intensive nature of performing multiple A-C_i curves across a range of temperatures, only the most common species, *P. brachiata*, was used for V_{cmax} and J_{max} measurements. Pre-warming measurements were collected in July 2015 and post-warming measurements were taken July–August 2017, after approximately 9 months of warming (Figure 2 and Supplementary Table S2). A-C_i curves were measured on the same leaf at 20, 25, 30, 35, 40°C using a LI6400XT (Li-COR Inc.) fitted with an Expanded Temperature Control Kit (6400-88, Li-COR Inc). V_{cmax} and J_{max} were extracted from each curve constructed from 400, 300, 200, 100, 50, 0, 450, 600, 850, 1000, and 1200 ppm CO₂ concentrations, at saturating irradiance (800 μmol m⁻² s⁻¹) and a flow adjusted to control chamber VPD from 1 to 2 kPa (A-C_i curves at different temperatures shown in Supplementary Figure S1). V_{cmax} and J_{max} were extracted from the net assimilation rate (A_{net}) response to internal CO₂ concentration (C_i) using the “Ecophys” package (Duursma, 2015) in R version 3.5.0 (R Core Team, 2018), which implements the Farquhar, von Caemmerer, and Berry model (Farquhar et al., 1980; von Caemmerer and Farquhar, 1981). Biochemical parameters were extracted by fitting the J_{max} and V_{cmax} vs. temperature response curves to a peaked Arrhenius function (Medlyn et al., 2002):

$$(T_k) = (k_{opt}) \frac{H_d \exp\left(\frac{H_a(T_k - T_{opt})}{(T_k R T_{opt})}\right)}{H_d - H_a 1 - \exp\left(\frac{H_d(T_k - T_{opt})}{(T_k R T_{opt})}\right)} \quad (2)$$

where T_k is the measured leaf temperature in Kelvin, (k_{opt}) is the value of J_{max} or V_{cmax} at the optimum temperature (μmol m⁻² s⁻¹), H_a is the activation energy, or exponential increase, in an Arrhenius function (kJ mol⁻¹), H_d is the decrease in J_{max} or V_{cmax} after T_{opt} and was held constant at 200 kJ mol⁻¹ (Medlyn et al., 2002), and R is the universal gas constant (8.314 JK⁻¹mol⁻¹). Equation 2 was fit as one curve for all measurements made within a single plot; therefore, all measurements collected in one plot were calculated as one sample.

Leaf Dark Respiration

Foliar dark respiration (R_d) was measured on the same leaves that we used to measure net photosynthesis whenever possible. R_d measurements were conducted using a LI6400XT fitted with the 6400-05 conifer chamber head wrapped in aluminum foil and a water jacket (Expanded Temperature Control Kit 6400-088 Li-COR Inc., Lincoln, NE, United States). We used this chamber because it fit larger leaf areas, providing more accurate detection of the low R_d rates. For each measurement, a single leaf

was rolled or folded loosely to fit in the chamber and to allow adequate air mixing. Whether or not entire leaves fit inside of the chamber, respiration rates were corrected by the actual leaf area inside the chamber. For respiration – temperature response curves, we measured CO₂ efflux rates at 25, 30, 35, 37, and 40°C for each leaf. Measurements began at least 1 h after sunset, from approximately 7:00 pm to no later than 2:00 am. Chamber reference CO₂ was controlled at 400 ppm. Each curve took approximately 25–35 min to complete.

Leaf respiration generally declines at temperatures higher than 50°C (O'Sullivan et al., 2017), but shows an exponential increase with temperature below this threshold. We measured the rate of respiration at temperatures up to 40°C; therefore, each respiratory response curve was fitted to the exponential, non-linear equation as in Cavaleri et al. (2008) and Slot et al. (2014):

$$R_d = \beta_0 \times \exp(T_{leaf} \times \beta_1) \quad (3)$$

where R_d is the respiration rate ($\mu\text{mol m}^{-2} \text{s}^{-1}$) at T_{leaf} and β_0 and β_1 are model parameters. The change in respiration rate with every 10°C (Q_{10}) is calculated as:

$$Q_{10} = \exp(10 \times \beta_1) \quad (4)$$

R_{25} was calculated using:

$$R_{25} = \frac{R_{T_{leaf}}}{(T_{leaf}-25)/10} \quad (5)$$

where $R_{T_{leaf}}$ is the respiration rate at T_{leaf} . R_{25} was calculated for each measurement temperature and then averaged to obtain one value for each leaf.

R:A Ratio

To assess leaf carbon balance, the ratio of leaf respiration to photosynthesis (R:A) was calculated by dividing R_{25} by the photosynthetic rate at 25°C (A_{25}). A_{25} was extracted from Eq. 1 by setting T_{leaf} equal to 25. For the eight curves that would not fit Eq. 1, the actual photosynthetic rate measured at 25°C was used for the values of A_{25} . When respiration and photosynthesis were measured on separate leaves, measurements were matched from the same individual plant.

Stomatal Traits

To provide further mechanistic insight to photosynthetic responses, we measured stomatal size and density during August 2016 (pre-warming) and late June 2017 (post-warming) for *P. brachiata* only. Stomatal traits were collected on fully expanded leaves that developed during the warming treatment. Stomatal impressions were collected by applying clear nail varnish to the abaxial side of the leaf. Clear cellophane tape was used to remove the dried varnish and mounted to glass microscope slides. Photos of the slides were taken under 20× magnifications using a compound light microscope (Eclipse 400, Nikon Instruments Inc., Melville, NY, United States) and camera (DFC295, Leica Microsystems Inc., Buffalo Grove, IL, United States) fitted with a 55X coupler. Stomatal density was calculated as the number

of stomata within the 20× magnified area and divided by total visible area using ImageJ v.1.50. Stomatal size was calculated by multiplying the length and width, including guard cells, of each stoma visible within the magnified area.

Detecting Acclimation by Analyzing Gain Score Treatment Effects

We found differences between plant temperature response parameters in the heated and control plots prior to the initiation of the warming treatment (Figures 3, 4). To account for this natural variation and to avoid spurious treatment effects, the changes in measured physiological parameters (A_{opt} , T_{opt} , Ω , g_{s-opt} , Q_{10} , R_{25} , R:A) in response to warming were analyzed using a gain score analysis, rather than simply comparing treatment versus control plants after warming initiation. Gain scores were calculated as the difference between post-warming and pre-warming plot averages, and were analyzed using two-way ANOVAs by treatment, season, and their interaction. Stomatal size and density gain scores were calculated as the difference between post-warming and pre-warming plot averages, measured during summer only, using Student's *t*-tests to compare treatments. J_{max} and V_{cmax} data were also only collected once pre- and post- warming (Figures 2B–D); therefore, Student's *t*-tests were used to compare differences between the optimum temperature of V_{cmax} (T_{optV}), the optimum temperature of J_{max} (T_{optJ}), the rate of V_{cmax} at T_{optV} (V_{opt}), the rate of V_{cmax} at 25°C, the rate of J_{max} at T_{optJ} (J_{opt}), and the rate of J_{max} at 25°C. V_{cmax} and J_{max} parameters were successfully extracted for two control and three heated plots; therefore, control plot gain scores were analyzed using two of the three control plots.

Gain scores were also used to analyze how plot environmental variables [daily maximum (T_{vegMAX}), mean ($T_{vegMEAN}$), and minimum (T_{vegMIN}) surface vegetation and soil volumetric water content at 10 cm (VWC_{10})] differed between treatment and season, calculated using dates of measurement campaigns only. Student's *t*-tests were used to compare VPD between the heated and control plots using hourly means, then comparing mean VPD between heated and control plots during daytime hours.

Acclimation Effect Size

To assess acclimation potential of the physiological parameters in a synthetic meta-analytic way, we calculated effect size of each gain score for all gas exchange and stomatal traits. Effect size was calculated as in Gurevitch et al. (1992) and Hedges and Olkin (1985):

$$d = \frac{Y_e - Y_c}{s} \times J_m \quad (6)$$

where d is the effect size, Y_e is the mean of the experimental gain score, Y_c is the mean of the control gain score, and s is the standard deviation calculated as:

$$s = \sqrt{\frac{((N_e - 1) \times s_e^2) + ((N_c - 1) \times s_c^2)}{N_e + N_c - 2}} \quad (7)$$

where N_e is the experimental gain score sample size, N_c is the control gain score sample size, s_e is the standard deviation of the

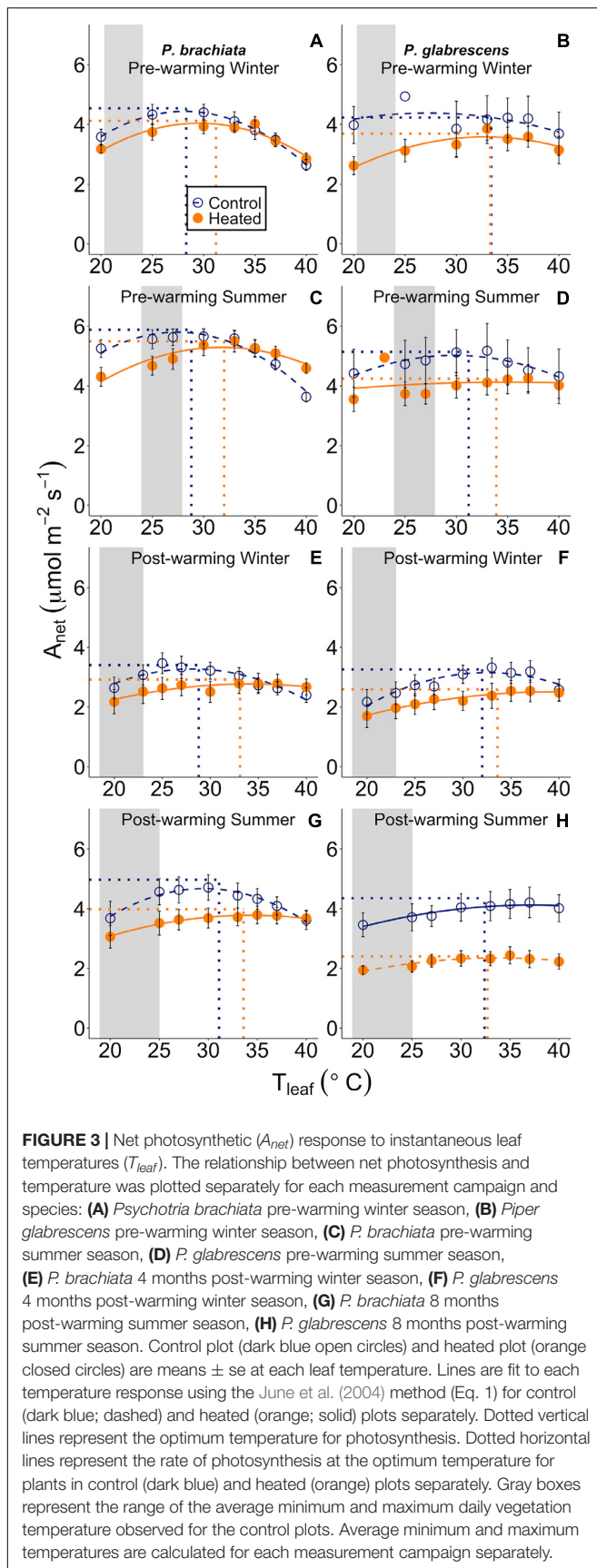


FIGURE 3 | Net photosynthetic (A_{net}) response to instantaneous leaf temperatures (T_{leaf}). The relationship between net photosynthesis and temperature was plotted separately for each measurement campaign and species: (A) *Psychotria brachiata* pre-warming winter season, (B) *Piper glabrescens* pre-warming winter season, (C) *P. brachiata* pre-warming summer season, (D) *P. glabrescens* pre-warming summer season, (E) *P. brachiata* 4 months post-warming winter season, (F) *P. glabrescens* 4 months post-warming winter season, (G) *P. brachiata* 8 months post-warming summer season, (H) *P. glabrescens* 8 months post-warming summer season. Control plot (dark blue open circles) and heated plot (orange closed circles) are means $\pm$ se at each leaf temperature. Lines are fit to each temperature response using the June et al. (2004) method (Eq. 1) for control (dark blue; dashed) and heated (orange; solid) plots separately. Dotted vertical lines represent the optimum temperature for photosynthesis. Dotted horizontal lines represent the rate of photosynthesis at the optimum temperature for plants in control (dark blue) and heated (orange) plots separately. Gray boxes represent the range of the average minimum and maximum daily vegetation temperature observed for the control plots. Average minimum and maximum temperatures are calculated for each measurement campaign separately.

experimental gain score, and s_c is the standard deviation of the control gain score.

J_m is a correction for small sample size biases and is calculated as:

$$J_m = 1 - \frac{3}{4m - 1} \quad (8)$$

Where m is calculated as:

$$m = N_e + N_c - 2 \quad (9)$$

Seasons were pooled for effect size analysis.

Detecting Acclimation by Analyzing Responses to Environmental Variables

The large variation in actual environmental conditions across space and time represented in “warmed” and “control” plots of a large-scale *in situ* manipulative experiment and the nature of these time-intensive physiological measurements resulted in rather low statistical power for the gain score analysis (i.e., using only binary categorical data). Additionally, we compared plots rather than individual plants across time when calculating gain scores because we did not always measure the same plants in each campaign, which decreased our sample size further for this analysis. To increase our statistical power for detecting acclimation, we took advantage of our rich datasets of actual environmental conditions experienced by these plants and investigated physiological plasticity in response to environmental variables (i.e., using continuous data). First, mixed effects models were used to investigate how photosynthetic temperature response parameters responded to daily values of temperature (T_{vegMAX} , $T_{vegMEAN}$), and soil volumetric water content at depths of 0–10 and 20–30 cm (VWC_{10} , and VWC_{20} , respectively) for each season. Respiratory temperature response parameters were analyzed in response to T_{vegMIN} , $T_{vegMEAN}$, VWC_{10} , and VWC_{20} because respiration was measured at night, when minimum temperatures occur. Models were run for each species separately with plot as the random variable, where individual plant was the sampling unit. In cases where multiple leaves were measured for a single plant, the mean of the dependent parameter of the two leaves were used as a single sample. Two Q_{10} values > 3 standard deviations from the mean were removed. Environmental variables on the day prior to gas exchange sampling were used because the heaters were turned off for safety during sampling days. Mixed effects models were analyzed using the “lmer” function in the “lme4” package (Bates et al., 2015). P -values were extracted for the mixed effect model using the “Anova” function from the “car” package in R (Fox and Weisberg, 2011), which calculates p -values based on Wald chi-square tests. In addition, we used a hierarchical partitioning approach to quantify the explanatory power of each environmental variable on parameter variance using the “heir-part” package in R (Walsh and Mac Nally, 2013). By using these additional approaches to assess acclimation potential of our study species, we were not only able to capture responses to the high spatial and temporal variability in environmental conditions that were obscured when only comparing binary treatment effects, but we also were able to tease apart the most important environmental drivers.

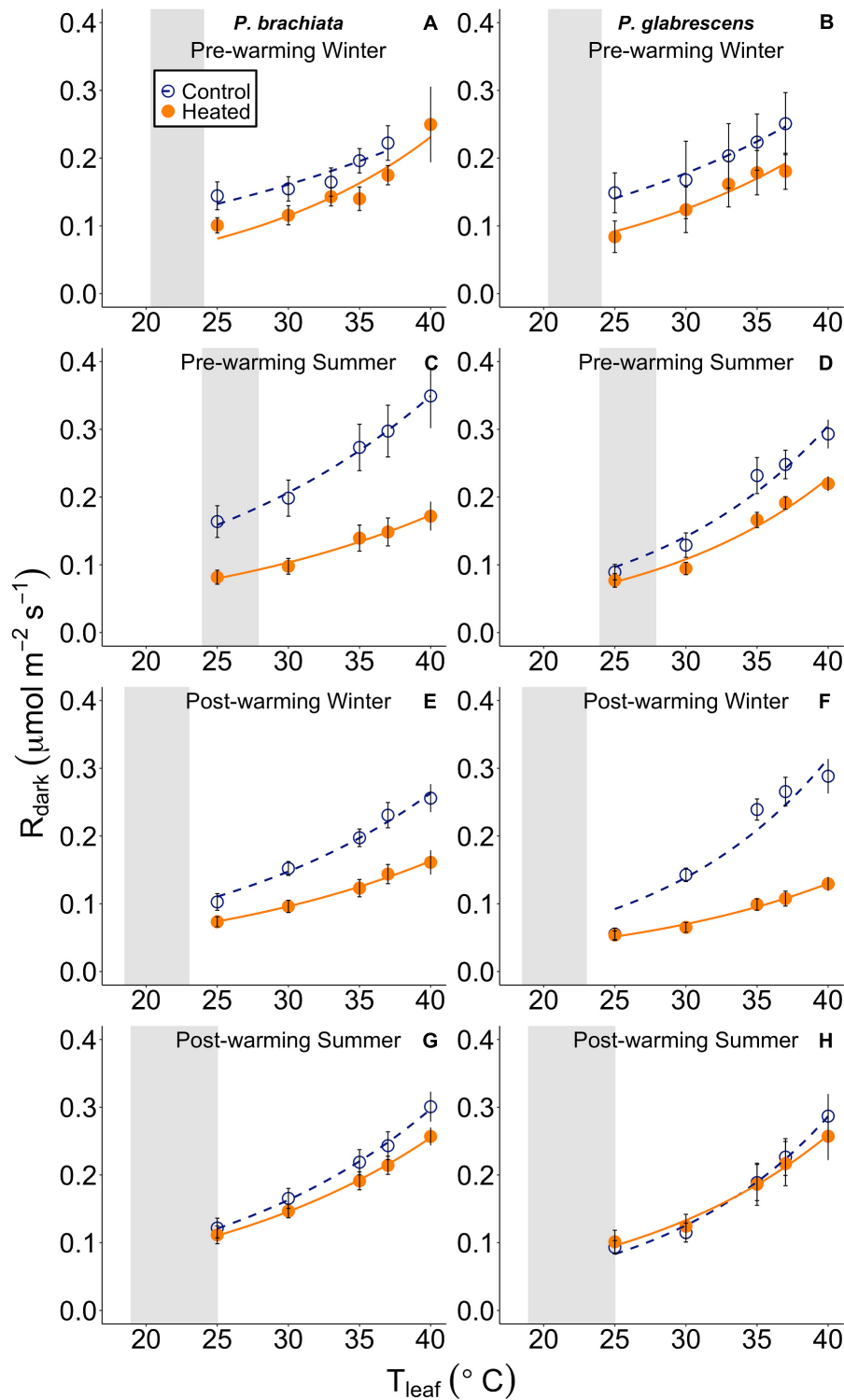


FIGURE 4 | Leaf respiratory (R_{dark}) response to instantaneous leaf temperatures (T_{leaf}). The respiratory response to temperature was plotted separately for each measurement campaign and species: **(A)** *Psychotria brachiata* pre-warming winter season, **(B)** *Piper glabrescens* pre-warming winter season, **(C)** *P. brachiata* pre-warming summer season, **(D)** *P. glabrescens* pre-warming summer season, **(E)** *P. brachiata* 4 months post-warming winter season, **(F)** *P. glabrescens* 4 months post-warming winter season, **(G)** *P. brachiata* 8 months post-warming summer season, **(H)** *P. glabrescens* 8 months post-warming summer season. Control plot (dark blue open circles) points and heated plot (orange; closed) points are means $\pm$ se at each leaf temperature. Exponential fit lines were fit to control (dark blue; dashed) and heated (orange; solid) plots separately. Gray boxes represent the range of the average minimum and maximum daily vegetation temperature for the control plots. Average minimum and maximum temperatures are calculated for each measurement campaign separately.

RESULTS

Environmental Variables

On average, the summer sampling campaigns were both hotter and rainier than the winter campaigns. Average daily precipitation was 3–6 times higher, and average daily below-canopy air temperature (T_{air}) was $\sim 3.5^{\circ}\text{C}$ warmer in summer than winter (Supplementary Table S1 and Figure 2A). Daily average minimum T_{air} showed slightly less variability between seasons ($\sim 1\text{--}4^{\circ}\text{C}$), while maximum T_{air} showed a greater difference between summer and winter ($\sim 3\text{--}7^{\circ}\text{C}$; Supplementary Table S1). Similarly, control plot mean daily vegetation temperature ($T_{vegMEAN}$) ranged from 23.5 to 25.3°C during summer and 20.7 to 21.9°C during winter campaigns (Figure 2B).

The infrared warming treatment resulted in hotter vegetation and drier soils compared to the control plots during both seasons. Daily mean, maximum, and minimum vegetation temperature gain scores (i.e., the difference between post- and pre-warming) showed a treatment effect (Supplementary Table S3), where heated leaf T_{vegMAX} was $\sim 4^{\circ}\text{C}$ greater, heated leaf $T_{vegMEAN}$ was $\sim 3^{\circ}\text{C}$ greater, and heated leaf T_{vegMIN} was $\sim 2^{\circ}\text{C}$ greater compared to the control plots for both seasons (Supplementary Figures S2A–C). We did find a “seasonal” effect for $T_{vegMEAN}$, T_{vegMIN} , and VWC_{10} (Supplementary Figures S2B–D), however, this does not indicate environmental parameter differences between summer and winter. Gain scores measure the change from pre- to post-warming; therefore, a “seasonal” effect indicates that there is more inter-annual variation during one of the seasons compared to the other. Additionally, soil volumetric water content at 0–10 cm (VWC_{10}) gain score was $\sim 34\%$ lower in the heated plots than the control (Supplementary Table S3 and Supplementary Figure S2D), showing that the warming treatment did significantly alter the heated plants’ growth environment through both higher vegetation temperatures and lower soil moisture (Figures 2B,C) and this treatment effect was consistent across seasons. Environmental gain scores were only calculated during our campaigns; therefore, differences outside of this timeframe could have had higher or lower gain score results.

Vapor pressure deficit (VPD) was not measured in the heated and control plots prior to warming, so we could not calculate gain scores for this variable. From data that coincided with our final measurement campaign only (late summer 2017), heated plots showed a mean daytime VPD of 0.36 ± 0.03 kPa, which was higher than the control plots (control VPD = 0.19 ± 0.03 kPa, $p = 0.002$, Supplementary Figure S3).

Treatment Effects on Net Photosynthesis and Foliar Respiration

While *Piper glabrescens* did not show treatment effects for any photosynthetic parameters, *Psychotria brachiata* did shift to a broader photosynthetic thermal niche under the warming treatment. Gain score analysis showed the optimum temperatures of photosynthesis (T_{opt}) and the rates of both photosynthesis and stomatal conductance at that optimum temperature (A_{opt} and g_{s-opt}) were not statistically

different between warmed and control plots for either species (Supplementary Table S4 and Supplementary Figures S4A–D,G,H). However, the photosynthetic thermal niche (Ω) gain score of *P. brachiata* was $\sim 6^{\circ}\text{C}$ wider in the heated plots compared to the control plots (Supplementary Table S4 and Supplementary Figure S4E), while *P. glabrescens* Ω did not differ between treatments (Supplementary Figure S4F).

For both species, photosynthetic optimum temperatures exceeded maximum daily vegetation temperatures in both heated and control plots during all measurement campaigns. T_{opt} values ranged from 30 to 32°C in control plots and 32 to 34°C in heated plots for both species (Supplementary Table S5). Control plot T_{opt} was $\sim 7^{\circ}\text{C}$ higher than maximum vegetation temperature for *P. brachiata* and $\sim 6\text{--}9^{\circ}\text{C}$ higher for *P. glabrescens*, with greater differences during the winter (Supplementary Table S5 and Figure 3).

We found no evidence of a warming treatment effect on foliar respiration temperature response or the ratio between respiration and photosynthesis at 25°C ($R:A$) for either species. Neither *P. brachiata* nor *P. glabrescens* showed significant treatment, season, or interaction effects on the gain scores of Q_{10} , R_{25} , or $R:A$ (Supplementary Table S4 and Supplementary Figures S5A–F).

Treatment Effects on Component Processes of Net Photosynthesis and Stomata

We investigated underlying mechanisms of photosynthetic thermal acclimation by exploring the shifts in temperature responses of component processes of net photosynthesis, including the maximum rates of Rubisco carboxylation (V_{cmax}) and the maximum rates of electron transport (J_{max}) (Supplementary Table S6). Consistent with our analyses of net photosynthesis, we detected no warming treatment effects for the temperature responses of J_{max} or V_{cmax} . Neither the optimum temperature of V_{cmax} (T_{optV} ; Student’s t -test $p = 0.304$), rate of V_{cmax} at T_{optV} (V_{opt} ; $p = 0.824$), the rate of V_{cmax} at 25°C ($p = 0.503$), the optimum temperature of J_{max} (T_{optJ} ; $p = 0.546$), the rate of J_{max} at T_{optJ} (J_{opt} ; $p = 0.747$), nor the rate of J_{max} at 25°C ($p = 0.468$) gain scores differed between treatments (Supplementary Figure S6).

Psychotria brachiata leaves had slightly smaller stomata in the heated ($-23.80 \pm 26.75 \mu\text{m}^2$) compared to the control ($67.28 \pm 31.70 \mu\text{m}^2$) plot gain scores after 8 months of warming ($p = 0.095$). There was no alteration in stomatal density ($p = 0.443$), where control density was $-0.57 \pm 42.24 \text{ m m}^{-2}$ and heated density was $54.29 \pm 49.50 \text{ m m}^{-2}$.

Acclimation Effect Size for All Variables

Effect size calculations were used to show acclimation responses of gas exchange and stomatal trait gain scores in a standardized meta-analytic framework. Photosynthetic acclimation indicates an up-regulation of photosynthesis with experimental warming; therefore, photosynthetic parameter (T_{opt} , A_{opt} , Ω , g_{s-opt} , T_{optV} , T_{optJ} , V_{opt} , J_{opt}) effect size greater than zero indicates acclimation. With an effect size of 1.36 ± 0.41 (effect size $\pm$ variance), *P. brachiata* thermal niche (Ω) was the only photosynthetic

parameter that was not close to overlapping zero (Figure 5). Evidence of respiratory acclimation would result in Q_{10} and R_{25} , and $R:A$ parameter effect sizes lower than zero (i.e., down-regulation). Agreeing with the gain score analyses, neither species showed high magnitude respiratory effect sizes (Figure 5). Adjustment of stomatal traits can result in either increasing or decreasing stomatal size and/or density (e.g., Shen et al., 2017), however, the only stomatal trait showing acclimation was *P. brachiata* stomatal size. With an effect size of -1.43 ± 0.84 , *P. brachiata* adjusted to have slightly smaller stomata under experimental warming (Figure 5).

Gas Exchange Parameter Responses to Environmental Variables

While categorical gain score analysis of treatment and season only revealed photosynthetic thermal niche treatment effects, we did find significant responses to continuous environmental variables across plots. Overall, photosynthetic optimum temperatures and thermal niches increased for *P. brachiata* as vegetation became warmer, but *P. glabrescens* photosynthesis and stomatal conductance declined as temperatures rose. *P. brachiata* T_{opt} increased significantly with daily mean and maximum vegetation temperatures (Supplementary Table S7, Figure 6A and Supplementary Figure S7A). *P. glabrescens* T_{opt} showed no response with $T_{vegMEAN}$ or T_{vegMAX} (Supplementary Table S7, Figure 6B, and Supplementary Figure S7B). *P. brachiata* A_{opt} did not respond to surface vegetation temperature (T_{veg} ; Figure 6C and Supplementary Figure S7C); although, A_{opt} intercept was higher during winter for $T_{vegMEAN}$ analysis but higher during summer for the T_{vegMAX} analysis (Supplementary Table S7). *P. glabrescens* A_{opt} declined as $T_{vegMEAN}$, but not T_{vegMAX} , increased (Figure 6D and Supplementary Figure S7D) and was higher during the summer (Supplementary Table S7). Thermal niche (Ω) broadened for *P. brachiata* as T_{veg} rose (Figure 6E and Supplementary Figure S7E), but *P. glabrescens* Ω did not respond to T_{veg} (Figure 6F and Supplementary Figure S7F). *P. brachiata* showed no relationship between g_{s_Opt} and T_{veg} (Figure 6G and Supplementary Figure S7G). *P. brachiata* g_{s_Opt} followed similar trends to A_{opt} , where g_{s_Opt} decreased as $T_{vegMEAN}$ increased (Figure 6H), however, the relationship was not significant for T_{vegMAX} (Supplementary Table S7 and Supplementary Figure S7H).

Temperature sensitivity of respiration did not respond to vegetation temperatures for either species, while respiratory rates at a standard temperature showed evidence of down regulation only for *P. brachiata*. Neither *P. brachiata* nor *P. glabrescens* Q_{10} responded to temperature (Figures 7A,B and Supplementary Figures S8A,B). *P. brachiata* R_{25} decreased with increasing $T_{vegMEAN}$ during the winter months only (Supplementary Table S7 and Figure 7C) and had no response to T_{vegMIN} (Supplementary Figure S8C). *P. glabrescens* R_{25} did not respond to $T_{vegMEAN}$ (Figure 7D) or T_{vegMIN} (Supplementary Figure S8D).

As soils dried, both species' optimum temperatures increased, and *P. brachiata* photosynthesis declined. *P. brachiata* and *P. glabrescens* T_{opt} rose as deeper (20–30 cm; VWC_{20}) but

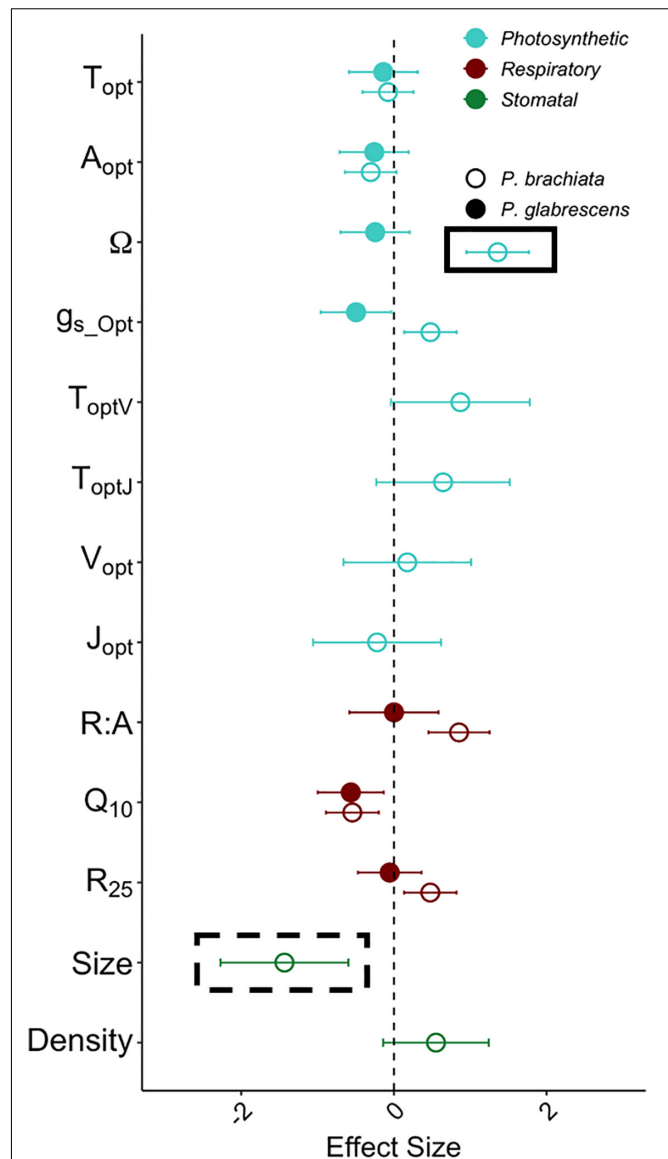


FIGURE 5 | Acclimation effect size with warming for the photosynthesis, respiration, and stomatal size and density variables. Variables represented are the optimum temperature of photosynthesis (T_{opt}) ($^{\circ}\text{C}$), the photosynthetic rate at T_{opt} (A_{opt}) ($\mu\text{mol m}^{-2} \text{s}^{-1}$), the width of the photosynthetic – temperature response curve (Ω) ($^{\circ}\text{C}$), the rate of stomatal conductance at T_{opt} (g_{s_Opt}) ($\text{mol m}^{-2} \text{s}^{-1}$), optimum temperature of the maximum rate of Rubisco carboxylation (T_{optV}) ($^{\circ}\text{C}$), optimum temperature of the maximum photosynthetic electron transport (T_{optJ}) ($^{\circ}\text{C}$), rate of the maximum rate of Rubisco carboxylation at the optimum temperature (V_{opt}) ($\mu\text{mol m}^{-2} \text{s}^{-1}$), rate of maximum photosynthetic electron transport at the optimum temperature (J_{opt}) ($\mu\text{mol m}^{-2} \text{s}^{-1}$), respiration increase with every 10°C (Q_{10}), the rate of leaf dark respiration at 25°C (R_{25}) ($\mu\text{mol m}^{-2} \text{s}^{-1}$), the ratio between R_{25} and photosynthesis at 25°C ($R:A$), stomatal density (Density) (m m^{-2}) and stomatal size (Size) (μm^2). Colors of data points depict the variable type, where light blue represents photosynthetic variables, dark red represents dark respiration variables, and green represents stomatal size and density. Symbols depict whether the sample is *Psychotria brachiata* (open) or *Piper glabrescens* (closed). Black boxes around *P. brachiata* Ω and stomatal size indicate significant (solid box) or marginally significant (dashed box) acclimation as calculated through the gain score analysis. Error bars depict variance of effect size as calculated from Gurevitch et al. (1992).

not shallow (0–10 cm; VWC_{10}) soils dried (Supplementary Table S7, Figures 8A,B, and Supplementary Figures S9A,B). *P. brachiata* A_{opt} declined as shallow soils dried during the summer but not during the winter (Supplementary Figure S9C) and declined with decreasing VWC_{20} (Figure 8C). *P. glabrescens* A_{opt} did not respond to VWC_{10} (Supplementary Figure S9C). As shown by the significant interaction (Supplementary Table S7), *P. glabrescens* A_{opt} declined as deep soils dried during the summer, however, A_{opt} was overall lower and did not respond to soil moisture during the winter (Figure 8C). Ω and g_{s_opt} did not respond to soil moisture for either species (Supplementary Table S7, Figures 7E–H, and Supplementary Figures S9E–H).

Both species showed down-regulation of respiratory basal rates with drying soils. Neither species Q_{10} responded to soil moisture (Supplementary Table S7, Figures 9A,B, and Supplementary Figures S10A,B). *P. brachiata* R_{25} did not respond to VWC_{10} but R_{25} declined as the 20 cm soils dried (Figure 9C and Supplementary Figure S10C). *P. glabrescens* R_{25} decreased with decreasing VWC_{10} and VWC_{20} (Figure 9D and Supplementary Figure S10D).

Hierarchical partitioning revealed that most variation in photosynthesis and respiratory parameters was controlled, unexpectedly, by deeper soil moisture, rather than vegetation temperature. Variance in T_{opt} , A_{opt} , and g_{s_opt} were all strongly controlled by VWC_{20} for both species (Supplementary Figure S11). Variance of thermal niche (Ω) for *P. brachiata* was strongly driven by VWC_{20} (Supplementary Figure S11A), however, *P. glabrescens* Ω variance was better explained by vegetation temperature (Supplementary Figure S11B). Q_{10} variance was relatively evenly explained by vegetation temperature and soil moisture (Supplementary Figure S11), particularly for *P. glabrescens*, while VWC_{20} explained most of the variance in R_{25} (Supplementary Figure S11).

DISCUSSION

Photosynthetic and Stomatal Responses to Temperature

Our hypothesis that tropical woody shrubs would not photosynthetically acclimate after 8 months of experimental warming was supported for *P. glabrescens*, but not *P. brachiata*. While we did not detect a warming treatment effect on the optimum temperature for photosynthesis (T_{opt}) from the gain score analysis, *P. brachiata* T_{opt} up-regulated as mean daily temperature increased (Figure 6A). *P. glabrescens* T_{opt} showed no acclimation to either experimental warming or daily temperature variations. Other studies have found that tropical species have some ability to photosynthetically acclimate to warmer temperatures (Cheesman and Winter, 2013; Slot and Winter, 2017; but see Fauset et al., 2019) or stimulate photosynthesis (Krause et al., 2013), however, acclimation was limited and had, until now, only been found in greenhouses or growth chambers. This emphasizes the importance of whole-plant *in situ* studies, such as this one, which may provide a more comprehensive representation of how plants will respond to climate warming.

Fewer warming studies investigate thermal niche acclimation, however, Slot and Winter (2017) found that thermal niche narrowed in experimentally warmed tropical seedlings where these plants shifted to maintain carbon gain under a wider temperature range. *P. brachiata* revealed a broadened photosynthetic thermal niche (Ω) following extended experimental warming (Figure 5 and Supplementary Figure S4) and in response to warmer daily temperatures (Figure 6E). Acclimation to both the experimental treatment and daily temperature variation suggests that this species' thermal niche has a particularly plastic response to warming. Similar to T_{opt} results, *P. glabrescens* showed no thermal niche acclimation potential. *P. brachiata* thermal niche acclimation occurred alongside slightly declined stomatal size with prolonged warming (Figure 5). Warming can either increase (Hill et al., 2014; Becker et al., 2017; Jumrani et al., 2017; Shen et al., 2017) or decrease (Rodrigues et al., 2016; Shen et al., 2017) stomatal density or size. For example, Wu et al. (2018) found that subtropical tree species with smaller stomata are better able to maintain rates of stomatal conductance and photosynthesis under high temperature-induced water deficits. Within our study, smaller size potentially allowed *P. brachiata* stomata to lose less water as the apertures remain open under wider temperature ranges (i.e., broadened thermal niche). Due to sampling restriction, we were unable to assess alterations in stomatal size for *P. glabrescens*; therefore, we cannot speculate if a lack of stomatal plasticity limited acclimation for this species. The co-occurring acclimation toward a wider thermal niche and slightly smaller stomata may give *P. brachiata* a slight advantage under climate warming, however, with mean values above 17°C, both of these understory species already had relatively wide photosynthetic thermal niches (Supplementary Table S5 and Figure 3).

Conversely, our hypothesis was supported for *P. glabrescens*, as this species showed no positive shift in photosynthetic temperature response with experimental warming (Figure 5 and Supplementary Figure S4). Instead of acclimating, *P. glabrescens* experienced slightly declined photosynthesis with warmer daily temperatures (Figure 6D). The negative correlation between A_{opt} and vegetation temperature in *P. glabrescens* corresponded with a decline in stomatal conductance (g_s) at T_{opt} (Figure 6H). Many model simulations of tropical forests have predicted that temperature will negatively affect carbon gain through lowered g_s (Doughty and Goulden, 2008; Lloyd and Farquhar, 2008; Galbraith et al., 2010), rather than more direct effects to photosynthetic machinery. For our species, *P. glabrescens* experienced declines in g_s as vegetation warmed (Figure 6H), while *P. brachiata* did not (Figure 6G). *P. brachiata* leaves shifted toward lower stomatal size in the heated leaves (Figure 5), suggesting that *P. brachiata* may be able to more readily maintain plant water status. While we did not measure plot-level VPD during most of the physiological measurement campaigns, VPD was higher in the heated plots during late summer 2017 (Supplementary Figure S3). Higher temperatures increase VPD, further increasing transpirational drive, and the shift to smaller stomata may have allowed *P. brachiata* to lose less water during warmer days. Smaller size allows stomata to close more quickly (Aasamaa et al., 2001), allowing plants to have more

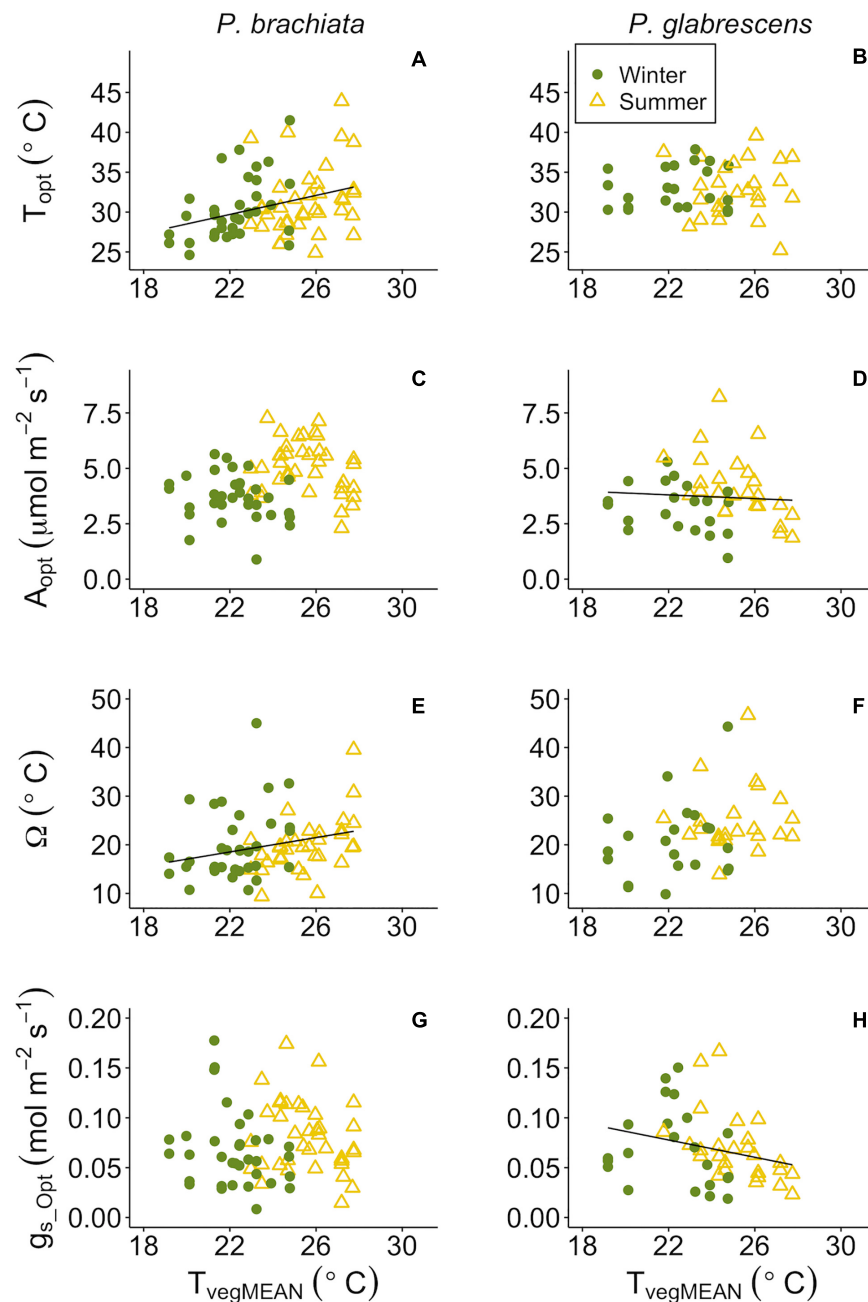
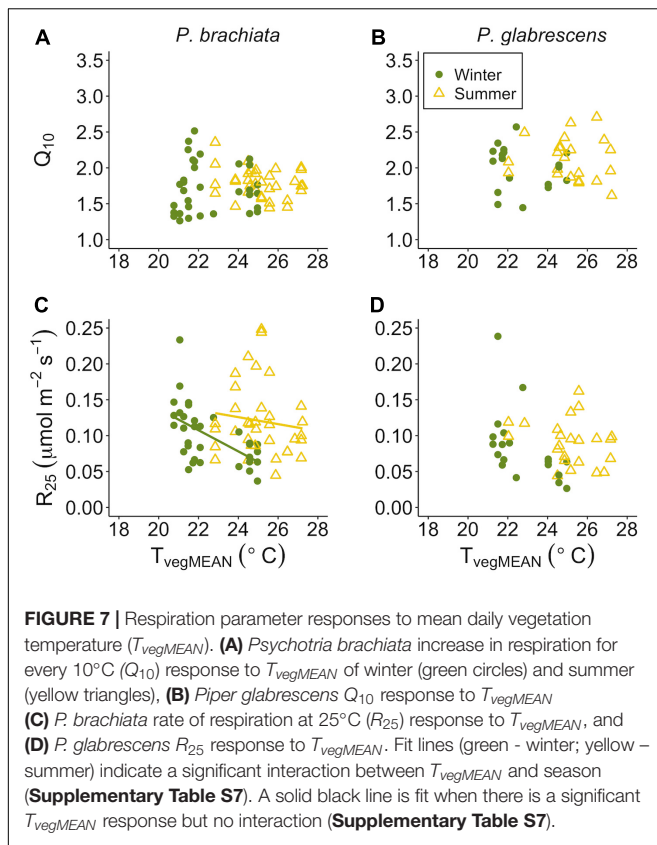


FIGURE 6 | Photosynthetic parameter responses to mean daily vegetation temperature ($T_{vegMEAN}$). **(A)** *Psychotria brachiata* optimum temperature of photosynthesis (T_{opt}) response to $T_{vegMEAN}$ of winter (green circles) and summer (yellow triangles), **(B)** *Piper glabrescens* T_{opt} response to $T_{vegMEAN}$, **(C)** *P. brachiata* rate of photosynthesis at T_{opt} (A_{opt}) response to $T_{vegMEAN}$, **(D)** *P. glabrescens* A_{opt} response to $T_{vegMEAN}$, **(E)** *P. brachiata* photosynthetic thermal niche (Ω) response to $T_{vegMEAN}$, **(F)** *P. glabrescens* Ω response to $T_{vegMEAN}$, **(G)** *P. brachiata* rate of stomatal conductance at T_{opt} (g_{s_Opt}) response to $T_{vegMEAN}$, and **(H)** *P. glabrescens* rate of stomatal conductance at T_{opt} (g_{s_Opt}) response to $T_{vegMEAN}$. Fit lines (green - winter; yellow - summer) indicate a significant interaction between $T_{vegMEAN}$ and season (**Supplementary Table S7**). A solid black line is fit when there is a significant $T_{vegMEAN}$ response but no interaction (**Supplementary Table S7**).

dynamic responses to environmental conditions (Hetherington and Woodward, 2003). This, in conjunction with overall lower water loss from smaller stomata, could further contribute to maintained photosynthesis with higher daily temperatures. Although we did find that *P. glabrescens* A_{opt} declined with

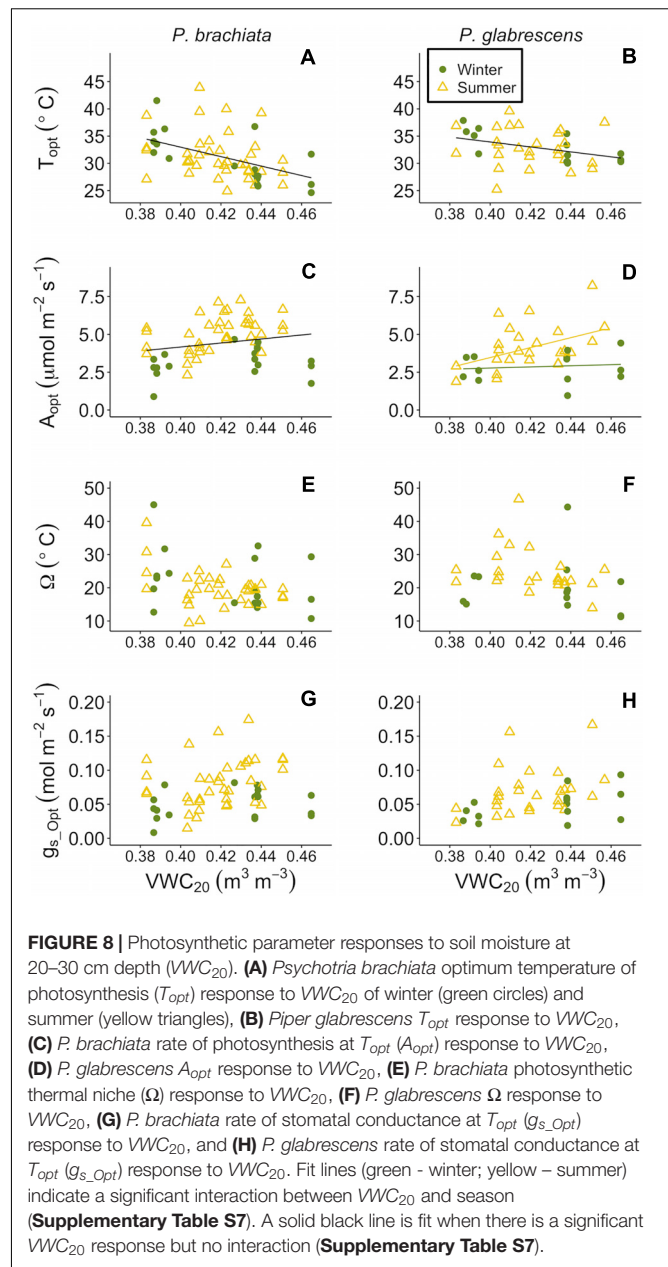
increasing temperature, *P. glabrescens* A_{opt} was up-regulated during the warmer, wetter summer months, suggesting that this species does show some seasonal plasticity.

While we found indications of *P. brachiata* A_{net} acclimation potential, we did not detect evidence of thermal acclimation

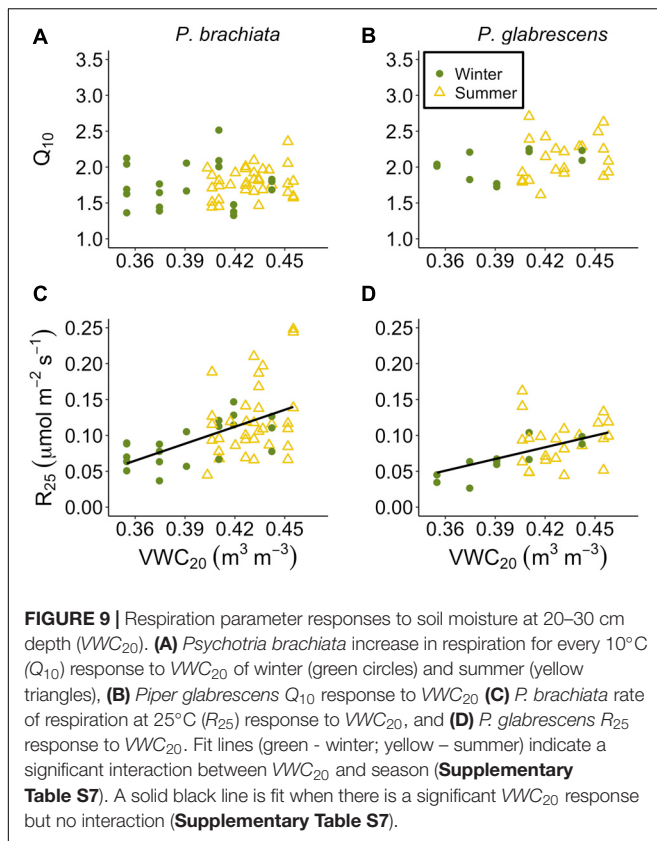


of the biochemical components of net photosynthesis (J_{max} and V_{cmax}). While both J_{max} and V_{cmax} have been found to acclimate within days (Smith and Dukes, 2017), it is possible that longer-term warming would be required to detect a shift of these processes within our study system or, perhaps, acclimation more readily occurs in the controlled environment of a growth chamber than for *in situ* studies (Gunderson et al., 2000). In addition, for both the photosynthetic and respiratory acclimation results, some of the study leaves were already fully developed prior to warming initiation. Leaf age, however, was not tracked, leading to uncertainty in whether leaves developed under the treatment (with the exception of the stomatal size and density measurements). Leaves that developed prior to initiation of warmer temperatures are less likely to acclimate (Loveys et al., 2003); therefore acclimation potential could have been limited for the fully developed leaves.

Our study site is in an area with frequent hurricanes, which can rapidly increase the light and temperature environment experienced by understory species. The higher plasticity in response to elevated temperature of *P. brachiata* may allow this species to respond more quickly to new environments (Figures 6A,E, 7C). Early successional species such as *P. brachiata* are often associated with higher plasticity (Valladares et al., 2000, 2002). There is some evidence that thermal acclimation is more likely to occur in faster growing, early successional species (e.g., Slot and Winter, 2018), however, this



is not consistent across all species (reviewed in Atkin et al., 2006). More shade tolerant species, such as *P. glabrescens*, are generally adapted to thrive in relatively stable environmental conditions (Valladares et al., 2002; Niinemets and Valladares, 2004), which is supported by *P. glabrescens*' wide thermal niches (Supplementary Table S5 and Figure 3). As a result, these two species may respond differently to the greater hurricane intensity and frequency predicted to occur with climate change (Knutson et al., 2015; Bacmeister et al., 2018). While higher *P. brachiata* plasticity may allow quicker responses to both warming and disturbance, the wider thermal niches found in *P. glabrescens* could also potentially mitigate negative effects of climate warming.



Respiratory Responses to Temperature

There is currently a general consensus that, unlike photosynthesis, tropical plants will be able to acclimate their rates of respiration (Slot and Kitajima, 2015). Contrary to this expectation, we found little to no respiratory thermal acclimation in either species. *P. brachiata* showed no acclimation response to experimental warming, however, there was slight evidence for acclimation to daily environmental variables. *P. brachiata* showed physiological plasticity by down-regulating the basal rate of respiration (R_{25}) as vegetation warmed, but that trend only occurred during the cooler, drier winter months (Figure 7C). *P. brachiata* R_{25} decreased as soils dried (Figure 9C); therefore, R_{25} might have been more sensitive to warming temperatures, and the corresponding drier soils, during the drier winter season. *P. glabrescens* respiration did not acclimate to daily changes in temperature or experimental warming (Figure 5 and Supplementary Figures S5, S7B,D). While there are few studies to corroborate respiratory acclimation through *in situ* tropical warming studies, Slot et al. (2014) found that canopy leaves exposed to only one week of nighttime warming were able to acclimate through a down-regulation of basal respiration rate (R_{25}). Studies on juvenile tropical species have found evidence of both decreased slope (Cheesman and Winter, 2013; Krause et al., 2013) or both decreased basal rate and slope (Cheesman and Winter, 2013; Drake et al., 2015) of respiratory acclimation. Meta-analyses suggest that plants across all biomes will acclimate their rates of respiration in response to warmer temperatures (Way and Oren, 2010; Slot and Kitajima, 2015) and, our study

adds an important set of results suggesting that respiratory acclimation might not be universal across tropical species.

There were no treatment differences in the ratio of respiration to photosynthesis ($R:A$; Figure 5 and Supplementary Figure S5E,F). Because these species were operating well below their optimum temperatures for photosynthesis (Supplementary Table S5), +4°C of current conditions would likely not negatively affect leaf carbon balance for these two species. It is possible, however, that maximum vegetation temperatures inside the plots were underestimated (Supplementary Table S5). The infrared thermometer reads temperature over multiple surfaces; therefore, measured temperature was likely lower than the actual temperature experienced for some individual leaves if the thermometer was measuring multiple leaves experiencing different temperatures. Photosynthesis and respiration were measured on different days; therefore, we were unable to investigate variability in $R:A$ with daily fluctuations of temperature and soil moisture.

Soil Moisture: A Stronger Driver Than Temperature

Our study aimed to investigate how plants specifically respond to elevated temperature, however, along with heating plant tissues, our warming treatment caused soil drying (Supplementary Figure S2D). Changing precipitation patterns and soil drying are predicted to have large effects on ecosystem carbon balance (Ciais et al., 2005; Phillips et al., 2009; Kao and Ganguly, 2011; Sherwood and Fu, 2014). Periodic drought and lowered soil moisture cause declines in tropical forest productivity (Santos et al., 2018; van Schaik et al., 2018). Importantly, hierarchical partitioning revealed that deeper soil moisture, even more so than vegetation temperature or shallow soil moisture, was the most influential climate variable determining variation in many gas exchange parameters (Supplementary Figure S11). Deeper, rather than shallow, soil moisture may have been a stronger driver because it was less variable in general (Figure 2D; Kimball et al., 2018) likely because of less surface evaporation and fewer roots present to rapidly draw water from that soil depth (Yaffar and Norby, 2020). T_{opt} increased as soil moisture decreased (Figures 8A,B), suggesting that neither species' T_{opt} will likely be negatively affected by a drying climate. T_{opt} increasing with drying soil in both of our species is somewhat surprising and the increase may not be a direct effect of soil drying. If increases in vegetation temperature (T_{veg}) coincided with soil drying, the negative relationship between T_{opt} and soil moisture could be due to the indirect influence of higher T_{veg} , however, T_{opt} only increased with T_{veg} for *P. brachiata*. We may lack the data to fully explain T_{opt} responses to soil moisture.

While drier soils may not negatively affect T_{opt} , we did observe decreased A_{opt} for *P. brachiata* as soils dried (Figure 8C). This negative association between photosynthesis and drier soil also occurred for *P. glabrescens* during the summer (Figure 8D), suggesting that drying soil has the potential to negatively affect both species' carbon uptake. Similarly, a long-term study found that photosynthesis declined as soil moisture decreased in 11 boreal and temperate species, likely due to

stomatal conductance restrictions (Reich et al., 2018). R_{25} also decreased as soil moisture decreased (Figures 9C,D), perhaps following the trend of A_{opt} , where decreased substrate may have limited the rate of respiration in drier soils. Soil drying often occurs with ecosystem-level warming experiments, and is also expected to occur with climate warming (e.g., Jarvi and Burton, 2013; Rich et al., 2015). Accordingly, the role soil moisture played in our study and the likelihood that, in the absence of precipitation changes, higher temperatures will result in reduced soil moisture, reinforces the importance of investigating how both temperature and moisture affect plant responses to climate change.

Assessing Physiological Responses by Treatment Versus Actual Environmental Conditions

We assessed thermal acclimation of physiological responses in two ways: (1) by assessing treatment effects using a binary comparison of control vs. warmed plots, and (2) by exploring responses to actual environmental conditions experienced by the plants in the day prior to each measurement. Although the experiment was initially designed to assess thermal acclimation through treatment effects, we detected stronger responses when looking at the responses to values of environmental variables across all treatments and time points. Several factors may have contributed to the disparity between the two assessments. First, *in situ* studies contend with many sources of variation across space and time. Gunderson et al. (2010) found a higher capacity for acclimation in seedlings grown in a growth chamber than saplings grown in the field. The authors suggest factors such as uncontrolled VPD and lower soil moisture could have limited acclimation capacity *in situ*.

The differences between plots prior to initiation of the warming treatment necessitated the gain score analysis comparing pre- and post- warming responses. This approach required all leaf-level measurements to be pooled at the plot level, thus dramatically reducing our statistical power compared to the regression-style analyses, which included plant-level data. We found no differences in temperature, soil moisture, or available nutrients between treatments that could easily explain pre-warming physiological variation between heated and control plots (Kimball et al., 2018; Reed et al., 2020). In January 2016, prior to initiation of the warming treatment, there were no differences in foliar nitrogen per unit leaf area for either *P. brachiata* (Student's *t* test, $p = 0.122$) nor *P. glabrescens* ($p = 0.520$). Our experience here highlights not only the value of pre-treatment measurements, but also the importance of high-resolution measurements of actual environmental conditions when designing large-scale *in situ* experiments.

CONCLUSION AND IMPLICATIONS

We assessed physiological acclimation potential of two Puerto Rican understory shrub species +4°C in an *in situ* field-scale warming experiment in a tropical forest and, overall, found different acclimation potential between species. Only one of our two study species, *P. brachiata*, acclimated to

warmer temperatures through a wider photosynthetic thermal niche and higher photosynthetic optimum temperatures. In contrast, our other study species, *P. glabrescens*, showed no evidence photosynthetic acclimation, and in fact showed declines in both photosynthesis and stomatal conductance with warmer temperatures. Surprisingly, we found little evidence for leaf respiratory acclimation, *P. brachiata* R_{25} decreased with increasing vegetation temperature during the drier winter season, but *P. glabrescens* showed no acclimation to experimental warming or daily temperature variations. These data provide important insight into how tropical understory plants respond to warming, as there are limited examples and, to our knowledge, no other large-scale *in situ* assessments of tropical forest understory plant physiological acclimation.

Expanding *in situ* experimental warming studies to include more species across different tropical forest types is essential, as a lack of acclimation could indicate altered tropical forest carbon uptake as global temperatures rise. Of our two study species, *P. brachiata* may be more resilient to climate warming due to higher plasticity in traits that conserve water and promote carbon gain. *P. glabrescens* experienced lower rates of photosynthesis at higher daily vegetation temperatures, perhaps due to reduced stomatal conductance. If a trend of higher physiological plasticity in early successional species and reduced photosynthesis with higher shade tolerance is conserved across more tropical understory species, we could see shifts in understory composition with climate warming. Soil moisture played an important role in determining the variation of many gas exchange variables and thus should be carefully considered *in situ* warming studies.

This study was conducted on understory species, yet upper canopy foliage may respond similarly to temperature as shaded layers (Slot et al., 2019) and, due to high heat stress, leaves in the canopy are *already* operating at or above thermal thresholds for photosynthesis (Doughty and Goulden, 2008; Mau et al., 2018). Thus, canopy foliage may not have the plasticity to up-regulate physiology to the same degree as understory plants. If so, the understory could become an even more dominant component of tropical forests' carbon cycle. In conclusion, our study addresses a critical gap in our understanding of how tropical plants may respond to warming and suggests that species' plasticity may play an important role in their ability to respond to climate change.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article doi: 10.2737/RDS-2020-0055.

AUTHOR CONTRIBUTIONS

MC, TW, and SR designed the TRACE experiment and acquired financial support for the overall study. KC and MC designed the physiological study. KC collected the photosynthesis, respiration, and leaf trait data. ES collected the stomatal size and density data. KC carried out all of the statistical analysis and wrote the first draft of the manuscript. MC provided additional advice and contributed to revisions of the text. All authors provided input and revisions to the final draft.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/ffgc.2020.576320/full#supplementary-material>

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Supplementary Figure 1 | $A-C_i$ curves at different temperatures.

Supplementary Figure 2 | Gain score analysis of heated and control plot vegetation temperature and soil moisture.

Supplementary Figure 3 | VPD of heated and control plots during summer 2017.

Supplementary Figure 4 | Photosynthetic parameter gain scores between pre-warming and post-warming measurements.

Supplementary Figure 5 | Dark respiration parameter gain scores between pre-warming and post-warming measurements.

Supplementary Figure 6 | Biochemical parameter responses to leaf temperature.

Supplementary Figure 7 | Photosynthetic parameter responses to maximum daily vegetation temperature.

Supplementary Figure 8 | Respiratory parameter responses to minimum daily vegetation temperature.

Supplementary Figure 9 | Photosynthetic parameter responses to soil moisture at 0–10 cm depth.

Supplementary Figure 10 | Respiratory parameter responses to soil moisture at 0–10 cm depth.

Supplementary Figure 11 | Hierarchical partitioning results of gas exchange parameter variances explained by environmental variables.

Supplementary Table 1 | Mean daily vegetation temperature of the control plots, mean, minimum, and maximum daily air temperature and rainfall during each measurement campaign.

Supplementary Table 2 | Count of total individual trees and leaves measured for each campaign, species, treatment, and gas exchange measurement.

Supplementary Table 3 | P -values and degrees of freedom from ANOVA results of gain score of vegetation temperature and soil moisture.

Supplementary Table 4 | P -values and degrees of freedom from ANOVA results of gain score of leaf gas exchange variables.

Supplementary Table 5 | Mean daily maximum vegetation temperature and physiological parameter means of the heated and control plots for each species.

Supplementary Table 6 | Temperature response parameters estimated for the maximum rate of Rubisco carboxylation (V_{cmax}) and the maximum rate of electron transport (J_{max}).

Supplementary Table 7 | Summary of mixed effects model results of gas exchange parameter responses to environmental variables.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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