Species interactions mediate thermal evolution

M. Tseng | Joey R. Bernhardt | Alexander E. Chila

Abstract
Understanding whether populations and communities can evolve fast enough to keep up with ongoing climate change is one of the most pressing issues in biology today. A growing number of studies have documented rapid evolutionary responses to warming, suggesting that populations may be able to persist despite temperature increases. The challenge now is to better understand how species interactions, which are ubiquitous in nature, mediate these population responses to warming. Here, we use laboratory natural selection experiments in a freshwater community to test hypotheses related to how thermal evolution of Daphnia pulex to two selection temperatures (12 and 18°C) is mediated by rapid thermal evolution of its algal resource (Scenedesmus obliquus) or by the presence of the zooplankton predator Chaoborus americanus. We found that cold-evolved algae (a high-quality resource) facilitated the evolution of increased thermal plasticity in Daphnia populations selected at 12°C, for both body size and per capita growth rates (r). Conversely, warm-evolved algae facilitated the evolution of increased r thermal plasticity for Daphnia selected at 18°C. Lastly, we found that the effect of selection temperature on evolved Daphnia body size was more pronounced when Daphnia were also reared with predators. These data demonstrate that trait evolution of a focal population to the thermal environment can be affected by both bottom-up and top-down species interactions and that rapid temperature evolution of a resource can have cascading effects on consumer thermal evolution. Our study highlights the importance of incorporating species interactions when estimating ecological and evolutionary responses of populations and communities to ongoing temperature warming.

KEYWORDS
adaptation, algae, FlowCam, rapid evolution, species interactions, temperature, thermal evolution, thermal reaction norm, zooplankton

1 | INTRODUCTION

Understanding how populations and communities are responding to ongoing rapid climate warming is one of the most pressing issues facing biologists today. Fortunately, in both ecology and evolution there has been a long history of investigating the effects of temperature on phenotypes, populations and communities (Angilletta, Steury, & Sears, 2004; Bennett, Dao, & Lenski, 1990; Bernhardt, Sunday, & O’Connor, 2018; Bochdanovits, Jong, & Jong, 2003; Kingsolver & Huey, 2008; Loboda, Savage, Budde, Schmidt, & Haye, 2018; Mousing, Ellegaard, & Richardson, 2014; Newman, 2004; O’Connor et al., 2007; Patrick, 1971; Régnière & Nealis, 2002; Tseng et al.,...
2018; Tseng & O’Connor, 2015). These and many other ecological and evolutionary studies of temperature responses have collectively demonstrated that temperature affects all levels of organization in the natural world and that evolutionary responses of focal populations to the thermal environment can occur at ecological time scales.

Although evolutionary responses to temperature have been documented in a range of taxa including fruit flies, stickleback, mustard plants, zooplankton, lizards, fish parasites and phytoplankton (Anderson, Inouye, McKinney, Colautti, & Mitchell-Olds, 2012; Barrett et al., 2010; Bochdanovits et al., 2003; Campbell-Staton, Edwards, & Losos, 2016; Hendry, 2017; James & Partridge, 1995; Mazé-Guilmot, Blanchet, Rey, Canto, & Loot, 2016; Padfield, Yvon-Durocher, Buckling, Jennings, & Yvon-Durocher, 2016; Schaum et al., 2017; Tseng & O’Connor, 2015), most estimates of thermal evolution have been made on populations in the absence of any species interactions. Consequently, although biotic interactions such as those between predators and prey, consumers and resources, competitors, and hosts and parasites are ubiquitous in nature, we have little data showing the importance of these interactions for how a focal population responds to the thermal environment.

Bottom-up species interactions, such as those between resources and their consumers, can affect thermal responses of consumers if the resources themselves are rapidly evolving to temperature. For example, resources that occupy the lower rungs of aquatic and terrestrial food chains (e.g., phytoplankton, microbes and insects) are also the organisms mostly likely to evolve rapidly, and these trait changes may have important consequences for thermal evolution of higher trophic levels. Of particular global importance are the trophic consequences of phytoplankton thermal evolution. These microscopic organisms fix over 50% of the planet’s carbon dioxide, they produce over 50% of the Earth’s oxygen, and they are the primary source of nutrients such as proteins and essential fatty acids, which cascade up the aquatic food web and that are ultimately consumed by humans (Finkel et al., 2010).

Conversely, top-down species interactions, such as those between predators and prey (for consistency, here we will refer to the prey as consumers), can also affect thermal evolution if predators depress the consumer population below which evolutionary responses are possible, or if predators preferentially eat consumers that happen to be particularly bad, or good, at withstanding the change in temperature (Osmond et al., 2017). Given the speed with which climate is warming around the world, there is a pressing need to better understand how these common species interactions affect thermal evolution. Here, we use a cosmopolitan freshwater food web to investigate the role of bottom-up (phytoplankton thermal evolution) and top-down (presence of a voracious predator) species interactions on rapid thermal evolution in the zooplankton Daphnia pulex (Cladocera: Daphniidae, Leydig 1860).

Our tri-trophic study system consists of the phytoplankton Scenedesmus obliquus (Chlorococcales: Scenedesmaceae, Kützing, 1833, also known as Tetraselmis obliquus and Acutodesmus obliquus), Daphnia pulex, and larvae of the predatory midge Chaoborus americanus (Diptera: Chaoboridae; De Haan 1849). We use controlled laboratory experiments to assess the effects of a) S. obliquus thermal evolution, and b) predator presence, on Daphnia thermal reaction norm (TRN) evolution in response to two temperature selection treatments (12 and 18°C).

1.1 | Predictions

Given their rapid doubling time (1–2 days) and large population size during the experiment (~500,000), we expected S. obliquus to show rapid thermal evolution.

This temperature evolution manifests as shifts in cell size, whereby cell size decreases as temperature increases (Chen, Li, Dai, Sun, & Chen, 2011; Margalef, 1954). Scenedesmus obliquus also produce fewer essential fatty acids at warmer temperatures, and warm-reared S. obliquus sustain smaller populations of Daphnia compared to cold-reared algae (Breuer, Lamers, Martens, Draaisma, & Wijffels, 2013; Sikora, Dawidowicz, & Elert, 2014; Vigeolas et al., 2012). We thus expect warm-evolved S. obliquus to be a lower-quality food resource for Daphnia, and we propose two hypotheses for how phytoplankton thermal evolution may affect ecological and evolutionary responses of Daphnia to temperature-mediated selection (Box, Hypotheses 1–2). Cold-evolved (higher quality) algae could facilitate Daphnia thermal reaction norm (TRN) evolution if this food type sustains higher populations of Daphnia, thus maintaining the standing genetic variation on which thermal selection can act (Frankham, 1996). Concomitantly, if a lower-quality food source (warm-evolved algae) imposes selection for Daphnia genotypes that can better withstand this food environment, the stressful environment may erode the available genetic variation (Charmantier & Garant, 2005; Kause et al., 2001), resulting in a muted response of Daphnia to temperature selection (Box, Hypothesis 1 (a) versus (b)). Conversely, because zooplankton and phytoplankton tend to be exposed to similar abiotic environments in nature, it is also possible that we will see local adaptation, whereby cold-selected Daphnia evolve higher performance when they are fed with phytoplankton from a similar environment, and vice versa for warm-reared Daphnia (Box, Hypothesis 2 (c) versus (d)). Here, we have assayed thermal reaction norm evolution so that we can quantify the relative roles of phenotypic plasticity versus evolutionary change in Daphnia thermal responses.

We also examined whether the presence of predators mediated evolutionary responses of Daphnia to the thermal environment (Box). Here, we propose two possible outcomes (Hypothesis 3.1, 3.2). In the presence of predators, classical life-history theory predicts that consumers should evolve higher overall per capita growth rate (Abrams & Rowe, 1996) (Box, Hypothesis 3.1 (a) versus (b)). Secondly, Chaoborus larvae are known to be voracious predators of Daphnia, and thus, predator presence may reduce Daphnia population size (and thus genetic variation) to levels below which thermal evolution is feasible (Hypothesis 3.2 (a) versus (c)).

2 | MATERIALS AND METHODS

Approximately 5,000 Daphnia pulex and 100 third-instar Chaoborus americanus were collected from the University of British Columbia
Research Ponds in early November 2016 (average temperature 8°C). *Daphnia* and *Chaoborus* were brought back to UBC and held at 12°C in a walk-in environmental chamber. *Daphnia* were fed a laboratory culture of *Scenedesmus obliquus*, and *Chaoborus* were fed wild-collected *Daphnia*. *Scenedesmus obliquus* (CPCC 5) was obtained from the Canadian Phycological Culture Centre in 2015 and maintained in axenic cultures at 20°C, under light- and nutrient-saturated conditions in Bold's Basal Medium (Bold, 1949). After 4 weeks of acclimation to the laboratory environment, 30 *Daphnia* were haphazardly allocated to each of 160 glass jars (750 ml) filled with 700 ml COMBO media (Kilham, Kreeger, Lynn, Goulden, & Herrera, 1998). One third-instar *Chaoborus* was placed into half of the jars (n = 80). Half of the 160 jars were then placed in 12°C water baths (40 jars with predators, 40 without), and the other half were placed in 18°C water baths (40 jars with predators, 40 without). For the 18°C treatment, the temperature of the water bath was gradually raised from 12 to 18°C over a period of 1 week. To minimize environmental variation across water baths, each week all *Daphnia* population jars were haphazardly moved among water baths of the same temperature.

### 2.1 Algae treatments and maintenance

Sixteen replicate populations of algae were seeded using the laboratory culture of *Scenedesmus obliquus*. Populations were grown in COMBO medium in 1.9-L glass jars. Jars were capped and aerated using filtered air (0.3-µm in-line HEPA-VENT filter) from a large aquarium air pump. Eight replicate jars were maintained at 12°C (cold-algae treatment), and eight were maintained at 18°C (warm-algae treatment).

Three times per week, approximately 3/5 of the contents of each of the eight algal populations per temperature treatment were poured out into a single bucket. Individual algal jars were refilled with COMBO and returned to the water bath. The pooled contents from each temperature treatment were fed to the appropriate *Daphnia* populations. The pooling of algae across the eight populations per temperature treatment for *Daphnia* feeding was purposeful because we wanted *Daphnia* to be exposed to generic "cold-reared" or "warm-reared" algae. Algal replicate jars remained separate for quantification of experimental evolution.

At four occasions during the experiment, we quantified algal cell size, and the number of algal cells after dilution, and immediately prior to the following dilution (typically 2 days later), using a FlowCam imaging particle counter (FlowCam® VS Series; Fluid Imaging Technologies), at a flow rate of 0.3 ml/min. We calculated population growth rate during each period as cell number at time $t + 1$ divided by cell number at time $t$. There was no difference in population growth rate between algae maintained at the two temperature treatments (growth rate = 0.43 cells/day; temperature × date
F_{1,124} = 0.17, p = 0.68). This lack of difference in growth rate is perhaps surprising because we typically believe that algae grow faster at warmer temperatures. We have since confirmed this result by growing the same strain of *S. obliquus* across multiple temperatures without dilution and found that differences in growth rate are not detectable until after the third day of continuous growth (data available upon request). Finally, at the beginning of the experiment we also used haemocytometer counts to confirm density estimates generated by the FlowCam, and found little difference between the estimates of cell density obtained using the two methods (FlowCam: 402,366 cells/ml; haemocytometer: 454,000 cells/ml).

2.2 | *Daphnia* maintenance

*Daphnia* were fed either cold- or warm-reared algae for the duration of the experiment. The number of algal cells per feeding was quantified using FlowCam, and we converted cells per feeding to carbon per feeding using published biovolume to biomass conversion equations (Montagnes, Berges, Harrison, & Taylor, 1994). Algal carbon per feeding (16 mg C/L per day) exceeded the amount typically fed to *Daphnia* to maintain saturating levels of algae (1–2 mg C/L) (Horton, Rowan, Webster, & Peters, 1979; Koch, Martin-Creuzburg, Grossart, & Straile, 2012; Weider, Jeyasingh, & Looper, 2008; West & Post, 2016). *Daphnia* populations were given a 100 ml COMBO change once per week (~14% media change). Overall, this was a three-factor experiment, with two levels of predators (present/absent), two levels of *Daphnia* rearing temperature (12°C, 18°C) and two levels of algal food type (cold, warm). There were twenty replicate *Daphnia* populations per treatment combination for a total of 160 replicate populations.

2.3 | Data collection

2.3.1 | *Daphnia* population size

*Daphnia* population size was censused on days 1, 15, 45, 75, 105 and 150 of the experiment. The contents of each replicate jar were poured into a sorting tray, and all *Daphnia* within each jar were manually counted by returning individual *Daphnia* to replicate jars using a plastic disposable pipette. Approximately 76,000 individual *Daphnia* were counted over the duration of the experiment.

2.3.2 | Common garden environment

On day 150 of the experiment, all predators were removed, and the temperature of all *Daphnia* and algae populations was set to 15°C (common garden environment). Day 150 represents approximately 10–15 rounds of reproduction in *Daphnia* and ~65 cell division cycles for *S. obliquus*. *Daphnia* have overlapping generations. It takes 10–15 days to go from neonate to neonate at the two temperatures used in this experiment, and mature adults continue to produce a new brood every 3–5 days until death. *Daphnia* were fed the laboratory stock of *S. obliquus* from day 150 onward.

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**Box 2** Potential top-down effects of predators (*Chaoborus* larvae) on thermal reaction norm (TRN) evolution of consumers (*Daphnia*).

### Hypothesis 3. Predators select for higher overall consumer growth rate

| No predators | Predators | Predators |
|--------------|-----------|-----------|
| ![Graph](image) | ![Graph](image) | ![Graph](image) |

**Predicted results:**

3.1 Consumers evolve higher \( r \) at warmer selection temp.; predators select for even higher \( r \) (a vs b).

3.2 Direct predation depresses population size below which evolutionary responses to the thermal selection environment are able to occur (a vs c).
2.3.3 | Assaying evolved differences in algal respiration and photosynthesis

After two rounds of algal replication in the common garden environment, we assayed cell volume and rates of photosynthesis and respiration, across a range of temperatures. We used the FlowCam to quantify average individual algal cell volume from a 0.3 ml sample from each replicate population. *Scenedesmus obliquus* exhibits morphological variation, whereby cells can either exist as unicells or colonies (coenobia, a group of daughter cells that share a parent cell wall) (Lürling & Van Donk, 1996). Our measure of cell volume does not distinguish between these two forms; we define “volume” as the total volume of material within the cell wall.

Rates of photosynthesis and respiration from subsamples of each replicate jar were assayed using a sealed glass microplate in combination with a microplate oxygen sensor. The microplate contained 24 x 200 µl chambers each equipped with an oxygen sensor spot (Loligo Systems). The microplate was connected to a 24-channel optical fluorescence oxygen system (SDR SensorDish Reader; Presens). The reader and microplate were placed in a temperature-controlled incubator (Panasonic MIR-154). Algal photosynthesis and respiration were assayed at five temperatures (10, 12, 16, 20 and 24°C). Rates of photosynthesis were estimated by measuring rates of oxygen evolution in the light and consumption in the dark. 200 µl of well-mixed *Scenedesmus* populations from each replicate population was transferred to each well on the microplate, and measurements of oxygen concentration were taken every 15 s over 3-hr periods, first in darkness and then in light. Wells were sealed with transparent PCR Film (Thermo Scientific). Prior to oxygen flux measurements, sensor spots were calibrated with air-saturated water (100% oxygen) and water containing 2% sodium sulphite (0% oxygen) at each experimental temperature. Algal cells were acclimated to the assay temperature for 1 hr in the dark prior to measurements. Three blank wells with COMBO medium were run at the same time as the phytoplankton at each temperature, and the average rate of oxygen flux in these wells was subtracted from the experimental wells to account for background microbial respiration. Gross photosynthesis was estimated as GP = net photosynthesis + respiration at each temperature. We assumed that net photosynthesis was directly proportional to oxygen production in the light. We mass-normalized rates of photosynthesis using the mean population biovolume (mean cell volume × cell density) from the source populations, measured using the FlowCam (flow rate = 0.3 ml/min) on the day of the respirometry assays. Cell volume (µm³) was determined with area-by-diameter estimation (ABD) in each replicate population.

2.3.4 | Assaying evolved differences in *Daphnia* life-history traits

After 3 weeks in the common garden environment, four juvenile *Daphnia* from each replicate jar were placed in individual 50-ml Falcon brand centrifuge tubes. Once one of the four *Daphnia* matured and produced at least four neonates, the remaining three *Daphnia* were discarded. These four neonates were transferred to individual 50-ml Falcon tubes filled with 40 ml COMBO and placed in one of four assay temperatures: 12, 18, 22 or 26°C. Because these offspring were produced parthenogenetically, each of the 160 replicate *Daphnia* populations thus contributed four genetically identical individuals to the evolution assay. We chose to use one *Daphnia* per replicate population in order to avoid pseudoreplication. To be statistically conservative, we used “jar” as the unit of replication for the experiment. We included 20 replicate jars (selection lines) per treatment, which is at the highest end of the number of replicates typically used for this type of experimental evolution study (Bochdanovits et al., 2003; Huang, Tran, & Agrawal, 2016; Padfield et al., 2016; Schaum et al., 2017; Van Doorslaer, Stoks, Jeppesen, & Meester, 2007).

*Daphnia* in the evolution assay were maintained on laboratory *S. obliquus* stock cultures. We checked *Daphnia* daily for eggs, and once eggs were visible, *Daphnia* were removed and preserved in 95% ethanol and photographed at 10× using an inverted microscope (Leica MC120-HD). We recorded the number of days to first clutch (development time) and measured adult size and offspring number. The latter two traits were measured using the measuring tool in the Leica Application Suite software (version 4.4). *Daphnia* adult size (mm) was measured as the length of the body from the top of the head above the eyespot to the base of the tail spine. To measure egg number, we counted the number of eggs visible from a side-view photograph of the ethanol-preserved *Daphnia*. This measure is likely an underestimate of the actual egg number as we were not able to count any eggs that were hidden under the top visible layer of eggs. We used egg number to calculate *Daphnia* per capita growth rate, r, using the following equation: \( r = \ln(\text{viable eggs in first clutch}/\text{time to first clutch}) \) (Trubetskova & Lampert, 2002).

2.4 | Data analysis

2.4.1 | Algal evolution

We used linear mixed-effects models (R package lmer) to evaluate whether *S. obliquus* photosynthesis or respiration rates evolved in response to rearing temperature. In these statistical models, selection temperature (12°C, 18°C) and assay temperature (10, 12, 16, 20, 24°C) were included as fixed effects, and “replicate” was included as a random effect. We assessed the statistical significance of the treatments and interactions using likelihood ratio tests, following Winter (2013). For the statistical analysis of algal cell volume, each replicate contributed one mean value to the analysis, and we used ANOVA to test whether rearing temperature explained variation in evolved cell volume.

2.4.2 | *Daphnia* population size and evolution

We used ANOVA to assess whether selection temperature (12°C, 18°C), predators (present/absent) and food type (cold vs. warm algae) explained significant variation in mean *Daphnia* population size.
We used linear mixed-effects models (R package lmer) to examine whether rearing temperature, predators, food type and assay temperature (12, 18, 22, 26°C) explained variation in *Daphnia* evolved size at maturity, development time and per capita growth rate, *r*. Assay temperature was modelled as a continuous variable so that we could compare slopes of the thermal reaction norms among treatments. Because each replicate population jar contributed one *Daphnia* genotype (4 individuals) to the evolution assays, we included a “family” random effect in each of the models. We incorporated trait variation arising from drift among replicates by including random slopes and intercepts for the “family” term. This approach allows us to examine the variance in traits attributable to treatment effects after among-replicate variance is accounted for (Dutilleul et al., 2014).

We included all main effects and interaction terms in the full model for each dependent variable (main effects: food type, selection temperature, assay temperature, predator). We assessed the statistical significance of interaction terms using the ANOVA command in R (e.g., ANOVA model), and we removed interactions terms if they were not statistically significant. We kept nonsignificant two-way interactions in the model if a three-way interaction that incorporated the two-way interaction was statistically significant. We had no a priori reason to expect significant statistical interactions between food type and predator presence, and none of these interactions were significant. All statistical analyses were conducted in R v.3.4.1 (R Core Team, 2017), and all raw data are available on Dryad (https://doi.org/10.5061/dryad.d6f6q7p).

3 | RESULTS

3.1 | Algal responses to temperature

3.1.1 | Cell size

When assayed after two generations in the common garden environment (15°C), algal populations that had been maintained for the duration of the experiment at the warmer temperature were 23% smaller in volume compared to algae maintained at the cooler temperature (12°C: 1,065 µm³ ± 91 SD; 18°C: 821 µm³ ± 145; *F*₁,₁₁₄ = 16, *p* < 0.001). We observed a similar pattern of cell volume during the experiment, with algae reared at the warmer temperature exhibiting a 25% decrease in cell size (776 µm³ ± 93) compared to algae reared at the cooler temperature (1,041 µm³ ± 104; average of sampling days 25, 75, 105). Algal cell volumes recorded after the common garden period were not statistically different from the average cell volume during the experiment (*F*₁,₆₀ = 1.3, *p* = 0.26).

3.1.2 | Metabolism

Rates of *S. obliquus* photosynthesis showed plastic increases with assay temperature, but there was no difference in photosynthesis between the temperature selection lines (Figure 1a; assay temperature: *χ*²(4) = 100.3, *p* < 0.0001; selection temperature: *χ*²(1) = 1.1,

![Figure 1](https://example.com/figure1.png)

**FIGURE 1** Thermal reaction norms for *Scenedesmus obliquus* (a) photosynthesis and (b) respiration. There was no effect of selection temperature on photosynthesis. Algae reared at 18°C evolved higher rates of respiration than those reared at 12°C. Both photosynthesis and respiration increased with assay temperature. See Section 3.1.2 for statistics. Blue symbols denote algae selected at the cooler temperature, and red symbols represent algae selected at the warmer temperature. Error bars represent ± 1 standard error.*

![Figure 1](https://example.com/figure1.png)

Algal respiration also increased plastically with assay temperature (Figure 1b; *χ*²(4) = 66.7, *p* < 0.0001), and *S. obliquus* populations selected at the warmer temperature evolved higher rates of respiration compared to populations selected at the colder temperature (Figure 1b; selection temperature: *χ*²(1) = 4.1, *p* = 0.04). The effect of selection temperature did not depend on assay temperature (*χ*²(4) = 5.98, *p* = 0.2).

3.2 | *Daphnia* population size

Both food type and predator treatment explained significant variation in *Daphnia* population size (Figure 2; predators: *F*₁,₁₅₂ = 33.6,
3.3 | Daphnia thermal reaction norms and evolution

3.3.1 | Body size

Algal food type altered Daphnia body size evolution in response to selection temperature (Figure 3a and Table 1). Daphnia fed cold-evolved algae evolved steeper thermal reaction norms when selected at 12°C, but there was no difference in TRNs for body size in Daphnia fed warm-evolved algae. There was a phenotypically plastic effect of assay temperature on Daphnia body size (Table 1), with Daphnia assayed at warmer temperatures showing slightly larger body sizes.

The presence of predators modified how Daphnia body size evolved in response to the thermal environment (Figure 3b and Table 1). The effect of temperature selection was more pronounced when Daphnia were also reared with predators, with Daphnia evolving overall larger bodies in the 12°C versus 18°C selection treatments.

3.3.2 | Time to maturity

Variation in Daphnia development time was primarily explained by assay temperature, with Daphnia assayed at warmer temperatures maturing several days earlier than those assayed at cooler temperatures (Figure 4a and Table 1). There was no effect of selection temperature or algal food type on TRN evolution. There was also no effect of predator presence on Daphnia development time TRNs (Figure 4b).

3.3.3 | Per capita growth rate, r

Algal food type altered how TRNs for Daphnia r evolved in response to selection temperature (Figure 5a and Table 1). When fed cold-evolved algae, Daphnia evolved steeper TRNs in the 12°C selection treatment, but when fed warm-evolved algae, Daphnia evolved steeper TRNs in the 18°C selection treatment (Figure 5a)
Assay temperature explained a significant fraction of variation in *Daphnia* (*r*), with *Daphnia* exhibiting faster *r* when grown in warmer assay temperatures. There was no effect of predators on TRN evolution in *Daphnia* (*Figure 5b and Table 1*).

### DISCUSSION

The goal of this study was to investigate the effect of bottom-up and top-down species interactions on thermal evolution of a focal zooplankton population. We documented increases in phytoplankton cell volume in response to the colder rearing environment from early on the experiment and confirmed that these changes had an evolutionary basis at the end of the experiment. We demonstrated that *Daphnia* fed cold-evolved algae maintained larger population sizes compared to populations fed warm-evolved algae. *Daphnia* reared in the presence of predators evolved overall larger bodies. Lastly, the evolution of TRNs for *Daphnia* per capita growth rate was driven by bottom-up species interactions, with *Daphnia* fed cold-evolved algae exhibiting steeper TRNs in the cold *Daphnia* selection environment, and vice versa for *Daphnia* fed warm-evolved algae. We discuss these results in further detail below.

#### 4.1 Algal evolution

Algae exhibit cell size plasticity in response to temperature (Chalifour & Juneau, 2011; Chen et al., 2011; Margalef, 1954; Rhee & Gotham, 1981), and thus, differences in cell size between the two temperature selection experiments were likely a function of both plasticity and evolution. The data collected here do now allow us to quantify exactly when in the experiment algal evolution occurred. However, using population sizes less than those used in this experiment (~100,000 cells/ml), we have since demonstrated that *S. obliquus* can...
Two recent studies on thermal reaction norm evolution in other species of phytoplankton also reported the evolution of elevated respiration in warmer selection environments, but in both of these studies, they also saw the evolution of increased photosynthesis (Chlorella vulgaris: Padfield et al., 2016, Chlamydomonas reinhardtii: Schaum et al., 2017). The duration of selection was 100 generations, and 10 years, for Chlorella, and Chlamydomonas, respectively, suggesting that the 70 generations used here may have been too short to observe photosynthesis evolution. However, the evolved increase in resource consumption (respiration) but not resource acquisition (photosynthesis) at warmer temperatures is consistent with the observed evolved decrease in cell size. Finally, although not directly tested here, cell size is positively correlated with lipid content in other algal species (Bono, Garcia, Sri-Jayantha, Ahner, & Kirby, 2015). This pattern suggests a possible mechanistic link between cold temperature, cell size, increased algal fatty acids and increased nutritional quality of cold-reared algae.

4.2 | Effect of resource thermal evolution on Daphnia population size

Our results demonstrated that algal thermal evolution had significant effects on Daphnia population size. Daphnia maintained on cold-reared algae attained a 12% higher population size during the experiment compared to Daphnia maintained on warm-reared algae. These results are consistent with a previously published study showing that cold-reared Scenedesmus obliquus are more nutritious and sustain higher Daphnia population sizes, compared to warm-reared S. obliquus (Breuer et al., 2013; Sikora et al., 2014; Vigeolas et al., 2012).

4.3 | Effect of resource evolution on Daphnia thermal evolution: “bottom-up effects”

From previous studies demonstrating a positive correlation between environmental quality and genetic variation/heritability (Bochdanovits et al., 2003; Charmantier & Garant, 2005), we predicted that Daphnia reared long term on cold-evolved (higher quality) phytoplankton would show an elevated evolutionary response to temperature-mediated selection (Box, Hypothesis 1). However, we also predicted that because zooplankton and phytoplankton often encounter the same abiotic environment in nature, a “better” environment might be one in which the consumer environment matches that of the resource (Box, Hypothesis 2). Our data were consistent with both hypotheses. (a) Daphnia body size evolution in response to selection temperature was more pronounced in populations that were fed the cold-evolved/higher quality algae, consistent with Hypothesis 1 (Figure 3a); (b) conversely, Daphnia evolution exhibited patterns indicative of local adaptation between the consumer and resource thermal environment, consistent with Hypothesis 2 (Figure 5a). What we did not predict at the outset of this study was that rapid algal evolution could potentially change the slope of the evolved TRN. A steeper TRN slope reflects an overall increase in thermal plasticity. It is unclear why Daphnia in these “matched” (e.g., cold-evolved algae + cold-selected Daphnia) environments would show elevated thermal plasticity, compared to Daphnia in the “mismatched” environments. However, these patterns are consistent with previous studies showing the expression of steeper TRNs when organisms were fed higher quality food (Álvarez & Nicieza, 2002; Jang, Rho, Koh, & Lee, 2015; Kingsolver, Shlichta, Ragland, & Massie, 2006).

4.4 | Potential for maladaptation

One would predict that organisms with rapid generation times (e.g., phytoplankton or microbes) will evolve much more quickly.
to warming, relative to their consumers (e.g., zooplankton), which have longer generation times and smaller population sizes. As temperatures in nature continue to warm, the mismatch in rates of thermal evolution between resources and consumers could result in slower-evolving consumers interacting with resources that have already evolved to a changed environment. This hypothetical situation is not totally dissimilar from the “warm-evolved algae + cold-selected Daphnia” treatment combination used in this study. For both body size and r, Daphnia in this treatment combination evolved muted thermal plasticity compared to Daphnia in the “cold-evolved algae + cold-selected Daphnia” combination. If thermal plasticity confers fitness benefits, then rapid thermal adaptation in resources could lead to maladaptation in consumers.

4.5 | Effect of predators on Daphnia thermal evolution: “top-down” effects

We predicted that the presence of predators would modify Daphnia TRN evolution (Box 1, Hypothesis 3). We did not observe the predicted increase in Daphnia r in response to predators, but Daphnia did evolve larger bodies, especially those in the 12°C selection treatment. These results are consistent with a previous study showing the evolution of larger bodies and earlier reproduction in the same predator-prey system (Spitze, 1991). However, another study that used the same study system, but with food limitation, did not observe the evolution of larger Daphnia body size in the presence of Chaoborus predators (Tseng & O’Connor, 2015).

5 | CONCLUSIONS

Using laboratory experiments on a tri-trophic aquatic community, we have demonstrated that species interactions can play an important role in determining how a focal population response to temperature-mediated selection. The challenge moving forward is to apply these results to natural settings and to assess how frequently species interactions hinder or help focal populations adapt to warming climates. There are multiple cases in nature where interacting species may respond to the same selection pressure but at very different speeds (e.g., insects and their coevolved viruses (Cory & Myers, 2003), small mammals and their prey (hare/lynx: Krebs, Boonstra, Boutin, & Sinclair, 2001), trees and their seed predators (Mezquida & Benkman, 2014), fish and zooplankton (Vanni, 1986)), and thus, we expect the “bottom-up” results of this study to be widely applicable across natural food webs.

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CONFLICT OF INTEREST

None declared.

DATA ACCESSIBILITY

All raw data are available in Dryad (https://doi.org/10.5061/dryad.d6f6q7p) (Tseng, Bernhardt, & Chila, 2019).

ORCID

Michelle Tseng https://orcid.org/0000-0002-4306-507X

Joey R. Bernhardt https://orcid.org/0000-0003-1824-2801

REFERENCES

Abrams, P. A., & Rowe, L. (1996). The effects of predation on the age and size of maturity of prey. Evolution, 50, 1052–1061.

Álvarez, D., & Nicieza, A. G. (2002). Effects of temperature and food quality on anuran larval growth and metamorphosis. Functional Ecology, 16, 640–648.

Anderson, J. T., Inouye, D. W., McKinney, A. M., Colautti, R. I., & Mitchell-Olds, T. (2012). Phenotypic plasticity and adaptive evolution contribute to advancing flowering phenology in response to climate change. Proceedings of the Royal Society B: Biological Sciences, 279(1743), 3843–3852.

Angillette, M. J., Steury, T. D., & Sears, M. W. (2004). Temperature, growth rate, and body size in ectotherms: Fitting pieces of a life-history puzzle. Integrative and Comparative Biology, 44, 498–509.

Barrett, R. D. H., Paccard, A., Healy, T. M., Bergek, S., Schulte, P. M., Schulte, D., & Rogers, S. M. (2010). Rapid evolution of cold tolerance in stickleback. Proceedings of the Royal Society B: Biological Sciences, 278, 233–238.

Bennett, A. F., Dao, K. M., & Lenski, R. E. (1990). Rapid evolution in response to high-temperature selection. Nature, 346, 79–81.

Bernhardt, J. R., Sunday, J. M., & O’Connor, M. I. (2018). Metabolic theory and the temperature-size rule explain the temperature dependence of population carrying capacity. American Naturalist, 192, 687–697.

Bochdanovitz, Z., de Jong, G., & De Jong, G. (2003). Experimental evolution in Drosophila melanogaster: Interaction of temperature and food quality selection regimes. Evolution, 57, 1829–1836.

Bold, H. C. (1949). The morphology of Chlamydomonas chlamydogama, sp. nov. Bulletin of the Torrey Botanical Club, 76, 101–108.

Bono, M. S., Garcia, R. D., Sri-Jayantha, D. V., Ahner, B. A., & Kirby, B. J. (2015). Measurement of lipid accumulation in Chlorella vulgaris via flow cytometry and liquid-state 1H NMR spectroscopy for development of an NMR-traceable flow cytometry protocol. PLoS ONE, 10, e0134846.

Breuer, G., Lamers, P. P., Martens, D. E., Draaisma, R. B., & Wijffels, R. H. (2013). Effect of light intensity, pH, and temperature on tri-acylglycerol (TAG) accumulation induced by nitrogen starvation in Scenedesmus obliquus. Bioresource Technology, 143, 1–9.

Campbell-Staton, S. C., Edwards, S. V., & Losos, J. B. (2016). Climate-mediated adaptation after mainland colonization of an ancestrally subtropical island lizard, Anolis carolinensis. Journal of Evolutionary Biology, 29, 2168–2180.

Chalifour, A., & Juneau, P. (2011). Temperature-dependent sensitivity of growth and photosynthesis of Scenedesmus obliquus, Navicula pelliculosa and two strains of Microcystis aeruginosa to the herbicide atrazine. Aquatic Toxicology, 103, 9–17.

Charmantier, A., & Garant, D. (2005). Environmental quality and evolutionary potential: Lessons from wild populations. Proceedings of the Royal Society B: Biological Sciences, 272, 1415–1425.
Chen, M., Li, J., Dai, X., Sun, Y., & Chen, F. (2011). Effect of phosphorus and temperature on chlorophyll a contents and cell sizes of Scenedesmus obliquus and Microcystis aeruginosa. Limnology, 12, 187–192.

Cory, J., & Myers, J. (2003). The ecology and evolution of insect bacte‐
vorises. Annual Review of Ecology, Evolution, and Systematics, 34, 239–272.

Dutilleul, M., Bonzom, J. M., Lecomte, C., Goussen, B., Daian, F., Galas, S., & Réale, D. (2014). Rapid evolutionary responses of life history traits to different experimentally‐induced pollutions in Caenorhabditis elegans. BMC Evolutionary Biology, 14, 1–14.

Finkel, Z. V., Beardall, J., Flynn, K. J., Quigg, A., Rees, T. A. V., & Raven, J. A. (2010). Phytoplankton in a changing world: Cell size and elemental stoichiometry. Journal of Plankton Research, 32, 119–137.

Frankham, R. (1996). Relationship of genetic variation to population size in wildlife. Conservation Biology, 10, 1500–1508.

Hendry, A. (2017). Eco‐evolutionary dynamics. Princeton, NJ: Princeton University Press.

Horton, P. A., Rowan, M., Webster, K. E., & Peters, R. H. (1979). Browsing Eco‐evolutionary dynamics.

Kilham, S. S., Kreeger, D. A., Lynn, S. G., Goulden, C. E., & Herrera, L. (2001). What Thermal reaction norms for caterpillars depend on diet. Evolutionary Ecology Research, 10, 251–268.

Kingsolver, J. G., & Huey, R. B. (2008). Size, temperature, and fitness: Three rules. Evolutionary Ecology Research, 10, 251–268.

Kingsolver, J., Shlitcha, J. G., Ragland, G. J., & Massie, K. R. (2006). Thermal reaction norms for caterpillars depend on diet. Evolutionary Ecology Research, 8, 703–715.

Koch, U., Martin-Creuzburg, D., Grossart, H.-P., & Straile, D. (2012). Differences in the amino acid content of four green algae and their impact on the reproductive mode of Daphnia pulex With 4 figures. Fundamental and Applied Limnology, 1814, 327–336.

Krebts, C. J., Boonstra, R., Boutin, S., & Sinclair, A. R. E. (2001). What drives the 10‐year cycle of snowshoe hares? BioScience, 51, 25.

Loboda, S., Savage, J., Biddle, C. M., Schmidt, N. M., & Haye, T. T. (2018). Declining diversity and abundance of High Arctic fly assemblages over two decades of rapid climate warming. Ecography, 41, 265–277.

Lürling, M., & Van Donk, E. (1996). Zooplankton‐induced unicell‐colony transformation in Scenedesmus acutus and its effect on growth of herbivore Daphnia. Oecologia, 108, 432–437.

Margalef, R. (1954). Modifications induced by different temperatures on the cells of Scenedesmus obliquus (Chlorophyceae). Hydrobiologia, 6, 83–91.

Mazé-Guilmo, E., Blanchet, S., Rey, O., Canto, N., & Lott, G. (2016). Local adaptation drives thermal tolerance among parasite populations: A common garden experiment. Proceedings of the Royal Society B: Biological Sciences, 283, 20160587.

Mezquida, E. T., & Benkman, C. W. (2014). Causes of variation in biotic interaction strength and phenotypic selection along an altitudinal gradient. Evolution, 68, 1710–1721.

Montagnes, D. J. S., Berges, J. A., Harrison, P. J., & Taylor, F. J. R. (1994). Estimating carbon, nitrogen, protein, and chlorophyll‐a from volume in marine‐phytoplankton. Limnology and Oceanography, 39, 1044–1060.

Mousing, E. A., Ellegaard, M., & Richardson, K. (2014). Global patterns in phytoplankton community size structure‐evidence for a direct temperature effect. Marine Ecology Progress Series, 497, 25–38.

Newman, J. A. (2004). Climate change and cereal aphids: The relative effects of increasing CO2 and temperature on aphid population dynamics. Global Change Biology, 10, 5–15.

O’Connor, M. I., Bruno, J. F., Gaines, S. D., Halpern, B. S., Lester, S. E., Kinlan, B. P., & Weiss, J. M. (2007). Temperature control of larval dispersal and the implications for marine ecology, evolution, and conservation. Proceedings of the National Academy of Sciences, 104, 1266–1271.

Osmond, M. M., Otto, S. P., Klausmeier, C. A., Station, K. B., Otto, S. P., & Klausmeier, C. A. (2017). When predators help prey adapt and persist in a changing environment. American Naturalist, 190, 83–98.

Padfield, D., Yvon‐Durocher, G., Buckling, A., Jennings, S., & Yvon‐Durocher, G. (2016). Rapid evolution of metabolic traits explains thermal adaptation in phytoplankton. Ecology Letters, 19, 133–142.

Patrick, R. (1971). The effects of increasing light and temperature on the structure of diatom communities. Limnology and Oceanography, 16, 405–421.

R CoreTeam (2017). R: A language and environment for statistical comput‐
ing. Vienna, Austria: R foundation for Statistical Computing.

Régnière, J., & Nealis, V. (2002). Modelling seasonality of gypsy moth, Lymantria dispar (Lepidoptera: Lymantriidae), to evaluate probability of its persistence in novel environments. Canadian Entomologist, 134, 805–824.

Rhee, G.-Y., & Gotham, I. J. (1981). The effect of environmental factors on phytoplankton growth: Temperature and the interactions of temperature with nutrient limitation 1. Limnology and Oceanography, 26, 635–648.

Schaum, C.-E., Barton, S., Breston, E., Buckling, A., Garcia-Carreras, B., Lopez, P., ... Yvon-Durocher, G. (2017). Adaptation of phytoplankton to a decade of experimental warming linked to increased photosynthesis. Nature Ecology & Evolution, 1, 94.

Sikora, A. B., Davidowicz, P., & Von Elert, E. (2014). Daphnia fed algal food grown at elevated temperature have reduced fitness. Journal of Limnology, 73, 421–427.

Spitze, K. (1991). Chaoborus predation and life-history evolution in Daphnia pulex: Temporal pattern of population diversity, fitness, and mean life history. Evolution, 45, 82–92.

Trubetskova, I., & Lampert, W. (2002). The juvenile growth rate of Daphnia: A short-term alternative to measuring the per capita rate of increase in ectotoxicology? Archives of Environmental Contamination and Toxicology, 42, 192–198.

Tseng, M., Bernhardt, J., & Chila, A. (2019). Data from: Species interactions mediate thermal evolution. Dryad: doi:https://doi.org/10.5061/dryad.d6f6q7p

Tseng, M., Kaur, K. M., Soleimani Pari, S., Sarai, K., Chan, D., Yao, C. H., ... Fograser, K. (2018). Decreases in beetle body size linked to climate change and warming temperatures. Journal of Animal Ecology, 87, 647–659.

Tseng, M., & O’Connor, M. I. (2015). Predators modify the evolution‐
ary response of prey to temperature change. Biology Letters, 11, 20150798.

Van Doorslaer, W., Stoks, R., Jeppesen, E., & De Meester, L. (2007). Adaptive microevolutionary responses to simulated global warming in Simocephalus vetulus: A mesocosm study. Global Change Biology, 13, 878–886.

Vanni, M. J. (1986). Fish predation and zooplankton demography: Indirect effects. Ecology, 67, 337–354.
Vigeolas, H., Duby, F., Kaymak, E., Niessen, G., Motte, P., Franck, F., & Remacle, C. (2012). Isolation and partial characterization of mutants with elevated lipid content in *Chlorella sorokiniana* and *Scenedesmus obliquus*. *Journal of Biotechnology*, 162, 3–12.

Weider, L. J., Jeyasingh, P. D., & Looper, K. G. (2008). Stoichiometric differences in food quality: Impacts on genetic diversity and the coexistence of aquatic herbivores in a *Daphnia* hybrid complex. *Oecologia*, 158, 47–55.

West, D. C., & Post, D. M. (2016). Impacts of warming revealed by linking resource growth rates with consumer functional responses. *Journal of Animal Ecology*, 85, 671–680.

Winter, B. (2013). Linear models and linear mixed effects models in R with linguistic applications. *arXiv*, 1308, 5499.

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