Leaf-cutting ants’ critical and voluntary thermal limits show complex responses to size, heating rates, hydration level, and humidity

Cleverson Lima1,3 · André Frazão Helene1 · Agustín Camacho1,2

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Abstract
Thermal variation has complex effects on organisms and they respond to these effects through combined behavioral and physiological mechanisms. However, it is less clear how these traits combine in response to changes in body condition (e.g., size, hydration) and environmental factors that surround the heating process (e.g., relative humidity, start temperatures, heating rates). We tested whether these body conditions and environmental factors influence sequentially measured Voluntary Thermal Maxima (VTmax) and Critical Thermal Maxima (CTmax) in leaf-cutting ants (Atta sexdens rubropilosa, Forel, 1908). VTmax and CTmax reacted differently to changes in body size and relative humidity, but exhibited similar responses to hydration level, start temperature, and heating rate. Strikingly, the VTmax of average-sized workers was closer to their CTmax than the VTmax of their smaller and bigger sisters, suggesting foragers maintain normal behavior at higher temperatures than sister ants that usually perform tasks within the colony. Previous experiments based on hot plate designs might overestimate ants’ CTmax. VTmax and CTmax may respond concomitantly or not to temperature rises, depending on body condition and environmental factors.

Keywords Voluntary thermal maximum · Body size · Dehydration · Heating rate · Relative humidity · Critical thermal maximum

Introduction
Accurately predicting ecological responses to climate change requires a thorough understanding of how organisms perform under thermal stress in different contexts. Traditionally, this problem has been approached by comparing thermal limits under different conditions that may alter tolerance levels (e.g., Angilletta et al. 2007; Christian and Morton 1992). However, since organisms integrate behavioral and physiological thermal tolerance to deal with temperature rises (Williams et al. 2008), detailed information on behavioral responses to temperature is needed to accurately predict responses to climate warming.

The physiological performance of ectotherms in response to temperature change is described with a Gaussian thermal performance curve (Angilletta 2014; Camacho et al. 2018; Huey and Stevenson 1979—although often skewed from the standard original model, see Sinclair et al. 2016). Within voluntarily tolerated thermal levels, different aspects of physiological performance are optimized at different temperatures (e.g., stamina may be more optimized at lower temperatures than sprint speed, Huey et al. 1984). Nonetheless, if body temperatures rise excessively, locomotor and neural processes eventually stop, and the animals reach their Critical Thermal Maximum (CTmax; Cowles and Bogert 1944; Jørgensen et al. 2020), which can kill them almost immediately (Angilletta et al. 2007; Christian and Morton 1992; Ribeiro et al. 2012).

At temperatures close to CTmax, animals often move away from heat sources, exhibiting their Voluntary Thermal Maximum (VTmax; Camacho et al. 2018; Cowles and...
Bogert 1944). This trait represents how organisms try to keep their body temperatures below too costly or dangerous levels (Martin and Huey 2008). VT\textsubscript{max} may remain invariable across populations separated by millions of years (Wiens et al. 2019), and, despite being a behavioral trait, VT\textsubscript{max} has been observed to change relatively little compared with CT\textsubscript{max} in lizards (e.g., Camacho and Rusch 2017). Moreover, preferred temperatures correlate with CT\textsubscript{max} across species (Huey and Kingsolver 1993; Sinervo et al. 2010), suggesting that thermal preference and thermal tolerance may vary being positively correlated (but not always, see Huey and Bennett 1987). In contrast, VT\textsubscript{max} and CT\textsubscript{max} have been found to vary independently within populations (e.g., lizards, Camacho et al. 2018). In this sense, behavioral and physiological traits of thermal tolerance, such as VT\textsubscript{max} and CT\textsubscript{max}, might respond differently to internal and external influences, such as different body conditions and environmental factors.

Physiological thermal tolerance is influenced by both environmental conditions and inherent characteristics of the organism. Among these characteristics, body size affects many physiological (Hurlbert et al. 2008; Jensen and Nielsen 1975; Ribeiro et al. 2012) and ecological traits (Johnson 2008; Kaspari 1993). Larger body size often raises the CT\textsubscript{max} of ectothermic animals (Angilletta et al. 2007; Christian and Morton 1992, Ribeiro et al. 2012, but see Oyen and Dillon 2018). Also, larger bodies tend to store more water and present lower rates of water loss (Edney 1977). In small arthropods (i.e., ants), longer limbs also increase the distance of the separation of the body from heated surfaces, which greatly reduces the heat load within millimeters (Cerdá and Retana 2000). Ant castes can be differentiated by morphological aspects (size), but individuals of overlapping size often specialize in different tasks within and out of the nest (called temporal castes, Wilson 1980). Accordingly, it could be expected that ants of different sizes may show different thermal tolerance as a result of exposure to different temperatures (Baudier and O’donnel 2018). Few studies have related thermal tolerance and body size in insects (e.g., Clémencet et al. 2010; Verble-Pearson et al. 2015; Baudier et al. 2015; Johnson and Stahlschmidt 2020) and none have integrated VT\textsubscript{max} and CT\textsubscript{max} before. One potential problem to explore such integration is that traditional approaches with linear models might not accurately describe the relationship of size and thermal tolerance (see Ribeiro et al. 2012 Fig. 1C, where a non-linear response of CT\textsubscript{max} to size is suggested by the data). Thus, the shapes of these relationships remain unknown.

The level of body hydration (HL) can be another body condition regulating thermal tolerance among ectothermic animals. Water loss rates increase with rising body temperatures (Edney 1977; Lighton and Bartholomew 1988), and dehydration stress lowers the VT\textsubscript{max} and CT\textsubscript{max} of some ectotherms (e.g., Anurans, Anderson and Andrade 2017). Small arthropods also react to water stress. Messor pergandei ants with lower water reserves forage closer to their nests (Lighton et al. 1994), Atta Columbica ants select resources with higher water content (Bowers and Porter 1981), and individuals with specific smaller size from Atta sexdens rubropilosa are recruited to transport water to the colony when it is in hydric stress (Ribeiro and Navas 2008). Thus, dehydrated individuals might have their VT\textsubscript{max} lower to protect themselves from lower CT\textsubscript{max}, especially if they are experiencing heat stress in drier environments (i.e., environments with lower relative humidity, RH). Yet, to our knowledge, such responses have not been evaluated in arthropods.

Other environmental traits are typical elements of the experimental setup. These are the temperature at which the animal starts to be heated (the start temperature, ST), and the rate at which temperature rises (the heating rate, HR, sometimes called ramping rates, Overgaard et al. 2012). Both may alter the risks of overheating by changing the time at which the animal is exposed to stressful hot temperatures. They also can change the animal’s perception of risk of overheating. For example, faster heating rates could lead an organism to seek thermal refuge at lower temperatures (i.e., present lower VT\textsubscript{max}), in anticipation of a higher risk of exceeding their CT\textsubscript{max}. In contrast, slower heating rates involving longer exposures to sublethal temperatures might induce higher physiological stress (Rezende et al. 2020). If such conditions lower an organism’s CT\textsubscript{max}, a decrease in VT\textsubscript{max} might be expected, under the premise that VT\textsubscript{max} will parallel CT\textsubscript{max} variation. These scenarios can be tested by manipulating heating rates (HR) and start temperatures (ST) in heat tolerance experiments (Terblanche et al. 2007; Camacho et al. 2018).

Ants play indispensable roles in mediating ecosystems’ services and disservices (Del Toro et al. 2012). This makes important to understand how these animals are affected by environmental changes that may be harmful, such as climate change trends. The small size and the availability of countless individuals make ants appropriate experimental models for thermal tolerance studies. Also, the leaf-cutting ant Atta sexdens rubropilosa (Forel 1908), which is the focus of this study, varies in body size both within and across castes (Hölldobler and Wilson 1990), which allowed us to test hypotheses regarding body size and thermal tolerance.

Based on the topics presented above, we tested if leaf-cutting ants’ VT\textsubscript{max} and CT\textsubscript{max} (1) increased in parallel with body size, (2) increased or decreased similarly with both relative humidity and hydration levels, and (3) increased or decreased similarly with variations in start temperatures and heating rates.

Quantifying the effects of body conditions and environmental factors on VT\textsubscript{max} and CT\textsubscript{max} will help us better understand the integration of behavioral and physiological...
thermal tolerance. In addition, these results will inform how experimental setups influence measures of behavioral and physiological thermal tolerance.

**Materials and methods**

**Study animals**

The ants used in the experiments came from five colonies collected at Rio Claro, SP (Brazil), near the Laboratório de Formigas Urbanas (Center of Social Insect Studies, CEIS—Universidade Estadual Paulista). They were maintained in laboratory at 24 °C ± 1 and 55–65% relative humidity. These colonies were brought and maintained for 1–2 years at the Laboratório de Ciências da Cognição (Department of Physiology, Instituto de Biociências, Universidade de São Paulo), where the experiments were conducted. The animals were fed every day with leaves of Acalypha spp. These conditions have been used for over 30 years by researchers of the CEIS, a reference institution of social insect studies (more details in Bueno et al. 2002 and Ribeiro et al 2012).

**The thermal tolerance meter**

We developed a device capable of sequentially measuring the VT\textsubscript{max} and CT\textsubscript{max} of four ants in about 15 min, the Thermal Tolerance Meter. In this device, ants are simultaneously heated in five chambers immersed in a thermal bath. This thermal bath consists of a transparent plastic box (15 × 10 × 8 cm) filled with water (1200 ml) and heated by a Magnetic Hot Plate Stirrer (Quimis Q261). These identical individual chambers are 5-ml polystyrene tubes, with 3.5 mm diameter. The chambers are horizontally inserted in the thermal bath, with the tip left outside to provide a temporal thermal refuge. The heated part of each tube has 8 cm length, while the refuge is 3 cm in length. Both openings of each tube are closed during the experiments by glass rods whose diameter fit the opening gap. Yet, they leave a very thin space which allows the thin thermocouple to pass through and record the body temperature at the model ant. Apart from closing the entrances, these rods allowed to push ants out of the thermal refuge, or the tube if necessary.

During the tests, four ants (one in each chamber) had their VT\textsubscript{max} and CT\textsubscript{max} measured in each run. Ants’ body temperatures were represented by that of an individual of similar size which had been killed immediately before. For that purpose, we inserted a thermocouple in this fifth individual’s thorax and placed it in the (middle) chamber after it was dead. This way, we accounted for ants’ body size and shape during the measurement of VT\textsubscript{max} and CT\textsubscript{max}, and avoided changing its mass, as it could have happened during freezing euthanasia. For each trial, we used a different model ant, which always had a similar size to the ants being tested. The body temperatures were monitored by a T-type thermocouple (1 mm diameter, Omega ®), connected to a computer through a datalogger (Picolog® TC H8). The temperature inside the thermal refuge was monitored to assess the differences with the heated part of the chamber. The thermal dynamics and thermal heterogeneity within the heating system were assessed, and they showed satisfactorily low thermal heterogeneity (See Fig. S2 in the supplementary file https://doi.org/10.6084/m9.figshare.14414243.v2). The RH of the room ranged from 55–65%, except during the experiments which had been designed to alter hydration level and relative humidity.

**Sequential measures of ants’ VT\textsubscript{max} and CT\textsubscript{max} using the thermal tolerance meter**

Before each experiment, the ants were acclimated to room temperature (25 °C) for 1 h (with water ad libitum). Each individual’s VT\textsubscript{max} was registered as the temperature at which they entered and remained in the thermal refuge for at least 7 s. In a pilot experiment, the ants could quickly visit the thermal refuge (always less than 3–4 s). Thus, by waiting 7 s before registering the VT\textsubscript{max}, we ensured the ants were avoiding the heating chamber. After recording the ants’ VT\textsubscript{max}, the refuge was occupied by the glass rod, closing it and pushing each ant back to the heating chamber, preventing it to come back into the thermal refuge. In the heating part, ants kept moving inside the chamber until their legs became paralyzed, causing disorganized locomotion (which was visible for the observer because of the fully transparent thermal bath). The CT\textsubscript{max} was then recorded and the ants were taken to a Petri dish for cooling down and observation. The ants that did not survive for at least two hours after the experiment had their VT\textsubscript{max} and CT\textsubscript{max} disregarded in the analyses (8% of 186 subjects tested (15) were disregarded from the analyses). At the end of each day of experiments, all tested ants were killed by decapitation and disposed.

**The dehydration treatment**

First, four groups of ten ants were isolated from the colonies for a few hours in recipients with water available ad libitum. Assuming that the animals were fully hydrated, they were weighted in a semi-analytical balance (readability: 0.0001 g), which was recorded as the Initial Weight (100% of Hydration Level). Then, the ants were placed in perforated vials (1.5 ml Eppendorf tubes, with one ant inside each), which were placed within four 7 × 7 × 8 cm sealed recipients. The bottoms of two of these recipients were filled with silica gel to create a dehydrating atmosphere within the recipients. The other two recipients had water-soaked silica gel, creating a very humid atmosphere. The vials containing the ants...
were separated from the silica by paper towels, preventing any direct contact with ether the silica or the water. After a few hours, the final weight of each animal was recorded and its Hydration Level (HL) was obtained as the difference between the initial and final weight (in %). This procedure provided ants with a range of differently dehydrated bodies (~75–100%).

Experiments

Measuring the effects of body size on VT$_{\text{max}}$ and CT$_{\text{max}}$

We measured the VT$_{\text{max}}$ and CT$_{\text{max}}$ of 49 individuals, ranging from 1.5 to 4 mm in head width. Head width is widely used to represent leaf-cutting ants’ size (see Wilson 1980). Although some smaller individuals (0.1–1.4 mm) can be found outside the nest, most of them aid the larvae inside the fungal gardens. The 1.5–4 mm range in head width thus represents the individuals found at the foraging trails. The sizes were measured with an analogic caliper (accuracy: 0.01 mm). For this experiment, the ST was 25°C, but the heating rate varied on an individual basis (i.e., heating rate was calculated based on how much each ant’s body temperature increased per minute during the experiment). The initial RH inside the thermal chambers during this experiment was 55–65%. Potential observer bias was evaluated in the following manner: three different observers collected the measures of the VT$_{\text{max}}$ and CT$_{\text{max}}$, independently, and then compared their measures. The observed differences between their independent measures remained below one degree across the 49 measures.

Measuring the effects of hydration level and relative humidity conditions on VT$_{\text{max}}$ and CT$_{\text{max}}$

The VT$_{\text{max}}$ and CT$_{\text{max}}$ of individuals with HL ranging 75–100% were tested at ambient RH of 50% (N = 15) and 85% (N = 20), always using individuals of about 2.3 mm head width, ST = 25 °C, and HR ~ 1 °C/min. The experiments were carried out in a controlled climatized room (FITOTRON EL011—Eletrolab). The initial RH of the air and inside thermal chambers was similar to the ambient RH, as checked using a hygrometer (HT-600 Instrutherm). Measuring humidity within the thermal chambers was not possible, but even if some unnoticed variation in RH might happen during heating, it can be safely assumed that tubes at 50% remained always much drier than the ones at 85%.

Measuring the effects of start temperature and heating rates on VT$_{\text{max}}$ and CT$_{\text{max}}$

We assessed these effects in two series of trials that we later pooled together for analysis. First, we measured the VT$_{\text{max}}$ and CT$_{\text{max}}$ of 57 individuals varying the HR between 0.5 and 3 °C/min but keeping the start temperature at 25 °C. Next, we measured 45 additional individuals, varying the ST between 23 and 32.5 °C, and heating rates between (0.6–2.6 °C/min). In this way, we ensured that ST and HR would vary independently across the full set of 102 ants, and considered both factors as continuous variables. For these trials, we used ants with about 2.3 mm in head width, taken directly from the colonies and kept for 1 h with access to water, acclimating to the room temperature (25 °C). The initial RH inside the thermal chambers was always in the range 55–65%.

Data analyses

We fitted Linear Mixed Models (Bates et al. 2014) relating the VT$_{\text{max}}$ or the CT$_{\text{max}}$, separately to the described predictors (body size, heating rate, start temperature, hydration level, and relative humidity conditions). In each fitted model, either the VT$_{\text{max}}$ or CT$_{\text{max}}$ was the response variable, and the corresponding predictors entered as fixed factors. Ants’ colony (categorical factor with five levels) entered as grouping variable (random effect) to control the lack of independence in traits among ants of the same colony.

Model selection

Before estimating fixed and random effects, we selected the best models describing the relationships between each response trait and its most relevant predictive factors. We used the Akaike Information Criterion (AIC, Akaike 1974), which penalizes the addition of parameters more than the Bayesian Information Criterion (Wang and Liu 2006), to choose the model that best describes such relationships. One model can be considered as having a better fit than another if its AIC value is lower by a difference of at least 2 (Wang and Liu 2006). Among models with a similar fit (difference below 2 AIC units), we chose the one with the lowest AIC, but considered the result given by the second-lowest. Given the important effect of heating rates (HR), we included this factor in all the models fitted.

To flexibly identify the relationships between body size and VT$_{\text{max}}$ and CT$_{\text{max}}$ in ants, we compared four models in total, ranging from first-order equations (linear) up to a fourth-order one (non-linear). As models become more complex, having a low number of categories in the fixed factor may lead to overfitting because parameters are calculated within each level of the random effects. In this case,
dropping terms and changing the random effect by a fixed effect (Barr et al. 2013). Thus, we repeated this model selection procedure twice, based on simpler, generalized least squares with either colony as fixed effect, or not including colony as factor. The procedure and results can be found in the supplementary material (Table S1).

When testing for the relative importance of hydration level (HL) and relative humidity (RH) on VT\textsubscript{max} and CT\textsubscript{max}, we used AIC to select between four competing mixed models in which colony was again the grouping factor. These models were: (1) a model containing only HL, (2) a model including HL and RH, (3) a model including both predictive variables and HR, without interactions, and (4) a model including the three terms and an interaction between HL and RH. To describe the fit of the selected model, we calculated a conditional pseudo-R\textsuperscript{2} value developed for mixed models (Nakagawa et al. 2017).

When testing the effect of ST and HR on VT\textsubscript{max} and CT\textsubscript{max}, we selected four competing models. These models were: (1) a null model containing only the intercept, (2) a model including HR alone, (3) including HR and ST independently, and (4), a model including their interaction. We also added the conditional pseudo-R\textsupersquared value for the selected model to describe this model’s fit.

All analyses were performed in the R environment (R Development Core Team 2018) using the package nlme (Pinheiro et al. 2017). Fixed Pseudo R\textsupersquared was obtained using the MuMin package (Barton 2009).

Results

Effects of body size (BS) on VT\textsubscript{max} and CT\textsubscript{max}

BS did not show linear correlations with VT\textsubscript{max} (DF 42, B 0.130, SD 0.294, t 0.451, p 0.657) or CT\textsubscript{max} (DF 42, B 0.185, SD 0.369, t 0.503, p 0.621), although heating rate did affect both measures (VT\textsubscript{max}; DF 42 B 3.466, SD 0.546, t 6.344, p 0.000; CT\textsubscript{max}; DF 42 B 1.014, SD: 0.368, t 2.755, p 0.008). Instead, a third-order polynomial explained the relationship between VT\textsubscript{max} and size better than both simpler and more complex models (AIC difference was > 2 with second best model). The observed non-linear trend in VT\textsubscript{max} was created by average-sized workers (2–2.6 mm head width) from four different colonies, whose VT\textsubscript{max} reached closer to their CT\textsubscript{max}, compared to larger and smaller individuals of the same and other colonies (Fig. 1). This trend was also indicated by the 4th order polynom, which had the second best fit.

In turn, the CT\textsubscript{max} response was not better described by more complex models compared to a straight line (Table 1). These models show that both traits respond differently to body size in leaf-cutting ants. More detailed results can be

![Fig. 1 Relationships between size, VT\textsubscript{max} and CT\textsubscript{max}](image)

### Table 1 Fit of different models describing the relationship of VT\textsubscript{max} and CT\textsubscript{max} with different factors

| Predictors               | DF | VT\textsubscript{max} | CT\textsubscript{max} |
|--------------------------|----|-----------------------|-----------------------|
| BS linear + HR           | 5  | 202.2                 | 128.89                |
| BS 2nd Order + HR        | 6  | 202.02                | 128.96                |
| BS 3rd Order + HR        | 7  | 199.67                | 129.27                |
| BS 4th Order + HR        | 8  | 201.21                | 131.19                |
| RH                       | 4  | 174.8                 | 160.08                |
| RH + HR                  | 4  | 176.32                | 172.27                |
| RH + HL + HR             | 6  | 159.84                | 144.46                |
| RH*HL + HR               | 7  | 157.24                | 143.54                |
| Intercept                | 3  | 431.64                | 336.53                |
| HR                       | 4  | 430.05                | 293.31                |
| HR + ST                  | 5  | 422.24                | 294.61                |
| HR*ST                    | 6  | 422.75                | 295.88                |

*BS body size, ST start temperature, HR heating rate, RH relative humidity, HL hydration level, DF parameters estimated*

*Indicates independent terms

*Indicates interaction between terms*
found in the supplementary material (Table S1 A for VT\textsubscript{max} and B for CT\textsubscript{max}).

**Combined effects of hydration level (HL) and relative humidity (RH) on VT\textsubscript{max} and CT\textsubscript{max}**

The competing models that we compared here were: (1) HL, (2) RH plus HL (3) the three terms, independent, (4) the three terms, with the interaction between HL and RH. Regarding the VT\textsubscript{max}, model 4 was the most likely by a difference > 2 in AIC values with the second best (Table 1). The model detects a positive but very weak interaction between HL and RH, where individuals were more reactive to HL within the humid treatment (N 40, B 0.106, SD 0.050, t 2.110, p 0.042). In general, VT\textsubscript{max} increased with HL (N 40, B 0.079, SD: 0.035, t 2.264, p 0.031) and HR (N 40, B 7.414, SD 3.118, t 2.377, p 0.023), but the effect of RH was not statistically significant (N 40, B −8.761, SD 3.936, t −194, p 0.061). See observations and resulting trends in Fig. 2.

The full output of this model can be found in Table S2 A (Table 2).

Regarding CT\textsubscript{max}, models 3 and 4 exhibited a similar fit (difference in AIC value < 1). Both models indicated the effects of the three independent terms with no interaction (Fig. 2). Model 4 suggested a lower effect for RH, with respect to model 3 (RH: N 40, B 1.345, SD 0.424, t 3.1, p 0.003; HL: N 40, B 0.081, SD: 0.023, t 3.44, p 0.001), and HR (N 40, B 9.011, SD 2.069, t 4.354, p 0.001). Full results can be found in the supplementary material (Table S2 B).

**Fig. 2** Effects of hydration state and relative humidity on CT\textsubscript{max} (above) and VT\textsubscript{max} (below). Ants’ VT\textsubscript{max} was more reactive to hydration level when heated in a humid environment (85% relative humidity, grey line, lozenges), compared to that of ants heated in a drier environment (50% relative humidity, black line, circles). Meanwhile, ants’ CT\textsubscript{max} was similarly reactive to hydration level in either a more humid (85% relative humidity, black line, squares) or drier environment (50% relative humidity, triangles)

| Table 2 | Model coefficients table |
|---------|---------------------------|
| **VT\textsubscript{max}** | Trait | Effect | DF | B  | SD  | t    | p value |
| BS on VT\textsubscript{max} | 42 | 0.13 | 0.294 | 0.451 | 0.657 |
| BS and HR on VT\textsubscript{max} | 42 | 3.466 | 0.546 | 6.344 | 0 |
| HL and RH on VT\textsubscript{max} | 40 | 0.106 | 0.05 | 2.11 | 0.042 |
| HL on VT\textsubscript{max} | 40 | 0.079 | 0.035 | 2.264 | 0.031 |
| HL and HR on VT\textsubscript{max} | 40 | 7.414 | 3.118 | 2.377 | 0.023 |
| RH on VT\textsubscript{max} | 40 | −8.761 | 3.936 | −194 | 0.061 |
| ST on VT\textsubscript{max} | 101 | 0.134 | 2.467 | 1.47 | 0.143 |
| HR on VT\textsubscript{max} | 101 | 1.313 | 0.373 | 3.516 | 0.001 |
| **CT\textsubscript{max}** | Trait | Effect | DF | B  | SD  | t    | p value |
| BS on CT\textsubscript{max} | 42 | 0.185 | 0.369 | 0.503 | 0.621 |
| BS and HR on CT\textsubscript{max} | 42 | 1.014 | 0.368 | 2.755 | 0.008 |
| HL on CT\textsubscript{max} | 40 | 0.081 | 0.023 | 3.44 | 0.001 |
| HL and HR on CT\textsubscript{max} | 40 | 9.011 | 2.069 | 4.354 | 0.001 |
| RH on CT\textsubscript{max} | 40 | 1.345 | 0.424 | 3.1 | 0.003 |
| ST on CT\textsubscript{max} | 101 | 0.041 | 0.048 | 0.85 | 0.396 |
| HR on CT\textsubscript{max} | 101 | 1.655 | 0.212 | 7.807 | 0 |
Effects of start temperature (ST) and heating rate (HR) on VT_{max} and CT_{max}

The models compared were: (1), intercept, (2) HR, (3) HR plus ST, and (4) HR plus ST, interacting. Among the three models with similarly low AIC (Table 1), the one with lowest AIC indicated that ST had no effects on either VT_{max} (N 102, B 0.134, SD 2.467, t 1.47, p 0.143) or CT_{max} (N 102, B 0.041, SD 0.048, t 0.85, p 0.396) (Fig. S1). On the contrary, raising the HR increased both ants’ VT_{max} (N 102, B 1.313, SD 0.373, t 3.516, p 0.001) and CT_{max} (N 102, B 1.655, SD 0.212, t 7.807, p 0.000), making VT_{max} range from 30 °C to 37.8 °C and CT_{max} from 40.6 to 48 °C across the range of heating rates (Fig. 3, Table S3). The model including an interaction found no effects at all, but we stay with the effect of heating rate, since it was observable in all experiments.

Discussion

Our measurement system allowed us to observe how physiological and behavioral thermal tolerance may combine in response to different factors. We discovered that the VT_{max} and CT_{max} of ants may describe different responses (linear vs non-linear) to changes in body size. No previous data relating VT_{max} and size from other ants or arthropod species are available to compare with ours. Lizards have shown a negative relationship of VT_{max} with body size (Camacho et al. 2018), possibly related to age. The body size of active leaf-cutting ants may relate less to age and mostly to their physical caste, where ants of different sizes dedicate more time to specific tasks. Small workers (i.e., head width up to 1.4 mm) often perform tasks within the nest (e.g., hyphae and larvae care, gardening, nest defense, Hölldobler and Wilson 1990; Wilson 1980) and related to water transportation (Ribeiro and Navas 2008). In turn, average-sized workers (head width about 2.2 mm) are most often involved in tasks that require more time outside the nest (temporal castes, Wilson 1980). That is, they explore, forage and recruit more often (Wilson 1980). Since workers of such sizes (2–2.6 mm head width) exhibited a higher VT_{max}, we propose the hypothesis that within morphological castes, determined by body size, temporal castes might be formed due to their predisposition to accept higher temperatures. This seems to come at the expense of exposing themselves to higher thermal risk, since their CT_{max} did not increase in parallel. Future experiments, designed within a context of division of labor, may test the two hypotheses by comparing the VT_{max} and thermal tolerance (e.g., CT_{max} or survival time at VT_{max}) of ants specifically selected when performing different tasks. Our nests were originated from wild queens and kept at room temperature. Therefore, our studied ants were never exposed to sunrays or particularly hot ant’s trails, as it would happen in the wild. In this sense, it seems unlikely that the observed increases in VT_{max} arise from adaptation or acclimation due to specific exposure to heat in captivity. Therefore, it remains to be determined how some average-sized Atta ants come to be more “thermally daring”.

Among hymenopterans, physiological differences are often found within morphologically defined castes (e.g., ability to follow pheromones, Robinson 2009). The separation of reproductive and working castes may be achieved by feeding the animals with different substances (Dussoutour and Simpson 2009; Markin 1970; Petralia and Vinson 1978). However, no inter-castes or intra-castes differences have been reported in behavioral thermal tolerance for any hymenopteran. Yet, another example of thermally daring “special forces” might be the self-heating warrior bees of the species Apis cerana and Apis mellifera, which are able to kill Asian wasps Vespa velutina (Ken et al. 2005). Having thermally daring workers that maintain normal behavior at higher temperatures might benefit nests by extending foraging times during hotter periods or larger foraging areas, while the rest of the colony occupies cooler spaces (Cerdá and Retana 2000). More thermotolerant species have been found to be more abundant among Mediterranean species (Cerdá et al. 2002), and in at least one species, larger and
more thermotolerant individuals forage at hotter hours of the day (Cerdá et al. 1998a, b). For leaf-cutting ants, as a consequence of spending longer times foraging at hotter temperatures, the lifespan of average-sized workers might be shortened (Mirhosseini et al. 2017; Rezende et al. 2014), compared to other workers specialized in bringing water (the smallest in the colony, Ribeiro and Navas 2008) or defending the colony (the largest ones, Powell and Clark 2004).

Our results on the $CT_{\text{max}}$ raise considerations for the design of studies of $CT_{\text{max}}$-size relationships, for leaf-cutting ants and other species. The homogeneous warming system used in this study (see Fig. S2 in the supplementary material for further details) prevented ants from creating large thermal gradients by raising over their legs to avoid the heating hotplate, where the temperature is often measured in studies on ants’ thermal tolerance. The existence of these gradients in laboratory assays may explain why leaf-cutting ants’ $CT_{\text{max}}$ correlates with body size in studies using a hot plate (e.g., Ribeiro et al. 2012, Whitford and Ethershank 1975), while not in ours. Longer legs might well protect larger ants by distancing them from heating surfaces (Cerdá and Retana 2000; Sommer and Wehner 2012), but our results suggest that size does not pose further protection against rapid homogeneous heating, at least within leaf-cutting ants. Yet, interspecific effects of body size on ants’ $CT_{\text{max}}$ have been found when using a setup more similar to ours (homogeneously and slowly heated vials, Baudier et al. 2015; Baudier and O’Donnell 2020). To better understand the implications of body size on the $CT_{\text{max}}$ of ants and other arthropods, future studies might combine slow and dynamic methods for calculating the $CT_{\text{max}}$ (Lutterschmidt and Hutchison 1997). In this way, they could evaluate how behavior ($VT_{\text{max}}$), critical limits ($CT_{\text{max}}$) and morphology (size) interact with the time spent at stressful/sublethal temperatures (see Castañeda et al. 2015; Rezende et al. 2020).

Hot plates are widely used for estimating the $CT_{\text{max}}$ of arthropods but this technique might overestimate this trait. This methodological problem is difficult to evaluate when using large global databases of thermal tolerance data, such as the GlobTherm (Bennett et al. 2018), resulting in a potential underestimation of their vulnerability to high temperatures. For instance, Pogonomyrmex desertorum is considered one of the most thermophilic species known, with critical limits ($CT_{\text{max}}$) up to 53 °C (Marsh 1985). In addition, other species of desert ant, heated from below (a heating mantle with a variable transformer), presented $CT_{\text{max}} > 50 ^\circ C$ (Whitford and Ethershank 1975). Using a similar hot plate procedure, Ribeiro et al. (2012) measured $CT_{\text{max}}$ up to 53 °C in Atta sexdens rubropilosa, which is not a thermophilic species, despite presenting $CT_{\text{max}}$ similar to $P. \text{ desertorum}$. Meanwhile, the maximum $CT_{\text{max}}$ of A. sexdens rubropilosa in this study was 48 °C. These cases exemplify the importance of considering the heating method during the determination of thermal tolerance in arthropods.

Hydration increased both the $VT_{\text{max}}$ and $CT_{\text{max}}$ of leaf-cutting ants. Both traits can also increase with HL across different ectothermic vertebrates (e.g., Anurans, Anderson and Andrade 2017; Guevara-Molina et al. 2020, and lizards, Camacho et al. 2018). Instead, the humidity had only observable effects on the $VT_{\text{max}}$ of ants. The humidity did increase the $CT_{\text{max}}$ of termites (Woon et al. 2018), suggesting that the physiological thermal tolerance of leaf-cutting ants is less susceptible to large changes in environmental humidity. We know of no previous studies comparing the strength of body condition (hydration level) and external (relative humidity) cues on voluntary maximum temperatures, but our results suggest ants may integrate both in their behavior. These responses agree with reports of water-stressed ants selecting leaves with higher water content (Bowers and Porter 1981). However, RH did not affect the $CT_{\text{max}}$ during heating, and ants’ $VT_{\text{max}}$ changed most in the humid treatment. Thus, we hypothesize that this response was not anticipating higher thermal risks due to lower $CT_{\text{max}}$, or risks derived from long-term exposures to high temperatures in a drier environment. Instead, they might simply be reacting to a better heat transmission within a more humid environment.

Ants’ $VT_{\text{max}}$ and $CT_{\text{max}}$ were always strongly dependent on heating rate but not start temperature. Heating rates increase the $VT_{\text{max}}$ of bullfrogs (Guevara-Molina et al. 2020) and some lizard species (e.g., skinks), but not of other lizards (Phrynosomatids, Camacho et al. 2018). In these cases, start temperatures only marginally affected the $VT_{\text{max}}$ of lizards, and did not affect the $VT_{\text{max}}$ of Anurans. These observations suggest that the total heating time does not affect their behavioral responses. Yet, all of them increased their $VT_{\text{max}}$ with heating rates, indicating that ants and other organisms might react to the time spent at stressful temperatures only. In the future, this could be evaluated using start temperatures closer to the $VT_{\text{max}}$, and very different heating rates.

Regarding physiological thermal tolerance, heating rate may increase anurans’ $CT_{\text{max}}$ while being detrimental for flies (Nyangonkodiwa and Terblanche 2010). While heating rates have been often kept constant when measuring the $CT_{\text{max}}$ (Lutterschmidt and Hutchison 1997), varying them still allows detecting the effects of other variables and bringing the experiments closer to the natural conditions. In nature, heating rates should vary importantly according to where individuals are (e.g., think of a sun-hit surface compared to a nest underground), and species’ ecology (e.g., terrestrial versus aquatic organisms) and physiology (e.g., more or less thermotolerant, or prone to lose water). Thus, further observations of $CT_{\text{max}}$ are needed to find general patterns.

Concluding, the $VT_{\text{max}}$ and $CT_{\text{max}}$ of leaf-cutting ants may respond differently to some factors (changes in body size, humidity) and in parallel to others (start temperatures,
heating rates, and hydration). Our results help understanding how behavior and thermal tolerance integrate in different situations, and also show how CT\textsubscript{max} estimates may be affected by experimental design. We propose that leaf-cutting ants may have more “thermally daring” workers of average size. To continue understanding these integrative responses of organisms to temperature rises, further studies could compare the effects of these or other factors, combining dynamic and static heating systems to better understand how behavioral and physiological limits face off the temporal dimension of thermal tolerance. With few adjustments, our system could aid the observation of these limits in other small arthropods (e.g., bees, caterpillars, spiders, etc.).

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Author contributions CSL: Execution, methodological development, data obtention, analysis, writing. AHF: Discussion of experiments and results, infrastructural and institutional support. ACG: Conception, methodological design and development, analysis, writing.

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Declarations

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 competing interest.

Ethics approval According to Brazilian law, an ethics committee is not necessary for biological experiments with invertebrates.

Consent to participate and publication of data Does not apply to invertebrates as experimental subjects. All authors consent to participate in this manuscript.

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