Bacterial Symbionts Confer Thermal Tolerance to Cereal Aphids *Rhopalosiphum padi* and *Sitobion avenae*

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**Abstract:** High-temperature events are evidenced to exert significant influence on the population performance and thermal biology of insects, such as aphids. However, it is not yet clear whether the bacterial symbionts of insects mediate the thermal tolerance traits of their hosts. This study is intended to assess the putative association among the chronic and acute thermal tolerance of two cereal aphid species, *Rhopalosiphum padi* (L.) and *Sitobion avenae* (F.), and the abundance of their bacterial symbionts. The clones of aphids were collected randomly from different fields of wheat crops and were maintained under laboratory conditions. Basal and acclimated CTmax and chronic thermal tolerance indices were measured for 5-day-old apterous aphid individuals and the abundance (gene copy numbers) of aphid-specific and total (16S rRNA) bacterial symbionts were determined using real-time RT-qPCR. The results reveal that *R. padi* individuals were more temperature tolerant under chronic exposure to 31 °C and also exhibited about 1.0 °C higher acclimated and basal CTmax values than those of *S. avenae*. Moreover, a significantly higher bacterial symbionts’ gene abundance was recorded in temperature-tolerant aphid individuals than the susceptible ones for both aphid species. Although total bacterial (16S rRNA) abundance per aphid was higher in *S. avenae* than *R. padi*, the gene abundance of aphid-specific bacterial symbionts was nearly alike for both of the aphid species. Nevertheless, basal and acclimated CTmax values were positively and significantly associated with the gene abundance of total symbiont density, *Buchnera aphidicola*, *Serratia symbiotica*, *Hamilton defensa*, *Regiella insecticola* and *Spiroplasma* spp. for *R. padi*, and with the total symbiont density, total bacteria (16S rRNA) and with all aphid-specific bacterial symbionts (except *Spiroplasma* spp.) for *S. avenae*. The overall study results corroborate the potential role of the bacterial symbionts of aphids in conferring thermal tolerance to their hosts.
Keywords: wheat aphids; thermal traits; critical thermal maxima; chronic temperature tolerance; aphid endosymbionts; bacterial gene abundance

1. Introduction

Global warming is primarily manifested not only by a gradual rise in the Earth’s average temperature, but also by the occurrence of frequent extreme high temperature events. Such high temperature events can influence population performances and the demographic parameters of insects, both in temporal and spatial scales [1–3].

Inferring the influence of extreme climate events on insects necessitates a better comprehension of their thermal tolerance limits and the mechanisms behind it [4–6]. Previous works have shown the significant impact of extreme high temperatures, both under acclimated and chronic exposures, on the physiology and thermal biology of invertebrates, including aphids [1,7–13]. Some studies, for instance, have demonstrated that different aphid species respond differently to chronic and acclimated temperature exposures [3,9–13]. However, the underlying mechanisms for such thermal impacts on aphids’ biology and ecology are largely unknown.

Aphids have been model systems to study insect–microbial symbiont interactions. They harbor many symbiotic bacteria within their bodies, including primary or obligate (Buchnera aphidicola) and secondary or facultative (Serratia symbiotica, Hamiltonella defensa, Regiella insecticola, Rickettsia spp. and Spiroplasma spp.) symbionts [14–18]. These aphid-specific symbiotic bacteria perform various functions, such as obligatory B. aphidicola provides nutritional supplementation and other facultative endosymbionts confer their hosts resistance to natural enemies and environmental extremities [19–24]. Previous studies have demonstrated the potential mediation of heat tolerance in aphids by their obligate and facultative bacterial symbionts [25–28].

In this paper, we address the putative effects of high temperature exposure on the thermal tolerance of aphids and their bacterial symbionts’ abundance. We intend to understand if short-term heat acclimation and chronic exposure to high temperatures would mediate any effect on the thermal tolerance, survival and abundance of symbiotic bacterial of aphids and if the thermal tolerance traits of aphids correlate to the abundance of their respective bacterial symbionts. To this end, individuals of two cereal aphids, i.e., bird cherry-oat aphid, Rhopalosiphum padi (L.), and English grain aphid, Sitobion avenae (F.), were chronically exposed to 31 °C (until death) and were acclimated to 34 °C for 3 h, before subjecting them to basal and acclimated critical thermal maxima (CTmax) determination. The abundance of aphid-specific bacterial endosymbionts was assessed using qPCR and their correlation with host thermal traits was worked out.

2. Materials and Methods

2.1. Collection and Rearing of Aphids

In this study, cereal aphids R. padi and S. avenae were studied as model species because of their differential performance under extreme temperature regimes [1,3]. About 100 wild clones of R. padi and S. avenae were randomly collected from winter wheat (Triticum aestivum L.) fields near the Henan (35°59′26.0″ N 114°31′37.9″ E) and Hebei (39°30′36.4″ N 115°55′58.2″ E) provinces of China. These aphid clones were transferred separately under cool conditions in plastic tubes. These clones were reared separately on wheat seedlings. Rearing was conducted up to F3 generations under standard conditions, i.e., at 65 ± 5% relative humidity, 22 ± 1 °C temperature and under 16 h:8 h light:dark photoperiod.

2.2. Experiment of Chronic Thermal Exposure

In order to determine the association between chronic heat tolerance and the gene abundance of aphid-specific bacterial symbionts, we collected three batches of aphids from the three generations reared in the laboratory. There were 33 apterous (5 days old) active
and healthy aphid individuals in each batch. The tested aphids were reared individually on wheat leaves plugged in a moist sponge fixed in plastic tubes (30 mm diameter and 100 mm length) and the leaves were changed on alternate days. The three batches of aphids were exposed until death to 31 °C in a climate chamber (RXZ-280B; Jiangnan Ltd., Ningbo, China) set at 55–70% relative humidity and under 16 h: 8 h light:dark photoperiod. The observations were made at regular intervals of 3–6 h until the end of experiment and dead aphids were transferred immediately in vials containing 95% ethanol and were preserved at −20 °C in a freezer for the extraction of DNA. For the comparison of aphid-specific and total bacterial symbiont communities, dead aphid individuals were categorized into four mortality time periods as per their tolerance to chronic temperature (31 °C). For R. padi, mortality time period 1–4 refer to the individuals that died within 6–24, 24–48, 48–72 and 72–96 h, respectively, while for S. avenae, mortality time period 1–4 refer to the individuals that died within 6–18, 18–36, 3–54 and 54–66 h, respectively (Figure 1).

2.3. Basal and Acclimated Critical Thermal Maxima Determination

To determine basal critical thermal maxima (CTmax), we collected 3 batches of aphids (each batch with 33 individuals) from the 3 generations of aphids reared in laboratory. The tested aphids were placed individually in a multi-well transparent plastic arena, which was then hanged in a vertical position in the middle of a double-layered glass container (20 × 30 cm) of a programmed glycol bath with an accuracy of ±0.01 °C (Ministat 230-cc-NR; Huber Ltd., Berching, Germany). The temperature in the container was first maintained at 21 °C for 20 min and then was augmented gradually by 0.1 °C min⁻¹ until the death of all test aphid individuals. Panasonic HDC-HS700 HD Camcorder (Panasonic, Osaka, Japan) was employed for recording the entire behavior of aphids in the arena plate during whole heating process. The temperature at which an aphid individual showed body spasms and lost its ability to move was noted as its CTmax value [29]. At the end of experiments, dead
aphids were transferred immediately in vials containing 95% ethanol and were preserved at −20 °C in a freezer for the extraction of DNA.

To determine acclimated CTmax, we collected 3 batches of aphids (each batch with 33 individuals) from the 3 generations of aphids reared in laboratory. The tested aphids were first acclimated at 34 °C for 3 h (this acclimation duration was selected as aphids did not lose their fitness up to 3 h in a pilot test; see Figure S1), and then were subjected to acclimated CTmax determination using same protocol as mentioned above for basal CTmax.

2.4. Quantification of Aphid Bacterial Symbionts

After surface sterilization with 1.0% sodium hypochlorite, the total genomic DNA was extracted from the preserved aphid individuals using TIANamp® genomic DNA Kit (Tiangen Biotech, Beijing, China), according to the protocol provided by the manufacturer. Purified DNA was eluted from each sample using 200 μL of Tris-EDTA buffer provided in the DNA extraction kit, and after quantification by NanoDrop™ spectrophotometer (ND-1000, Thermo Fisher Scientific, Waltham, MA, U.S.A.), it was preserved at −20 °C in a freezer until downstream molecular analysis. Diagnostic PCR tests were carried out with MyCycler™ thermal cycler (Bio-Rad Laboratories Inc., Hercules, CA, U.S.A.) for the detection and subsequent optimization of annealing temperatures of primers. Primer pairs along with their sequences used for the diagnostic PCR amplifications of obligate and facultative bacterial symbionts of aphid individuals are detailed in Table 1.

Table 1. Primer sequences used for the PCR amplification of total and aphid-specific bacterial symbionts of cereal aphids.

| Symbiont Category | Type     | Taxonomic Name | Bacterial GROUP | Gene Length (kb) | Primer Code  | Real Time Quantitative PCR Primer Sequence |
|-------------------|----------|----------------|-----------------|-----------------|--------------|------------------------------------------|
| Total Bacterial Community (16S rRNA) | Eubacteria | 1.47 | 16S rRNAF | CCTACGGGAGGCAGCAG |
| Primary/Obligate Symbiont | P-type | Buchnera aphidicola | Gammaproteobacterim | 1.5 | BaF | TGAGAGGATAACGACGCCACAC |
| Secondary/Facultative Symbionts | R-type/PASS | Serratia symbiotica | Gammaproteobacterim | 1.46 | BsF | CCGTGACAAAAACTGACGCC |
| T-type/PABS | Hamiltonella defensa | Gammaproteobacterim | 1.3 | HdF | CCTCTAAACAGTAACTGAGC |
| U-type/PAUS | Regiella insecticola | Gammaproteobacterim | 1.3 | RIF | AGCCACACTGGAACCTAGAAAC |
| S-type/PAR | Rickettsia spp. | Alphaproteobacterium | 1.46 | RspF | GCTGCCAAAATTGAACCTGTCATCT |
| Spiroplasma spp. | Mollicutes | 1.4 | SspF | GCGTATACATAGTGGGCAAAC |

For RT-qPCR amplifications, linearized recombinant plasmids (pGEM-T Easy Vector; Promega) were prepared using pGM-T Cloning Kit (Tiangen Biotech, Beijing, China) having a standard sequence inserts of target genes. Standard curves were created using serial dilutions (10-fold) of purified linearized plasmids containing 101 to 109 copies of the targeted bacterial genes. Each RT-qPCR reaction mixture of 20 μL was constituted of 10 μL 2× SuperReal PreMix Plus (Tiangen, Beijing, China), 0.6 μL of each of the 10μM forward and reverse primers, 8.4 μL of ddH2O and 1 μL of 10 ng μL−1 of DNA template. The thermal protocol used for RT-qPCR amplifications included an enzyme activation step of 94 °C for 600 s, 40 cycles with denaturation, annealing and extension steps, respectively.
at 94 °C for 30 s, 55 °C for 30 s and 72 °C for 60 s. For each sample, three independent biological and technical replicates were run.

2.5. Statistical Analysis

Data were statistically analyzed using Statistix V. 8.1® (Tallahassee, FL, USA) analytical software. For each aphid species, the comparison of basal and acclimated CTmax indices was conducted by Student’s paired t-test at \( p \leq 0.05 \). Similarly, Student’s paired t-test was employed to compare the gene abundance of aphid-specific bacterial symbionts between different critical thermal maxima (CTmax) treatments or between both aphid species. Spearman’s rank correlation analysis was worked out to explore the potential association among the gene abundance of aphid-specific endosymbionts and thermal indices of aphids.

3. Results

3.1. Symbionts Gene Abundance and Chronic Thermal Tolerance

When \( R. \) padi were exposed to 31 °C, mortality was less than 5% within 36 h, then increased rapidly and reached a high level of 25% between 36 to 66 h, and then maintained a low level of less than 10% between 66–96 h (Figure S2). The proportion of tested \( R. \) padi individuals that died within the mortality time periods 1–4 when exposed chronically to temperature of 31 °C was 3.6, 49.4, 40.9 and 6.1%, respectively (Figure 1). For \( S. \) avenae, when the tested aphids were exposed to 31 °C, mortality was less than 10% within 18 h, then it increased dramatically and reached a peak of ca. 40% between 18 to 36 h, and then decreased gradually from 15% to very low levels between 36–66 h (Figure S2). The tested \( S. \) avenae individuals that died within the mortality time periods 1–4 when exposed chronically to temperature of 31 °C were 7.2, 72.5, 17.8 and 2.4%, respectively (Figure 1).

Furthermore, qPCR determinations showed that tolerant \( R. \) padi individuals (of mortality time periods 3 and 4) harbored significantly higher gene copy numbers of total (16S rRNA) bacterial symbionts and of all aphid-specific bacterial symbionts, i.e., \( B. \) aphidicola, \( S. \) symbiotica, \( H. \) defensa, \( R. \) insecticola and \( R. \)ickettsia spp. (Figure 2). For \( S. \) avenae, the difference was significant only for 16S rRNA, \( S. \) symbiotica and \( R. \)ickettsia spp. (Figure 3).

![Figure 2](image_url)

Figure 2. Abundance (mean gene copy numbers ± SD; \( n = 20 \)) of bacterial symbionts of aphids (\( R. \)hopalosiphum \( p \)adi) under chronic exposure to 31 °C. Dead aphid individuals were divided into four mortality time periods, i.e., 1–4, representing susceptible-to-tolerant thermal threshold levels. 16S rRNA = total eubacterial rDNA gene; \( B. \) aphidicola; \( S. \) symbiotica; \( H. \) defensa; \( R. \) insecticola; \( R. \)ickettsia spp.; \( S. \)pp. Different letters at the tops of the treatment bar show significant differences between treatments (one-way ANOVA; \( p \leq 0.05 \)).
3.2. Symbionts Gene Abundance and Acute Thermal Tolerance

Mean acclimated and basal CTmax values were 38.82 ± 0.44 °C and 37.41 ± 0.48 °C for *R. padi* and 37.53 ± 0.51 °C and 36.79 ± 0.46 °C for *S. avenae* (Figure 4). Acclimated individuals of both aphid species exhibited significantly higher CTmax values than the non-acclimated aphids (basal CTmax) (Figure 4). On average, CTmax values of *S. avenae* were about 1.0 °C less than those of *R. padi*.

Both aphid species had higher absolute gene copy numbers of total bacteria (16S rRNA) in non-acclimated (basal) aphids than the acclimated ones, but it did not reach the significant level. However, the gene copy numbers of *B. aphidicola*, *S. symbiotica*, *H. defensa* and *R. insecticola* in *R. padi* were significantly higher in the acclimated than in the non-acclimated (basal) aphid individuals (Figure 5). Likewise, in *S. avenae*, the gene abundance of *B. aphidicola*, *S. symbiotica* and *H. defensa* were moderately, but significantly, higher in the acclimated than non-acclimated (basal) aphid individuals (Figure 6). The mean gene abundance of total bacterial (16S rRNA) and aphid-specific bacterial symbionts was higher

![Figure 2](image-url) Abundance (mean gene copy numbers ± SD; n = 20) of bacterial symbionts of aphids (*Sitobion avenae*) under chronic exposure to 31 °C. Dead aphid individuals were divided into four mortality time periods, i.e., 1–4, representing susceptible-to-tolerant thermal threshold levels. 16S rRNA = total eubacterial rDNA gene; *Ba* = Buchnera aphidicola; *Ss* = *Serratia symbiotica*; *Hd* = Hamiltonella defensa; *Ri* = Regiella insecticola; *Rsp* = Rickettsia spp.; *Ssp* = *Spiroplasma* spp. Different letters at the tops of the treatment bar show significant differences between treatments (one-way ANOVA; *p* ≤ 0.05).

![Figure 4](image-url) Basal and acclimated critical thermal maxima (CTmax) (mean ± SD; n = 99) of cereal aphids *Rhopalosiphum padi* and *Sitobion avenae*. Asterisks indicate significant difference among acclimated and basal CTmax for each aphid species (Student’s paired *t*-test; *p* ≤ 0.05).
in *S. avenae* than *R. padi*, except the abundances of *B. aphidicola* and *R. insecticola* gene copy numbers, which were similar in both species (Figure S3).

![Figure 5](image)

**Figure 5.** Abundance (mean gene copy numbers ± SD; *n* = 99) of bacterial symbionts of *Rhopalosiphum padi* aphids in acclimated and basal treatments. 16S rRNA = total eubacterial rDNA gene; *Ba* = *Buchnera aphidicola*; *Ss* = *Serratia symbiotica*; *Hd* = *Hamiltonella defensa*; *Ri* = *Regiella insecticola*; *Rsp* = *Rickettsia* spp.; *Ssp* = *Spiroplasma* spp. Asterisks signify significant difference between acclimated and basal treatments (Student’s paired *t*-test; *p* ≤ 0.05).

![Figure 6](image)

**Figure 6.** Abundance (mean gene copy numbers ± SD; *n* = 99) of bacterial symbionts of *Sitobion avenae* aphids in acclimated and basal treatments. 16S rRNA = total eubacterial rDNA gene; *Ba* = *Buchnera aphidicola*; *Ss* = *Serratia symbiotica*; *Hd* = *Hamiltonella defensa*; *Ri* = *Regiella insecticola*; *Rsp* = *Rickettsia* spp.; *Ssp* = *Spiroplasma* spp. Asterisks signify significant difference between acclimated and basal treatments (Student’s paired *t*-test; *p* ≤ 0.05).

Nevertheless, a significant rank correlation was recorded between the thermal tolerance indices and the abundance (gene copy numbers) of total symbiont density, *B. aphidicola*, *S. symbiotica*, *H. defensa*, *R. insecticola* and *Spiroplasma* spp. for *R. padi*, and with the total symbiont density, total bacteria (16S rRNA) and with all aphid-specific bacterial symbionts (except *Spiroplasma* spp.) for *S. avenae* (Table 2).
Table 2. Correlation among the critical thermal maxima (CTmax) of cereal aphids with their bacterial symbiont gene abundance.

| Bacterial Symbionts          | Rhopalosiphum padi (n = 99) | Sitobion avenae (n = 99) |
|------------------------------|------------------------------|--------------------------|
| Total eubacterial community  | 0.298                        | 0.670 **                 |
| Buchnera aphidicola          | 0.529 **                     | 0.939 **                 |
| Serratia symbiotica         | 0.481 **                     | 0.850 **                 |
| Hamiltonella defensa         | 0.435 *                      | 0.658 **                 |
| Regiella insecticola        | 0.414 *                      | 0.618 **                 |
| Rickettsia spp.             | 0.248                        | 0.390 *                  |
| Spiroplasma spp.            | 0.473 **                     | 0.048                    |
| Total symbionts density     | 0.514 **                     | 0.350 *                  |

Spearman’s Rank Correlation Coefficients (rho); * = correlation is significant at the 0.05 level (2-tailed); ** = correlation is significant at the 0.01 level (2-tailed).

4. Discussion

Extreme high temperature events exert significant effects on the thermal tolerance of aphids as simulated through chronic and acclimated exposures to high temperatures [7,9–13]. However, do the bacterial symbionts of aphids mediate the chronic and acute thermal tolerance of their hosts? This question remains to be cleared. This study was performed to determine the association between the chronic and acute thermal tolerance of R. padi and S. avenae aphids, and the gene abundance of their total (16S rRNA), secondary or facultative (S. symbiotica, H. defensa, R. insecticola, Rickettsia spp. and Spiroplasma spp.) and primary or obligate (B. aphidicola) endosymbiotic bacteria [17,18].

The findings of this study corroborated the fact that the R. padi species is more heat tolerant and exhibits higher evolutionary potential to high temperature events as compared to S. avenae [10,13,30]. Moreover, the acclimation-induced enhanced thermal threshold observed in both aphid species is in line with previous studies demonstrating the greater thermal plasticity induced by the acclimation to elevated temperatures and by heat-hardening in Trichogramma wasps [31], mites [32] and in other organisms, such as in the aquatic hydra/algae holobiont system [33].

Furthermore, a significantly higher gene abundance of bacterial symbionts, particularly of B. aphidicola, S. symbiotica, R. insecticola and Rickettsia spp., were recorded in the cohorts of temperature-tolerant aphid individuals (i.e., of mortality time periods 3 and 4) as compared to susceptible ones (i.e., of mortality time periods 1 and 2) for both aphid species. Likewise, the gene abundance of B. aphidicola, S. symbiotica and H. defensa were significantly higher in the acclimated than non-acclimated (basal) aphid individuals of both aphid species and for R. insecticola for R. padi. More interestingly, a significant and positive correlation was found among the thermal tolerance indices and gene abundance of total symbionts density, B. aphidicola, S. symbiotica, H. defensa, R. insecticola and Spiroplasma spp. for R. padi, and with the total symbionts density, total bacteria (16S rRNA) and with all aphid-specific bacterial symbionts (except Spiroplasma spp.) for S. avenae.

These results validate the potential role of aphid symbiotic bacteria in conferring ecological fitness and thermal tolerance to their host aphids [24–26,28,34,35]. Insect–microbial symbiont interactions play a vital role in the evolutionary adaptation of host insects to ecological stresses, such as extreme thermal exposures [36]. Many previous studies have demonstrated the significance of endosymbionts S. symbiotica and H. defensa in improving aphid tolerance to extreme temperature exposures [26,34–38]. Russell and Moran [35], Montllor et al. [34] and Dunbar et al. [39] demonstrated that bacterial the symbionts S. symbiotica, H. defensa and B. aphidicola ameliorate the thermal tolerance of their hosts and confer tolerance to high temperature exposures. However, how short-term acclimation boosted the gene abundance in the individuals of both species compared to basal (non-acclimated) needs further investigation. In-large, the results of this study corroborate that...
the interactions of aphids and their bacterial symbionts may play an important role in aphids’ thermal adaptation to high temperature exposures or events.

5. Conclusions

In short, a significantly higher abundance of bacterial symbionts was harbored by the individuals of the *R. padi* and *S. avenae* species tolerant to chronic thermal exposures than the susceptible ones. Short-term acclimation to 34 °C considerably enhanced the CTmax (thermal tolerance) for both aphid species. Furthermore, interestingly, the critical thermal maxima values of both species were positively associated with the gene abundance of *B. aphidicola*, *S. symbiotica*, *H. defensa* and *R. insecticola* signifying their putative role in conferring thermal tolerance to their host aphids. In future studies, the diversity and community structure of bacterial symbionts associated with these aphids should be conducted by Illumina deep sequencing of 16S rRNA.

**Supplementary Materials:** The following are available online at https://www.mdpi.com/article/10.3390/insects13030231/s1, Figure S1: Effect of different acclimation times on critical thermal maxima (CTmax) of 5-day-old apterous adults of cereal aphid *Sitobion avenae*; Figure S2: Cumulative percent mortality of cereal aphids *Rhopalosiphum padi* and *Sitobion avenae* under 31 °C for different exposure times; Figure S3: Gene copy numbers (mean ± SD) of total (16S rRNA) and aphid-specific bacterial symbionts in basal and acclimated 5-day-old apterous adults of cereal aphids *Rhopalosiphum padi* (RP) and *Sitobion avenae* (SA).

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