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# **Elevated Temperature May Afect Nectar Microbes, Nectar Sugars, and Bumble Bee Foraging Preference**

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#### **Abstract**

Floral nectar, an important resource for pollinators, is inhabited by microbes such as yeasts and bacteria, which have been shown to infuence pollinator preference. Dynamic and complex plant-pollinator-microbe interactions are likely to be afected by a rapidly changing climate, as each player has their own optimal growth temperatures and phenological responses to environmental triggers, such as temperature. To understand how warming due to climate change is infuencing nectar microbial communities, we incubated a natural nectar microbial community at diferent temperatures and assessed the subsequent nectar chemistry and preference of the common eastern bumble bee, *Bombus impatiens*. The microbial community in foral nectar is often species-poor, and the cultured *Brassica rapa* nectar community was dominated by the bacterium *Fructobacillus*. Temperature increased the abundance of bacteria in the warmer treatment. Bumble bees preferred nectar inoculated with microbes, but only at the lower, ambient temperature. Warming therefore induced an increase in bacterial abundance which altered nectar sugars and led to signifcant diferences in pollinator preference.

**Keywords** Floral microbes · Pollination · Symbiosis · Fructophilic bacteria

### **Introduction**

Rapid increases in global temperature have the potential to disrupt many ecological processes. Warming due to climate change has caused an increase of 0.85 °C in global surface temperatures over the past century [\[1](#page-8-0)]. Models based on low  $CO<sub>2</sub>$  emissions estimate that there will be another 1.5 °C rise by the turn of the century [\[2](#page-8-1)]. Sixteen of the 17 hottest years in the past 138 years have occurred in the 2000s with 2016 and 2020 being the hottest years on record [\[1\]](#page-8-0). Extreme climatic events, including drought and heat waves, are also predicted to become more common [\[3\]](#page-8-2).

Most living organisms, including microorganisms, have an optimal living temperature at which they thrive [\[4](#page-8-3), [5](#page-8-4)]. As global temperatures increase, many species interactions will be disrupted [[6\]](#page-8-5). Although there is ample research on plant phenological and physiological change in the context of climate change, there currently is a lack of information on how climate change will infuence the interaction between fowers, pollinators, and their associated microorganisms. The center of this interaction web is foral nectar. Pollinators rely on nectar as a valuable source of energy [\[7](#page-8-6)]. However, nectar is more than a sugar-rich resource, and surveys of many wild plant species in varied ecological regions have revealed nectar is often inhabited by bacteria [[8,](#page-8-7) [9](#page-8-8)] and fungi (mainly yeast) [\[10\]](#page-8-9).

Nectar-inhabiting microorganisms have been found to alter nectar chemistry and infuence pollinator behavior [[11\]](#page-8-10). Specialized nectar-inhabiting microbes can tolerate the harsh nectary environment that flters out non-specialist microbes [[12\]](#page-8-11). These specialized microbes utilize resources within foral nectaries and change nectar chemistry in many ways. For example, yeasts and bacteria change nectar sugar concentrations as they metabolize nutrients [\[13](#page-9-0)] as well as release volatile organic compounds which afect pollinator preference for nectar [[14](#page-9-1), [15](#page-9-2)]. Although fungi and bacteria can reduce overall nectar sugars when compared to sterile nectar, bees tend to prefer nectar colonized with specifc microbes [[13](#page-9-0), [16\]](#page-9-3). However, microbe-mediated pollinator preference is dependent on pollinator species, the microbes present, and even gustatory and olfactory cues, which in some cases lead to pollinators showing indiference or even avoiding nectar colonized with less attractive microbes [[14,](#page-9-1)

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[15](#page-9-2), [17\]](#page-9-4). These microscopic changes in nectar lead to drastic changes in pollinator behavior, however, little is known about how climate change will afect this interaction.

With the looming prospect of rapid temperature increases, understanding the efects of climate change on plant-pollinator-microbial interactions is of great importance. We hypothesize that environmental temperature mediates microbial community structure in nectar because diferent microbes are likely to have diferent optimal growth temperatures. We predict that nectar-inhabiting microbial communities will difer between temperature treatments, leading to altered nectar chemistry and ultimately diferences in pollinator preference. Here we test the efects of three diferent temperature treatments, representing baseline temperatures and climate change predicted temperatures, on the same starting microbial community using synthetic nectar. We determine how temperature-mediated changes in microbial communities alter nectar chemistry and pollinator preference. Our results help tease apart the mechanisms within plant-pollinator-microbe interactions that will be afected by climate change.

### **Methods**

#### **Brassica rapa and Nectar Extractions**

*Brassica rapa* subsp. *oleifera*, Brassicaceae, is a wild mustard introduced to North America from Europe which is pollinated by many invertebrate species. We selected *B. rapa* as it is abundant in Southern California and is visited by many diferent pollinator species. We collected *B. rapa* fowers in the early morning (between 8:00 and 10:00am) from a meadow in Beaumont, California (33.933670,−117.002738) in April 2017 and brought the flowers back to the lab for nectar extractions. Although the fowers were not bagged, we collected them early in the morning before many pollinators were out foraging. While we collected nectar early in the morning to control for pollinator visitation and microbial degradation, the nectar could have already been inoculated with sugar-altering microorganisms. The collection site was a mature feld with all plants in the fowering stage. *Brassica rapa* inflorescences have 2–5 flowers open at a time, and we gently removed all unopened foral buds prior to nectar extraction. To avoid pollen and pollen microbes contaminating the nectar, we used sterile micro-dissecting scissors to carefully remove anthers and pollen from each fower. To extract nectar, we placed 2–5 fowers facing down in a sterile, modifed 1.5 mL Eppendorf tube and centrifuged the flowers at  $7500 \times g$  for 1 min to remove nectar. We repeated this process with about 20 *B. rapa* inforescences from fve diferent plants until we had 25µL of nectar. To prevent any debris or small invertebrates from falling into the nectar during centrifugation, we modifed the 1.5 mL Eppendorf tubes by gluing fne mesh halfway up the tube. We used centrifugation as the method to collect nectar as the *Brassica* fowers are too small to use capillary tubes. Although this method may introduce microbes from the petals or bracts it is unlikely that these microbes would thrive in the artifcial nectar as they are phyllosphere bacteria [\[18](#page-9-5)]. We sterilized all 1.5 mL collection tubes with mesh modifcations by UV (254 nm) sterilization in an AirScience UV-Box (Fort Meyers, FL) for 20 min prior to use. We pooled the nectar from the 20 *B. rapa* inforescences from fve diferent plants for chemical composition analysis and microbial community characterization.

#### **Nectar Analysis**

To quantify nectar sugars, we used the Megazyme Sucrose, D-Fructose, D-Glucose Assay Kit and followed the manufacturer's protocol. To identify the amino acid composition of the nectar samples, we sent *B. rapa* nectar to Texas A&M University Proteomics department where there is an established free amino acid assay for plant nectar using high-performance liquid chromatography (HPLC, Online Resource 1). Based on these two analyses of *B. rapa* nectar, we designed sterile, synthetic nectar by autoclaving a solution of 7.4% w/v molecular grade sucrose, 5.8% w/v molecular grade glucose, and 1.1% w/v molecular grade fructose for a roughly 7:5:1 ratio—which is similar to what was found in *B. rapa* nectar in Wykes (1952) [[19\]](#page-9-6). We then added 8 mL of Minimum Essential Medium (MEM) sterile Non-Essential Amino Acid Solution (100×) (Sigma-Aldrich M7145), which contains seven of the highest registered amino acids from *B. rapa* nectar. We added amino acids to replicate the nectar as closely as possible to promote feld-realistic microbial growth.

#### **Choice Experiment**

We inoculated synthetic nectar with nectar-inhabiting microbial communities from wild *B. rapa* flowers by adding 50 $\mu$ L of pooled, freshly extracted *B. rapa* nectar from roughly 45 inforescences (as described above) to 10 mL of synthetic nectar and allowed 24 h for growth at 25 °C. Then we gently vortexed this single pool of inoculated artifcial nectar and added 10µL to 110 individual 2 mL tubes of synthetic nectar. We then separated these tubes into two treatments of 55 tubes each and incubated each treatment at one of two temperature treatments. The frst temperature treatment (27 °C) represents the average spring-time high in Riverside, CA [[20](#page-9-7)], where the nectar microbes were collected, and the second treatment (32 °C) represents a climate change predicted temperature  $[21]$  $[21]$  $[21]$ . High CO<sub>2</sub> emission scenarios predict a 5 °C increase in global temperatures by the turn of the century [\[21\]](#page-9-8). We incubated treatments for 3 days, which is the longest that nectar will sit in the nectary of *Brassica* before being depleted or the flower senesces [\[22](#page-9-9)]. For controls, we also incubated 55 tubes of sterile synthetic nectar at each of the temperature treatments. After the 3-day incubation, we divided each volume of the mature microbial communities into two aliquots: one for characterization of the microbial community and nectar sugars, and one for the bumble bee choice assay.

We presented 1–5-day-old female worker *Bombus impatiens* with a choice assay to assess feeding preference. We purchased fve *B. impatiens* colonies from Koppert Biological Systems (Howell, MI) and maintained these colonies with pollen and 60% sterile sucrose water ad libitum in environmentally controlled rooms at the University of California Riverside, which were held at 27 °C. We assayed 10 bumble bees from each colony for a total of 50 bumble bees. To perform the choice assay, we put one bumble bee in a small foraging chamber (9.5 cm $\times$ 10.15 cm) that had access to 4 feeders each flled with 1 mL of one of four treatments: (1) synthetic nectar inoculated with microbes and incubated at 27 °C, (2) sterile synthetic nectar incubated at 27 °C as a control, (3) synthetic nectar inoculated with microbes and incubated at 32 °C, and (4) sterile synthetic nectar incubated at 32 °C as a control. We point out that the temperature treatments occurred before the choice assay; all 4 nectar treatments were offered to bees at the same temperature as the bees. The bees themselves were not exposed to any temperature treatments but instead held at a constant 27 °C.

To assess bumble bee preference, we carefully weighed each feeder before and after a 24-h foraging period and counted choice as amount in grams of nectar consumed by each bee. We set up fve control pots that had all four nectar treatments but no bumble bees. To account for evaporation, we weighed the feeders before and after the foraging period and subtracted the amount evaporated from the amount consumed by bees for each block. Each bee participated in only one choice test and was not returned to the colony after the assay was completed. To be sure that the experimental bees were not acclimated to any of the offered treatments, we did not train the bees to the assay arenas or to the artifcial nectar, thus the experimental bees were naive to all four ofered treatments in the choice assay. As experimental bees were reared in their respective colonies, it is possible that they were acclimated to hive microbial contaminants in the sugar water while in the colony. However, the lab colonies we used for these experiments were never exposed to environmental microbes and we regularly changed their sterilized sucrose solution. The shared sucrose source in the colonies would therefore only be exposed to the specialized bumble bee gut microbiota, which is closely related to the honey bee microbiota and is not known to grow in sucrose-only media [\[23](#page-9-10)]. However, previous work described the commonly found microbes in commercial bumble bee microcolonies [\[24](#page-9-11)] and none of these microbes were detected in our nectar treatment. Exposure of our experimental bees to nectar-inhabiting microbes before the choice trials is therefore highly unlikely if not impossible.

#### **Extreme Heat Experiment**

We conducted a second choice assay using the same methods as described above but with more extreme temperatures. Using the *B. rapa* synthetic nectar recipe, we inoculated the same wild *Brassica* fower microbial community and incubated the nectar for 3 days. In this experiment, we used 32  $\degree$ C and 42  $\degree$ C to incubate the nectar before offering it to 50 bumble bees from 5 colonies (these colonies difered from those used in experiment above) in a choice assay as described above. We selected these temperature treatments as they represent the average summer daytime high in the geographical region and a typical heatwave temperature increase.

#### **Post‑Assay Nectar and Microbiota Analysis**

To characterize microbial communities, we centrifuged the aliquoted sample at  $4500 \times g$  for 5 min to pellet out microbes. Once the pellet was formed, we pipetted nectar off for carbohydrate analysis, leaving the pellet for DNA extraction. To measure carbohydrate concentrations on a subset of samples, 10 samples from each treatment, we again used the Sucrose, D-Fructose, D-Glucose Assay Kit (Megazyme, Chicago, IL) to characterize diferences in nectar sugars according to temperature treatments.

We extracted DNA from the remaining pellet of all 100 samples using the DNeasy Blood and Tissue Kit (Qiagen, Valencia, CA). To control for possible reagent contaminants, we included  $N=1$  "blank" samples that contained no cells beyond those that may have occurred in the reagents or via possible contamination. We ran this blank sample through all of our library preparation and analysis pipeline. To prepare the samples for extraction, we used a Qiagen tissue lyser to bead-beat samples for 6 min at 30 Hz with two sterile 3.2 mm chrome-steel beads and roughly 100µL of 0.1 mm glass beads (Biospec, Bartlesville, OK), in 180µL of bufer ATL from the Qiagen extraction kit. We then added 20μL of Proteinase K, incubated the samples overnight at 57 °C, and followed the DNeasy standard extraction protocol.

To characterize the microbial communities within nectar, we used dual-index inline barcoding to prepare samples for sequencing on the MiSeq sequencer (Illumina), following the same protocols as detailed in McFrederick and Rehan (2016) [[25](#page-9-12)]. We used primers that included either the forward or reverse Illumina sequencing primer, a unique 8-nt-long barcode, and the forward or reverse genomic oligonucleotide  $[26]$  $[26]$ . We used the bacterial 16S rRNA sequence primers 799F-mod3 CMGGATTAGATACCCKGG [[27\]](#page-9-14) and 1115R AGGGTTGCGCTCGTTG [[26](#page-9-13)] and the fungal internal transcribed spacer (ITS) primers ITS1F (50- CTTGGTCATTTAGAGGAAGTAA-30) and ITS4R (50-TCCTCCGCTTATTGATATGC-30). We performed PCRs using  $10\mu L$  of  $2 \times$  Pfusion High-Fidelity DNA polymerase (New England Biolabs, Ipswich, MA), 10μL of ultrapure water, 0.5μL of each 10 μM primer stock, and 4μL of DNA, with an annealing temperature of 57 °C for 30 cycles. We cleaned this product using Ultraclean PCR cleanup kit (MoBio, Carlsbad, CA), to remove unincorporated primers and dNTPs. To complete the Illumina sequencing construct, we used 1μL of the clean PCR product as a template for a second PCR, using HPLC-purified primers: CAAGCAGAAGACGGCATA CGAGATCGGTCTCGGCATTCCTGC and AATGAT ACGGCGACCACCGAGATCTACACTCTTTCCCTA CACGACG  $[26]$  $[26]$ . We then normalized  $18\mu$ L of PCR product using SequalPrep Normalization plates (Thermo Fisher Scientific, Waltham, MA). We pooled 5μL of each sample and performed another Ultraclean PCR cleanup on this combined sample. We assessed library quality using a 2100 Bioanalyzer (Agilent, Santa Clara, CA). After quality control, we sequenced the libraries using a MiSeq sequencer (Illumina) and MiSeq Reagent kit, version 3 (Illumina), with  $2 \times 300$  cycles, at the IIGB Genomics Core, UC Riverside. We were unable to amplify fungi from our artificial nectar samples, and therefore do not consider fungi further.

#### **Quantifcation of the Microbial Community (qPCR)**

To determine the absolute abundance of bacterial cells in the artificial nectar samples of both temperature treatments, we used quantitative real-time PCR (qPCR) of the inoculated DNA extractions as above,  $N = 100$ ; however, 30 samples failed to amplify  $(N = 70)$ . Each PCR reaction consisted of 7.5μL SsoAdvanced master mix (Bio-Rad Laboratories, Los Angeles, CA), 3.6 mL molecular grade water, 0.45μL forward primer, 0.45μL reverse primer and 1.0 μl sample or standard DNA. We used the universal bacterial primers for the conserved 16S rRNA region Univ331F as our forward primer (5′- TCCTACGGGAGGCAGCAGT-3′) and Univ797R as our reverse primer (5′-GGACTACCAGGGTATCTA ATCCTGTT-3′) [[28\]](#page-9-15). We used this 16S qPCR primer set instead of 799F-1115R due to its established use in previous microbiome studies [[28–](#page-9-15)[31](#page-9-16)]. The reaction conditions were an initial heating at 95 °C for 3 min, followed by 39 cycles of 95 °C for 10 s, 59 °C for 30 s on a BioRad C1000 Touch thermal cycler. We compared

our samples to a standard curve of  $1 \times 10^2 - 1 \times 10^8$  copies of the 16S rRNA gene cloned into a TOPO-TA plasmid (Invitrogen, Carlsbad, CA), with all qPCR efficiencies between 90 and  $100\%$  and  $R^2$  above 0.99.

#### **Statistical Analysis and Microbiome Bioinformatics**

We used a two-way analysis of variance (ANOVA) and TukeyHSD post hoc pairwise comparison to assess whether temperature treatment affects individual nectar sugar concentrations and a Kruskal–Wallis chi-square test to test for diferences in total sugars. We used a linear mixed model with Gaussian error distribution (GLMMs; package lme4) [[32\]](#page-9-17) to assess differences in nectar consumption by bumble bees during the choice assay. We used nectar consumption as the response variable, temperature treatment and presence or absence of a microbial community as fxed efects, and colony of origin as random intercept. We used package lmerTest to compare coefficients of fixed effects  $[33]$  $[33]$  $[33]$ . To test if 16S rRNA gene copy numbers (qPCR results) were diferent between temperature treatments, we used Welch's *t*-test. We performed all the above statistical analyses in R 3.4.4 [[34\]](#page-9-19). We used QIIME2-2018.6 [\[35](#page-9-20)] to process the 16S rRNA gene sequence libraries. We trimmed the low-quality ends off the reads with QIIME2's default settings. Next, we binned our sequences into amplicon sequence variants (ASVs) using DADA2 [[36](#page-9-21)], followed by chimera removal using the default settings of the DADA2 Pipeline. To assign taxonomy to the ASVs we used the QIIME2 q2-featureclassifer [[37](#page-9-22)] trained to the 799–1115 region of the 16S rRNA gene and conducted local BLASTn searches against the NCBI 16S microbial database (July 2017). We cleaned the data by fltering out ASVs from the resulting feature table that corresponded to contaminants of reagents as identifed in our blanks (Online Resource 2). As the artifcial nectar in which the foral microbiomes were incubated contained no plant material, we found no plant plastid contamination in our sequencing reads. To generate a phylogenetic tree of our sequences, we used the MAFFT aligner [[38\]](#page-9-23) and FastTree v2.1.3 [[39\]](#page-9-24). We used rarefaction analysis to determine a standardized coverage of bacterial species diversity to be used in alpha and beta diversity analyses of the DNA extracted from synthetic nectar after incubation period. At 2600 reads per sample we found that the rarefaction curves levelled off, and 63 samples could be included in the subsequent analyses. We used this tree and ASV table for alpha diversity analysis and to calculate unweighted UniFrac distance matrices in QIIME2 using qiime diversity core metrics. We used the Shannon Diversity Index and the Kruskal–Wallis test in QIIME2 to analyze alpha diversity. We analyzed diferences in beta diversity with temperature treatment as a fxed factor using Adonis (type II sum of squares) with the vegan package in R [[40](#page-9-25)].

#### **Results**

#### **Sugar Changes with Temperature and Microbes**

After the 3-day incubation period, artifcial nectar sugars were signifcantly reduced by 9.5% when microbes were present (Kruskal–Wallis chi-square = 13.391; *N* = 50, *p*<0.0001). As microbes consume nectar resources, overall nectar sugars decreased in the inoculated treatments compared to sterile nectar. Temperature treatment and presence of microbes infuenced individual sugar concentrations. There was no signifcant diference in sucrose levels between any of the four treatments (Kruskal–Wallis chisquare = 2.7792, df = 3,  $p = 0.4269$ ; Fig. [1\)](#page-5-0). There was an interaction effect of temperature and treatment on fructose levels  $(F_{1,36} = 38.8, p < 0.0001;$  $(F_{1,36} = 38.8, p < 0.0001;$  $(F_{1,36} = 38.8, p < 0.0001;$  Fig. 1). There was roughly 50% more fructose in the in the 27 °C compared to 32 °C (Tukey HSD,  $p < 0.0001$ ), and 75% more in the sterile nectar compared to nectar inoculated with microbes (Tukey HSD,



<span id="page-5-0"></span>**Fig. 1** Boxplot indicating amounts of the three sugars after a 3-day incubation. There was more fructose in the in the 27 °C compared to 32 °C and signifcantly more in the sterile nectar compared to nectar with microbes. There were diferences in glucose amounts between 32 °C sterile treatment, 27 °C microbe-inoculated treatment, and 32 °C microbe-inoculated treatment. There was no signifcant diference in amount of sucrose between treatments. Statistical diference is indicated by letters, NS, no signifcance

 $p < 0.0001$ ). There was also an interaction effect between temperature and treatment on glucose levels  $(F_{1,36}=10.505,$  $p=0.002$ ; Fig. [1](#page-5-0)) with significant differences in glucose amounts between 32 °C sterile treatment (Tukey HSD,  $p = 0.003$ ), 27 °C microbe-inoculated treatment (Tukey HSD,  $p = 0.001$ ), and 32 °C microbe-inoculated treatment (Tukey HSD, *p*<0.0001).

#### **Bumble Bee Preference**

*Bombus impatiens* consumed significantly more nectar inoculated with microbes (GLMM;  $t = 6.854$ ,  $df = 207$ ,  $p < 0.0001$ , Fig. [2](#page-5-1)) and incubated at 27 °C (GLMM; *t* = −4.190, *df* = [2](#page-5-1)07, *p* < 0.0001, Fig. 2), than all other nectar choices. Bumble bees preferred nectar inoculated with microbes and incubated at 27 °C 1.5 $\times$  more than synthetic nectar without microbes or nectar with microbes but incubated at 32 °C. We found no effect of evaporation on overall choice assay results. Similarly, in our "extreme heat" study, bumble bees preferred nectar incubated at lowest of the two temperature treatments (Online Resource 3). In both studies, bumble bees preferred nectar with a microbial community yet reared at the lower of the two temperatures.

#### **Microbial Community**

There was a total of 397,885 quality-fltered reads with an average of 5604 reads per sample (*N*=63) that clustered into 205 fltered ASVs for bacterial sequencing. We found that there was no signifcant diference in alpha diversity, using the Shannon Diversity Index, between temperature treatments (Shannon's  $H = 1.3532$ ,  $p = 0.244$ ). Non-metric Multidimensional Scaling (NMDS) analysis on the unweighted UniFrac distance matrix (Fig. [3](#page-6-0)) showed that there was no obvious



<span id="page-5-1"></span>**Fig. 2** Mass of nectar consumed by bumble bees during the 24-h choice assay. Bumble bees were given the choice between four treatments. For the two incubation temperatures, there were a sterile control and a treatment (inoculated with a microbial community). Asterisk (\*\*\*) indicates statistical signifcance



<span id="page-6-0"></span>**Fig. 3** Nonmetric multidimensional scaling plot of the unweighted UniFrac distance matrices of synthetic nectar inoculated with a microbial community and incubated at two temperature treatments. Blue squares indicate 27 °C (ambient) treatments and red circles denote 32 °C (heated) treatments. Colored ellipses designate 95% confdence intervals around the centroid median of the points

clustering by treatment. We analyzed the unweighted UniFrac distance matrix of our samples with the Adonis function in the R package vegan [[40](#page-9-25)] (999 permutations PerMANOVA) using temperature as an explanatory variable and found no signifcant difference between temperature treatments  $(F=1.0562)$ ,  $R^2$ =0.02344, *p*=0.32).

Across all samples, a *Fructobacillus* (Leuconostocaceae) ASV was the most abundant bacterium and dominated the communities regardless of temperature treatment (Online Resource 4). As sequencing data revealed that *Fructobacillus* spp*.* dominated microbial communities in all samples, we used 16S rRNA gene qPCR to determine if the absolute abundance of bacteria difered by temperature treatment. We found signifcantly higher total abundance of bacteria in the 32 °C treatment compared to the 27 °C treatment (*t*= −3.804, df=43.97, *p*<0.0001; Fig. [4\)](#page-6-1)*.*

### **Discussion**

Warming afected overall bacterial density within nectar which in turn afected nectar sugar composition and ultimately pollinator preference. Nectar sugars decreased when



<span id="page-6-1"></span>**Fig. 4** Quantitative PCR (qPCR) results showing absolute abundance of bacterial cells in each nectar sample of the diferent temperature treatments; there was an increase in bacterial abundance in the heated treatment. Ambient=27 °C and heated=32 °C. Asterisk (\*\*\*) indicates statistical signifcance

microbes were present, but overall sugar levels by themselves did not explain bumble bee preference. Fructose levels were lowest with bacterial inoculation at the warmest temperature, which agrees with our microbial community data. Our synthetic nectar bacterial communities were dominated by *Fructobacillus,* which were more abundant in the warmer (32 °C) incubation temperature. As its name implies, *Fructobacillus* spp. are fructophilic lactic acid bacteria that use fructose as their main carbohydrate source [[41\]](#page-9-26), indicating that increasing *Fructobacillus* abundance drives the decrease in fructose at the warmest temperature. *Fructobacillus* spp*.* are commonly isolated from fructose-rich environments, such as fowers and fruits, and are occasionally isolated from nectar although not as the dominant member of nectar microbial communities [\[41–](#page-9-26)[43,](#page-9-27) [45\]](#page-9-28).

Ours is the frst study to characterize the microbial community of *B. rapa* nectar. While *Fructobacillus fructosus* and *F. tropaeoli* are known to inhabit fowers [[44](#page-9-29)], a recent review of nectar-associated microbes did not classify *Fructobacillus* as flower specialists [[45\]](#page-9-28). Most of our nectar samples also contained *Acinetobacter* and lactobacilli*,* although these bacteria accounted for fewer than 5% of sequence reads. *Acinetobacter* is frequently isolated from the nectar of many diferent plant species [[46](#page-9-30)] and is considered a fower specialist [[45\]](#page-9-28). Lactobacilli are common bee- and fower-associated bacteria that may play a role in pollinator health [[47\]](#page-9-31). For example, *Apilactobacillus kunkeei* (formerly *Lactobacillus kunkeei*) has been isolated from flowers and is commonly associated with honey bee bread and may potentially aid in honey bee immunity [[43](#page-9-27), [48](#page-9-32)]. *Apilactobacillus micheneri* (formerly *Lactobacillus micheneri*) associates with fowers and non-apis bees [\[49,](#page-9-33) [50\]](#page-10-0). *Apilactobacillus* and *Acinetobacter* are also fructophilic bacteria and may therefore utilize fructose in our artifcial nectar similarly to *Fructobacillus*. Although not detected here, the only other bacterial genus besides *Acinetobacter* that has been classifed as a fower specialist is *Rosenbergiella* [[45\]](#page-9-28). *Rosenbergiella nectarea* can utilize fructose and various other sugars [\[51\]](#page-10-1) and is not considered a fructophilic bacterium. Nectar communities dominated by *Rosenbergiella* may therefore show different effects on floral chemistry and pollinator attraction under warming, warranting further study.

In contrast to the fructophilic bacteria that are often found in foral nectar, nectar-inhabiting yeasts tend to decrease overall sucrose while increasing fructose and glucose [\[16](#page-9-3)]. Although yeasts are commonly found in nectar their presence is not universal [[52](#page-10-2)], and we did not detect them in our samples, suggesting that *B. rapa nectar* may not harbor yeasts. Dispersal of nectar microbes depends on many mechanisms including flower visitor [[45](#page-9-28)], and lack of the appropriate vector or the chemistry of our artifcial nectar may explain the absence of yeast in our samples. Nectarinhabiting yeasts have the potential to warm flowers in cooler climates, which attracts pollinators [[53](#page-10-3)]. However, little is known about the efects of increased environmental temperatures on foral yeasts. Future studies should examine the effects of warming on nectar microbial communities that include yeast.

The communities that we studied were dominated by a single bacterial genus, as fower microbiomes are known for low species richness and evenness [\[45](#page-9-28)]. Floral microbial communities are often less diverse than that of leaves and other plant parts [[18](#page-9-5)]. Nectar microbial communities are filtered by nectar properties  $[12]$  $[12]$  making this environment hospitable mainly for a small group of flower specialists [\[45\]](#page-9-28). Research on priority effects indicates that the initial microbial colonist may persist and outcompete other nectar microbes later in the season [[54\]](#page-10-4). However, patterns of foral microbial diversity and richness vary with geographical location [\[18](#page-9-5)]. The dominance of *Fructobacillus* in our lab experiment represents a "snapshot" of the *B. rapa* microbial community and may refect a lack of diverse plant, pollinator, and microbial communities in the immediate area. As human-modifed landscapes now dominate much of our planet [\[55\]](#page-10-5), our low-diversity community may be common. Broad surveys of nectar microbial communities are still lacking [\[45](#page-9-28)], and continued research on this topic is needed to untangle the efects climate change has on plant-pollinatormicrobe interactions.

When given the choice, bumble bees preferred nectar with bacteria at ambient temperature, suggesting that either the loss of fructose or an overabundance of microbial metabolites infuenced bumble bee foraging choices. Nectar yeast metabolites, such as volatile organic compounds (VOCs), act as informative cues of nectar rewards, including indicating the presence of higher sugar concentrations [\[56](#page-10-6)]. Nectar with dissolved secondary metabolites from yeast elicited an enhanced gustatory response in bumble bees,

indicating the importance of these chemicals [[15\]](#page-9-2). Bumble bees have a strong preference towards sucrose-rich nectar [[57\]](#page-10-7). As sucrose levels remained unchanged but fructose levels decreased, our results suggest that bumble bee preferences are either also afected by fructose and glucose levels or by the correct blend of sugars and microbial metabolites. It could also be that as fructose was depleted in the highdensity *Fructobacillus* samples, the bacteria switched to a less preferred biochemical pathway that resulted in the formation of repulsive metabolites.

Whether sugars, bacterial metabolites, or interactions between the two drive bumble bee foraging choices needs further study. It is important to note that only the nectar bacterial communities were exposed to temperature treatments; bumble bees were kept at a constant temperature throughout the study, and we therefore do not consider changes in bee energetics with temperature. Overall, our data indicate that the microbial community within nectar is important for pollinator choice and is mediated by abiotic factors such as temperature. As temperatures increase due to climate change, alterations to nectar microbiomes may have adverse efects on pollinator choice.

As pollinators forage for resources, they use many mechanisms to choose high-quality pollen and nectar. Although high sugar concentrations are important for optimal foraging, the microbial component of nectar is also a signifcant factor for pollinator choice. For example, honey bees have been known to avoid nectar colonized with the bacteria *Asaia astilbes*, *Erwinia tasmaniensis*, and *Apilactobacillus kunkeei* [[58\]](#page-10-8). However, honey bees are not deterred by nectar colonized with *Metschnikowia reukaufi*, a commonly found nectar-inhabiting yeast [\[58\]](#page-10-8). Recent studies have revealed that these yeasts have a positive impact on bumble bee colony growth [\[59\]](#page-10-9). Our study follows a similar pattern to this previous work, as bumble bees consumed more nectar with a bacterial community than nectar with no microbes but a higher sugar concentration. The preference for microbeinoculated nectar indicates that either these microbes are potentially advantageous, perhaps giving nutritional benefts to the bee or that the bees are making sub-optimal foraging decisions.

We are the first to show that temperature can affect nectarmicrobe-pollinator interactions in vitro. With an increase in temperature, we saw an increase in the absolute abundance of *Fructobacillus* spp*.* in the nectar. This greater density of *Fructobacillus* altered nectar chemistry and ultimately pollinator preference, connecting climate change to pollinator behavior as mediated by nectar microbes. Although we only compared two temperatures in this study there is a clear difference in bacterial abundance with temperature in vitro, and future studies should investigate the efects of a gradient of natural temperatures on this system in vivo. Previous studies have shown that climate change is afecting plant-pollinator

mutualisms by causing plant phenological shifts that can disrupt pollinator mutualisms [\[60](#page-10-10)]. As temperatures change and precipitation decreases, plant metabolism is likely to respond to warming, altering nectar properties [\[61](#page-10-11)]. Alteration in nectar properties and plant response may select for diferent microbial communities, which in turn, may diferentially afect pollinator preference. We point out that as the climate changes nectar microbes and plant physiology may continue to adapt to the new environmental conditions. Bee preference and consumption may also adapt to these potentially new nectar microbial communities and nectar chemistries, such that the results we found here may not apply to the future climate. However, bees evolve more slowly than the climate is changing [\[62](#page-10-12)], which suggests that disruption of these plant-pollinator-microbial interactions is likely.

### **Conclusion**

Our data show that warming afects the density of nectarinhabiting microbes, which in turn alter nectar chemistry and pollinator preference. We thereby elucidate a connection between climate change, plant- and pollinator-associated microbes, and pollinator behavior. Field studies on these interactions can shed light on whether changes to nectar-inhabiting microbiomes mediated by climate change will infuence pollination success and if plants are able to select nectar microbial communities under climate change stress. Future studies looking at the effects of temperature on nectar-inhabiting microbes, changes in nectar composition *in planta,* and plant ftness will be especially valuable.

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**Author Contribution** KAR and QSM designed this experiment. KAR conducted the experiment and analyzed the data. KAR wrote the frst draft of the manuscript and KAR and QSM edited further drafts.

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**Data Accessibility** Metabarcoding amplicon data and associated metadata are available on the NCBI Sequence Read Archive (SRA accession number PRJNA758089).

#### **Declarations**

**Conflict of Interest** The authors declare no competing interests.

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