

1 **Interplay of diet, heat stress, and the microbiome shapes health and escape behavior in**
2 **amphibian larvae**

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19 Running title: Diet and heat shape tadpole microbiomes

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22 **Data availability statement**

23

24 Raw data and R scripts are deposited in FigShare
25 (<https://doi.org/10.6084/m9.figshare.29447390>). Raw sequences are deposited in the NCBI
26 (BioProject PRJNA1304763).

27

28 **Acknowledgements**

29

30 We are thankful to Miguel Vences, Sven Gippner, and Janina Rudolph for field assistance,
31 Maileen Weidner, Maline Türk, Fabian Bartels, and Ben Oetken for help with animal
32 husbandry, and Christoph Reisdorff for support in the isotope analyses conducted in his
33 laboratory. We also thank Frank Suhling for identification of the dragonfly naiads, Selma
34 Vieira and Johannes Sikorski for helpful advice during bioinformatic analyses, and Robin
35 Schmidt for providing the picture of the mating pair of *Rana temporaria* used in Fig. 1. We
36 especially thank Richard Wassersug for enlightening conversations about amphibian larvae
37 behavior and its analysis. We acknowledge financial support from the Open Access
38 Publication Fund of the Leibniz Institute DSMZ - German Collection of Microorganisms and
39 Cell Cultures GmbH. DNA analyses were supported by the Deutsche
40 Forschungsgemeinschaft (DFG; GZ: CA 3427/2-1, project number: 546565602 granted to

41 PCE). Experimental work at the Technical University of Braunschweig was funded by the
42 DFG (Project number: 459850971; granted to KR). KR was supported by Marie-Curie
43 Actions (Grant Number: 101151070-AMPHISTRESS).

44

45 **Conflict of interest statement**

46 Not applicable.

47

48 **Ethics statement**

49

50 Permits for the experiments were obtained from the Niedersächsisches Landesamt für
51 Verbraucherschutz und Lebensmittelsicherheit, Germany (Gz. 33.19-42502-04-20/3590 and
52 33.19-42502-04-22-00274). Fieldwork was carried out with permits from the Stadt
53 Braunschweig (Stadt Braunschweig - Fachbereich Umwelt und Naturschutz, Willy-Brandt-
54 Platz 13, 38102 Braunschweig; Gz. 68.11-11.8-3.3).

55

56 **Author contributions**

57 P.C.E. and K.R. were responsible for funding acquisition, conceptualization, methodology,
58 investigation, data curation, project administration, writing —review and editing; P.C.E. was
59 also responsible for formal analysis, validation, visualization, writing—original draft. J. G., F.
60 B. and J. O. contributed to investigation. All authors participated in writing —review and
61 editing.

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65

66 **Abstract**

67

68 Diet influences animal health and their microbiomes, potentially affecting how they cope with
69 environmental stressors such as rising temperatures and altered food quality associated with
70 climate change. Using a multifactorial experiment, larvae of the frog *Rana temporaria* were
71 reared on three diets differing in protein, fat, and animal-derived components (low-,
72 intermediate-, and high-quality), at two temperatures (18 °C and 24.5 °C), and either exposed
73 or not to a simulated heatwave (28 °C for 48 h). We examined how these treatments and
74 associated shifts in gut bacterial indicators and predicted microbial metabolic pathways
75 related to nutrient assimilation, host health (body condition and developmental rate), and
76 escape behavior. Larvae maintained body condition and developed faster at 24.5 °C, with
77 higher diet quality further accelerating development. An intermediate-quality diet reduced
78 responsiveness to an aversive stimulus at 24.5 °C, although this effect disappeared following
79 heatwave exposure. Heatwave conditions were associated with increased abundance of
80 *Klebsiella* and a predicted increase in the myo-inositol degradation pathway, which may
81 influence membrane dynamics and signaling and may increase attention levels. Despite
82 microbial shifts, host performance remained similar across most treatments, suggesting
83 substantial microbiome plasticity and the presence of functionally redundant enterotypes that
84 help buffer environmental stress.

85

86 *Key-words:* Food quality, thermal stress, bacteria, escape behavior, developmental plasticity,
87 behavioral plasticity, gut-brain-axis, *Rana temporaria*

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89

90 **Introduction**

91

92 What animals eat shapes their available energy, growth, and development, ultimately
93 affecting their likelihood of survival (e.g., Kupferberg, 1997; Wang et al., 2015; Llobat and
94 Marín-García, 2022). Beyond its nutritional value, food intake also influences the microbiome
95 - the diverse community of microorganisms (Archaea, Bacteria, Fungi, Protists, Viruses), their
96 metabolites, and interactions (Berg et al., 2020) - that inhabit animal guts (Tuddernham and
97 Sears, 2015) and contribute critically to nutrient assimilation and host health (McFall-Ngai et
98 al., 2013). Animals and their mutualistic or commensal microbial partners have likely been
99 co-evolving since the origin of the animal kingdom (McFall-Ngai et al., 2013). This long-
100 standing association expanded the metabolic potential of animals, enabling the use of
101 otherwise inaccessible food resources and tightly linking host and microbial genomes
102 (McFall-Ngai et al., 2013). The gut microbiome supports digestion and the assimilation of end
103 products by host cells (Perry et al., 2020) and may further influence the host's ability to cope
104 with environmental stress by regulating specific metabolic pathways (Fontaine and Kohl,
105 2023). Because microbiomes respond more rapidly to changing conditions than host genomes,
106 they act as key mediators of animal resilience to environmental stress (Voolstra and Ziegler,
107 2020). On the other hand, microbial communities may also respond to environmental
108 conditions in a way that compromises host health (Douglas and Werren, 2016), what is
109 usually associated to stressful conditions for their hosts (e.g., Fontaine and Kohl, 2023; Guo et
110 al., 2024). Thus, it is important to keep in mind that microbial communities have their own
111 dynamics with varying consequences for host health, and certain environmental conditions
112 can lead to unbalance of host-microbiome interactions or dysbiosis (Zaneveld et al., 2017).

113 Human activities and resulting climate change have created a world in which wildlife
114 faces multiple stressors that compromise individual health, drive population declines, and can
115 ultimately trigger species extinctions (Ruddiman, 2013; McCallum, 2015). Climate change
116 encompasses not only increasing mean temperatures but also transient temperature extremes,
117 altered precipitation patterns, droughts, and shifts in food webs, food quality, and food
118 availability (IPCC, 2023; Hardison and Eliason, 2024). Animals are exposed to both
119 prolonged elevated temperatures and short-term heatwaves, with responses varying according
120 to the intensity and duration of thermal stress (Carreira et al., 2016; Staniek et al., 2025; Xiao
121 and Wang, 2025). The increasing occurrence of heatwaves in Europe, Asia, and Australia
122 (IPCC, 2023) highlights their likely importance for the fate of species under climate change.

123 Emerging evidence suggests that gut microbes play a role in mediating heat tolerance in
124 ectotherms (Dallas et al., 2024). Conversely, impoverished microbiomes may reduce
125 ectotherm tolerance to thermal extremes (Fontaine et al., 2022; Fontaine and Kohl, 2023).
126 Shifts in microbiome composition can modify host thermal resilience by influencing
127 metabolic pathways, oxidative stress resistance, and energy balance. For example, pathways
128 linked to amino acid metabolism - often enriched in hosts with diverse microbiomes - may
129 allow hosts to use bacteria-derived amino acids as additional energy sources during thermal
130 stress (Fontaine and Kohl, 2023).

131 Even species with comparatively high warming tolerance may experience costs at higher
132 temperatures (Duarte et al., 2012). Temperature changes can alter predator-prey dynamics

133 (Seifert et al., 2014), affect key physiological processes, and influence nutrient assimilation
134 (Croll and Watts, 2004). This can lead to shifts in foraging behavior and food preferences
135 (Carreira et al., 2016) and ultimately influence food webs (Seifert et al., 2014). For example,
136 omnivorous amphibian larvae increase their consumption of plant material relative to animal
137 food at higher temperatures, improving growth and performance (Carreira et al., 2016). In
138 crayfish, increased temperatures reduce protein absorption but increase soluble carbohydrate
139 absorption (Croll and Watts, 2004), helping explain reduced consumption of animal-based
140 foods at high temperatures. Thus, diet preferences respond to temperature (Behrens and
141 Lafferty, 2007; Devries and Appel, 2014; Carreira et al., 2016), while the microbiome
142 responds to diet (Tuddernham and Sears, 2015) and can itself influence food intake and
143 behavior (Miri et al., 2023). Food quality and availability, as well as the abundance of key
144 microbial groups, are influenced by the same environmental stressors that affect host survival
145 and recruitment, making these interactions important determinants of species success or
146 failure (e.g., Manning and Sullivan, 2021; Yan et al., 2024; Videvall et al., 2023). Yet, the
147 combined effects of diet and temperature on the microbiome remain poorly understood
148 (Hardison and Eliason, 2024).

149 Behavioral changes influenced by the microbiome extend beyond shifts in foraging
150 behavior (Wong et al., 2015; Miri et al., 2023). Gut microbes produce and regulate numerous
151 neuroactive substances - hormones, neuropeptides, neurotransmitters, and many metabolites
152 that affect host metabolic pathways (Lynch & Hsiao, 2019). These microbial compounds
153 influence neuronal signaling and neural development (Bercik et al., 2012) and include
154 enzymes that synthesize key neuroactive molecules involved in behavioral regulation (Dinan
155 et al., 2015; Chen et al., 2013). This modulation is coordinated through the gut-brain axis - a
156 bidirectional network operating through neural (especially via the vagus nerve), endocrine,
157 and immune pathways (Miri et al., 2023; Silva et al., 2020). Short-chain fatty acids (SCFAs)
158 exemplify influential microbial metabolites that maintain gut integrity, modulate immune and
159 endocrine function, and cross the blood-brain barrier to affect neurotransmission,
160 neurotrophic factors, and microglial activity (Silva et al., 2020).

161 Much research on microbiome-driven behavior has focused on humans or mice as model
162 organisms (Sampson and Mazmanian, 2015), yet understanding the microbiome's role in
163 wildlife evolution and survival is urgently needed (Hird, 2017). In house sparrows,
164 microbiome diversity correlates with exploratory behavior, which in turn promotes greater
165 microbiome diversity (Florkowski and Yorzinski, 2023). The microbiome also influences
166 mate choice and social behavior, with implications for individual fitness and evolutionary
167 success (Sharon et al., 2010; Archie and Theis, 2011). Studies on microbiome-ectotherm
168 interactions are especially important given the sensitivity of ectotherms to climate change and
169 the potential role of their microbiome in mitigating associated stressors (Fontaine and Kohl,
170 2023).

171 Among ectotherms, amphibians are particularly vulnerable to climate change and other
172 stressors (Collins and Storfer, 2003; Hayes et al., 2010; Luedtke et al., 2023), making them
173 the most threatened vertebrate group globally (Wake & Vredenburg, 2008; Borzée et al.,
174 2025). They are therefore valuable model organisms for studying interactions among climate
175 change, diet, microbiome, and behavior. Amphibian diet shapes larval growth and

176 development (Kupferberg, 1997; Carreira et al., 2016; Ruthsatz et al., 2019), while the
177 microbiome affects larval thermal stress tolerance (Fontaine and Kohl, 2023). Altered
178 foraging behavior may reduce thermal stress impacts (Carreira et al., 2016), yet amphibian
179 larvae often exhibit lower thermal tolerance than their predators, potentially increasing their
180 vulnerability to predation (Bastiani, 2023). For instance, larvae of the treefrog *Pithecopus*
181 *rusticus* showed reduced thermal acclimation capacity and thermal tolerance compared to a
182 co-occurring dragonfly predator, losing locomotor capacity at temperatures at which predators
183 remained active (Bastiani, 2023). Because predation is a major source of mortality during
184 larval development (McDiarmid and Altig, 1999; Wells, 2019), the ability to avoid predators
185 is essential for survival. Predator avoidance behavior depends on both immobility in response
186 to predator cues (Relyea, 2001; Preston and Forstner, 2015; Eterovick et al., 2020) and rapid
187 escape responses once detected (Hébert et al., 2019). Diet can influence this behavior:
188 nutrient-rich diets enhance growth and escape performance (Kloh et al., 2024), whereas
189 ingestion of toxic cyanobacteria impairs locomotor performance (Moura et al., 2023). Low-
190 quality diets may therefore compromise escape responses, increasing predation risk.

191 Here, we investigated the interconnected and potentially synergistic effects of diet,
192 temperature, and the gut microbiome on the health and behavior of larvae of the European
193 Common Frog (*Rana temporaria*), an ectothermic model organism. Using a multifactorial
194 experimental design, we tested whether three diets differing in amounts of protein, fat, and
195 animal-derived components (low-, intermediate-, and high-quality), two rearing temperatures
196 (18 °C and 24.5 °C), and exposure to a heatwave (28 °C for 48 h) shaped the gut bacterial
197 communities of *R. temporaria* larvae. We then linked these experimental conditions - and the
198 resulting bacterial communities - to food assimilation (isotopic signatures), health biomarkers
199 (body condition and developmental rate), and behavior, focusing on escape responses to an
200 aversive stimulus as a proxy for predator avoidance.

201 We tested three hypotheses: (1) diet quality, sustained elevated rearing temperature,
202 and/or transient heatwaves affect gut bacterial diversity and composition; (2) diet, temperature
203 treatments, and/or altered gut bacterial communities influence larvae's carbon and nitrogen
204 isotopic signatures and affect health biomarkers; and (3) diet, temperature treatments, and/or
205 altered gut bacterial communities lead to differences in behavioral responses to a simulated
206 predator attack. Finally, we predicted metabolic pathways enriched in bacteria that increased
207 in abundance under each treatment to identify potential links between microbial activity and
208 amphibian larval performance.

209

210 **Materials and methods**

211

212 *Experimental design*

213

214 Five egg clutches of the European Common Frog (*Rana temporaria*) were collected on
215 25 March 2023 in the Kleiwiesen (52.328°N, 10.582°E; Braunschweig, Lower Saxony,
216 Germany) and transported to the Zoological Institute of the Technische Universität
217 Braunschweig. When hatched larvae reached developmental stage 25 (*sensu* Gosner, 1960)
218 they were distributed among three food treatments and two controlled-temperature rearing

219 environments (4 larvae per clutch × 5 clutches × 3 food treatments × 2 rearing temperatures =
220 120 larvae; Fig. 1).

221 The food treatments were prepared using soluble powdered foods that differed in protein
222 and fat content, as well as in the diversity of nutrient sources. The diet with the lowest protein
223 and fat levels and the lowest diversity of components (hereafter “low-quality”) consisted of an
224 organic grass powder (NaturaleBio®; *Hordeum vulgare*) containing 3% lipid, 11%
225 carbohydrate, and 32% protein. The diet with the highest protein and fat content and the
226 greatest diversity of components (hereafter “high-quality”) was Sera Micron Nature® fish
227 food, which contains 7.2% lipid, 10.3% carbohydrate, and 56.6% protein. The intermediate
228 diet (“intermediate-quality”) was a thoroughly blended 1:1 mixture of the powders used for
229 the low- and high-quality diets.

230 The energy content of each diet was determined by bomb calorimetry (6200 Isoperobol
231 Calorimeter, Parr Instruments, Moline, Illinois) at the laboratory for chemical analyses at the
232 University of Hamburg. Mean (\pm SD) caloric values were 17.13 ± 0.04 kJ/g ($n = 3$) for the
233 low-quality diet, 18.72 ± 0.03 kJ/g ($n = 3$) for the intermediate-quality diet, and 20.35 ± 0.06
234 kJ/g ($n = 4$) for the high-quality diet.

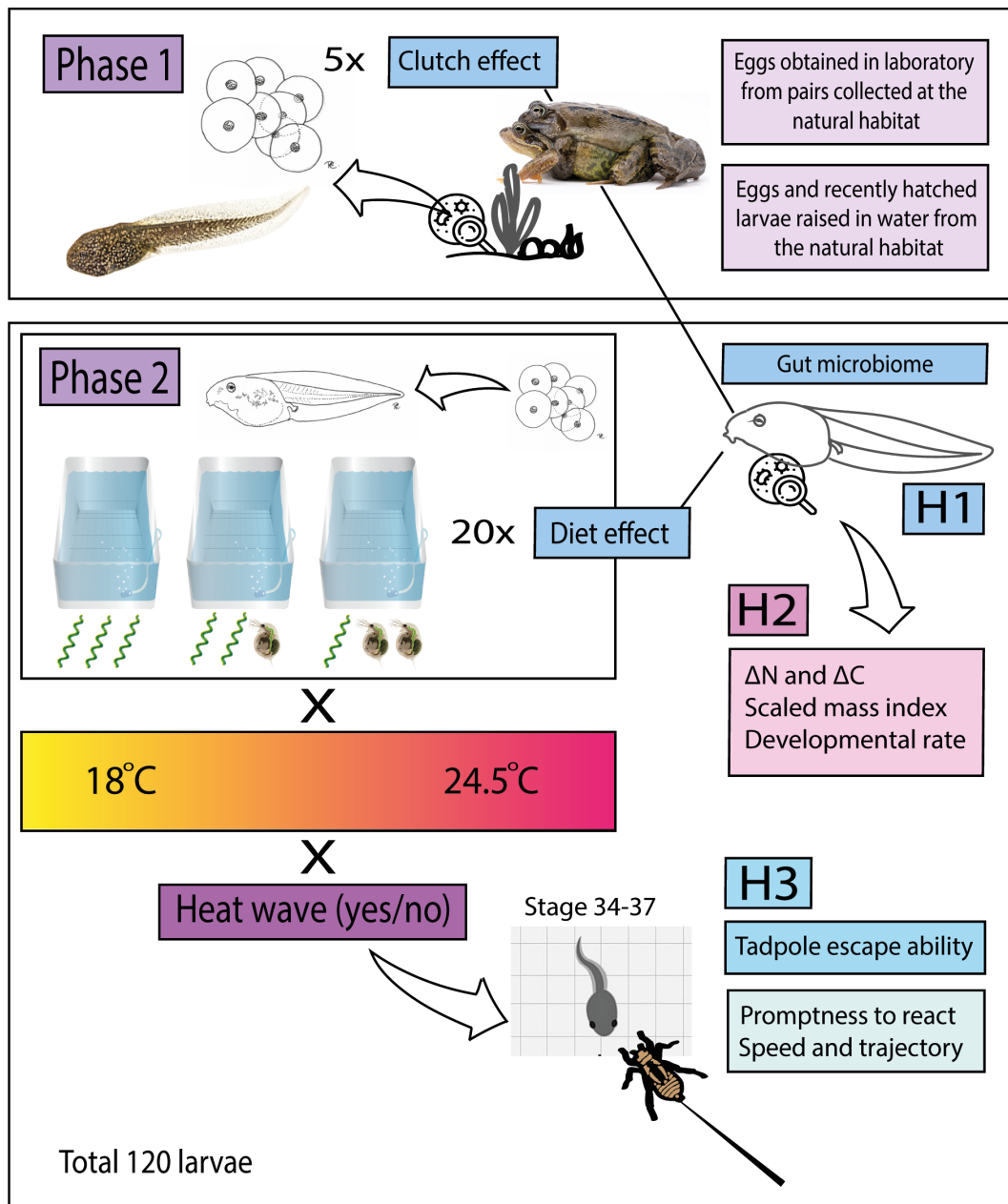
235 The lower temperature (18 °C) represented typical ambient conditions during *R.*
236 *temporaria* larval development. The higher temperature (24.5 °C) was chosen to fall within
237 the range of 22-26 °C, in which *R. temporaria* larvae exhibit elevated stress levels but can still
238 maintain body condition, likely supported by adjustments in their gut bacterial communities
239 (Eterovick et al., 2024).

240 When larvae reached developmental stages 34–37 (pre-metamorphic stages; digit
241 development in the hind limbs; *sensu* Gosner 1960), approximately half of the surviving
242 individuals from each treatment were exposed to a heatwave protocol to test the effects of
243 temperature extremes on escape behavior, as well as potential interactions with diet quality
244 and rearing temperature (Fig. 1). Larvae were kept at 28 °C for 48 h, after which temperature
245 was decreased at the same rate back to the original rearing temperature. Larvae remained in
246 their individual buckets throughout the procedure. Additional details on animal husbandry and
247 experimental setup are available in the supplementary material.

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251
 252 Fig. 1. Graphical summary of the experimental design representing acquisition of offspring (larvae)
 253 from five different egg clutches from *Rana temporaria* and the experiment itself. The experiment
 254 structure is shown based on three main hypotheses to be tested: whether diet and temperatures
 255 experienced during development affect assemblage of gut bacteria (H1), nutrient assimilation and
 256 biomarkers (body condition and developmental rate; H2), as well as escape ability of *R. temporaria*
 257 larvae (H3).

258
 259

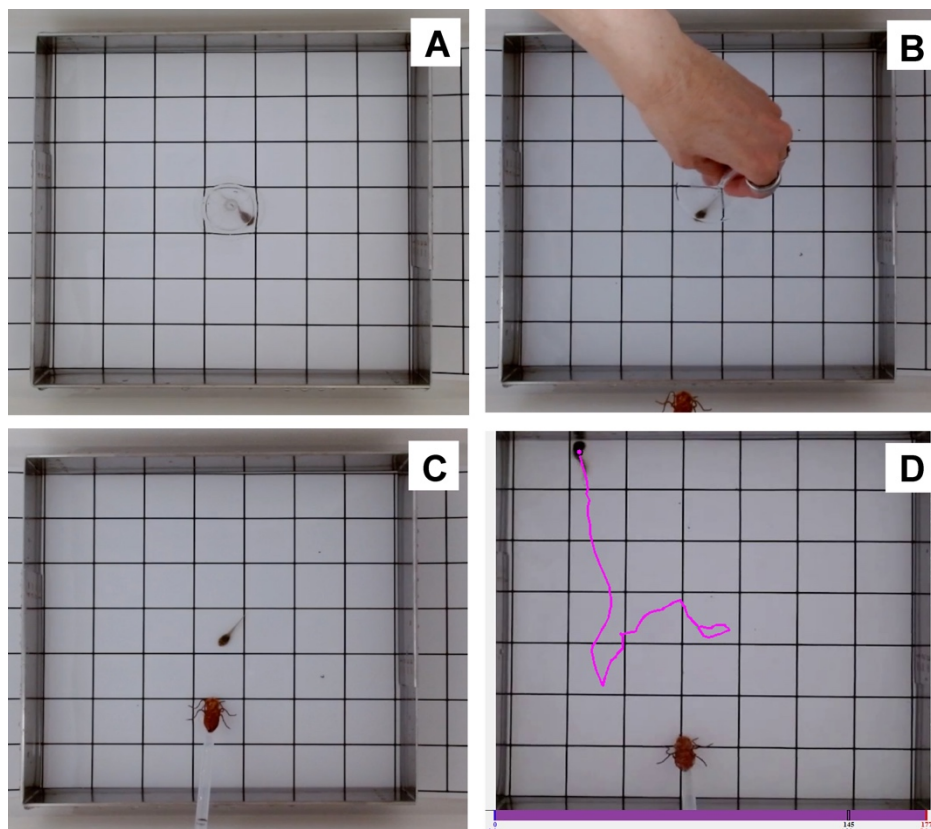
260 Behavioral trials

261
 262 One day after larvae exposed to the heatwave had returned to their original rearing
 263 temperature, behavioral trials were conducted with both heatwave and control (non-exposed)
 264 larvae. A white plastic tray (35 × 35 cm) was filled with 1.3 L of rested tap water at the
 265 rearing temperature of the tested larva (18 °C or 24.5 °C), reaching a water depth of 1 cm. A

266 laminated paper marked with 5×5 cm squares was placed at the bottom, and an adjustable
267 metal frame delineated the arena (Fig. 2). The tray was surrounded by white cardboard to
268 shield larvae from the experimenter, and a high-definition webcam (Logitech C920s HD Pro,
269 Logitech, Lausanne, Switzerland) was mounted on a tripod directly above the arena.

270 Each larva was gently captured from its bucket using a spoon, placed in the central square
271 of the arena, and covered with a glass funnel (Fig. 2A). Handling was minimized and
272 conducted as gently as possible. After a three-minute acclimation period (following Eterovick
273 et al., 2018), the funnel was removed, and a dragonfly naiad (*Libellula*, Libellulidae,
274 Anisoptera) model was presented as a potentially aversive stimulus. The stimulus consisted of
275 a transparent plastic pipette containing 4 mL of water assumed to hold chemical predator
276 cues. This water was obtained from a 500 mL container where ten dragonfly naiads (*Libellula*
277 *depressa*; returned to their habitat after use) from the same frog habitat had been held for 4 h.
278 Each stock of water was used for two hours after removal of the naiads, with water
279 temperatures matched to the larva's rearing temperature. A life-size predator model, made
280 from non-toxic modeling clay and ink, was attached to the pipette tip.

281



282

283

284 Fig. 2. Experimental setup for behavioral tests. Plastic trays filled up to 1 cm with rested tap water at
285 larvae rearing temperature were lined with a grid of 5 x 5 cm squares. A space of 35 x 35 cm was
286 delimited with a metal frame and the larva to be tested was placed at the central square, where it was
287 retained for 3 minutes under a glass funnel (A). After careful removal of the funnel without disturbing
288 the larva (B), a predator model was approached (C) and the reaction of the larva was filmed to
289 evaluate the escape response (see text for details). Fleeing trajectories of the larva were tracked with
290 the software AnimalTA (Chiara and Kim, 2023; D).

291

292 Immediately after funnel removal, the pipette was inserted at $\sim 45^\circ$ relative to the larva's
293 frontal direction, touching the water two grid squares (10 cm) away. Water containing
294 predator cues was slowly released, and the predator model was gradually moved toward the
295 larva until it elicited an escape response or gently touched it. Because amphibian larvae
296 perceive varied cues from predators (Melo et al., 2021), this combined stimulus was designed
297 to engage visual (model), mechanical (approach and water flow), and chemical (predator
298 exposed water) cues, as the most relevant cue for *R. temporaria* larvae is unknown. Trials
299 ended once the larva attempted to flee or if the model touched the larva without eliciting any
300 escape movement. Video recordings were captured using OBS Studio (Open Broadcaster
301 Software, Version 29.1; <https://obsproject.com/>). Larvae were tested in random order and
302 always by the same person, blind to their heatwave exposure and rearing conditions.

303 Videos were analyzed in random order and without treatment information using
304 AnimalTA software (Chiara and Kim, 2023). Occasionally, larvae moved during funnel
305 removal and had to be repositioned in the arena's center. We recorded the "number of
306 attempts" (times a larva was repositioned) as an additional variable, reflecting early
307 movements that could contribute to energy expenditure and stress. To ensure uniformity, we
308 quantified the elapsed time between funnel removal and the trial start (when the predator
309 model contacted the water) and found it to be strongly correlated with the number of
310 repositioning attempts (Spearman's $R_s = 0.9$, $p < 0.001$; Fig. S1), indicating no significant
311 variation in attempt durations.

312 Escape behavior was quantified using the following variables: (1) whether the larva
313 reacted (fleeing) or not, with no reaction defined as remaining stationary even when touched
314 by the model; (2) reaction time, measured from the moment the predator model touched the
315 water until the larva's flee response; (3) whether the larva reacted before or after contact with
316 the predator model; (4) average speed; and (5) trajectory linearity ("meander" function,
317 Chiara and Kim, 2023) during fleeing. Variables 2–5 were analyzed only for larvae exhibiting
318 escape responses. Speed and trajectory linearity were measured until the larva stopped or
319 touched a wall, as such a barrier would otherwise bias the metrics.

320 This behavioral test protocol was used to examine the effects of diet, rearing temperature,
321 and heatwave exposure on larval kinematics. We expected larvae fed higher-quality diets,
322 reared at 18°C , and not exposed to the heatwave to be more alert and reactive, fleeing earlier
323 and at higher speed. Escape trajectories were expected to be more curved, reflecting the
324 typical anti-predator strategy of anuran larvae, which rely on rapid turns with small radii
325 rather than straight-line swimming (Wassersug, 1989). Simply stated, tadpoles typically
326 escape from predatory attacks by turning away from the approaching predator rather than
327 trying to outrun it (Wassersug, 1989).

328 329 *Sample collection*

330
331 Within 12 hours after the behavioral trials, each tadpole was euthanized using $2\text{ g} \times \text{L}^{-1}$
332 tricaine methanesulfonate (MS-222; Ethyl 3-aminobenzoate methanesulfonate; Sigma-
333 Aldrich). The developmental stage of each larva was confirmed under a stereomicroscope
334 according to Gosner (1960). Snout-vent length (SVL) was measured to the nearest 0.5 mm

335 using a digital caliper. Larvae were then gently dry-blotted and weighed to the nearest 0.001 g
336 using an electronic balance (Sartorius A200 S, Germany).

337 A sterile scalpel was used to excise the tail for subsequent isotopic analysis. Using a
338 sterile scalpel and tweezers, the ventral skin was cut to remove the entire gut for bacterial
339 DNA extraction. The tail, gut, and remaining body were placed in three separate tubes, all
340 stored at -80 °C until further analysis.

341 342 *Isotope analyses*

343
344 Stable isotope analyses were conducted to assess differences in absorption and
345 incorporation of food components by larvae subjected to different diets, based on isotopic
346 signatures. Analyses were performed at the Biozentrum Klein Flottbek, University of
347 Hamburg, Germany, following the methods of Glos et al. (2020), as detailed in the
348 supplementary material.

349 350 *Body condition and developmental rate assessment*

351
352 Body condition was estimated using the scaled mass index (SMI), calculated from the
353 slope of the regression of log-transformed snout-vent length (SVL) and log-transformed body
354 mass (standardized major axis, SMA) as: $SMI = [\text{individual Mass} \times (\text{mean SVL of}$
355 $\text{population/individual SVL})^{SMA}]$ (Peig and Green, 2009; 2010).

356 This index has been previously applied to *R. temporaria* larvae (Dittrich et al., 2018;
357 Ruthsatz et al., 2020; Eterovick et al., 2024). In the present study, SMA was 2.742.

358 Developmental rate was calculated as the number of Gosner (1960) stages advanced by
359 each larva divided by the number of days from hatching to the end of the experiment.

360 361 *Bacterial 16S rRNA gene library preparation*

362
363 DNA was extracted from whole guts of larvae using the QIAamp Fast DNA Stool Mini
364 Kit (QIAGEN) following the manufacturer's instructions. Extractions were performed over
365 five days, with one negative control included per day to monitor for contamination. A
366 ZymoBIOMICS™ microbial community standard (Zymo Research Europe GmbH) was used
367 as a positive extraction control on the first and last days of the extraction process.

368 The V4 region of the 16S rRNA gene was amplified using the forward primer 515F (5'-
369 GTGCCAGCMGCCGCGGTAA-3') and reverse primer 806R (5'-
370 GGACTACHVGGGTWTCTAAT-3'; Caporaso et al., 2011). Each sample was tagged with a
371 unique combination of forward and reverse primers from a stock of 24 forward and 24 reverse
372 primer tags. Two PCR plates were prepared, each including one negative control. A positive
373 control consisting of ZymoBIOMICS™ microbial community DNA standard was also
374 included. The Zymo microbial community and DNA standards, which contain known species

375 compositions and abundances, were used to verify the precision of extraction and PCR
376 protocols, respectively.

377 PCR products were pooled and purified. Aliquots were electrophoresed on a 2% agarose
378 gel, and the desired 251 bp fragment was extracted using the Monarch DNA Gel Extraction
379 Kit (New England BioLabs, GmbH, Germany) following the manufacturer's protocol.
380 Purified DNA was quantified with a Qubit™ fluorometer (Invitrogen) and sequenced using
381 the MiSeq500 Illumina platform (paired-end 2 × 250 bp, v2 chemistry) at the Leibniz-Institut
382 DSMZ - German Collection of Microorganisms and Cell Cultures GmbH.

383

384 *Bioinformatic analyses*

385

386 Sequence denoising, filtering, and beta diversity calculations were performed in
387 QIIME2 (Bolyen et al., 2019), as detailed in the supplementary material.

388 Beta diversity was assessed using unweighted UniFrac distances (sensitive to low-
389 abundance microbes; Lozupone and Knight, 2005) and compared among treatments using
390 PERMANOVA with pairwise post hoc tests. Metagenomic predictions of metabolic pathways
391 of the gut microbiota were generated using PICRUSt2 (Douglas et al., 2020), based on the
392 MetaCyc database (Caspi et al., 2016), which provides robust predictions of metabolic
393 pathways related to genes contained in 16S rRNA gene sequence data.

394

395 *Statistical analyses*

396

397 Isotopic signatures were compared among diet treatments and between rearing
398 temperatures using the R package nicheROVER (Swanson et al., 2015; R Core Team, 2024).
399 This approach estimates the probability that the isotopic niches of individuals from one group
400 overlap with those of another, based on quantitative variables such as $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$. We ran
401 1,000 simulations to calculate niche breadth and overlap. Isotopic signatures of the three diets
402 were based on six replicate samples per food type (see Supplementary Material).

403 The effects of food treatment, rearing temperature, and heatwave exposure - including
404 all two- and three-way interactions - on larval mass, body condition, and developmental rate
405 were analyzed with GLMMs in the afex package (Singmann et al., 2024), with clutch identity
406 included as a random factor.

407 Before analyzing behavioral traits, we screened for outliers in the time elapsed between
408 the moment the predator model touched the water and when it touched the larva (where
409 applicable). Two outliers with unusually long times (Fig. S2) were removed. For the
410 remaining data, this interval averaged 7.44 ± 1.78 s. Mixed models were then built to test the
411 influence of food treatment, rearing temperature, and heatwave exposure (fixed variables),
412 including their interactions, on: (1) whether the larva reacted (binary), (2) reaction time, (3)
413 whether the reaction occurred before or after being touched (binary), (4) average speed, and
414 (5) trajectory linearity (see "Behavioral trials"). Trial day and clutch identity were included as
415 random effects nested within food treatment. When full models failed to converge due to
416 model complexity, we simplified random-effect structures or analyzed likely interactions

417 separately (Singmann et al., 2024). For binary outcomes, singular-fit warnings were expected,
418 but results were considered robust when outcomes were consistent across full and simplified
419 models (Singmann & Kellen, 2019; Singmann et al., 2024). Post hoc tests were performed
420 with emmeans (Lenth, 2017).

421 For each behavioral variable, we first tested whether larval mass, body condition, or
422 number of positioning attempts influenced results (Pearson or Spearman correlations for
423 quantitative variables; Wilcoxon tests for binary outcomes). When relevant, these variables
424 were incorporated into the models (e.g., number of attempts as a random factor). We expected
425 larvae in better condition to respond more rapidly and before being touched, and to escape
426 with higher speed and less linear trajectories. Positioning attempts were considered
427 problematic if they were associated with reduced responsiveness, delayed reactions, increased
428 likelihood of being touched, slower speeds, or more linear escapes, as these outcomes could
429 mean that additional positioning attempts reduced the responsive capacity of the larvae.

430 Microbiome bacteria alpha-diversity was calculated using Hill numbers with order of
431 diversity q1 (based on Shannon diversity index) (Alberdi and Gilbert, 2019) using the R
432 package iNext (version 4.5.1; R Core Team, 2025; Chao et al., 2014). Hill numbers represent
433 a number of taxonomic entities that would give the same diversity value if they all had the
434 same abundance (Hill, 1973; Alberdi and Gilbert, 2019). A GLMM was posteriorly built in
435 afex, using food treatment, rearing temperature, and heatwave exposure (and all interactions)
436 as fixed effects and clutch identity as a random effect to explain bacteria alpha diversity.

437 To assess microbiome composition, we constructed a phyloseq object (McMurdie &
438 Holmes, 2013) normalized via Total Sum Scaling (TSS) and tested for microbial markers that
439 differed across the 12 treatment combinations (3 diets \times 2 rearing temperatures \times heatwave vs.
440 no heatwave). Variance homogeneity among groups was evaluated with betadisper (vegan;
441 Oksanen et al., 2013), and ASV abundances were ordinated using PCoA. Microbiome
442 markers (i.e., taxa significantly increased in specific treatments) were identified through
443 LEfSe (Segata et al., 2011) using the R package microbiomeMarker (Cao et al., 2022), with
444 an LDA score threshold of 4. LEfSe identifies taxa (at all taxonomic levels) most likely to
445 explain group-level differences while accounting for statistical significance. We also searched
446 for microbiome markers at the genus and family levels using a probabilistic approach with
447 ALDEx2, which is more robust to variations in numbers of reads (Fernandes et al., 2014), for
448 confirmation purposes.

449 Predicted microbial metabolic pathways were compared among all 12 treatment
450 combinations using ggpicrust2 (Yang et al., 2023), applying the ALDEx2 method for multi-
451 group comparisons.

452

453 **Results**

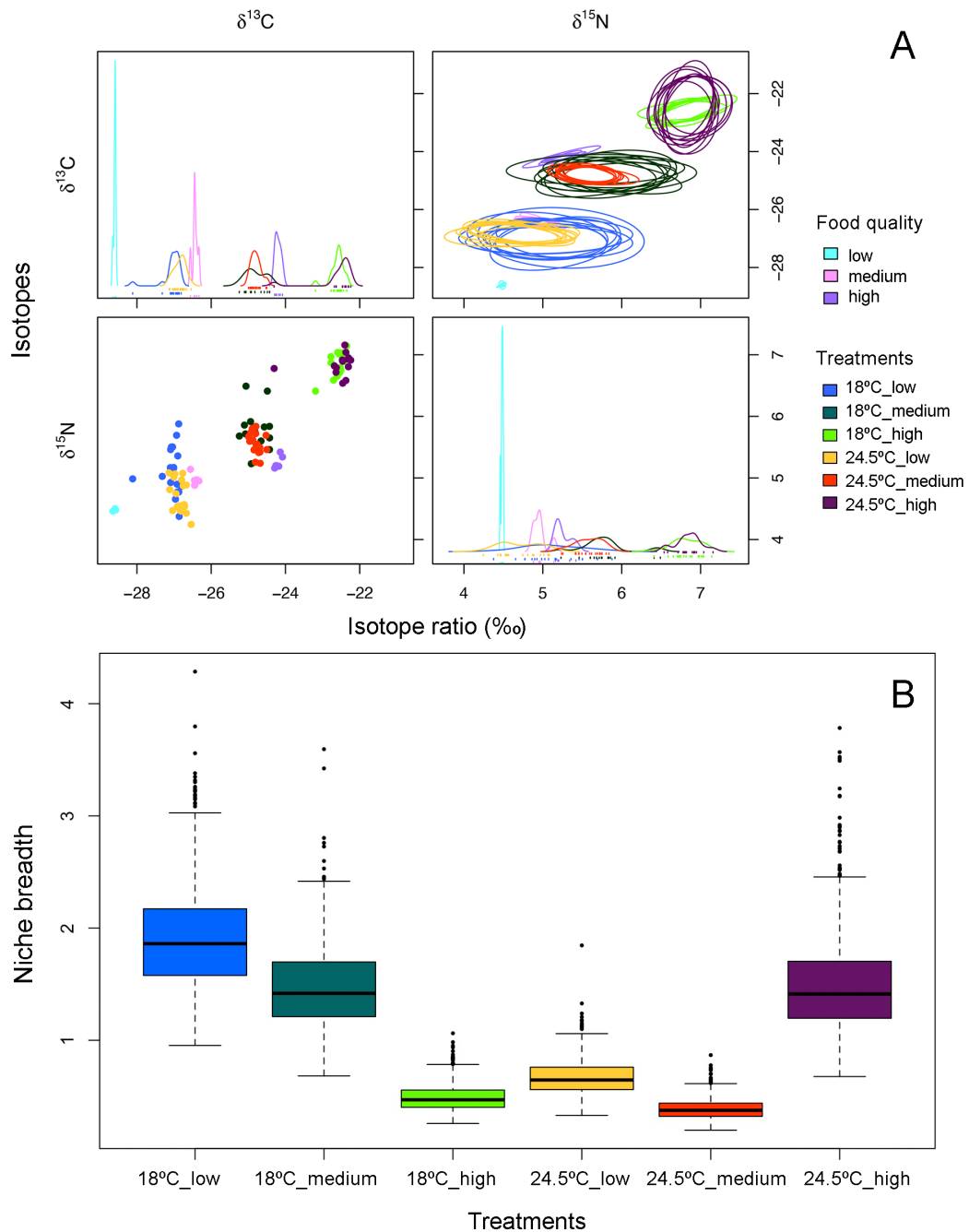
454

455 *Isotope analyses*

456

457 The three diets produced markedly different isotopic signatures in *Rana temporaria*
458 larvae, whereas isotopic niches of the two temperature treatments largely overlapped within

459 each diet (Figs. 3, S3). For the low- and intermediate-quality diets, higher temperature
 460 reduced isotopic niche breadth. In contrast, for the high-quality diet, niche breadth was
 461 narrower at 18 °C and increased at 24.5 °C (Fig. 3).
 462



463
 464 Fig. 3. Isotopic signatures (A) and niche breadth (B) of *Rana temporaria* larvae reared with three food
 465 treatments and two temperatures in a crossed experimental design. Food treatments correspond to diets
 466 with increasing levels of protein, fat, and components of animal origin (their isotopic signatures are
 467 also included in A). Rearing temperatures were 18 °C and 24.5 °C. The isotopic signatures are
 468 represented as lines in one-dimensional density plots (top left and bottom right graphs, A), two-
 469 dimensional scatterplots (bottom left graph; A) and ten random elliptical simulated projections of
 470 trophic niches corresponding to each diet and each treatment (top right graph; A).
 471

472 *Survivorship, development, and body condition*

473

474 Of the 120 larvae used in the experiment, 12 died: six in the 18 °C treatment (five with
475 intermediate- and one with high-quality food) and six in the 24.5 °C treatment (five with high-
476 and one with intermediate-quality food). Five of these deaths occurred during or after the
477 heatwave phase (three heatwave larvae and two controls). One larva developed hydrops and
478 was excluded.

479 Larval body condition (SMI) did not differ among food treatments, rearing
480 temperatures, or heatwave exposure; the same was true for body mass alone (Table 1, Figs.
481 S4, S5). In contrast, developmental rate was higher at 24.5 °C than at 18 °C (Table 1). At 24.5
482 °C, developmental rate also increased with intermediate-quality food (Kenward–Roger post-
483 hoc: estimate = -0.029, SE = 0.010, df = 91.5, $t = -2.874$, $p = 0.025$) and with high-quality
484 food (estimate = -0.054, SE = 0.011, df = 92.0, $t = -4.804$, $p < 0.001$) compared with low-
485 quality food. These effects were absent at 18 °C (Table 1; Figs. 4A, S6).

486

487 *Behavioral trials*

488

489 Escape-behavior trials were conducted with 102 *R. temporaria* larvae. Of the 108
490 surviving larvae, one showed hydrops and four displayed abnormal behavior (lethargy or
491 irregular swimming) and were therefore excluded. In addition, one video file was accidentally
492 lost. Of the 102 larvae tested, 81 responded to the aversive stimulus (61 before being touched
493 by the predator model and 20 upon contact), whereas 21 did not react even when gently
494 touched.

495

496 *Larvae likeliness to react*

497

498 Larval response (reacted vs. did not react) was unrelated to mass ($W = 711.5$, $p = 0.252$;
499 Fig. S7) or body condition ($W = 936$, $p = 0.482$; Fig. S8). The number of attempts needed to
500 position a larva before the trial differed between responders and non-responders ($W = 601$, p
501 $= 0.022$; Fig. S9); however, larvae requiring more positioning attempts were also more likely
502 to react, indicating that repositioning did not impair their ability to respond (Fig. S9). For this
503 reason, number of attempts was included as an additional random effect in the models
504 assessing reaction likelihood.

505 Reaction likelihood was not explained by any fixed factor alone but by interactions
506 among them (Table 2). The full mixed-effects model with random structure did not converge,
507 so we ran a model without random effect structure using the *glmer* function, which also did
508 not converge. Simpler models including only individual predictors and single interactions
509 yielded consistent results using the mixed function (Table 2).

510 Heatwave exposure increased larvae reaction likelihood at 24.5 °C (log odds scale post-
511 hoc tests: $z = -2.309$, $p = 0.021$), whereas at 18 °C it reduced the likelihood of reacting ($z =$
512 2.454 , $p = 0.014$). Heatwave exposure increased reaction likelihood at medium food quality (z
513 $= -2.017$, $p = 0.043$), but not in the other food treatments ($z = 0.475$, $p = 0.635$ for low quality
514 food and $z = 0.037$, $p = 0.970$ for high quality food; Fig. 4B, Table 2).

515

516 Table 1. Models built to explain variability in body condition (SMI), mass, developmental rate (dev_rate) and gut bacteria diversity of *Rana*
 517 *temporaria* larvae reared at two temperatures (either 18 °C or 24.5 °C) and receiving one of three food treatments considered as of low-, medium-,
 518 and high-quality (based on increasing content of protein, fat, and animal components), being exposed or not to a heatwave (HW) in a crossed
 519 experimental design. Developmental rate was calculated as the number of Gosner's (1960) developmental stages advanced during the experiment
 520 divided by the number of days from hatching to the end of the experiment. Significant effects are boldfaced and marked with an *. *mixed* refer to
 521 function employed to run the models.

Dependent variable / GLMM model	Fixed effects	df	F	p	n
Body condition (SMI)					
<i>mixed</i> (SMI ~ diet*temperature*HW + (1 Clutch))	diet	2; 91.72	2.196	0.117	107
	temperature	1; 91.66	0.067	0.797	
	HW	1; 94.99	0.236	0.628	
	diet:temperature	2; 93.03	1.001	0.372	
	diet:HW	2; 93.99	0.594	0.554	
	temperature:HW	1; 94.31	1.551	0.216	
	diet:temperature:HW	2; 90.66	0.190	0.827	
Mass (mg)					
<i>mixed</i> (mass ~ diet*temperature*HW + (1 Clutch))	diet	2; 91.72	2.291	0.107	107
	temperature	1; 91.66	0.124	0.725	
	HW	1; 94.99	0.106	0.745	
	diet:temperature	2; 93.03	0.322	0.272	
	diet:HW	2; 93.99	0.947	0.057	
	temperature:HW	1; 94.31	0.124	0.725	
	diet:temperature:HW	2; 90.66	1.705	0.188	
Developmental rate (dev_rate)					
<i>mixed</i> (dev_rate ~ diet*temperature*HW + (1 Clutch))	diet	2; 91.72	8.428	<0.001*	107
	temperature	1; 91.65	412.706	<0.001*	
	HW	1; 94.99	0.865	0.354	

diet:temperature	2; 93.03	4.404	0.015*
diet:HW	2; 93.99	0.281	0.756
temperature:HW	1; 94.31	3.364	0.070
diet:temperature:HW	2; 90.66	0.036	0.965

Gut bacteria diversity (Hill numbers)

<i>mixed</i> (Hill_q1 ~ diet*temperature*HW + (1 Clutch))	diet	2; 77.31	1.851	0.164
	temperature	1; 78.60	8.537	0.005*
	HW	1; 79.95	0.393	0.533
	diet:temperature	2; 78.43	3.905	0.024*
	diet:HW	2; 79.09	1.266	0.287
	temperature:HW	1; 77.97	0.077	0.783
	diet:temperature:HW	2; 69.18	3.347	0.103

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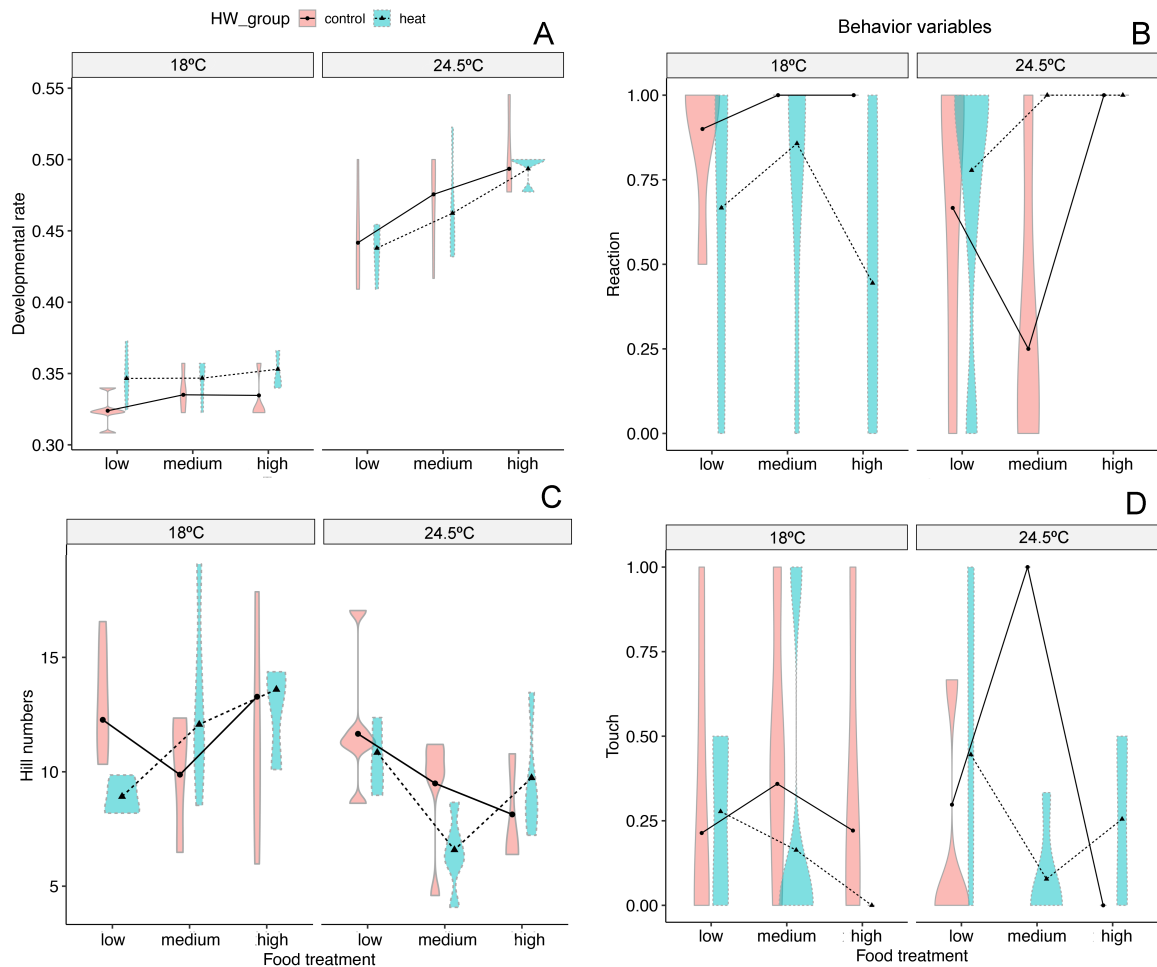
533 Table 2. Models built to explain variability in five dependent variables describing *Rana temporaria* larvae escaping behavior when exposed to an
 534 aversive stimulus consisting of an approaching transparent plastic pipette with a predator model glued to the top releasing 4 ml of water previously
 535 exposed to predators. Analyzed escape responses were: (1) whether the larva reacted or not (no reaction meant not moving even when touched by
 536 the model), (2) larvae reaction time (time elapsed from the moment the predator model touched the water to the fleeing response of the larva), (3)
 537 whether the larva reacted before or after being touched by the predator model, (4) average speed and (5) trajectory linearity while fleeing. *Rana*
 538 *temporaria* larvae were reared at two temperatures (either 18 °C or 24.5 °C) and received one of three food treatments considered as of low,
 539 medium, and high quality (based on increasing levels of protein, fat, and components of animal origin), being exposed or not to a heatwave (HW) in
 540 a crossed experimental design. Significant effects are boldfaced and marked with an *. *mixed* and *lmer* refer to functions employed to run the
 541 models.

Dependent variable / GLMM model	Fixed effects	df	F	p	n
Reaction to the aversive stimulus (binary)					
<i>lmer</i> (reaction ~ diet*temperature*HW + (1 day_filmed) + (1 Clutch) + (1 attempts))		-	-	-	102
<i>mixed</i> (reaction ~ diet*temperature + (1 day_filmed) + (1 Clutch) + (1 attempts))	diet	7; 2	3.010	0.222	
	temperature	8; 1	0.788	0.374	
	diet:temperature	7; 2	10.520	0.005*	
<i>mixed</i> (reaction ~ temperature*HW + (1 day_filmed) + (1 Clutch) + (1 attempts))	temperature	6; 1	0.309	0.580	
	HW	6; 1	0.212	0.654	
	temperature:HW	6; 1	16.366	<0.001*	
<i>mixed</i> (reaction ~ diet*HW + (1 day_filmed) + (1 Clutch) + (1 attempts))	diet	7; 2	3.483	0.175	
	HW	8; 1	1.116	0.290	
	diet:HW	7; 2	13.397	0.001*	
Reaction time					

<i>mixed</i> (reaction_time ~ diet*temperature*HW + (diet day_filmed+clutch))	diet	2; 0.29	0.087	0.934	81
	temperature	1; 1.24	0.290	0.670	
	HW	1; 64.31	0.880	0.352	
	diet:temperature	2; 0.55	0.148	0.888	
	diet:HW	2; 51.67	1.638	0.204	
	temperature:HW	1; 62.73	0.870	0.355	
	diet:temperature:HW	2; 63.25	1.078	0.346	
Touch by the predator model before reaction (binary)			Chisq		
<i>mixed</i> (touch ~ diet*temperature*HW + (1 day_filmed) + (1 Clutch))		-	-	-	81
<i>mixed</i> (touch ~ diet* HW + (1 day_filmed) + (1 Clutch))	diet	6; 2	2.125	0.345	
	HW	7; 1	0.478	0.489	
	diet:HW	6; 2	4.919	0.085	
Speed while fleeing (log)			F		
<i>mixed</i> (logspeed ~ diet*temperature*HW + (diet day_filmed+clutch))	diet	2; 0.86	0.364	0.768	81
	temperature	1; 0.85	0.079	0.831	
	HW	1; 61.65	0.087	0.769	
	diet:temperature	2; 0.84	0.268	0.813	
	diet:HW	2; 52.33	1.900	0.160	
	temperature:HW	1; 61.27	0.012	0.913	
	diet:temperature:HW	2; 61.42	1.697	0.192	
Trajectory non-linearity while fleeing or “meander” (log)					
<i>mixed</i> (logmeander ~ diet*temperature*HW + (diet day_filmed+clutch))	diet	2; 0.61	0.137	0.893	81
	temperature	1; 0.93	0.908	0.525	
	HW	1; 63.74	0.502	0.481	
	diet:temperature	2; 1.04	0.106	0.908	
	diet:HW	2; 51.23	0.614	0.545	

542

temperature:HW	1; 62.44	1.625	0.207
diet:temperature:HW	2; 63.22	1.203	0.307



543
 544 Fig. 4. Interactive effects among food quality, rearing temperature, and exposure to a heatwave in
 545 *Rana temporaria* larvae developmental rate (A), variables describing behavior (B, D) and gut bacteria
 546 diversity (C). Food quality refers to increasing levels of protein, fat, and components of animal origin.
 547 Rearing temperatures were 18 °C and 24.5 °C. The heatwave (HW) corresponded to increasing
 548 temperature at a ramping rate of 0.5 °C per hour until 28 °C, maintenance at 28 °C for 48 h and
 549 subsequent temperature decrease of 0.5 °C per hour until original rearing temperature. Variables
 550 describing behavior are larvae likeliness to react (fleeing) to an aversive stimulus (B) and to be
 551 touched by an approaching predator model before reacting (D). Graphs correspond to violin plots of
 552 estimated marginal means from the model including all fixed variables (see Table 1).

553
 554 *Larvae reaction time*

555 Reaction time, measured for the 81 larvae that responded to the stimulus, was not
 556 influenced by mass (Adjusted $R^2 = 0.030$, $F_{79} = 3.487$, $p = 0.066$; Fig. S10), body condition
 557 (Adjusted $R^2 = -0.013$, $F_{79} = 0.005$, $p = 0.946$; Fig. S11), or the number of positioning
 558 attempts before the trial ($\rho = -0.125$, $p = 0.263$; Fig. S12). Reaction time was also unaffected
 559 by any experimental factor - food treatment, rearing temperature, heatwave exposure - or by
 560 their interactions (Table 2).

561
 562 *Larvae likeliness of being touched*

563 Whether larvae reacted before or after being touched by the predator model was
564 unrelated to mass ($W = 577$, $p = 0.722$; Fig. S13), SMI ($W = 697$, $p = 0.343$; Fig. S14), or the
565 number of attempts needed to position them ($W = 533$, $p = 0.366$; Fig. S15). Reaction
566 likeliness tended to increase in larvae fed medium quality food when exposed to the heatwave
567 (Table 1, Figs. 2D).

568

569 *Larvae escape speed and trajectory*

570 Escape speed and movement non-linearity (“meander”; Chiara & Kim, 2023) were
571 quantified for the 81 larvae that fled, with both variables log-transformed to meet normality
572 assumptions. Neither metric was affected by mass, body condition, or the number of
573 positioning attempts (speed: Adjusted $R^2 = -0.013$, $F_{79} = 0.004$, $p = 0.949$; Fig. S16; Adjusted
574 $R^2 = -0.013$, $F_{79} = 0.009$, $p = 0.923$; Fig. S17; $\rho = -0.104$, $p = 0.354$; Fig. S18; meander:
575 Adjusted $R^2 = 0.003$, $F_{79} = 1.271$, $p = 0.263$; Fig. S19; Adjusted $R^2 = -0.009$, $F_{79} = 0.247$, $p =$
576 0.620 ; Fig. S20; $\rho = 0.050$, $p = 0.657$; Fig. S21). Food treatment, rearing temperature,
577 heatwave exposure, and their interactions likewise had no effect on larval escape speed or
578 trajectory (Table 2).

579

580 *Gut bacteria diversity and composition*

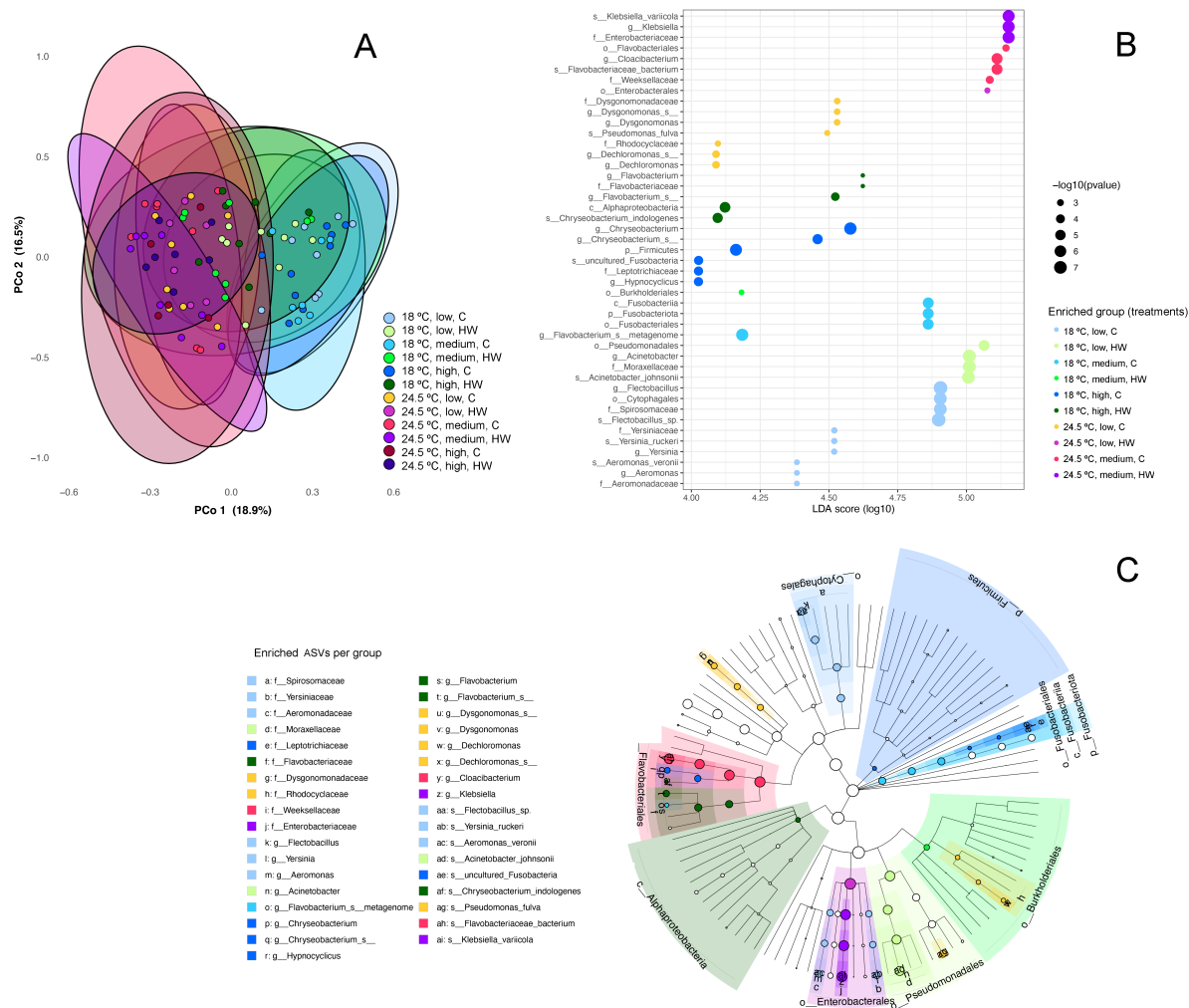
581

582 Gut bacterial diversity was influenced by rearing temperature and its interaction with
583 food treatment (Table 1). Both larvae fed medium-quality and high-quality food exhibited
584 reduced gut bacterial diversity when reared at 24.5 °C compared to 18 °C, with no difference
585 among temperatures for low-quality food (Fig. 4C, S22).

586 The two positive extraction controls (ZymoBIOMICS™ microbial community standard)
587 and the positive PCR control (ZymoBIOMICS™ microbial community DNA standard)
588 displayed identical species compositions but differed in the relative abundances of taxa
589 compared with the manufacturer’s expected profile (Fig. S23). The two extraction controls
590 yielded consistent results (Fig. S23), indicating that any deviations in relative abundances
591 were systematic rather than random.

592 In total, 207 Amplicon Sequence Variants (ASVs) were recovered from the gut
593 microbiomes of 92 *R. temporaria* larvae. The dominant phyla across treatments were
594 *Pseudomonadota* and *Bacteroidota* (Fig. S24). Most treatment pairs differed significantly in
595 gut bacterial community composition, with a few exceptions. No differences were detected
596 between medium-quality food with heatwave exposure and high-quality food without
597 exposure at 18 °C. At 24.5 °C, larvae fed low-quality food with heatwave exposure did not
598 differ from those fed medium-quality food (with or without heatwave exposure) or high-
599 quality food (with or without heatwave exposure) (Fig. 5A; Table S1).

600



601
 602 Fig. 5. Gut bacteria community composition (A) and enriched Amplicon Sequence Variants (ASVs; B)
 603 according to treatments imposed to larvae of *Rana temporaria*, corresponding to three diets with
 604 increasing levels of protein, fat, and components of animal origin (low-, medium-, and high-quality),
 605 two rearing temperatures (18 °C and 24.5 °C) and exposure or not to a heatwave (HW vs. C = control).
 606 Clustering of taxa with differences in abundance among treatments is also shown (C). Colors of ASVs
 607 correspond to colors of treatments in which they were the most abundant, cold colors (blue-green)
 608 correspond to 18 °C and warm colors (yellow-purple) to 24.5 °C rearing temperatures. Color intensity
 609 increases with food quality. Taxonomic levels are given before taxon names as species (s_), genus
 610 (g_), family (f_), order (o_), class (c_), and phylum (p_).
 611
 612

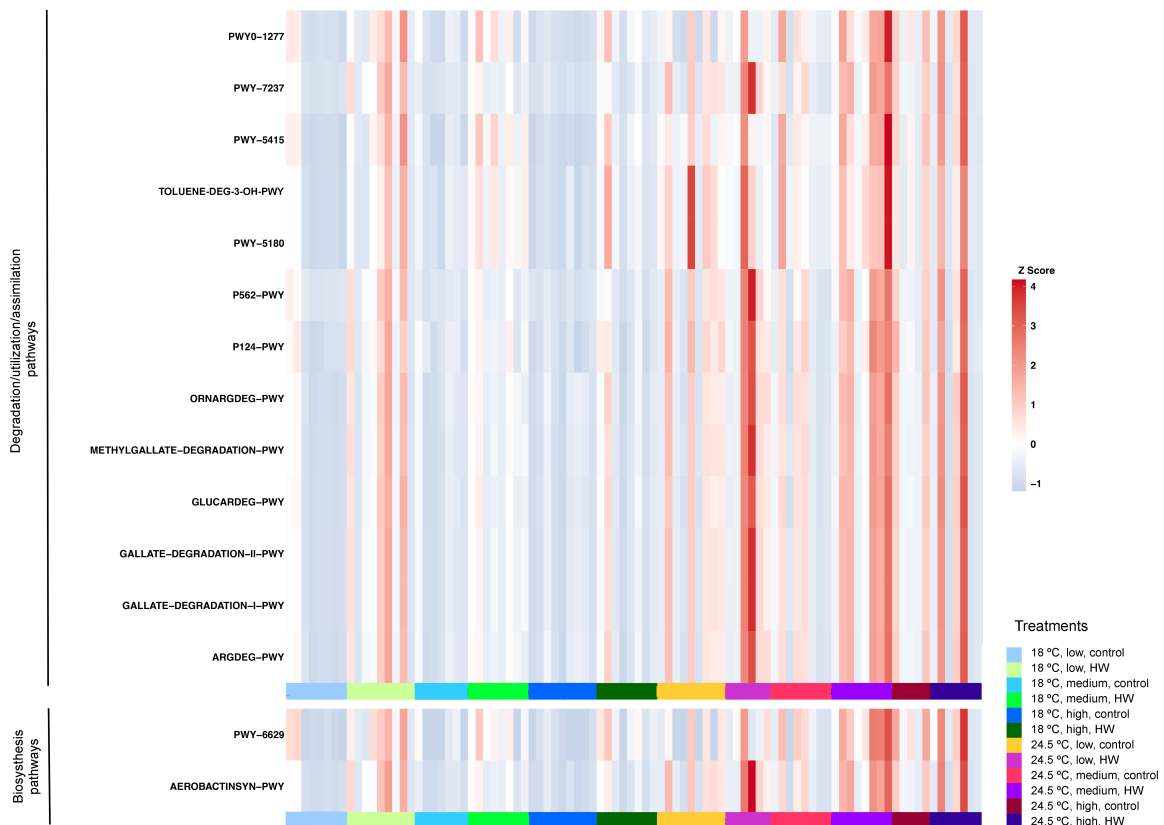
613 All treatment combinations except those involving high-quality food at 24.5 °C
 614 (regardless of heatwave exposure) had ASVs identified as indicators, totaling 45 ASVs (Fig.
 615 5). At low food quality, the main indicators identified through LefSe at 18 °C without
 616 heatwave exposure were *Flectobacillus* (*Spirosomaceae*, *Cytophagales*), *Yersinia ruckeri*
 617 (*Yersiniaceae*), and *Aeromonas veronii* (*Aeromonadaceae*). When exposed to a heatwave,
 618 *Acinetobacter johnsonii* (*Moraxellaceae*) was predominant. At 24.5 °C, *Dysgonomonas*
 619 (*Dysgonomonadaceae*), *Pseudomonas fulva*, and *Dechloromonas* (*Rhodocyclaceae*)
 620 dominated without heatwave exposure, whereas *Enterobacterales* predominated under
 621 heatwave exposure. The ALDEx2 analysis showed consistent results for all genera and
 622 families (results in the Supplementary Material).

623 At intermediate food quality, *Fusobacteriales* (*Fusobacteriia*, *Fusobacteriota*) and
 624 *Flavobacterium* were characteristic at 18 °C without heatwave exposure, while
 625 *Burkholderiales* dominated with heatwave exposure. At 24.5 °C, *Cloacibacterium*
 626 (*Weeksellaceae*, *Flavobacteriales*) predominated without heatwave exposure, whereas
 627 *Klebsiella variicola* (*Enterobacteriaceae*) was selected as an indicator under heatwave
 628 exposure.

629 At high food quality and 18 °C, *Chryseobacterium*, *Bacillota*, and *Hypnocyclicus*
 630 (*Leptotrichiaceae*) were indicators without heatwave exposure, and *Cryseobacterium*
 631 *indologenes*, *Flavobacterium* (*Flavobacteriaceae*), and *Alphaproteobacteria* predominated
 632 under heatwave exposure (Fig. 5).

633 In total, 357 unique metabolic pathways were predicted, of which 289 differed
 634 significantly among experimental treatments. The most significantly affected pathways
 635 included degradation of myo-inositol, D-glucarate, fructose, and various aromatic compounds
 636 (catechol, gallate, toluene, 3-phenylpropanoate, and 3-(3-hydroxyphenyl) propanoate), and
 637 conversion of amino acids into putrescine, as well as synthesis of L-tryptophan and
 638 aerobactin, (Fig. 6). The treatments with the highest expression of these pathways were those
 639 with larvae reared at 24.5 °C, exposed to a heatwave, or both (Fig. 6).

640



641 Fig. 6. The predicted 15 most significant pathways influenced by gut bacteria from larvae of *Rana*
 642 *temporaria* and their expression in treatments corresponding to a multifactorial experimental design of
 643 three diets with increasing levels of protein, fat, and components of animal origin (low-, medium-, and
 644 high-quality), two rearing temperatures (18 °C and 24.5 °C) and exposure or not to a heatwave (HW
 645

646 vs. C = control). Treatment colors are as in Fig. 5. Pathways follow the MetaCyc database (Caspi et
647 al., 2016). In the order of appearance: PWY0-1277 = pathway 3-phenylpropanoate and 3-(3-
648 hydroxyphenyl) propanoate degradation; PWY-7237 = pathway myo-, chiro- and scyllo-inositol
649 degradation; PWY-5415 = pathway catechol degradation I (meta-cleavage pathway); TOLUENE-
650 DEG-3-OH-PWY = toluene degradation II (aerobic) (via 4-methylcatechol); PWY-5180 = pathway
651 toluene degradation I (aerobic) (via o-cresol); P562-PWY = pathway myo-inositol degradation I;
652 P124-PWY = pathway *Bifidobacterium shunt*; ORNARGDEG-PWY = superpathway of L-arginine
653 and L-ornithine degradation; METHYLGALLATE-DEGRADATION-PWY = pathway methylgallate
654 degradation; GLUCARDEG-PWY = pathway D-glucarate degradation I; GALLATE-
655 DEGRADATION I and II-PWY = pathways gallate degradation I and II; ARGDEG-PWY =
656 superpathway of L-arginine, putrescine, and 4-aminobutanoate degradation; PWY-6629 =
657 superpathway of L-tryptophan biosynthesis; AEROBACTINSYN-PWY = pathway aerobactin
658 biosynthesis.

659

660 Discussion

661

662 The gut microbiome plays key roles in many aspects of animal biology, from nutrient
663 assimilation to immune defense and ultimately behavior (McFall-Ngai et al., 2013;
664 Tuddernham and Sears, 2015). Animals respond to environmental conditions and their gut
665 microorganisms are also expected to respond, potentially in ways that are adaptive and
666 enhance the animals' ability to cope with both natural environmental fluctuations (Park and
667 Do, 2024) and human-driven environmental challenges (Lynch and Hsiao, 2019; Fontaine and
668 Kohl, 2023). Under changing conditions, microbial taxa favored by the new environment may
669 increase in abundance and help maintain host metabolic functions, provided that the microbial
670 community has sufficient functional redundancy (Louca et al., 2018).

671 In our study, larvae of *R. temporaria* exposed to different temperatures and diets
672 exhibited shifts in gut bacterial diversity and composition, likely enabling them to maintain
673 body condition and still develop faster under higher temperatures. At elevated temperatures,
674 diet quality became a decisive factor for larval development and escape performance.

675

676 *Larvae nutrient assimilation, growth, and development*

677 Larvae of *Rana temporaria* exhibited clearly distinguishable stable isotope signatures
678 depending on food treatment, reflecting expected differences in nutrient acquisition from the
679 diets provided. However, body condition did not differ among food treatments, regardless of
680 rearing temperature. Development, on the other hand, was faster at 24.5 °C, and at this
681 temperature, higher food quality further increased developmental rate. Because temperature
682 determines the metabolic rate of ectotherms (Álvarez and Nicieza, 2002) and higher
683 metabolism requires more energy (Arendt, 1997), the improved food quality likely enabled *R.*
684 *temporaria* larvae to grow faster while maintaining good body condition.

685 The “macronutrient ratio hypothesis” predicts that ectotherms prefer increased
686 carbohydrate/protein ratios at higher temperatures to meet the energetic demands of elevated
687 metabolism, because excreting nitrogen from protein catabolism incurs a cost (Hardison and
688 Eliason, 2024). Similarly, the “temperature metabolic stoichiometry hypothesis” proposes that

689 ectotherms prefer diets with a higher carbon-to-nitrogen ratio under elevated temperatures
690 (Hardison and Eliason, 2024). Nitrogen excretion rates, however, usually increase with
691 temperature, reducing the cost of protein-rich diets (Hardison and Eliason, 2024). This
692 increase in nitrogen excretion may have allowed *R. temporaria* larvae to maintain body
693 condition and develop faster at higher temperatures when fed high-protein diets with
694 relatively constant carbohydrate content. A proportional increase in protein consumption at
695 higher temperatures has been observed in arthropods (Devries and Appel, 2014; Schmitz et
696 al., 2016), and invertebrates can maintain stable carbon-to-nitrogen ratios if food intake
697 increases with temperature (Anderson et al., 2017). In our study, this was likely the case
698 because larvae were fed *ad libitum*.

699 Niche breadth, estimated from stable isotope analyses, was higher at 18 °C with low to
700 intermediate food quality and at 24.5 °C with high food quality, but lower at 18 °C with high
701 food quality and at 24.5 °C with intermediate food quality. Because diet was uniform within
702 treatments, variation in niche breadth reflects individual differences in nutrient assimilation,
703 potentially mediated by the microbiome.

704 The microbiome, shaped by diet and host genetics, can influence nutrient absorption
705 and metabolism (Huda et al., 2022; Corbin et al., 2023). Thus, broader niche breadths may
706 indicate greater plasticity of the holobiont in adjusting nutrient assimilation at the individual
707 level. More diverse microbial communities often exhibit higher functional redundancy,
708 maintaining metabolic functions despite taxonomic shifts (Louca et al., 2018). In humans,
709 distinct host–microbial symbiotic states respond differently to diet (Arumugam et al., 2011),
710 suggesting that such plasticity can help meet host nutritional demands. As host genetic
711 variation was controlled across treatments (equal representation of five clutches), the larger
712 niche breadths observed likely reflect greater microbiome-mediated adaptive capacity,
713 potentially enhancing host performance.

714 If this hypothesis holds, higher efficiency in individual food assimilation could be
715 achieved at 24.5 °C when larvae consume high-quality food, as suggested by the observed
716 faster development without detriment to body condition or escape performance (discussed
717 below). However, in natural habitats, *ad libitum* access to the highest-quality food at elevated
718 temperatures may not be realistic. In such circumstances, herbivorous diets - which resulted in
719 broader niches than diets with intermediate animal components in our experiment - may
720 represent the best available solution. Therefore, dietary preferences toward herbivory under
721 heat stress could be subject to selection. In the wild, plant material has been associated with
722 higher nutritional value for fish at warmer temperatures and is thought to influence latitudinal
723 diversity gradients in herbivorous versus carnivorous fishes, with consumption of plant-based
724 food increasing with temperature (Behrens and Lafferty, 2007; González-Bergonzoni et al.,
725 2012). Choice experiments with ectotherms have similarly shown selection for more
726 herbivorous diets at higher temperatures (Vejříková et al., 2016; Zhang et al., 2020). Yet, in
727 some cases, herbivorous fish abundance did not increase with temperature in the southern
728 hemisphere (Trip et al., 2014), and grasshoppers increased preference for protein under higher
729 temperatures (Schmitz et al., 2016), indicating that increased plant consumption is not the
730 only strategy for coping with heat. The availability of suitable microorganisms to aid
731 digestion and assimilation of different nutrients, along with their own response to

732 temperature, may be critical for host success at varying temperatures and food qualities
733 (Vejříková et al., 2016).

734 Increasing temperatures can alter the diets of ectothermic animals by affecting both food
735 availability and quality or by triggering dietary shifts (Hardison and Eliason, 2024). For
736 instance, lipid content of algae decreases at higher temperatures (20-28 °C vs. 12 °C), which
737 also reduces the growth of *Daphnia* fed on them (Tseng et al., 2021). Altered temperatures
738 impose different nutrient demands, and species may adjust foraging behavior accordingly.
739 Thus, understanding the nutrients ectotherms can actually access in natural habitats is crucial
740 for interpreting laboratory results; otherwise, we risk overestimating their capacity to improve
741 performance based on animals kept in unrealistic conditions (Hardison and Eliason, 2024).

742
743 *Larvae escape behavior*

744
745 Larval ability to react was influenced by experimental conditions, whereas reaction
746 time, speed, and meander were not. At the higher rearing temperature (24.5 °C), not all diets
747 were sufficient to maintain an effective escape response in *R. temporaria* larvae. Diets with
748 high protein content and greater representation of animal-derived components, as well as an
749 herbivorous diet, resulted in efficient escape performance. Interestingly, the diet assumed to
750 be of lowest quality produced intermediate results in terms of larval reactivity, whereas larvae
751 receiving intermediate-quality food at 24.5 °C and not exposed to a heatwave exhibited the
752 poorest performance. These results align with observed patterns in larvae niche breadth,
753 suggesting a relationship between nutrient assimilation plasticity (i.e., broader isotopic
754 niches) and escape ability.

755 Using a combined visual, tactile, and chemical stimulus may have obscured cue-specific
756 differences in reaction time, as responses can vary by sensory modality (Melo et al., 2021).
757 We nevertheless combined cues to ensure stimulus detection by all larvae. Individuals that
758 failed to respond, or responded only upon contact, were considered less responsive, as
759 physical contact with a predator would likely result in capture in nature. *Rana temporaria*
760 larvae in the Kleiwiesen develop in small ponds with dragonfly naiads but without predatory
761 fish. High escape speed is adaptive against active predators such as fish, but less so against
762 ambush predators like Odonata (Teplitsky et al., 2005). Under these conditions, rapid threat
763 detection and initiation of escape likely have greater fitness consequences than speed or
764 escape trajectory (Staudinger et al., 2011).

765
766 *Gut bacteria, predicted metabolic pathways, and their potential influence on larvae*
767 *performance*

768
769 Variations in gut bacterial abundance and predicted metabolic pathways may have
770 contributed to differences in *R. temporaria* larvae performance under the experimental
771 conditions. Escape responses were markedly reduced in larvae reared at 24.5 °C with an
772 intermediate-quality diet and not exposed to a heatwave. In these larvae, *Cloacibacterium*
773 showed increased abundance. Interestingly, *Cloacibacterium* was also abundant in the control

774 group compared to elevated temperatures in rainbow trout (Zhou et al., 2022), although it
775 remains unclear whether this taxon contributed directly to the reduced reactivity in larvae.

776 In contrast, larvae exposed to a heatwave under the same dietary and rearing
777 temperature conditions showed improved escape performance and a higher abundance of
778 *Klebsiella* (*Enterobacteriaceae*, *Enterobacterales*). This suggests that the heatwave may have
779 triggered proliferation of *Klebsiella*, which in turn could have contributed to enhanced
780 performance. However, this shift in microbial composition came with additional reduction in
781 gut microbiome diversity, which was already low at medium quality food and 24.5 °C, and
782 may reduce host capacity to cope with additional stressors (Henry et al., 2021).

783 *Klebsiella* may influence host performance through multiple metabolic pathways.
784 Pathways such as P562-PWY and PWY-7237, involved in myo-inositol and related inositol
785 derivatives degradation (Berman and Magasanik, 1966a, 1966b; Anderson and Magasanik,
786 1971; Karp et al., 2019), were relatively increased in treatments with higher temperatures.
787 Myo-inositol is essential in eukaryotes for membrane phospholipids and cell signaling, and its
788 metabolism may help maintain membrane fluidity and protein activity - which are influenced
789 by temperature (Hazel, 1995) - under thermal stress. Additionally, *Klebsiella* may influence
790 behavior through neuromodulatory signals, as related species (*K. pneumoniae*) affect food
791 intake and attention in humans via serotonin and dopamine signaling (Miri et al., 2023). Other
792 increased pathways present in *Klebsiella*, such as GLUCARDEG-PWY (D-glucarate
793 degradation) and AEROBACTINSYN-PWY (aerobactin biosynthesis; Karp et al., 2019),
794 support bacterial growth by enabling carbon use and iron acquisition, which may indirectly
795 benefit host performance.

796 Other taxa also contributed to larvae performance under specific conditions. *Yersinia*
797 (*Yersiniaceae*, *Enterobacterales*) increased in abundance in larvae reared at low-quality food
798 and 18 °C under heatwave exposure, although performance did not differ from controls.
799 *Chryseobacterium*, associated with lipid absorption (Semova et al., 2012), predominated in
800 larvae fed high-quality food at 18 °C. In larvae fed high-quality food at 24.5 °C, no dominant
801 indicators were detected, yet these individuals developed the fastest and exhibited effective
802 escape responses, which could be linked to functional redundancy in the microbial
803 community.

804 Predicted metabolic pathways suggest that microbial plasticity may provide alternative
805 solutions for nutrient acquisition under different temperatures. For example, in larvae reared
806 at 24.5 °C with low-quality (herbivorous) diets, *Pseudomonas* and *Dysgonomonas* were
807 abundant in non-heatwave conditions, supporting aerobic aromatic catabolism pathways
808 (GALLATE-DEGRADATION-I-PWY, GALLATE-DEGRADATION-II-PWY,
809 METYLGALLATE-DEGRADATION) that enable degradation of plant lignin and tannins
810 (Karp et al., 2019). At the same time, increased abundance of *Enterobacterales* under
811 heatwave exposure likely allowed efficient carbon utilization and maintenance of membrane
812 function, supporting effective escape responses despite low-quality diets. In the fish
813 *Plectropomus leopardus* dominant gut bacterial taxa were shown to change within 12 h and
814 maintain estimated microbial functional capacity constant under different environmental
815 conditions (Mekuchi et al., 2018).

816 Protein absorption efficiency may decline with increasing temperature in ectotherms
817 (Croll and Watts, 2004). In fish, low-protein diets lead to gut microbiomes with altered
818 composition and reduced diversity, which are less efficient at absorbing protein—likely due to
819 the influence of specific bacterial strains on enterocyte protein uptake (Childers et al., 2025).
820 For instance, strains of *Acetivobacter*, *Aeromonas*, and *Pseudomonas* can reduce protein
821 absorption in the fish gut (Childers et al., 2025; Ye et al., 2019). Besides *Pseudomonas*,
822 *Dysgonomonas* may also be disadvantageous to the host at elevated temperatures. Members
823 of *Bacteroidales* (the order that includes *Dysgonomonas*) use putrescine to produce GABA
824 (gamma-aminobutyric acid), a molecule that modulates stress responsiveness in humans (Miri
825 et al., 2023). Thus, increased putrescine degradation may impair stress responses. In our
826 study, the superpathway of L-arginine, putrescine, and 4-aminobutanoate degradation
827 (ARGDEG-PWY) was activated in larvae reared at 24.5 °C on low-quality food and exposed
828 to a heatwave. However, these larvae also showed increased abundances of *Enterobacterales*
829 (the order that includes *Klebsiella*), which may have facilitated the degradation of diverse
830 carbon sources from the herbivorous diet and regulated membrane functions (as discussed
831 above), ultimately allowing larvae to maintain an effective escape response.

832 Some pathways also suggest potential benefits for coping with environmental
833 pollutants. PWY-5180 and TOLUENE-DEG-3-OH-PWY, corresponding to toluene
834 degradation, were associated with *Pseudomonas* (Fishman et al., 2004; Karp et al., 2019) and
835 may help larvae survive in polluted habitats. Additionally, PWY-6629, the L-tryptophan
836 biosynthesis pathway, increased under higher temperatures. In other ectotherms, dietary L-
837 tryptophan improves growth and thermic stress resistance (Akthar et al., 2013), suggesting
838 possible similar benefits mediated by the microbiome, although this pathway has only been
839 documented for *E. coli* due to limited ectotherm microbiome studies (Legrand et al., 2020;
840 Eterovick et al., 2024).

841 Overall, exposure to elevated temperatures - either long-term or as short-term
842 heatwaves - was associated with increases in the most significant metabolic pathways, though
843 not uniformly across treatments. This variability aligns with individual differences in
844 microbiome-host interactions and may underlie observed variation in larvae performance
845 under different environmental conditions.

846
847 *Concluding remarks*

848
849 At a temperature equivalent to that naturally experienced by *R. temporaria* (18 °C), food
850 quality - defined by high protein, fat, and animal component content - did not appear to be a
851 decisive factor for larval performance, including developmental rate and the ability to detect
852 and escape from threats. Under these conditions, food provided the necessary nutrients for
853 larvae performance and the gut bacterial community may have adjusted to variations in food
854 quality and exposure to short-term heat stress, maintaining nutrient assimilation for host
855 metabolic functions.

856 However, at elevated rearing temperatures, food quality became a key determinant of
857 developmental rate and interacted with additional temperature fluctuations, such as
858 heatwaves, to shape both the microbiome and behavioral outcomes. Larvae fed the diet richest

859 in protein, fat, and animal components developed the fastest and were among the most likely
860 to respond early to threats. Such traits would increase survival likelihood, allowing these
861 larvae to leave warming and potentially drying habitats quickly and to escape predators
862 efficiently. Interestingly, larvae fed an herbivorous diet - low in protein, fat, and component
863 diversity - also exhibited effective escape responses. The comparable outcomes of these
864 markedly different diets suggest that the different bacterial communities associated to them
865 may provide functional redundancy, supporting host performance.

866 Larvae receiving intermediate-quality diets, with moderate inclusion of animal
867 components, showed variable outcomes depending on heatwave exposure. This variability
868 indicates that a more herbivorous diet may represent a safer strategy in unpredictable
869 environments where high-quality animal food may not be consistently available. Temperature-
870 modulated microbial growth may further favor the consumption of specific food types, as
871 microbes play a key role in nutrient assimilation (Newsome et al., 2011; Vejříková et al.,
872 2016). Supporting this, studies across diverse ectotherms - from insects to vertebrates - have
873 often documented increased herbivory under elevated temperatures (Behrens and Lafferty,
874 2007; Carreira et al., 2016; Brankatschk et al., 2018; Zhang et al., 2020), although exceptions
875 exist (Trip et al., 2014; Schmitz et al., 2016). To better understand these patterns, future
876 research should investigate wild ectotherms' microbiomes, isotopic signatures, and health
877 biomarkers, linking diet composition, microbiome-mediated nutrient assimilation, and host
878 condition in natural habitats.

879 As human activities increase the intensity and frequency of environmental changes,
880 accelerating species extinction rates (IPCC, 2023), understanding the role of the microbiome
881 in animal resilience becomes increasingly important. Microbiomes are dynamic communities
882 (Louca et al., 2018) that respond to environmental fluctuations (Mekuchi et al., 2018).
883 Therefore, studies integrating multifactorial interactions among host, microbiome, and
884 environment, and collecting data from animals under natural conditions, are essential to
885 accurately interpret laboratory findings and predict ecological outcomes.

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References

- Akhtar, M. S., Pal, A. K., Sahu, N. P., Ciji, A., Meena, D. K., & Das, P. (2013). Physiological responses of dietary tryptophan fed *Labeo rohita* to temperature and salinity stress. *Journal of Animal Physiology and Animal Nutrition*, 97(6), 1075-1083. doi: 10.1111/jpn.12017
- Alberdi, A., & Gilbert, M. T. P. (2019). A guide to the application of Hill numbers to DNA-based diversity analyses. *Molecular Ecology Resources*, 19(4), 804-817. doi: 10.1111/1755-0998.13014
- Álvarez, D., & Nicieza, A. G. (2002). Effects of induced variation in anuran larval development on postmetamorphic energy reserves and locomotion. *Oecologia*, 131, 186-195. doi: 10.1007/s00442-002-0876-x
- Anderson, T. R., Hessen, D. O., Boersma, M., Urabe, J., & Mayor, D. J. (2017). Will invertebrates require increasingly carbon-rich food in a warming world?. *The American Naturalist*, 190(6), 725-742. doi: 10.1086/694122
- Anderson, W. A., & Magasanik, B. (1971). The pathway of myo-inositol degradation in *Aerobacter aerogenes*. Conversion of 2-deoxy-5-keto-D-gluconic acid to glycolytic intermediates. *The Journal of Biological Chemistry*, 246(18), 5662–5675. doi: 10.1016/S0021-9258(18)61857-5
- Archie, E. A., & Theis, K. R. (2011). Animal behaviour meets microbial ecology. *Animal Behaviour*, 82(3), 425-436. doi: 10.1016/j.anbehav.2011.05.029
- Arendt, J. D. (1997). Adaptive intrinsic growth rates: an integration across taxa. *The Quarterly Review of Biology*, 72(2), 149-177. doi:10.1086/419764
- Arumugam, M., Raes, J., Pelletier, E., Le Paslier, D., Yamada, T., Mende, D. R., ... & Bork, P. (2011). Enterotypes of the human gut microbiome. *Nature*, 473(7346), 174-180. doi: 10.1038/nature09944
- Bastiani, V. I. M. (2023). Anuran amphibians in highland grasslands in southern Brazil: Effects of habitat, degradation, sensitivity and thermal tolerance. PhD Dissertation, Universidade Federal de Santa Maria, Brazil.
- Behrens, M. D., & Lafferty, K. D. (2007). Temperature and diet effects on omnivorous fish performance: implications for the latitudinal diversity gradient in herbivorous fishes. *Canadian Journal of Fisheries and Aquatic Sciences*, 64(6), 867-873. doi: 10.1139/f07-063
- Bercik, P., Collins, S. M., & Verdu, E. F. (2012). Microbes and the gut-brain axis. *Neurogastroenterology & Motility*, 24(5), 405-413. doi: 10.1111/j.1365-2982.2012.01906.x
- Berg, G., Rybakova, D., Fischer, D., Cernava, T., Vergès, M. C. C., Charles, T., ... & Schloter, M. (2020). Microbiome definition re-visited: old concepts and new challenges. *Microbiome*, 8, 1-22. doi: 10.1186/s40168-020-00875-0
- Berman, T., & Magasanik, B. (1966a). The pathway of myo-inositol degradation in *Aerobacter aerogenes*. Dehydrogenation and dehydration. *The Journal of Biological Chemistry*, 241(4), 800–806. doi: 10.1016/S0021-9258(18)96836-5
- Berman, T., & Magasanik, B. (1966b). The pathway of myo-inositol degradation in *Aerobacter aerogenes*. Ring scission. *The Journal of Biological Chemistry*, 241(4), 807–813. doi: 10.1016/S0021-9258(18)96837-7

934 Bolyen, E., Rideout, J. R., Dillon, M. R., Bokulich, N. A., Abnet, C. C., Al-Ghalith, G. A., ...
935 & Caporaso, J. G. (2019). Reproducible, interactive, scalable and extensible microbiome
936 data science using QIIME 2. *Nature Biotechnology*, 37(8), 852-857. doi: 10.1038/s41587-
937 019-0209-9

938 Borzée, A., Prasad, V. K., Neam, K., Tarrant, J., Kosch, T. A., Barata, I. M., ... & Wren, S.
939 (2025). Conservation priorities for global amphibian biodiversity. *Nature Reviews*
940 *Biodiversity*. doi: 10.1038/s44358-025-00101-5

941 Brankatschk, M., Gutmann, T., Knittelfelder, O., Palladini, A., Prince, E., Grzybek, M., ... &
942 Eaton, S. (2018). A temperature-dependent switch in feeding preference improves
943 *Drosophila* development and survival in the cold. *Developmental Cell*, 46(6), 781-793.
944 doi: 10.1016/j.devcel.2018.05.028

945 Cao, Y., Dong, Q., Wang, D., Zhang, P., Liu, Y., & Niu, C. (2022). microbiomeMarker: an
946 R/Bioconductor package for microbiome marker identification and visualization.
947 *Bioinformatics*, 38(16), 4027-4029. doi: 10.1093/bioinformatics/btac438

948 Caporaso, J. G., Lauber, C. L., Walters, W. A., Berg-Lyons, D., Lozupone, C. A., Turnbaugh,
949 P. J., ... & Knight, R. (2011). Global patterns of 16S rRNA diversity at a depth of millions
950 of sequences per sample. *Proceedings of the National Academy of Sciences*,
951 108(supplement_1), 4516-4522. doi: 10.1073/pnas.1000080107

952 Carreira, B. M., Segurado, P., Orizaola, G., Gonçalves, N., Pinto, V., Laurila, A., & Rebelo, R.
953 (2016). Warm vegetarians? Heat waves and diet shifts in tadpoles. *Ecology*, 97(11), 2964-
954 2974. doi: 10.1002/ecy.1541

955 Caspi, R., Billington, R., Ferrer, L., Foerster, H., Fulcher, C. A., Keseler, I. M., ... & Karp, P.
956 D. (2016). The MetaCyc database of metabolic pathways and enzymes and the BioCyc
957 collection of pathway/genome databases. *Nucleic acids research*, 44(D1), D471-D480.
958 doi: 10.1093/nar/gkv1164

959 Chen, X., D'Souza, R., & Hong, S. T. (2013). The role of gut microbiota in the gut-brain axis:
960 current challenges and perspectives. *Protein & Cell*, 4, 403-414. doi: 10.1007/s13238-
961 013-3017-x

962 Chiara, V., & Kim, S. Y. (2023). AnimalTA: A highly flexible and easy-to-use program for
963 tracking and analysing animal movement in different environments. *Methods in Ecology*
964 *and Evolution* 14: 1699–1707. doi: 10.1111/2041-210X.14115

965 Childers, L., Park, J., Wang, S., Liu, R., Barry, R., Watts, S. A., ... & Bagnat, M. (2025).
966 Protein absorption in the zebrafish gut is regulated by interactions between lysosome rich
967 enterocytes and the microbiome. *eLife* 13:RP100611. doi: 10.7554/eLife.100611

968 Collins, J. P., & Storfer, A. (2003). Global amphibian declines: sorting the hypotheses.
969 *Diversity and Distributions*, 9(2), 89-98. doi: 10.1046/j.1472-4642.2003.00012.x

970 Corbin, K. D., Carnero, E. A., Dirks, B., Igudesman, D., Yi, F., Marcus, A., ... & Smith, S. R.
971 (2023). Host-diet-gut microbiome interactions influence human energy balance: a
972 randomized clinical trial. *Nature Communications*, 14(1), 3161. doi: 10.1038/s41467-
973 023-38778-x

974 Croll, S. L., & Watts, S. A. (2004). The effect of temperature on feed consumption and
975 nutrient absorption in *Procambarus clarkii* and *Procambarus zonangulus*. *Journal of the*
976 *World Aquaculture Society*, 35(4), 478-488. doi: 10.1111/j.1749-7345.2004.tb00113.x

977 Dallas, J. W., Kazarina, A., Lee, S. T., & Warne, R. W. (2024). Cross-species gut microbiota
978 transplantation predictably affects host heat tolerance. *Journal of Experimental Biology*,

979 227(1), jeb246735. doi: 10.1242/jeb.246735

980 Devries, Z. C. & Appel, A. G. (2014). Effects of temperature on nutrient self-selection in the
981 silverfish *Lepisma saccharina*. *Physiological Entomology* 39, 217–221. doi:
982 10.1111/phen.12064

983 Dinan, T. G., Stilling, R. M., Stanton, C., & Cryan, J. F. (2015). Collective unconscious: how
984 gut microbes shape human behavior. *Journal of Psychiatric Research*, 63, 1-9. doi:
985 10.1016/j.jpsychires.2015.02.021

986 Dittrich, C., Rodríguez, A., Segev, O., Drakulić, S., Feldhaar, H., Vences, M., & Rödel, M. O.
987 (2018). Temporal migration patterns and mating tactics influence size-assortative mating
988 in *Rana temporaria*. *Behavioral Ecology*, 29(2), 418-428. doi:10.1093/beheco/ax188

989 Douglas, G. M., Maffei, V. J., Zaneveld, J. R., Yurgel, S. N., Brown, J. R., Taylor, C. M., ... &
990 Langille, M. G. (2020). PICRUSt2 for prediction of metagenome functions. *Nature*
991 *Biotechnology*, 38(6), 685-688. doi: 10.1038/s41587-020-0548-6

992 Douglas, A. E., & Werren, J. H. (2016). Holes in the hologenome: why host-microbe
993 symbioses are not holobionts. *MBio*, 7(2), 10-1128. doi:10.1128/mBio.02099-15

994 Duarte, H., Tejedo, M., Katzenberger, M., Marangoni, F., Baldo, D., Beltrán, J. F., ... &
995 Gonzalez-Voyer, A. (2012). Can amphibians take the heat? Vulnerability to climate
996 warming in subtropical and temperate larval amphibian communities. *Global Change*
997 *Biology*, 18(2), 412-421. doi: 10.1111/j.1365-2486.2011.02518.x

998 Eterovick, P. C., Mendes, I. S., Kloh, J. S., Pinheiro, L. T., Václav, A. B. H. P., Santos, T., &
999 Gontijo, A. S. B. (2018). Tadpoles respond to background colour under threat. *Scientific*
1000 *Reports*, 8(1), 4085. doi: 10.1038/s41598-018-22315-8

1001 Eterovick, P. C., Kloh, J. S., Figueredo, C. C., Viana, P. I. M., Goulart, M., Milan, D. T., ... &
1002 Vences, M. (2020). Background choice and immobility as context dependent tadpole
1003 responses to perceived predation risk. *Scientific Reports*, 10(1), 13577. doi:
1004 10.1038/s41598-020-70274-w

1005 Eterovick, P. C., Schmidt, R., Sabino-Pinto, J., Yang, C., Künzel, S., & Ruthsatz, K. (2024).
1006 The microbiome at the interface between environmental stress and animal health: an
1007 example from the most threatened vertebrate group. *Proceedings of the Royal Society B*,
1008 291(2031), 20240917. doi:10.1038/s41598-018-22315-8

1009 Fernandes, A. D., Reid, J. N., Macklaim, J. M., McMurrough, T. A., Edgell, D. R., & Gloor,
1010 G. B. (2014). Unifying the analysis of high-throughput sequencing datasets:
1011 characterizing RNA-seq, 16S rRNA gene sequencing and selective growth experiments
1012 by compositional data analysis. *Microbiome*, 2(1), 15. doi: 10.1186/2049-2618-2-15

1013 Fishman, A., Tao, Y., & Wood, T. K. (2004). Toluene 3-monooxygenase of *Ralstonia pickettii*
1014 PKO1 is a para-hydroxylating enzyme. *Journal of Bacteriology*, 186(10), 3117-3123. doi:
1015 10.1128/jb.186.10.3117-3123.2004

1016 Florkowski, M. R., & Yorzinski, J. L. (2023). Gut microbiome diversity and composition is
1017 associated with exploratory behavior in a wild-caught songbird. *Animal Microbiome*,
1018 5(1), 8. doi: 10.1186/s42523-023-00227-x

1019 Fontaine, S. S., & Kohl, K. D. (2023). The microbiome buffers tadpole hosts from heat stress:
1020 a hologenomic approach to understand host–microbe interactions under warming. *Journal*
1021 *of Experimental Biology*, 226(1), jeb245191. doi:10.1242/jeb.245191

1022 Fontaine, S. S., Mineo, P. M., & Kohl, K. D. (2022). Experimental manipulation of microbiota
1023 reduces host thermal tolerance and fitness under heat stress in a vertebrate ectotherm.

1024 Nature Ecology & Evolution, 6(4), 405-417. doi: 10.1038/s41559-022-01686-2

1025 Glos, J., Ruthsatz, K., Schröder, D., & Riemann, J. C. (2020). Food source determines stable
1026 isotope discrimination factors ΔN and ΔC in tadpoles. *Amphibia-Reptilia*, 41(4): 501-
1027 507. doi:10.1163/15685381-bja10020

1028 González-Bergonzoni, I., Meerhoff, M., Davidson, T. A., Teixeira-de Mello, F., Baattrup-
1029 Pedersen, A., & Jeppesen, E. (2012). Meta-analysis shows a consistent and strong
1030 latitudinal pattern in fish omnivory across ecosystems. *Ecosystems*, 15, 492-503. doi:
1031 10.1007/s10021-012-9524-4

1032 Gosner, K.L., 1960. A simplified table for staging anuran embryos and larvae with notes on
1033 identification. *Herpetologica* 16: 183–190.

1034 Guo, J., Li, Z., Liu, X., Jin, Y., Sun, Y., Yuan, Z., ... & Zhang, M. (2024). Response of the gut
1035 microbiota to changes in the nutritional status of red deer during winter. *Scientific*
1036 *Reports*, 14(1), 24961. doi: 10.1038/s41598-024-76142-1

1037 Hardison, E. A., & Eliason, E. J. (2024). Diet effects on ectotherm thermal performance.
1038 *Biological Reviews*, 99(4), 1537-1555. doi: 10.1111/brv.13081

1039 Hayes, T. B., Falso, P., Gallipeau, S., & Stice, M. (2010). The cause of global amphibian
1040 declines: a developmental endocrinologist's perspective. *Journal of Experimental*
1041 *Biology*, 213(6), 921-933. doi: 10.1242/jeb.040865

1042 Hazel, J. R. (1995). Thermal adaptation in biological membranes: is homeoviscous adaptation
1043 the explanation?. *Annual Review of Physiology*, 57(1), 19-42.

1044 Hébert, M., Versace, E., & Vallortigara, G. (2019). Inexperienced preys know when to flee or
1045 to freeze in front of a threat. *Proceedings of the National Academy of Sciences*, 116(46),
1046 22918-22920. doi: 10.1073/pnas.191550411

1047 Henry, L. P., Bruijning, M., Forsberg, S. K., & Ayroles, J. F. (2021). The microbiome extends
1048 host evolutionary potential. *Nature Communications*, 12(1), 5141. doi: 10.1038/s41467-
1049 021-25315-x

1050 Hill, M. O. (1973). Diversity and evenness: a unifying notation and its consequences.
1051 *Ecology*, 54(2), 427-432. doi: 10.2307/1934352

1052 Hird, S. M. (2017). Evolutionary biology needs wild microbiomes. *Front Microbiol* 8: 725.
1053 doi: 10.3389/fmicb.2017.00725

1054 Huda, M. N., Salvador, A. C., Barrington, W. T., Gacasan, C. A., D'Souza, E. M., Deus
1055 Ramirez, L., ... & Bennett, B. J. (2022). Gut microbiota and host genetics modulate the
1056 effect of diverse diet patterns on metabolic health. *Frontiers in Nutrition*, 9, 896348. doi:
1057 10.3389/fnut.2022.896348

1058 IPCC (2023). Summary for Policymakers. In: *Climate Change 2023: Synthesis Report.*
1059 *Contribution of Working Groups I, II and III to the Sixth Assessment Report of the*
1060 *Intergovernmental Panel on Climate Change* [Core Writing Team, H. Lee and J. Romero
1061 (eds.)]. IPCC, Geneva, Switzerland, pp. 1-34. doi: 10.59327/IPCC/AR6-
1062 9789291691647.001

1063 Karp, P. D., Billington, R., Caspi, R., Fulcher, C. A., Latendresse, M., Kothari, A., Keseler, I.
1064 M., Krummenacker, M., Midford, P. E., Ong, Q., Ong, W. K., Paley, S. M., & Subhraveti,
1065 P. (2019). The BioCyc collection of microbial genomes and metabolic pathways.
1066 *Briefings in Bioinformatics*, 20(4), 1085–1093. doi: 10.1093/bib/bbx085

1067 Kloh, J. S., Figueredo, C. C., Calaça, P., & Eterovick, P. C. (2024). Pollen as food: effects of
1068 consumption on tadpole growth, development, and mobility. *Hydrobiologia*, 851(8),

1069 2071-2080. doi: 10.1007/s10750-023-05439-5

1070 Kupferberg, S. J. (1997). The role of larval diet in anuran metamorphosis. *American Zoologist*

1071 37, 146–159.

1072 Legrand, T. P., Wynne, J. W., Weyrich, L. S., & Oxley, A. P. (2020). A microbial sea of

1073 possibilities: current knowledge and prospects for an improved understanding of the fish

1074 microbiome. *Reviews in Aquaculture*, 12(2), 1101-1134. doi: 10.1111/raq.12375

1075 Lenth, R. (2017). emmeans: Estimated Marginal Means, aka Least-Squares Means. R package

1076 version 0.9.1. <https://CRAN.R-project.org/package=emmeans>

1077 Llobat, L., & Marín-García, P. J. (2022). Application of protein nutrition in natural ecosystem

1078 management for European rabbit (*Oryctolagus cuniculus*) conservation. *Biodiversity*

1079 and Conservation, 31(5), 1435-1444. doi: 10.1007/s10531-022-02426-5

1080 Louca, S., Polz, M. F., Mazel, F., Albright, M. B., Huber, J. A., O'Connor, M. I., ... & Parfrey,

1081 L. W. (2018). Function and functional redundancy in microbial systems. *Nature Ecology*

1082 & Evolution, 2(6), 936-943. doi: 10.1038/s41559-018-0519-1

1083 Lozupone, C., & Knight, R. (2005). UniFrac: a new phylogenetic method for comparing

1084 microbial communities. *Applied and environmental microbiology*, 71(12), 8228-8235.

1085 doi:10.1128/AEM.71.12.8228–8235.2005

1086 Luedtke, J. A., Chanson, J., Neam, K., Hobin, L., Maciel, A. O., Catenazzi, A., ... & Stuart, S.

1087 N. (2023). Ongoing declines for the world's amphibians in the face of emerging threats.

1088 *Nature*, 622(7982), 308–314. <https://doi.org/10.1038/s41586-023-06578-4>

1089 Lynch, J. B., & Hsiao, E. Y. (2019). Microbiomes as sources of emergent host phenotypes.

1090 *Science*, 365(6460), 1405-1409. *Science* 365: 1405–1409. doi: 10.1126/science.aay0240

1091 Manning, D. W., & Sullivan, S. M. P. (2021). Conservation across aquatic-terrestrial

1092 boundaries: Linking continental-scale water quality to emergent aquatic insects and

1093 declining aerial insectivorous birds. *Frontiers in Ecology and Evolution*, 9, 633160. doi:

1094 10.3389/fevo.2021.633160

1095 McCallum, M. L. 2015. Vertebrate biodiversity losses point to a sixth mass extinction.

1096 *Biodiversity and Conservation* 24, 2497–2519. doi: 10.1007/s10531-015-0940-6

1097 McDiarmid, R. W., & Altig, R. (Eds.). (1999). *Tadpoles: the biology of anuran larvae*.

1098 University of Chicago Press.

1099 McFall-Ngai, M., Hadfield, M. G., Bosch, T. C., Carey, H. V., Domazet-Lošo, T., Douglas, A.

1100 E., ... & Wernegreen, J. J. (2013). Animals in a bacterial world, a new imperative for the

1101 life sciences. *Proceedings of the National Academy of Sciences*, 110(9), 3229-3236. doi:

1102 10.1073/pnas.1218525110

1103 McMurdie, P. J., & Holmes, S. (2013). phyloseq: an R package for reproducible interactive

1104 analysis and graphics of microbiome census data. *PloS one*, 8(4), e61217. doi:

1105 10.1371/journal.pone.0061217

1106 Mekuchi, M., Asakura, T., Sakata, K., Yamaguchi, T., Teruya, K., & Kikuchi, J. (2018).

1107 Intestinal microbiota composition is altered according to nutritional biorhythms in the

1108 leopard coral grouper (*Plectropomus leopardus*). *PloS one*, 13(6), e0197256. doi:

1109 10.1371/journal.pone.0197256

1110 Melo, G. R., Solé, M., & Eterovick, P. C. (2021). Invisible or fearless: tadpole response to

1111 predator cues depends on color. *Ethology Ecology & Evolution*, 33(2), 99-107. doi:

1112 10.1080/03949370.2020.1830859

1113 Miri, S., Yeo, J., Abubaker, S., & Hammami, R. (2023). Neuromicrobiology, an emerging
1114 neurometabolic facet of the gut microbiome?. *Frontiers in Microbiology*, 14, 1098412.
1115 doi: 10.3389/fmicb.2023.1098412

1116 Moura, S., Kloh, J. S., Figueredo, C. C., & Eterovick, P. C. (2023). An empty stomach is not a
1117 good adviser: avoiding toxic Cyanobacteria can compromise tadpole antipredator
1118 defenses. *Amphibia-Reptilia*, 44(4), 457-465. doi: 10.1163/15685381-bja10153

1119 Newsome, S. D., Fogel, M. L., Kelly, L., & del Rio, C. M. (2011). Contributions of direct
1120 incorporation from diet and microbial amino acids to protein synthesis in Nile tilapia.
1121 *Functional Ecology*, 25(5), 1051-1062. doi: 10.1111/j.1365-2435.2011.01866.x

1122 Oksanen, J., Blanchet, F. G., Kindt, R., Legendre, P., Minchin, P. R., O'hara, R. B., ... &
1123 Wagner, H. (2013). Community ecology package. R package version, 2(0), 321-326.

1124 Park, J. K., & Do, Y. (2024). Combined effect of seasons and life history in an anuran
1125 strengthens the response and relationship between their physiology and gut microbiota.
1126 *Scientific Reports*, 14(1), 10137. doi: 10.1038/s41598-024-60105-7

1127 Peig, J., & Green, A. J. (2009). New perspectives for estimating body condition from
1128 mass/length data: the scaled mass index as an alternative method. *Oikos*, 118(12), 1883-
1129 1891. doi:10.1111/j.1600-0706.2009.17643.x

1130 Peig, J., & Green, A. J. (2010). The paradigm of body condition: a critical reappraisal of
1131 current methods based on mass and length. *Functional Ecology*, 24(6), 1323-1332.
1132 doi:10.1111/j.1365-2435.2010.01751.x

1133 Perry, W. B., Lindsay, E., Payne, C. J., Brodie, C., & Kazlauskaitė, R. (2020). The role of the
1134 gut microbiome in sustainable teleost aquaculture. *Proceedings of the Royal Society B*,
1135 287(1926), 20200184. doi: 10.1098/rspb.2020.0184

1136 Preston, D. B., & Forstner, M. R. (2015). Houston Toad (*Bufo (Anaxyrus) houstonensis*)
1137 tadpoles decrease their activity in response to chemical cues produced from the predation
1138 of conspecifics and congeneric (*Bufo (Incilius) nebulifer*) tadpoles. *Journal of*
1139 *Herpetology*, 49(2), 170-175. doi: 10.1670/13-059

1140 R Core Team. 2024. R: A language and Environment for Statistical Computing. R Foundation
1141 for Statistical Computing. Version 4.4.2, Vienna, Austria. <https://cran.r-project.org/>

1142 Relyea, R. A. (2001). Morphological and behavioral plasticity of larval anurans in response to
1143 different predators. *Ecology*, 82(2), 523-540. doi: 10.1890/0012-
1144 9658(2001)082[0523:MABPOL]2.0.CO;2

1145 Ruddiman, W. F. (2013). The anthropocene. *Annual Review of Earth and Planetary Sciences*,
1146 41(1), 45-68. doi: 10.1146/annurev-earth-050212-123944

1147 Ruthsatz, K., Dausmann, K. H., Paesler, K., Babos, P., Sabatino, N. M., Peck, M. A., & Glos,
1148 J. (2020). Shifts in sensitivity of amphibian metamorphosis to endocrine disruption: the
1149 common frog (*Rana temporaria*) as a case study. *Conservation Physiology*, 8(1),
1150 coaa100. doi:10.1093/conphys/coaa100

1151 Ruthsatz, K., Giertz, L. M., Schröder, D., & Glos, J. (2019). Chemical composition of food
1152 induces plasticity in digestive morphology in larvae of *Rana temporaria*. *Biology Open*,
1153 8(12), bio048041. doi: 10.1242/bio.048041

1154 Sampson, T. R., & Mazmanian, S. K. (2015). Control of brain development, function, and
1155 behavior by the microbiome. *Cell Host & Microbe*, 17(5), 565-576. doi:
1156 10.1016/j.chom.2015.04.011

1157 Schmitz, O. J., Rosenblatt, A. E. & Smylie, M. (2016). Temperature dependence of predation
1158 stress and the nutritional ecology of a generalist herbivore. *Ecology* 97, 3119–3130. doi:
1159 10.1002/ecy.1524

1160 Segata, N., Izard, J., Waldron, L., Gevers, D., Miropolsky, L., Garrett, W. S., & Huttenhower,
1161 C. (2011). Metagenomic biomarker discovery and explanation. *Genome Biology*, 12, 1-
1162 18. doi: 10.1186/gb-2011-12-6-r60

1163 Seifert, L. I., de Castro, F., Marquart, A., Gaedke, U., Weithoff, G., & Vos, M. (2014). Heated
1164 relations: temperature-mediated shifts in consumption across trophic levels. *PLoS One*,
1165 9(5), e95046. doi:10.1371/journal.pone.0095046

1166 Semova, I., Carten, J. D., Stombaugh, J., Mackey, L. C., Knight, R., Farber, S. A., & Rawls, J.
1167 F. (2012). Microbiota regulate intestinal absorption and metabolism of fatty acids in the
1168 zebrafish. *Cell Host & Microbe*, 12(3), 277-288. doi: 10.1016/j.chom.2012.08.003

1169 Sharon, G., Segal, D., Ringo, J. M., Hefetz, A., Zilber-Rosenberg, I., & Rosenberg, E. (2010).
1170 Commensal bacteria play a role in mating preference of *Drosophila melanogaster*.
1171 *Proceedings of the National Academy of Sciences*, 107(46), 20051-20056. doi:
1172 10.1073/pnas.1009906107

1173 Silva, Y. P., Bernardi, A., & Frozza, R. L. (2020). The role of short-chain fatty acids from gut
1174 microbiota in gut-brain communication. *Frontiers in Endocrinology*, 11, 508738. doi:
1175 10.3389/fendo.2020.00025

1176 Singmann H, Kellen D. (2019). An introduction to mixed models for experimental
1177 psychology. In *New methods in cognitive psychology* (eds DH Spieler, E Schumacher).
1178 Hove, UK: Psychology Press. doi:10.4324/9780429318405-2

1179 Singmann H, Bolker B, Westfall J, Aust F, Ben-Shachar M (2024). `afex`: Analysis of
1180 Factorial Experiments. doi: 10.32614/CRAN.package.afex
1181 <<https://doi.org/10.32614/CRAN.package.afex>>, R package version 1.4-1,
1182 <https://CRAN.R-project.org/package=afex>

1183 Staniek, M. A., Pansch, C., Shama, L. N., Mehler, K., Steinmann, A., Middelburg, J. J., &
1184 Meysick, L. (2025). Heatwave intensity drives eco-physiological responses in infaunal
1185 bivalves: a mesocosm experiment. *Limnology and Oceanography*, 70, S417-S431. doi:
1186 10.1002/lno.70012

1187 Staudinger, M. D., Hanlon, R. T., & Juanes, F. (2011). Primary and secondary defences of
1188 squid to cruising and ambush fish predators: variable tactics and their survival value.
1189 *Animal Behaviour*, 81(3), 585-594. doi:10.1016/j.anbehav.2010.12.002

1190 Swanson, H. K., Lysy, M., Power, M., Stasko, A. D., Johnson, J. D., & Reist, J. D. (2015). A
1191 new probabilistic method for quantifying n-dimensional ecological niches and niche
1192 overlap. *Ecology*, 96(2), 318-324. doi: 10.1890/14-0235.1

1193 Teplitsky, C., Plénet, S., Léna, J. P., Mermet, N., Malet, E., & Joly, P. (2005). Escape
1194 behaviour and ultimate causes of specific induced defences in an anuran tadpole.
1195 *Journal of Evolutionary Biology*, 18(1), 180-190. doi: 10.1111/j.1420-
1196 9101.2004.00790.x

1197 Trip, E. D. L., Clements, K. D., Raubenheimer, D., & Choat, J. H. (2014). Temperature-
1198 related variation in growth rate, size, maturation and life span in a marine herbivorous
1199 fish over a latitudinal gradient. *Journal of Animal Ecology*, 83(4), 866-875. doi:
1200 10.1111/1365-2656.12183

1201 Tseng, M., Di Filippo, C. M., Fung, M., Kim, J. O., Forster, I. P., & Zhou, Y. (2021).
1202 Cascading effects of algal warming in a freshwater community. *Functional Ecology*,
1203 35(4), 920-929. doi: 10.1111/1365-2435.13752

1204 Tuddenham, S., & Sears, C. L. (2015). The intestinal microbiome and health. *Current Opinion*
1205 *in Infectious Diseases*, 28(5), 464-470. doi: 10.1097/QCO.0000000000000196

1206 Vejřiková, I., Vejřík, L., Syvänta, J., Kiljunen, M., Čech, M., Blabolil, P., ... & Peterka, J.
1207 (2016). Distribution of herbivorous fish is frozen by low temperature. *Scientific*
1208 *Reports*, 6(1), 39600. doi: 10.1038/srep39600

1209 Videvall, E., Burraco, P., & Orizaola, G. (2023). Impact of ionizing radiation on the
1210 environmental microbiomes of Chernobyl wetlands. *Environmental Pollution*, 330,
1211 121774. doi: 10.1016/j.envpol.2023.121774

1212 Woolstra, C. R., & Ziegler, M. (2020). Adapting with microbial help: microbiome flexibility
1213 facilitates rapid responses to environmental change. *BioEssays*, 42(7), 2000004. doi:
1214 10.1002/bies.202000004

1215 Wake, D. B., & Vredenburg, V. T. (2008). Are we in the midst of the sixth mass extinction? A
1216 view from the world of amphibians. *Proceedings of the National Academy of Sciences*,
1217 105(supplement_1), 11466-11473. doi: 10.1073/pnas.0801921105

1218 Wang, W., Zhou, R., He, L., Liu, S., Zhou, J., Qi, L., ... & Hu, D. (2015). The progress in
1219 nutrition research of musk deer: Implication for conservation. *Applied Animal*
1220 *Behaviour Science*, 172, 1-8. doi: 10.1016/j.applanim.2015.09.006

1221 Wassersug, R. J. (1989). Locomotion in amphibian larvae (or "Why aren't tadpoles built like
1222 fishes?"). *American Zoologist*, 65-84.

1223 Wells, K. D. (2019). *The ecology and behavior of amphibians*. University of Chicago press.

1224 Wong, A. C. N., Holmes, A., Ponton, F., Lihoreau, M., Wilson, K., Raubenheimer, D., &
1225 Simpson, S. J. (2015). Behavioral microbiomics: a multi-dimensional approach to
1226 microbial influence on behavior. *Frontiers in Microbiology*, 6, 1359. doi:
1227 10.3389/fmicb.2015.01359

1228 Xiao, J., & Wang, W. X. (2025). Complex and lasting impacts of heatwaves on life-history
1229 traits and fitness in *Daphnia magna*. *Journal of Experimental Biology*, 228(18),
1230 jeb250837. doi:10.1242/jeb.250837

1231 Yan, K., Guo, F., Kainz, M. J., Li, F., Gao, W., Bunn, S. E., & Zhang, Y. (2024). The
1232 importance of omega-3 polyunsaturated fatty acids as high-quality food in freshwater
1233 ecosystems with implications of global change. *Biological Reviews*, 99(1), 200-218.
1234 doi: 10.1111/brv.13017

1235 Yang, C., Mai, J., Cao, X., Burberry, A., Cominelli, F., & Zhang, L. (2023). ggpicrust2: an R
1236 package for PICRUST2 predicted functional profile analysis and visualization.
1237 *Bioinformatics*, 39(8), btad470. doi:10.1093/bioinformatics/btad470

1238 Ye, L., Mueller, O., Bagwell, J., Bagnat, M., Liddle, R. A., & Rawls, J. F. (2019). High fat diet
1239 induces microbiota-dependent silencing of enteroendocrine cells. *Elife*, 8, e48479. doi:
1240 10.7554/eLife.48479.

1241 Zaneveld, J. R., McMinds, R., & Vega Thurber, R. (2017). Stress and stability: applying the
1242 Anna Karenina principle to animal microbiomes. *Nature microbiology*, 2(9), 1-8. doi:
1243 10.1038/nmicrobiol.2017.121

- 1244 Zhang, P., van Leeuwen, C. H. A., Bogers, D., Poelma, M., Xu, J. & Bakker, E. S. (2020).
1245 Ectothermic omnivores increase herbivory in response to rising temperature. *Oikos* 129,
1246 1–12. doi: 10.1111/oik.07082
- 1247 Zhou, C., Yang, S., Ka, W., Gao, P., Li, Y., Long, R., & Wang, J. (2022). Association of gut
1248 microbiota with metabolism in rainbow trout under acute heat stress. *Frontiers in*
1249 *Microbiology*, 13, 846336. doi: 10.3389/fmicb.2022.846336
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1253 **Supplementary material**

1254

1255 **Animal husbandry and experimental setup**

1256 The transport of egg clutches took approximately 30 minutes. Upon arrival, clutches were
1257 carefully transferred to separate trays containing about 10 L of water from the original habitat
1258 and equipped with aerators. Larvae hatched on 2 April 2023. Both clutches and newly hatched
1259 larvae were maintained in a large room with windows along two walls, which were kept open
1260 to expose the animals as closely as possible to natural light and temperature conditions.
1261 Approximately one third of the water was replaced every two days with fresh water from the
1262 original habitat. This replacement water was collected every three days and stored at 4 °C in
1263 buckets. Before use, buckets were placed in the same room as the animals until the water
1264 reached the same temperature as that in the rearing containers (14 ± 0.2 °C).

1265 Nine days after hatching, larvae reached developmental stage 25 (*sensu* Gosner, 1960),
1266 the point at which they deplete yolk reserves and begin feeding independently. At this stage,
1267 120 larvae were placed individually into 1.2-L buckets containing 1 L of filtered, rested tap
1268 water and kept under a 14:10 h light:dark cycle. Larvae were randomly assigned to three food
1269 treatments (Fig. 1). Buckets for the 18 °C treatment were placed in a climate chamber (Kälte-
1270 Klimatechnik-Frauenstein GmbH, Germany). For the 24.5 °C treatment, buckets were placed
1271 inside a water bath housed within large plastic boxes (Surplus Systems Eurobox, 60 × 40 × 22
1272 cm) in a different room, with temperature regulated by two adjustable heating elements (JBL
1273 PROTEMP S 25, 25 W, JBL GmbH & Co. KG, Germany). Water temperature in the buckets
1274 was gradually increased at a rate of 0.5 °C per hour until the target temperature was reached.

1275 Diet quality was classified based on component diversity, protein and fat levels, and
1276 caloric content. The organic grass powder contains only one plant species and has lower
1277 caloric, protein, and fat content, whereas the fish food contains a wide range of ingredients
1278 (algae, zooplankton, plant and animal products) and is higher in calories, protein, and fat.

1279 The powders used in all three diets have similar texture and solubility. They remain
1280 suspended in water for a short time before settling, ensuring that the feeding mechanisms of
1281 frog larvae - filtering and scraping surfaces - provide equal access to both powders when
1282 mixed at a 50:50 ratio. All diets were provided *ad libitum*. Buckets were cleaned at least every
1283 three days by completely replacing the water with rested tap water at the same temperature,
1284 during which each larva was briefly (<1 min) transferred to a sieve placed in a separate bucket
1285 of clean water.

1286 Buckets assigned to the heatwave treatment were placed in a water bath inside plastic
1287 boxes (60 × 40 × 22 cm) containing two adjustable heating elements. The setup was housed in
1288 a warmer room (29 °C air temperature). Prior to the experiment, the heating system was
1289 calibrated to ensure accurate temperature ramping, and water temperatures were monitored
1290 hourly. Water temperature in the buckets was increased at a rate of 0.5 °C per hour until
1291 reaching 28 °C. Because ramping protocols were identical and final temperatures (i.e.,
1292 original rearing temperatures) differed, larvae reared at 18 °C required more time to reach 28
1293 °C and return (20 h total) than larvae reared at 24.5 °C (7 h total). Buckets assigned to the

1294 control treatment (no heatwave) were also moved and returned to their original positions
1295 during treatment allocation so that handling was standardized across experimental groups.

1296

1297 **Methods for isotope analyses**

1298 Larval tails were dried in an oven at 60 °C for at least 24 hours. Subsequently, tail muscle
1299 tissue samples weighing 0.38-0.93 mg (mean = 0.76 mg) were taken in duplicate for each
1300 larva and placed in 4 × 6 mm tin cups (HEKAtech, Germany). The powdered foods
1301 corresponding to the three dietary treatments (NaturaleBio® grass powder, Sera Micron
1302 Nature® fish food, and a 50:50 mixture of both) were also analyzed, with six replicates per
1303 diet.

1304 Samples were combusted in a mass spectrometer (EURO-EA 3000, Euro Vector, Italy)
1305 using BBOT (2,5-Bis-(5-tert-butyl-2-benzoxazolyl)-thiophen; 6.51% N; 72.52% C;
1306 HEKAtech, Germany), KNO₃, and caffeine as standards. Isotope ratios are reported in δ
1307 notation (‰) relative to atmospheric nitrogen (AIR) for δ¹⁵N and Pee Dee Belemnite (PDB)
1308 for δ¹³C, following international reference standards (Fry, 2006).

1309

1310 **Sequence quality filtering, sample depth, and taxonomic assignment**

1311 Paired-end demultiplexed FASTQ files were imported into QIIME2 and denoised using
1312 the q2-deblur algorithm, which applies quality filtering based on Bokulich et al. (2013),
1313 associates erroneous sequences with their true biological sequences, and removes chimeras.
1314 Forward and reverse reads were paired, quality filtered, and trimmed to a high-quality length
1315 (median Illumina Q30), resulting in 250 bp sequences. Of the initial 2,737,481 reads, 148,401
1316 remained after filtering, with sequencing depths between 207 and 5,451 reads per sample. All
1317 negative controls (five extraction controls and two PCR controls) yielded zero reads after
1318 filtering.

1319 A phylogenetic tree was constructed using the Greengenes 16S rRNA backbone tree
1320 (version gg-13-8; McDonald et al., 2012). Taxonomic classification was performed using a
1321 custom-trained classifier built with reference sequences, taxonomy, and animal proximal gut-
1322 specific sequence weights (SILVA release 138.1, 515F/806R) from Kaehler et al. (2019;
1323 <https://github.com/BenKaehler/readytowear>). Positive controls were evaluated separately via
1324 BLAST (NCBI; Sayers et al., 2025) because they do not represent animal gut samples.

1325 Amplicon Sequence Variants (ASVs) represented by fewer than eight reads (~0.005% of
1326 total remaining sequences) were removed to minimize artifacts from amplification errors
1327 (Bokulich et al., 2013). The remaining reads were used to calculate Shannon entropy, which
1328 reached saturation at 556 reads. Thus, samples with fewer than 556 reads (15 samples, one to
1329 three per treatment) were excluded from further analyses. Because a very high number of
1330 samples was removed during quality filtering, we also estimated sample coverage and tested
1331 for curve stabilization using rarefaction in iNext (Chao et al., 2014). Sample coverage values
1332 were close to one and diversity curves stabilized for all maintained samples, indicating that
1333 our results were robust despite low sample depth.

1334 **Comparisons of bacterial genera and families among treatments using ALDEx2**

1335 All the genera (*Flectobacillus*, *Acetivibrio*, *Klebsiella*, *Cloacibacterium*,
1336 *Chryseobacterium*, *Dechloromonas*, *Hypnocyclicus*, *Yersinia*, *Dysgonomonas*, *Aeromonas*,
1337 *Pseudomonas*, and *Flavobacterium*) and families (*Spirosomaceae*, *Enterobacteriaceae*,
1338 *Moraxellaceae*, *Weeksellaceae*, *Rhodocyclaceae*, *Leptotrichiaceae*, *Dysgonomonadaceae*,
1339 *Yersiniaceae*, *Aeromonadaceae*, *Flavobacteriaceae*) identified with LEfSe were also
1340 identified with ALDEx2 as significantly different among treatments and their higher
1341 representativeness occurred in the same treatments to which they were assigned as indicator
1342 taxa (Figs. S25-S28).

1343

1344 **References**

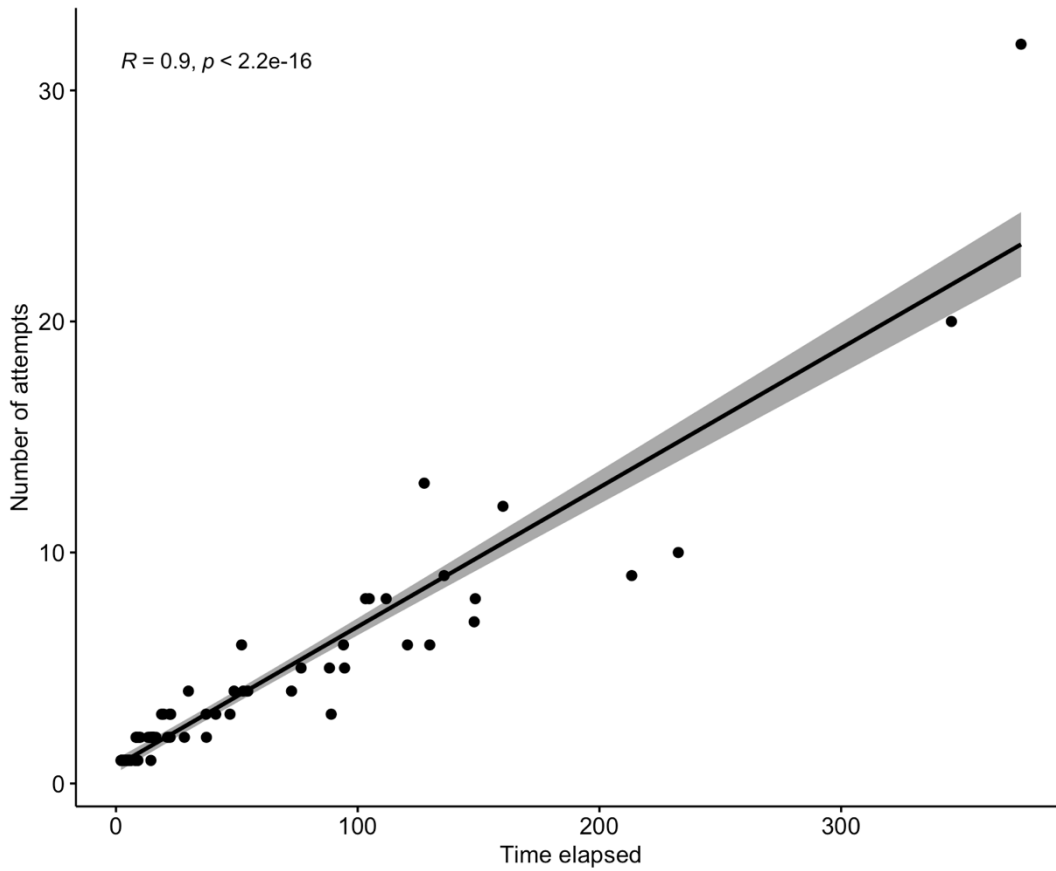
- 1345 Bokulich, N. A., Subramanian, S., Faith, J. J., Gevers, D., Gordon, J. I., Knight, R., ...
1346 Caporaso, J. G. (2013). Quality-filtering vastly improves diversity estimates from
1347 Illumina amplicon sequencing. *Nature Methods*, 10(1), 57-59. doi: 10.1038/nmeth.2276
- 1348 Chao, A., Gotelli, N. J., Hsieh, T. C., Sander, E. L., Ma, K. H., Colwell, R. K., & Ellison, A.
1349 M. (2014). Rarefaction and extrapolation with Hill numbers: a framework for sampling
1350 and estimation in species diversity studies. *Ecological Monographs*, 84(1), 45-67. doi:
1351 10.1890/13-0133.1
- 1352 Fry, B.G. (2006): *Stable Isotope Ecology*. Springer, New York.
- 1353 Gosner, K.L., 1960. A simplified table for staging anuran embryos and larvae with notes on
1354 identification. *Herpetologica* 16: 183–190.
- 1355 Kaehler, B. D., Bokulich, N. A., McDonald, D., Knight, R., Caporaso, J. G., & Huttley, G. A.
1356 (2019). Species-level microbial sequence classification is improved by source-
1357 environment information. *Nature Communications* 10: 4643. doi: 10.1038/s41467-019-
1358 12669-6
- 1359 McDonald, D., Price, M. N., Goodrich, J., Nawrocki, E. P., DeSantis, T. Z., Probst, A., ... &
1360 Hugenholtz, P. (2012). An improved Greengenes taxonomy with explicit ranks for
1361 ecological and evolutionary analyses of bacteria and archaea. *The ISME Journal*, 6(3),
1362 610-618. *ISME J.* 6, 610–618. doi: 10.1038/ismej.2011.139
- 1363 Sayers, E. W., Beck, J., Bolton, E. E., Brister, J. R., Chan, J., Connor, R., ... & Pruitt, K. D.
1364 (2025). Database resources of the National Center for Biotechnology Information in
1365 2025. *Nucleic Acids Research*, 53(D1), D20-D29. doi: 10.1093/nar/gkae979

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1368 **Supplementary figures**

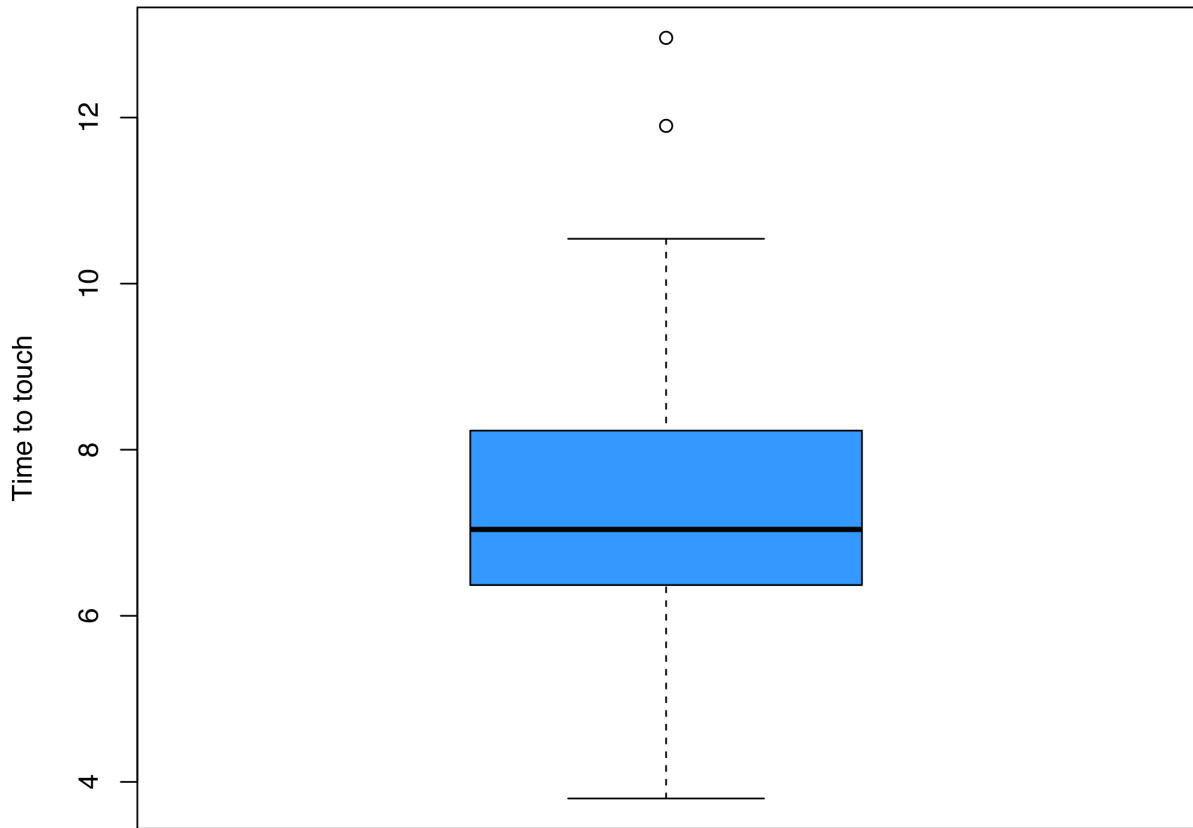
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1371 Fig. S1. Correlation between time elapsed from the end of the 3 minutes larvae remained
1372 under the funnel and the actual start of the behavioral trial (when the dragonfly naiad model
1373 touched the water) and number of attempts (number of times the larva had to be repositioned
1374 on the center of the tray). Refer to the section “Behavioral trials” for a detailed description of
1375 escape behavior trials of *Rana temporaria* larvae.

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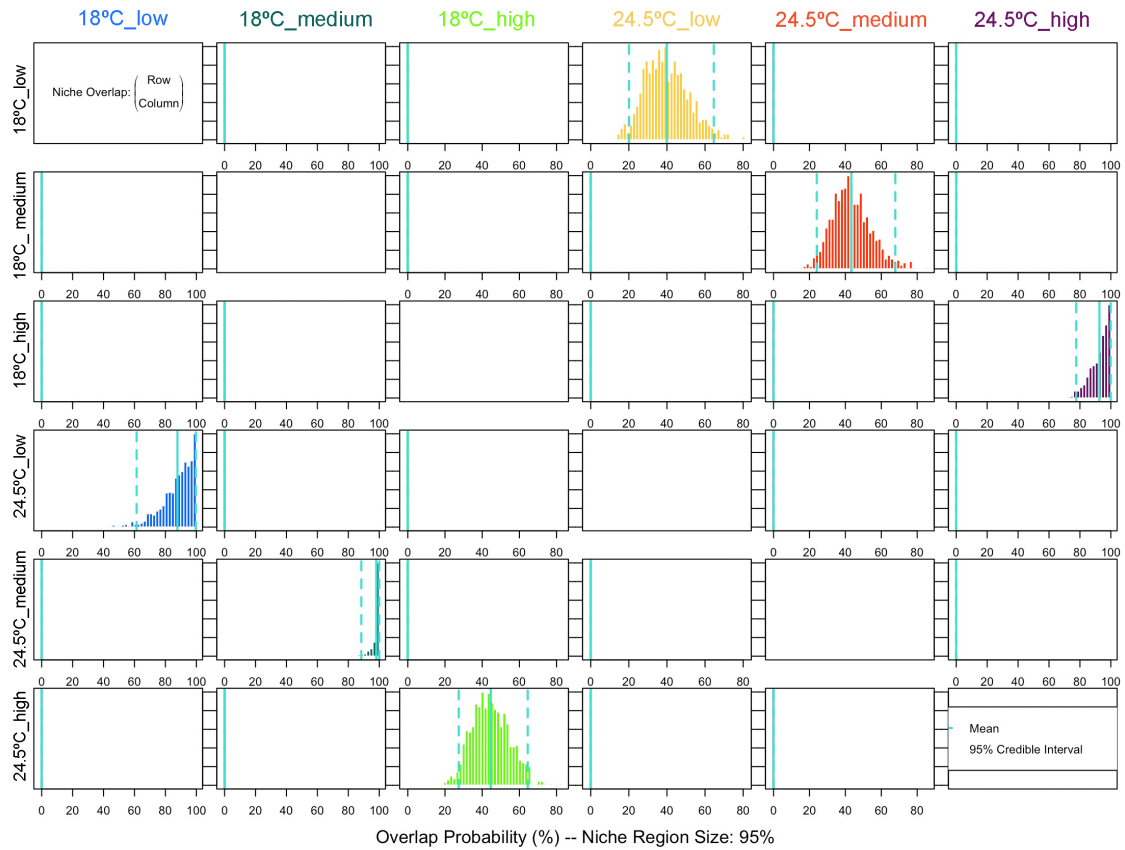
Larvae touched

1377

1378 Fig. S2. *Rana temporaria* larvae were tested for escape behavior to an aversive stimulus
1379 represented by an approaching transparent plastic pipette filled with 4 ml of water containing
1380 chemical predator cues to be released and a predator model glued to the top of the pipette. The
1381 graph shows the time elapsed from the moment the predator model touched the water to the
1382 moment it touched the larvae (when it happened) in behavioral trials (n = 102 trials). The two
1383 outliers above were excluded from posterior analyses.

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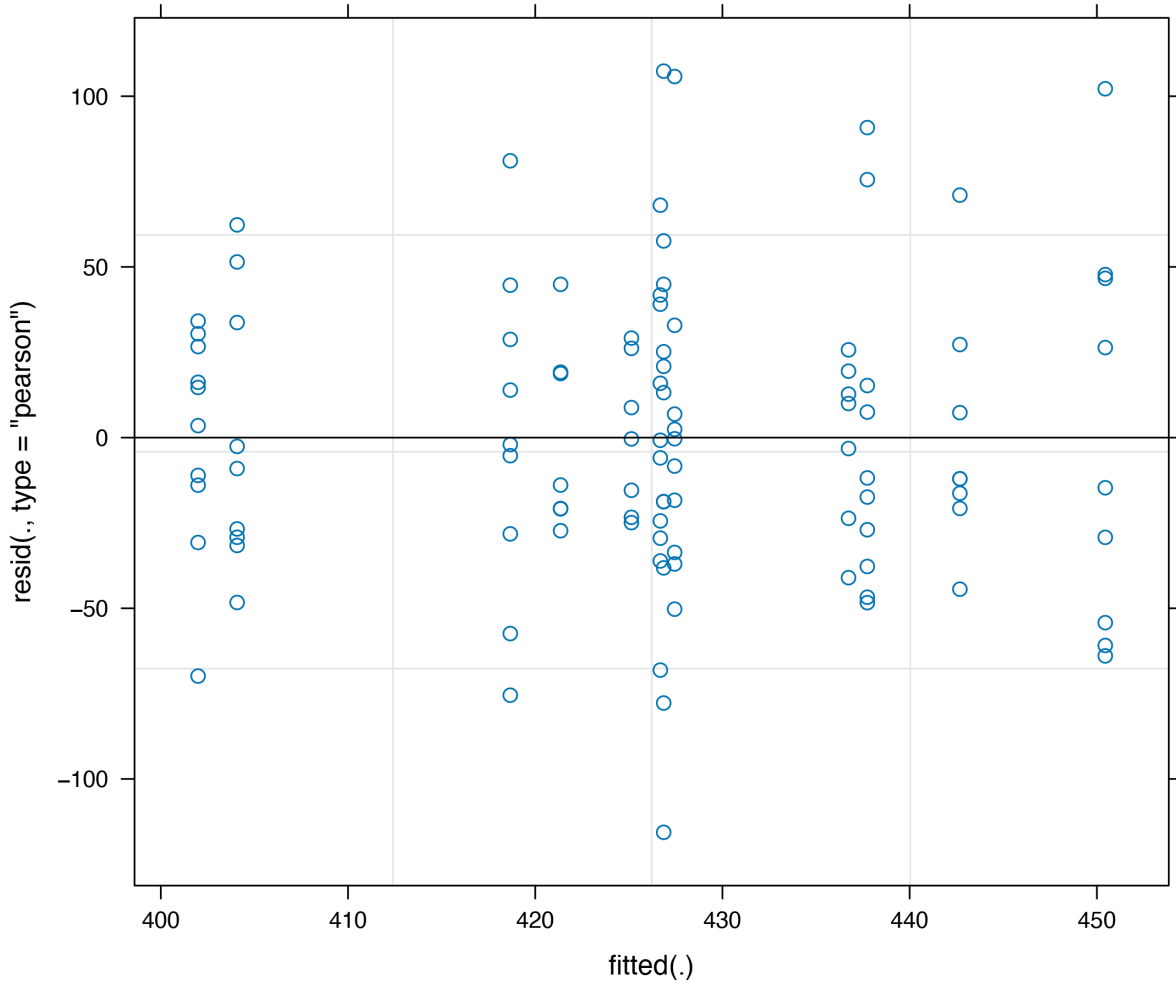


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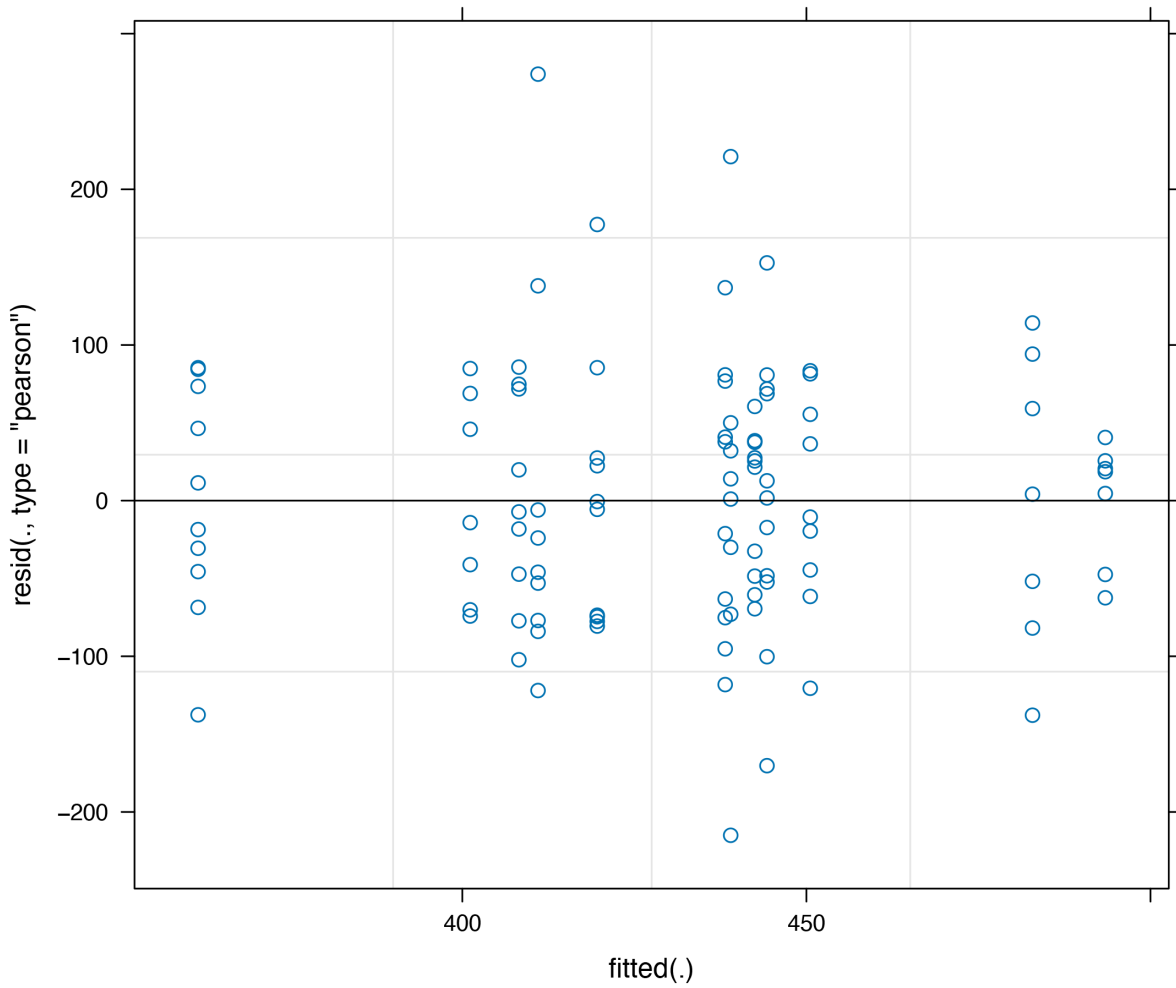
1388 Fig. S3. Posterior distribution of the probabilistic niche overlap of *Rana temporaria* larvae
 1389 reared with three different diets varying in nutritional quality and two temperatures in a
 1390 crossed experimental design (colors correspond to treatments on the columns). Niche overlap
 1391 metrics were generated by the package nicheROVER (Swanson et al., 2015). The probability
 1392 distribution of species displayed in rows overlapping onto those displayed in columns is
 1393 presented as well as posterior means (turquoise continuous lines) and 95% credible intervals
 1394 (turquoise dashed lines).

1395



1396
1397 Fig. S4. Residual distribution of the model testing the effects of food treatment, rearing
1398 temperature, and exposure or not to a heatwave on body condition (SMI) of *Rana temporaria*
1399 larvae (see Table 1 for model description).

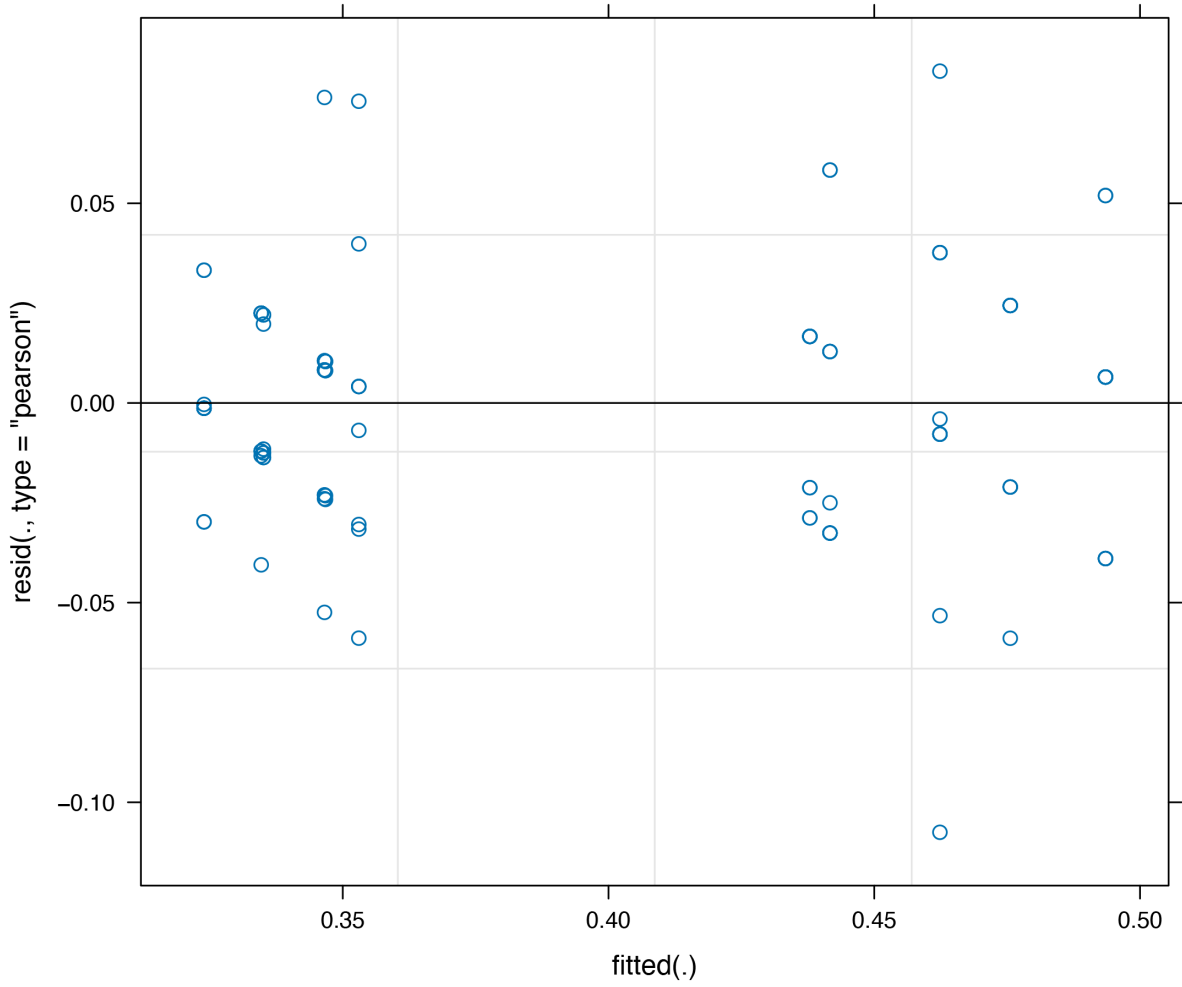
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1402 Fig. S5. Residual distribution of the model testing the effects of food treatment, rearing
1403 temperature, and exposure or not to a heatwave on mass of *Rana temporaria* larvae (see Table
1404 1 for model description).

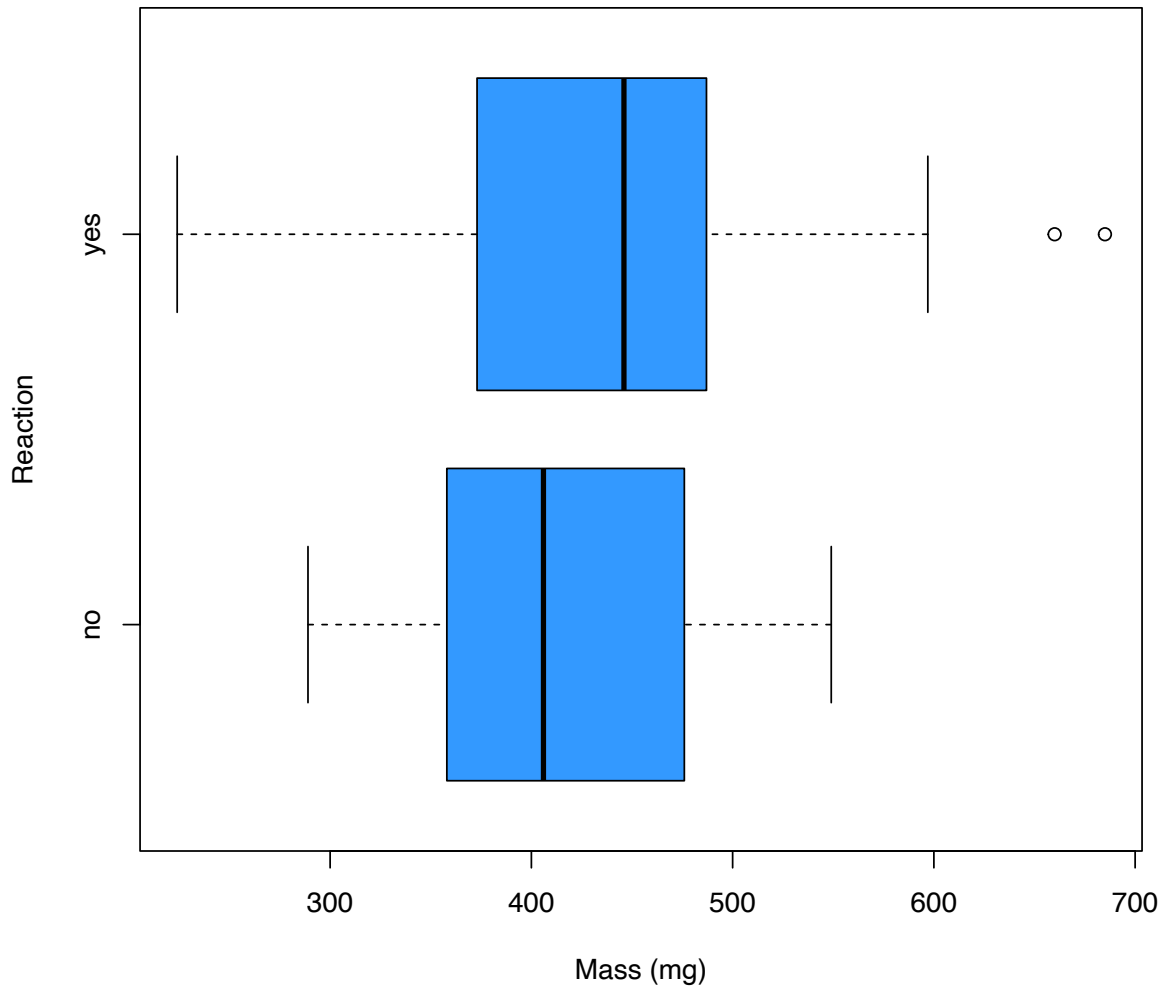
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1407 Fig. S6. Residual distribution of the model testing the effects of food treatment, rearing
1408 temperature, and exposure or not to a heatwave on developmental rate of *Rana temporaria*
1409 larvae (see Table 1 for model description).

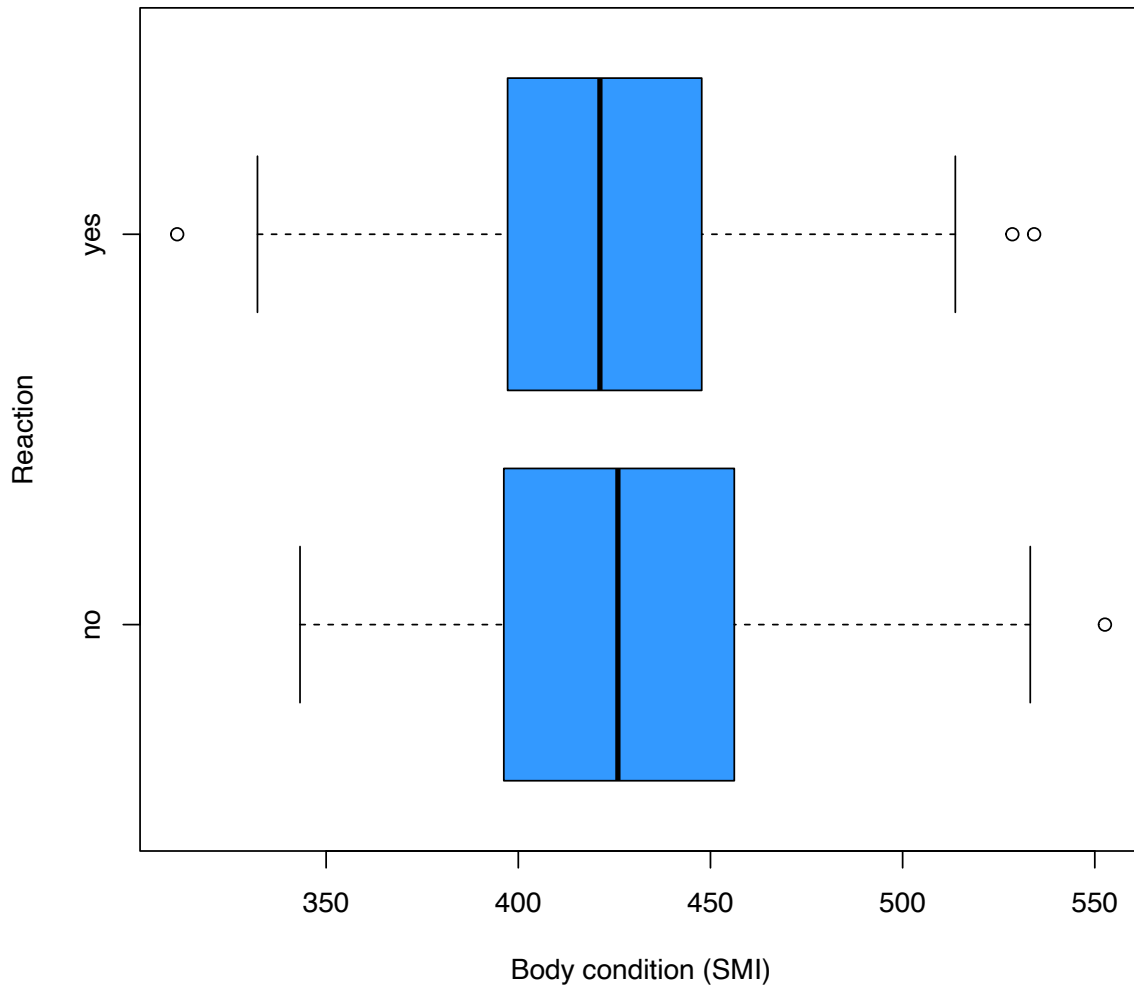
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1412 Fig. S7. Mass (mg) of *Rana temporaria* larvae that either reacted to the aversive stimulus
1413 presented in behavioral trials (1) or not (0). Wilcoxon-test: $W = 711.5$, $p = 0.252$.

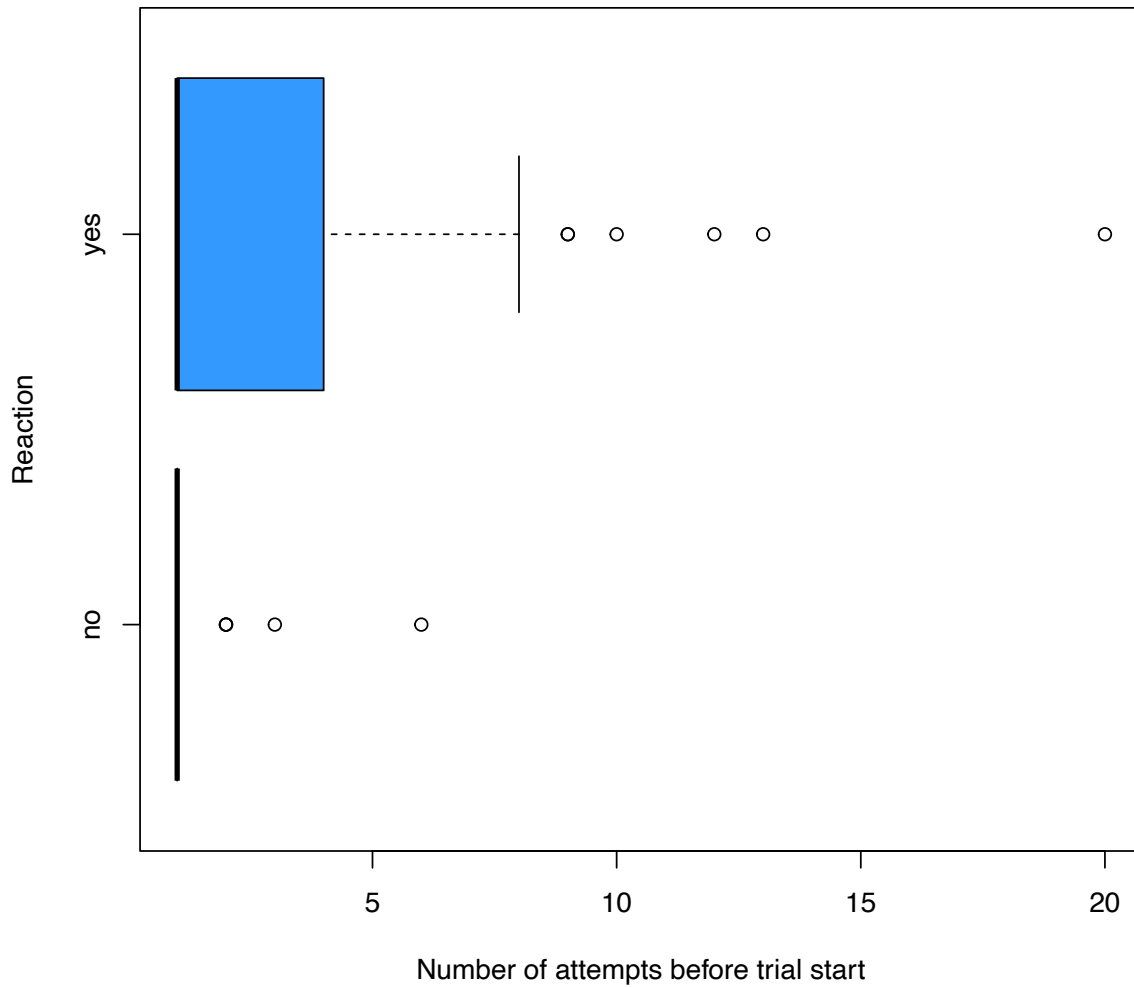
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1416 Fig. S8. Body condition (SMI) of *Rana temporaria* larvae that either reacted to the aversive
1417 stimulus presented in behavioral trials (1) or not (0). Wilcoxon-test: $W = 936$, $p = 0.482$.

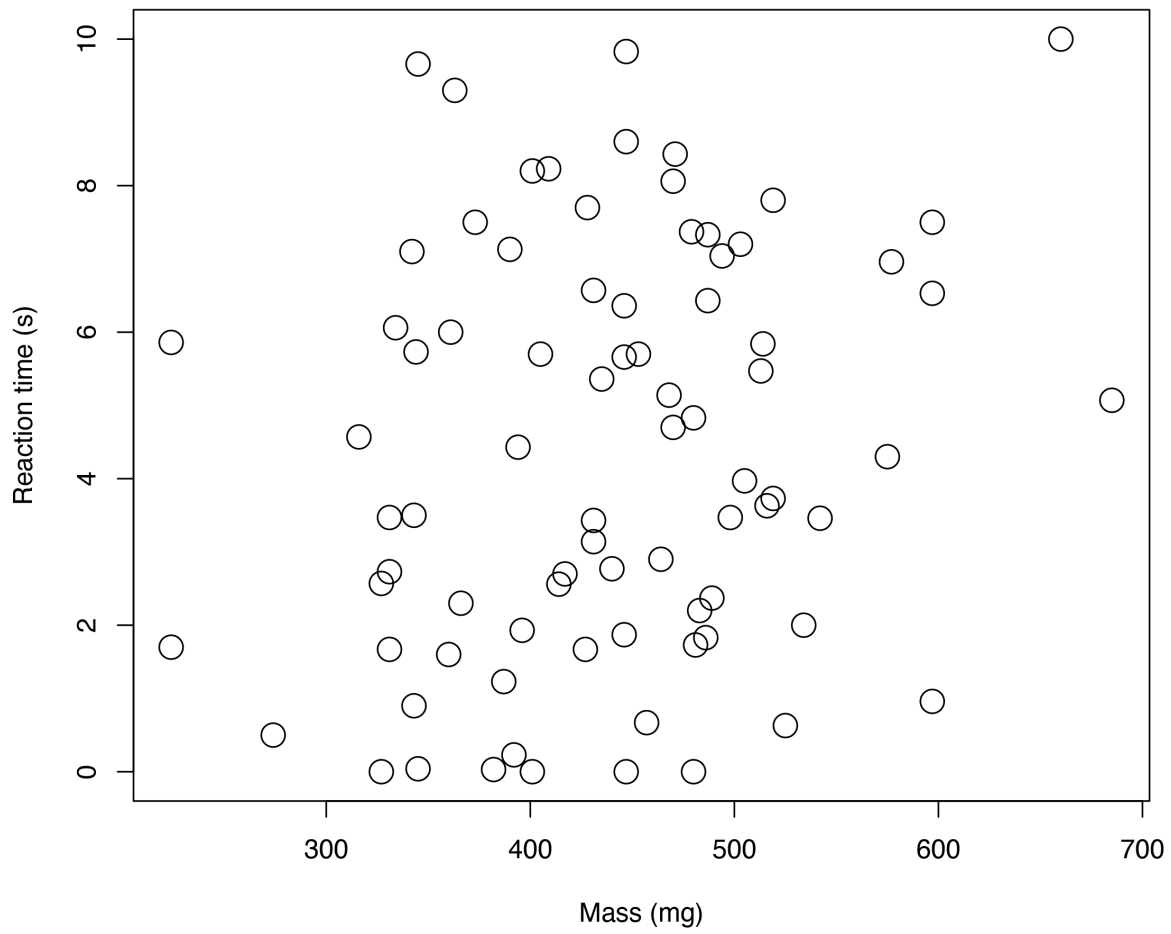
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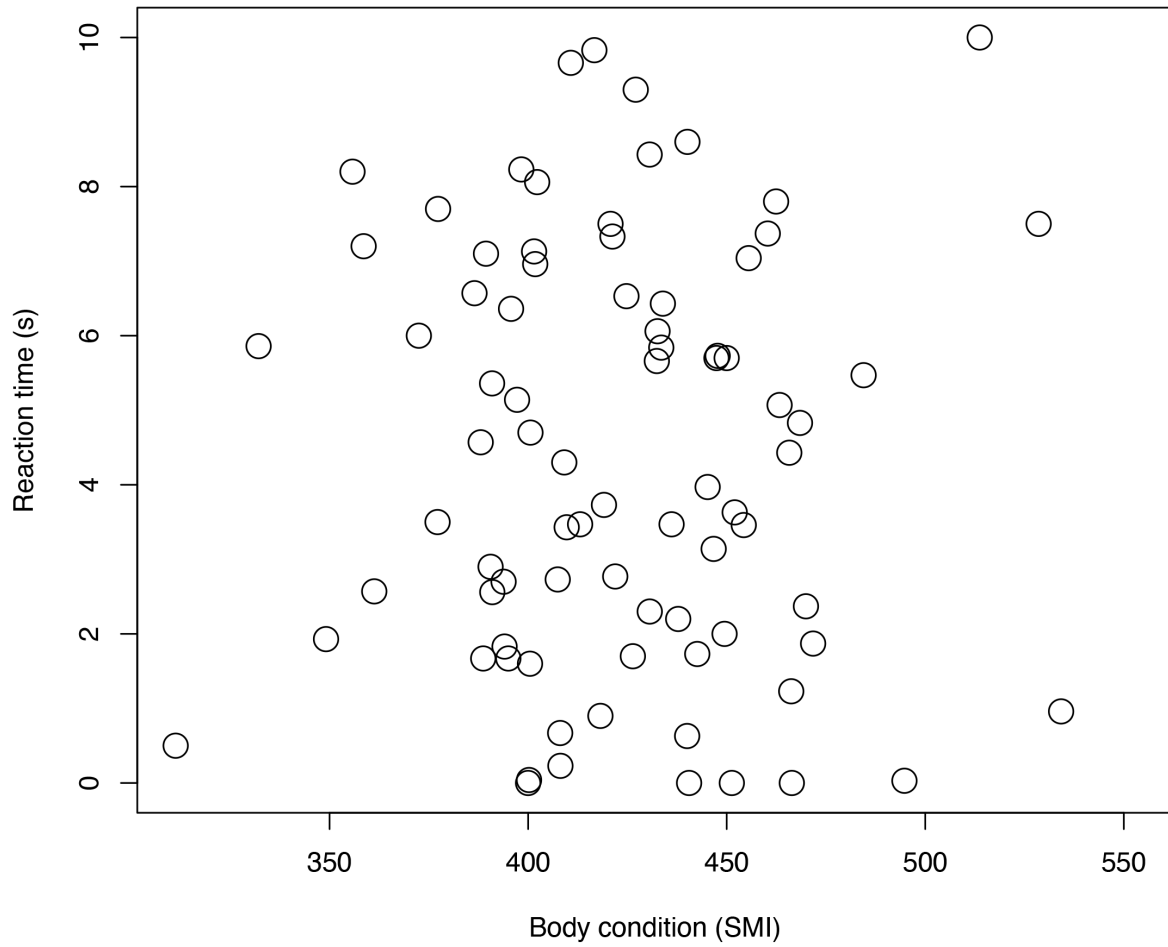
1420 Fig. S9. Number of attempts to position *Rana temporaria* larvae before the start of the
 1421 behavioral trials compared between larvae that either reacted to the aversive stimulus
 1422 presented in behavioral trials (1) or not (0). Wilcoxon-test: $W = 601$, $p = 0.022$.

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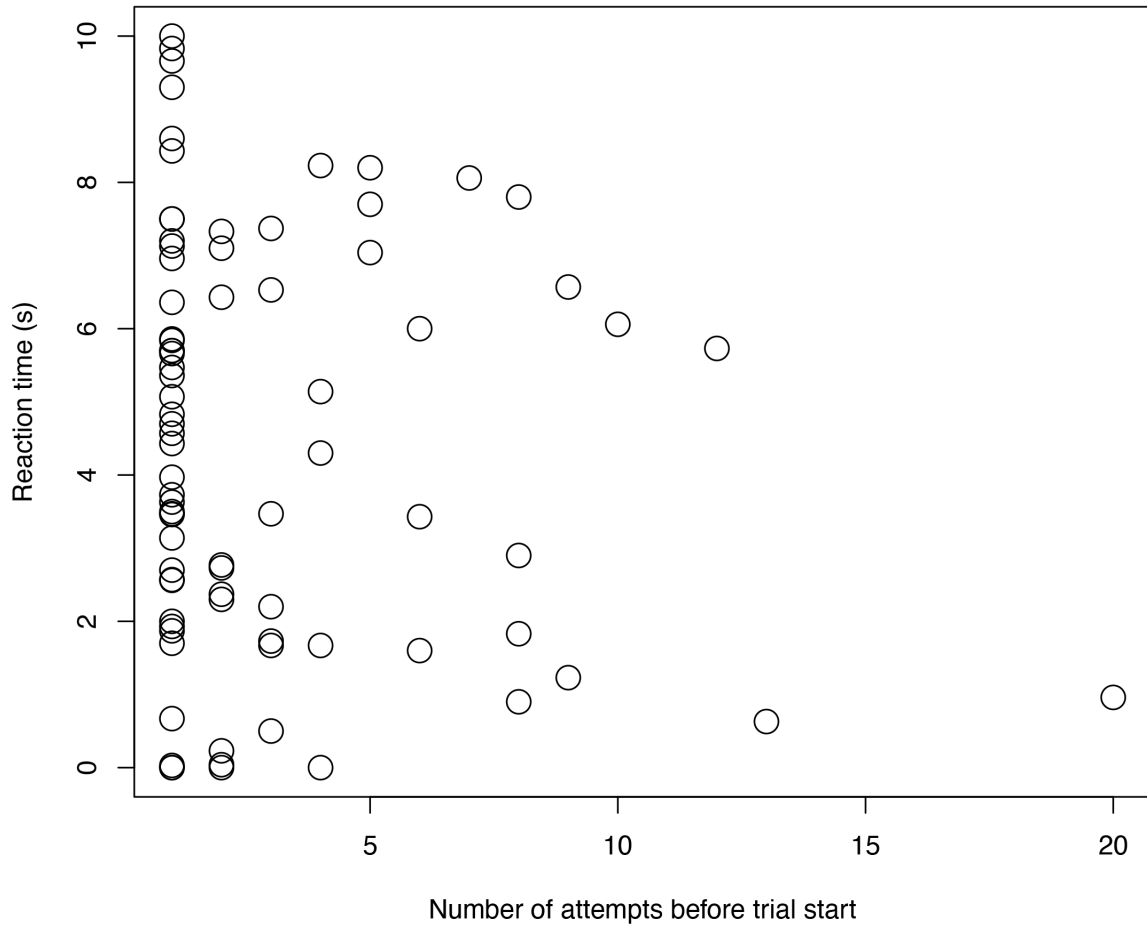
1425 Fig. S10. Relationship between mass (mg) of *Rana temporaria* larvae and time to react to the
1426 aversive stimulus presented in behavioral trials. Adjusted R-squared = 0.030, $F = 3.487$, $df =$
1427 79, $p = 0.066$.



1428

1429 Fig. S11. Relationship between body condition (SMI) of *Rana temporaria* larvae and time to
1430 react to the aversive stimulus presented in behavioral trials. Adjusted R-squared = -0.013, F =
1431 0.005, df = 79, p = 0.946.

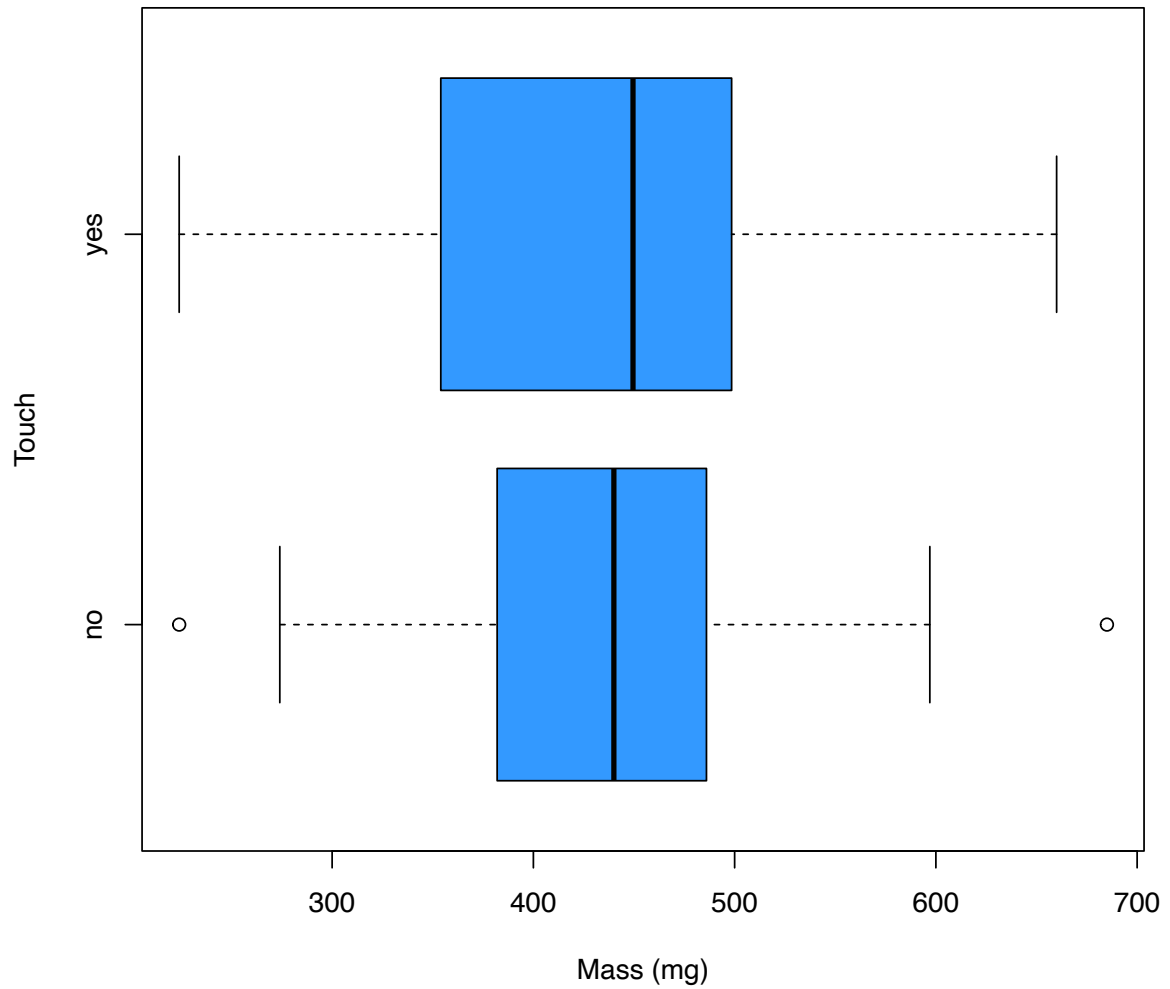
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1434 Fig. S12. Relationship between number of attempts to position *Rana temporaria* larvae before
1435 the start of the behavioral trials and time (s) the larvae took to react to the aversive stimulus
1436 presented. $\rho = -0.125$, $p = 0.263$.

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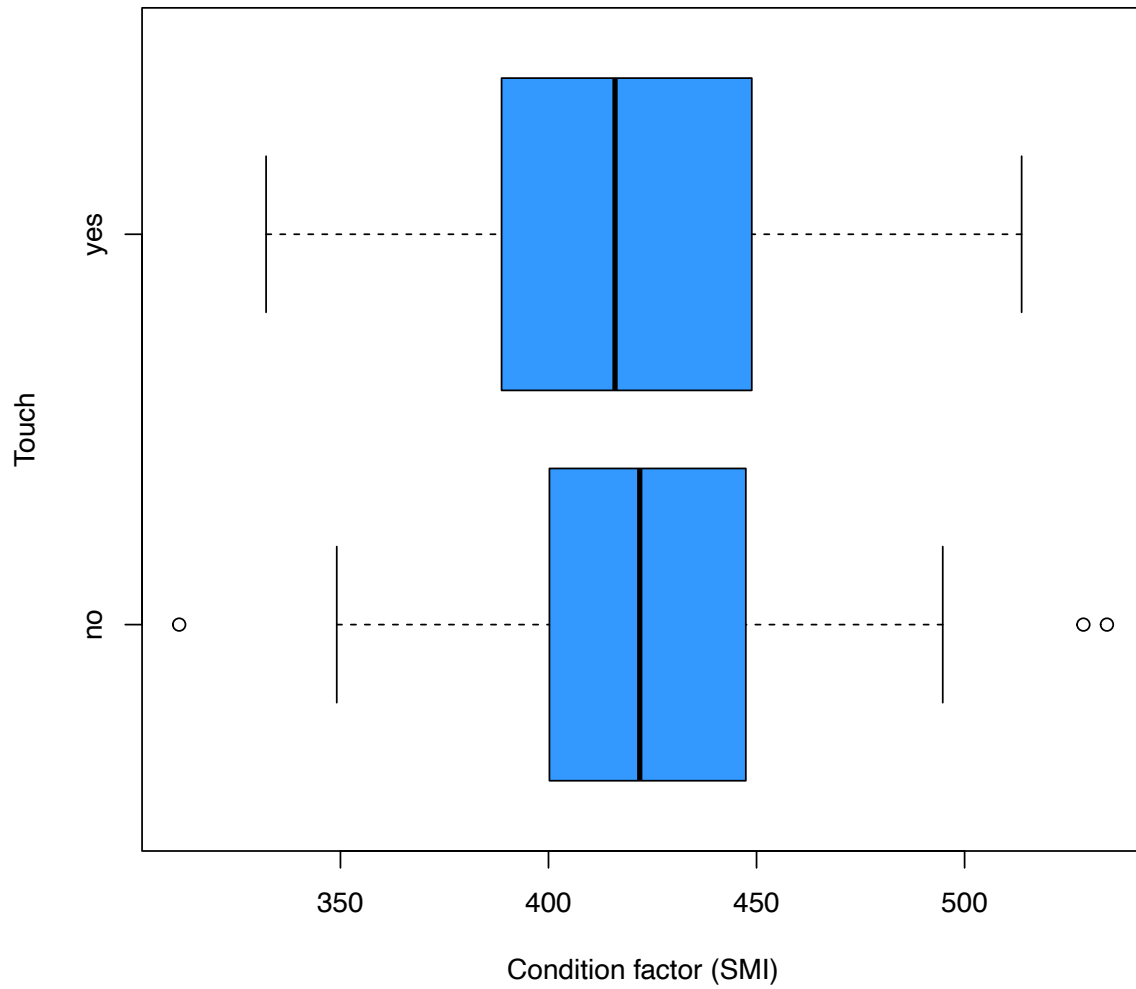


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1439 Fig. S13. Mass (mg) of reacting *Rana temporaria* larvae that either were touched by the
1440 predator model approached to them in behavioral trials (1) or not (0) before fleeing.

1441 Wilcoxon-test: $W = 577$, $p = 0.722$.

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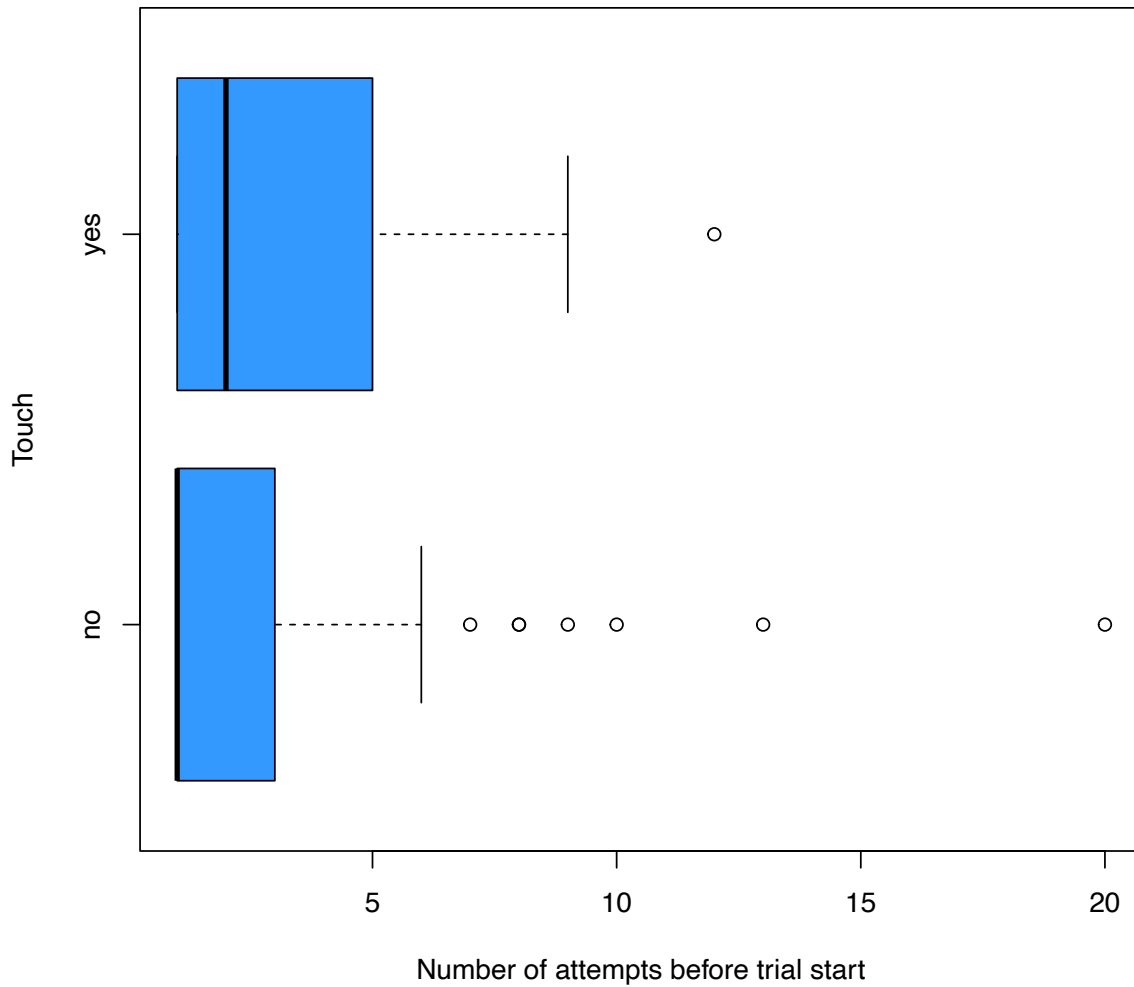


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1444 Fig. S14. Condition factor (SMI) of reacting *Rana temporaria* larvae that either were touched
 1445 by the predator model approached to them in behavioral trials (1) or not (0) before fleeing.

1446 Wilcoxon-test: $W = 697$, $p = 0.343$.

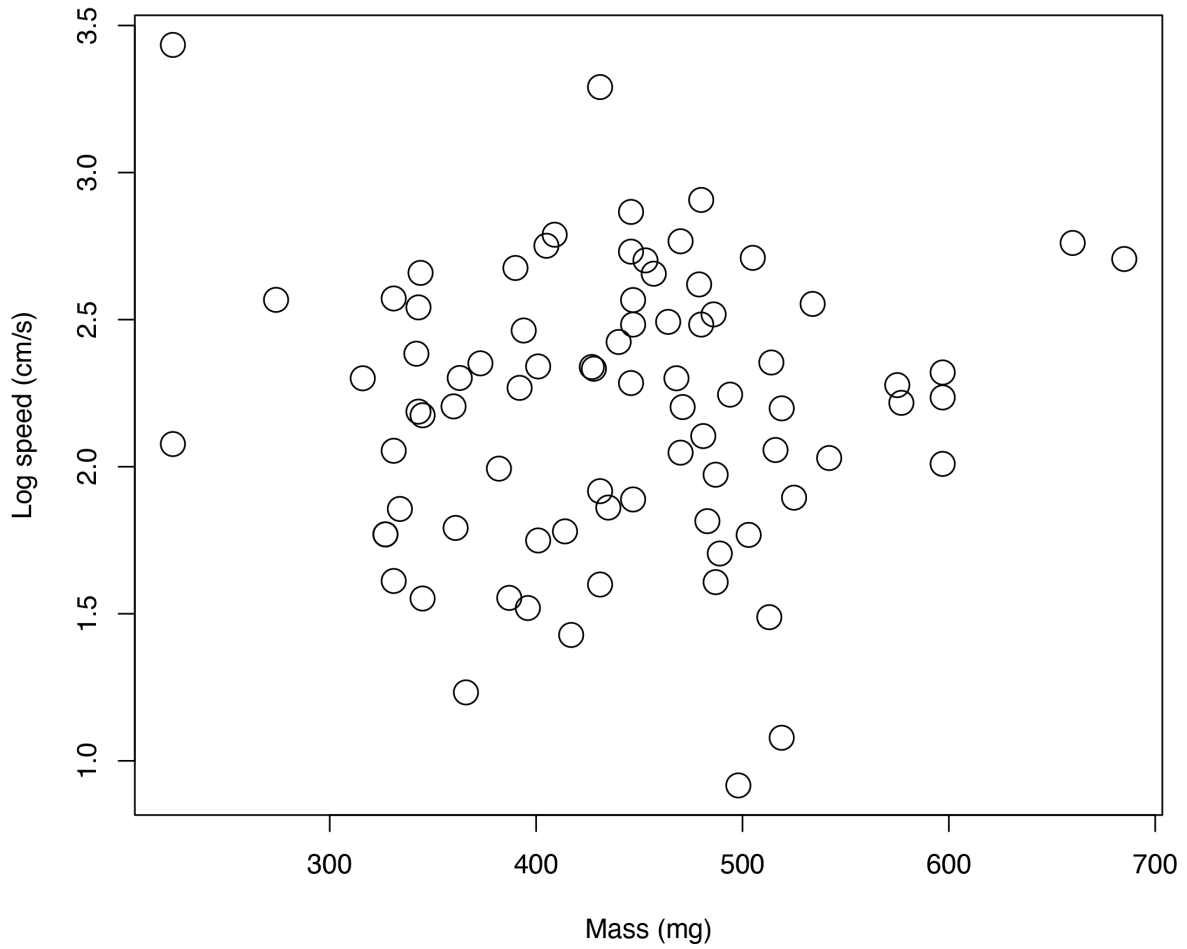
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1449 Fig. S15. Number of attempts to position *Rana temporaria* larvae before the start of the
 1450 behavioral trials compared between larvae that either were touched by the predator model
 1451 approached to them in the behavioral trials (1) or not (0) before fleeing. Wilcoxon-test: $W =$
 1452 533, $p = 0.366$.

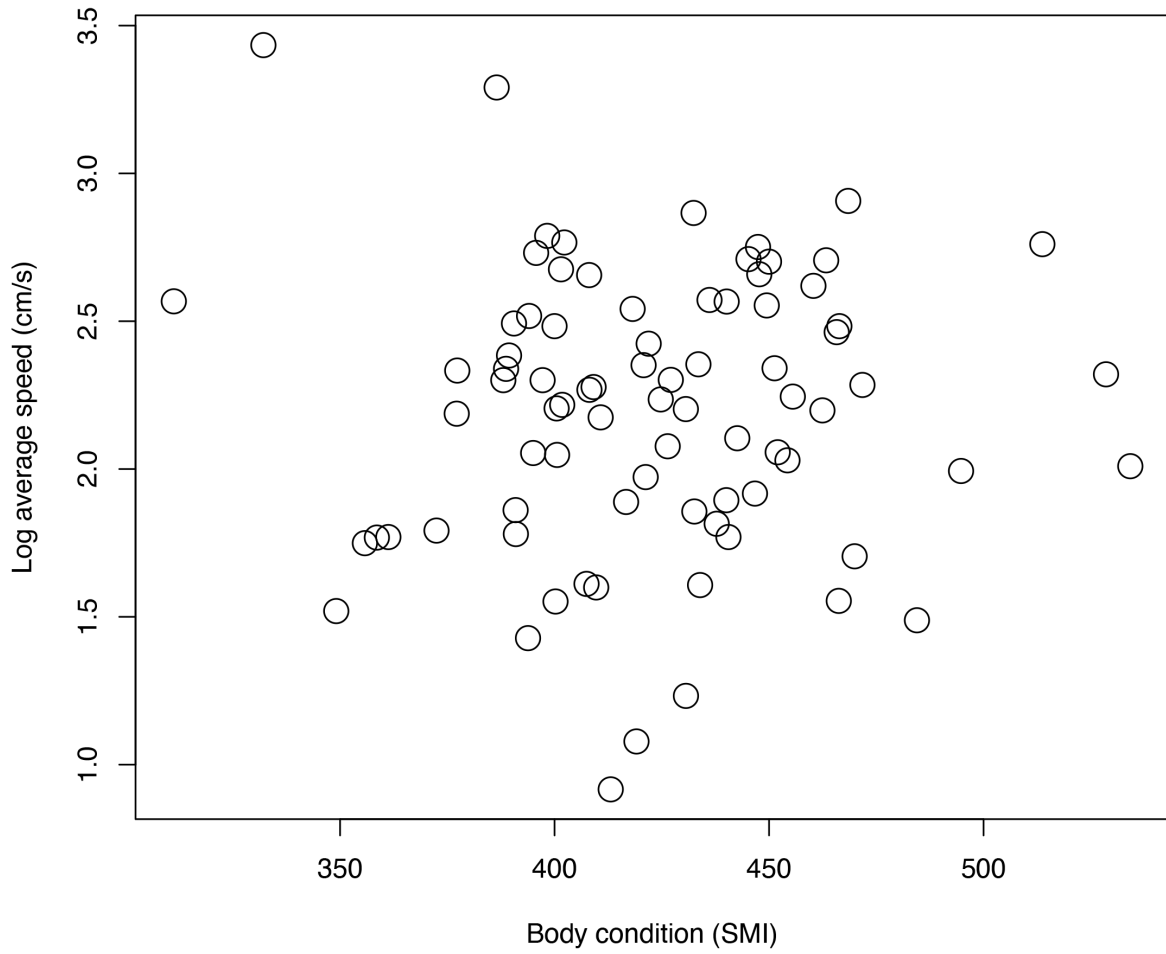
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1455 Fig. S16. Relationship between mass (mg) of *Rana temporaria* larvae and average speed (in
1456 cm/s, log transformed) while fleeing from the aversive stimulus presented in behavioral trials.
1457 Adjusted R-squared = -0.013, F = 0.004, df = 79, p = 0.949.

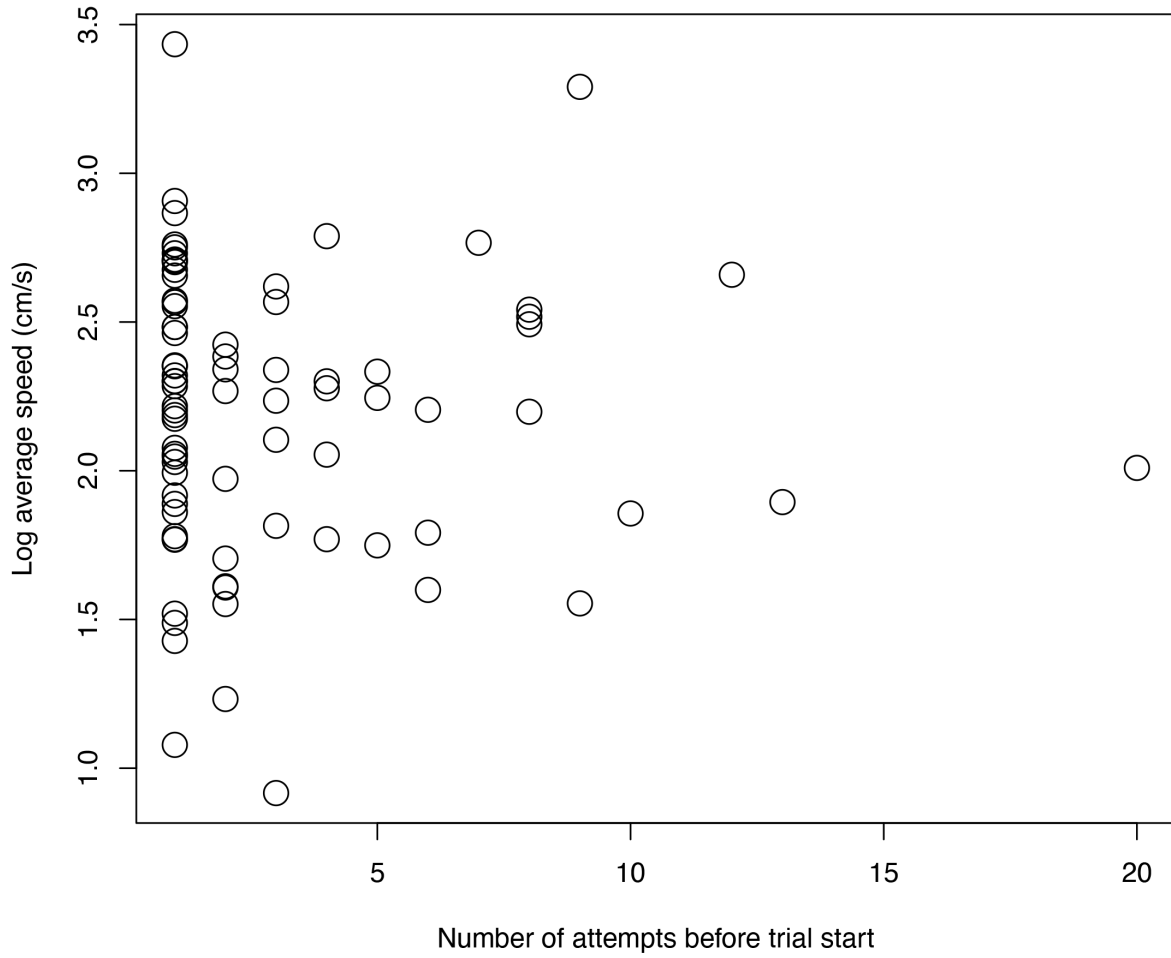
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1460 Fig. S17. Relationship between body condition (SMI) of *Rana temporaria* larvae and average
1461 speed (in cm/s, log transformed) while fleeing from the aversive stimulus presented in
1462 behavioral trials. Adjusted R-squared = -0.013, $F = 0.009$, $df = 79$, $p = 0.923$.

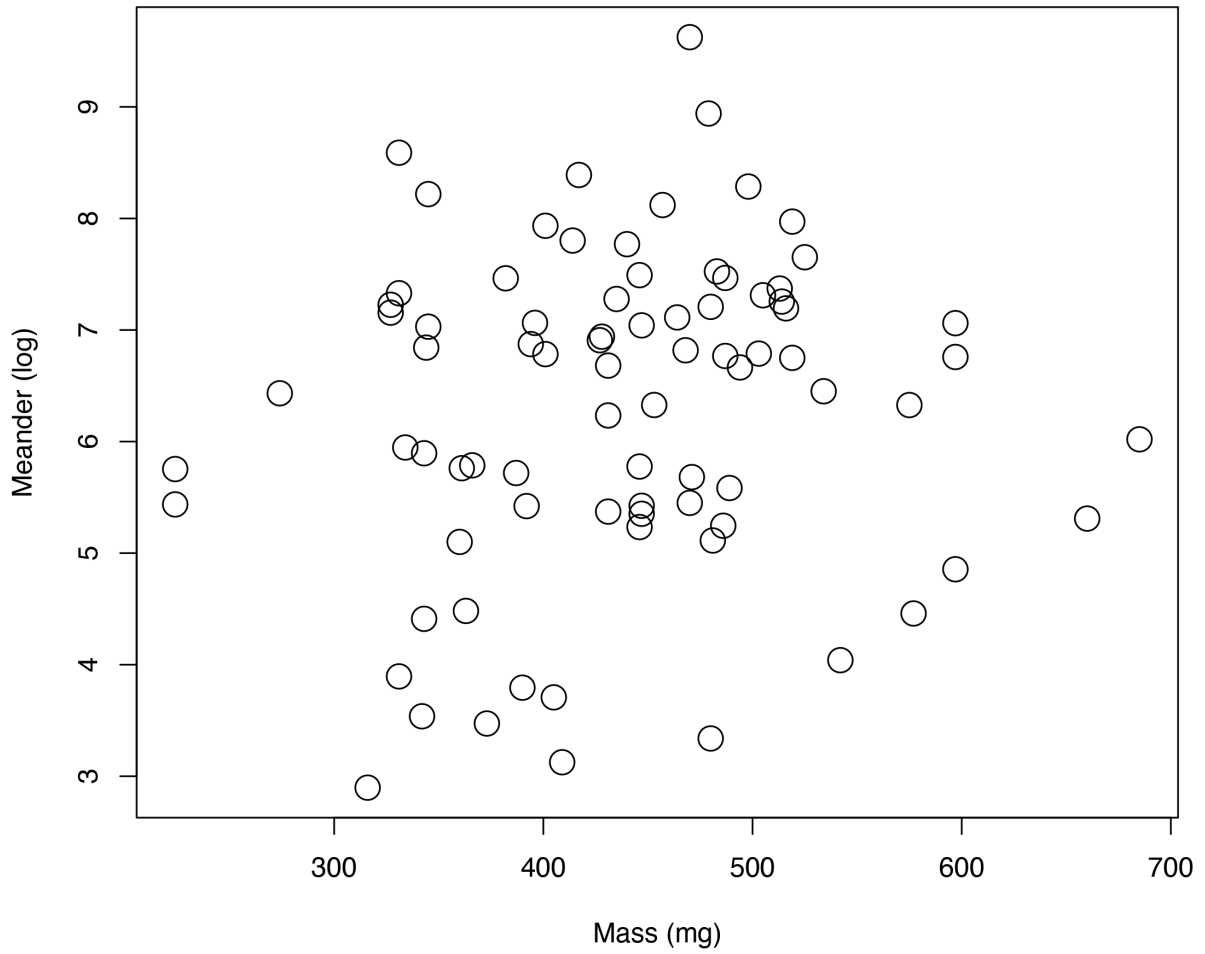
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1465 Fig. S18. Relationship between number of attempts to position *Rana temporaria* larvae before
1466 the start of the behavioral trials and average speed (in cm/s, log transformed) of the larvae
1467 while fleeing from the aversive stimulus presented. $\rho = -0.104$, $p = 0.354$.

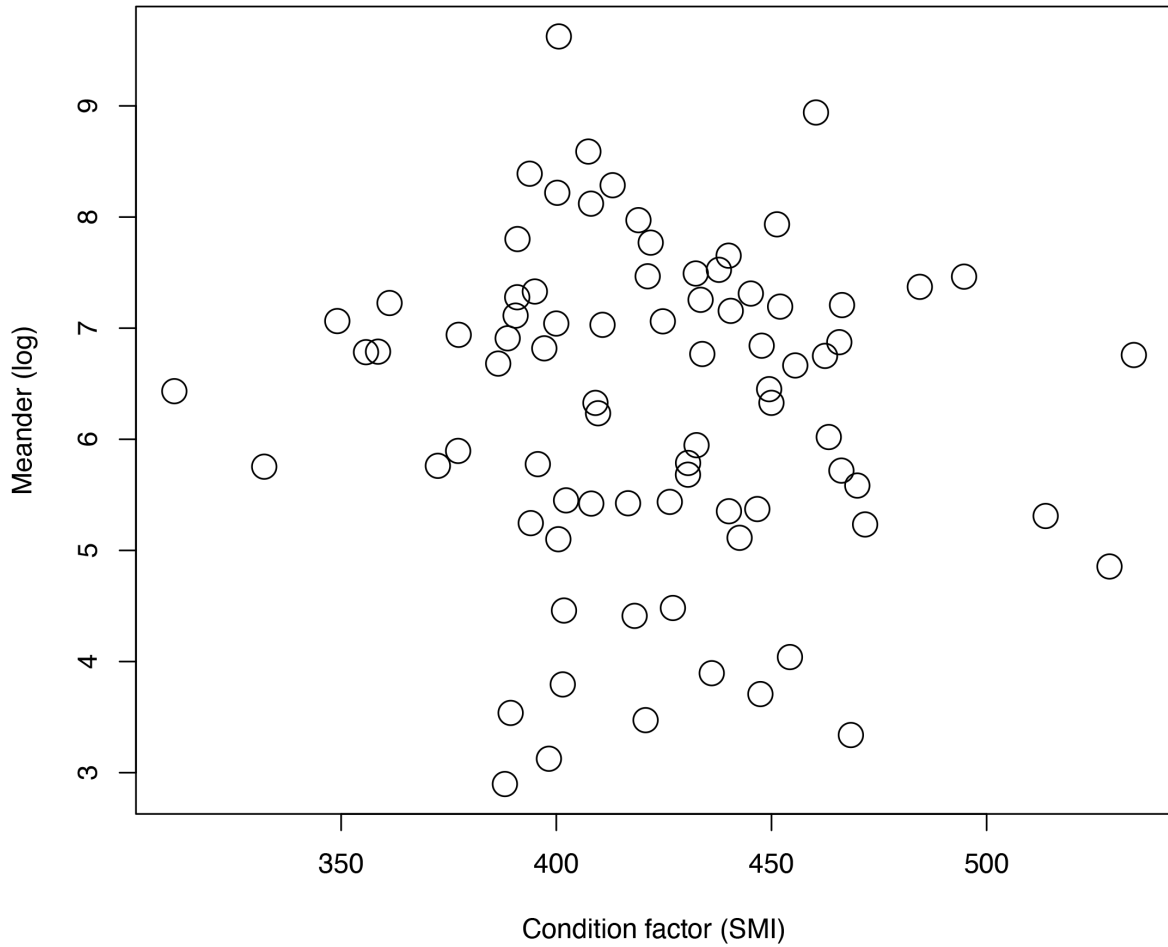
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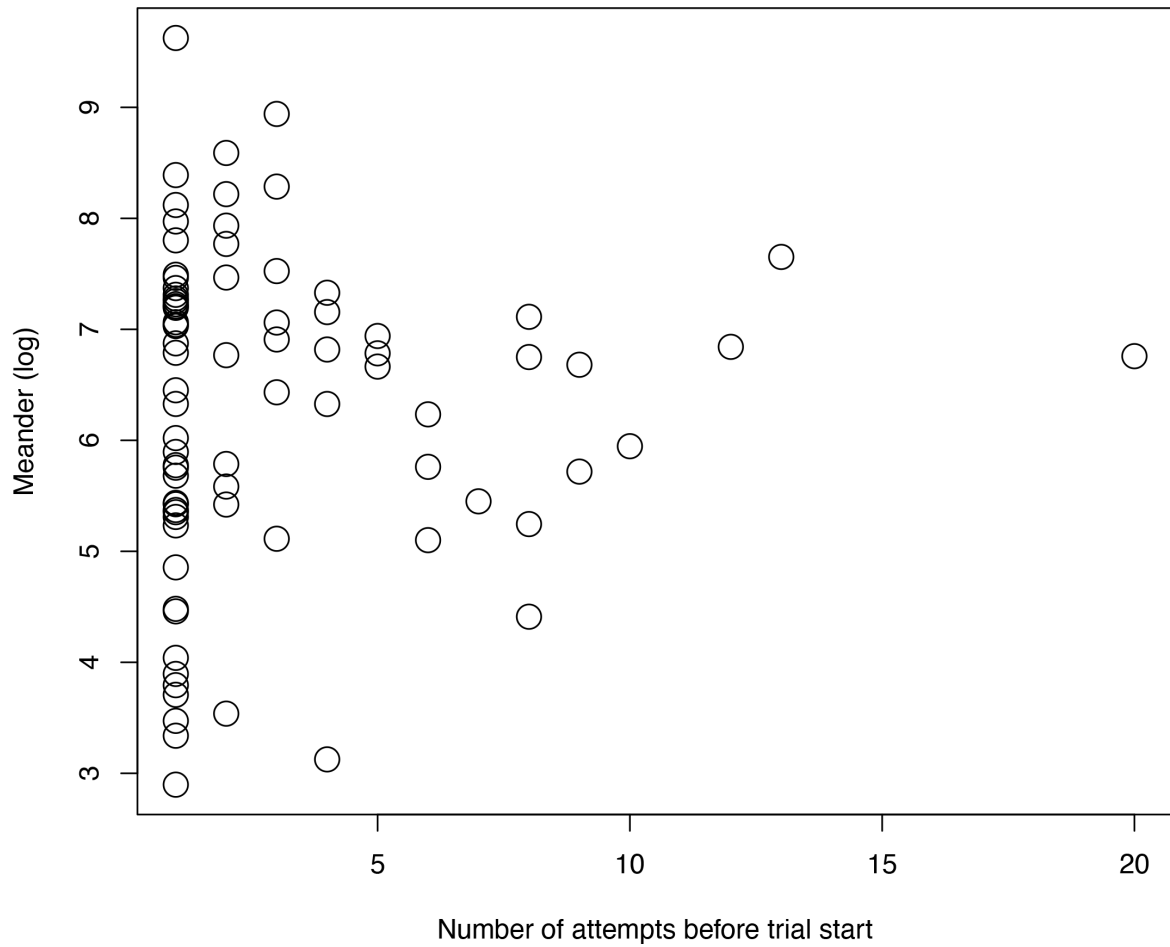
1470 Fig. S19. Relationship between mass (mg) of *Rana temporaria* larvae and trajectory non-
1471 linearity (“meander”, log transformed) while fleeing from the aversive stimulus presented in
1472 behavioral trials. Adjusted R-squared = 0.003, F = 1.271, df = 79, p = 0.263.

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1475 Fig. S20. Relationship between body condition (SMI) of *Rana temporaria* larvae and
1476 trajectory non-linearity (“meander”, log transformed) while fleeing from the aversive stimulus
1477 presented in behavioral trials. Adjusted R-squared = -0.009, F = 0.247, df = 79, p = 0.620.

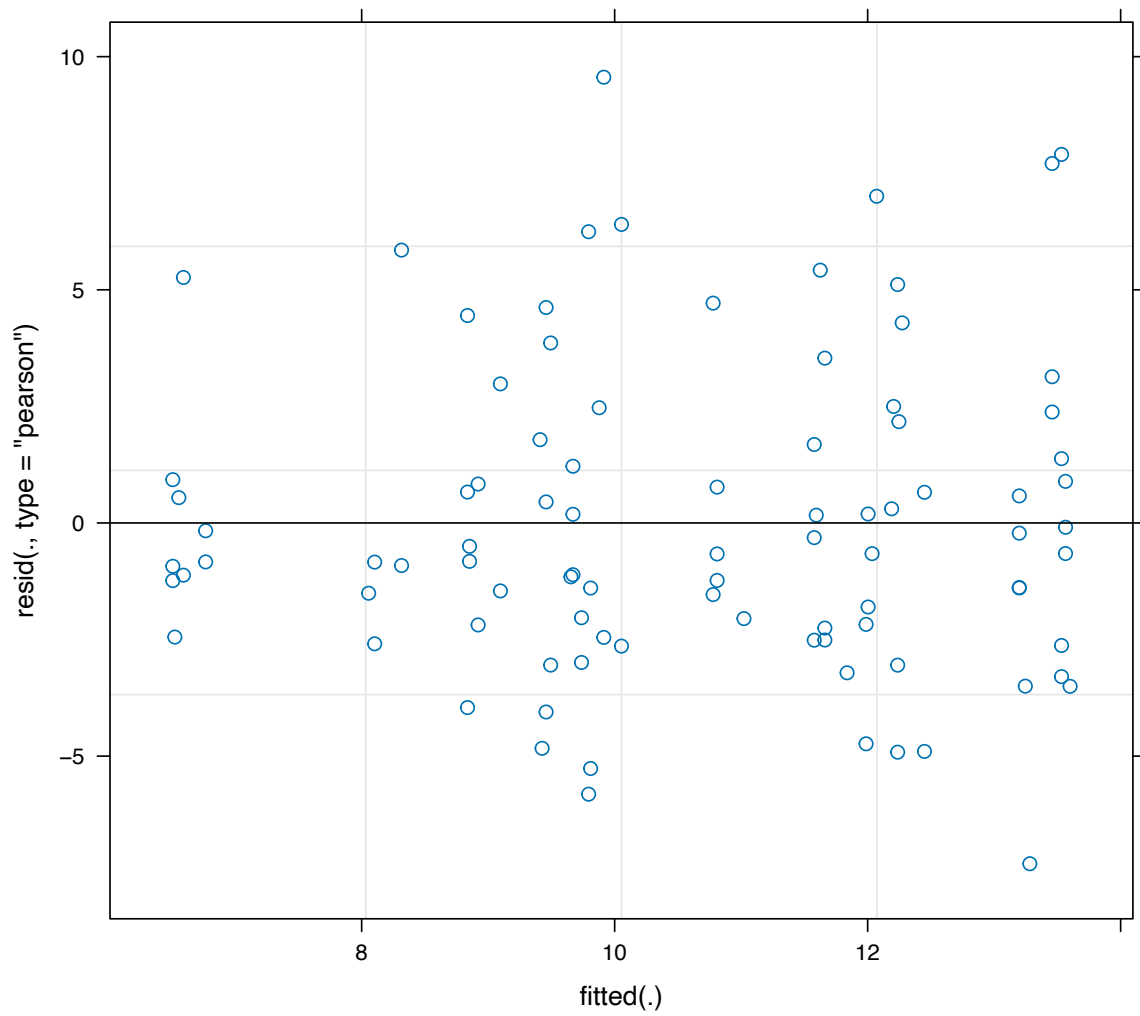


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1479 Fig. S21. Relationship between number of attempts to position *Rana temporaria* larvae before
 1480 the start of the behavioral trials and trajectory non-linearity (“meander”, log transformed) of
 1481 the larvae while fleeing from the aversive stimulus presented. $\rho = 0.050$, $p = 0.657$.

1482

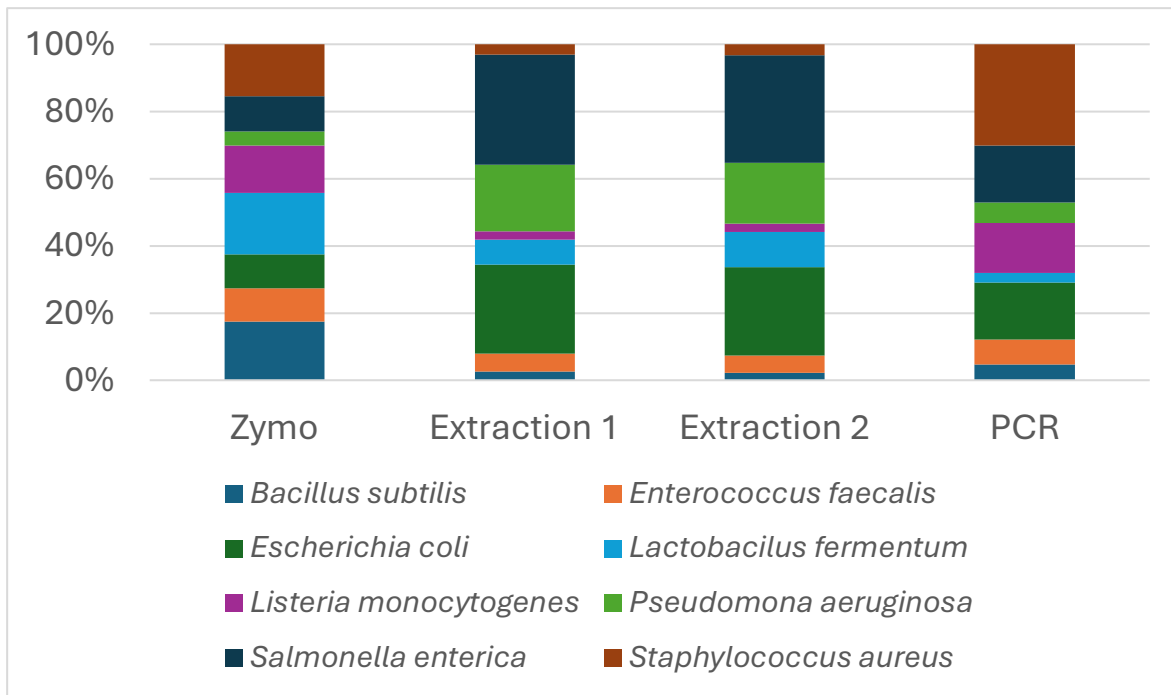
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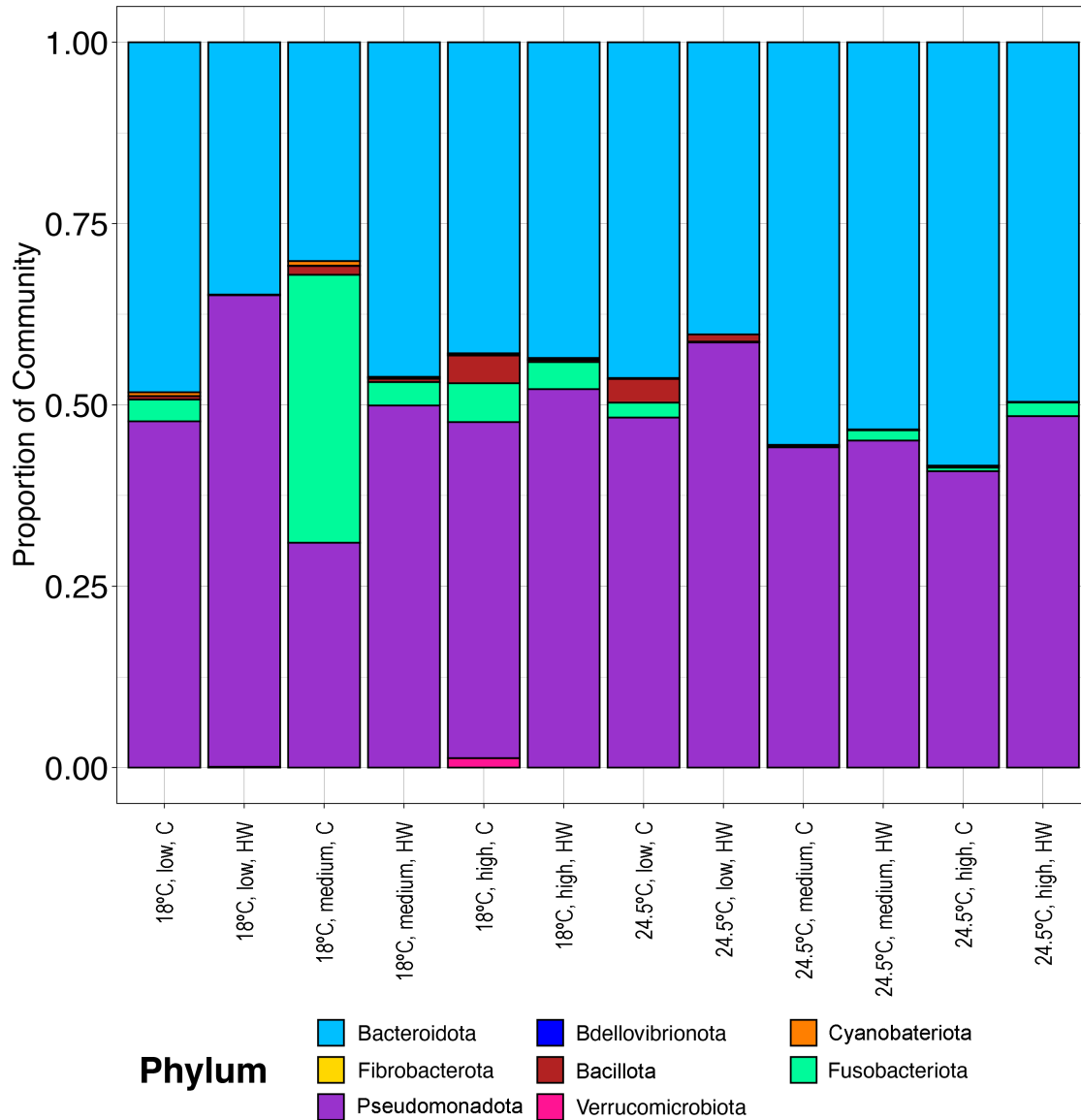
1485 Fig. S22. Residual distribution of the model testing the effects of food treatment, rearing
1486 temperature, and exposure or not to a heatwave on bacteria alpha diversity (Hill numbers) in
1487 the gut microbiome of *Rana temporaria* larvae (see Table 1 for model description).

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Fig. S23. Results of two positive controls for DNA extractions (ZymoBIOMICST[™] microbial community standard, Zymo Research Europe GmbH) and one positive PCR control (ZymoBIOMICST[™] microbial community DNA standard, Zymo Research Europe GmbH) in comparison with the expected community profile (Zymo), showing that taxonomic composition was precisely assessed, but not relative abundances. The similarity of the two extractions shows repeatability, meaning that bias in reflecting the real abundance of given taxa are consistent and, thus, comparable among samples.



1499

1500 Fig. S24. Community composition of gut bacteria based on phylum for *Rana temporaria*
 1501 larvae fed three diets with increasing levels of protein, fat, and animal components
 1502 (considered as low-, medium- and high-quality), reared at either 18 °C or 24.5 °C. and
 1503 exposed or not to a heatwave, in a crossed experimental design. The heatwave corresponded
 1504 to increasing temperature at a ramping rate of 0.5 °C per hour until 28 °C, maintenance at 28
 1505 °C for 48 h and subsequent temperature decrease of 0.5 °C per hour until original rearing
 1506 temperature.

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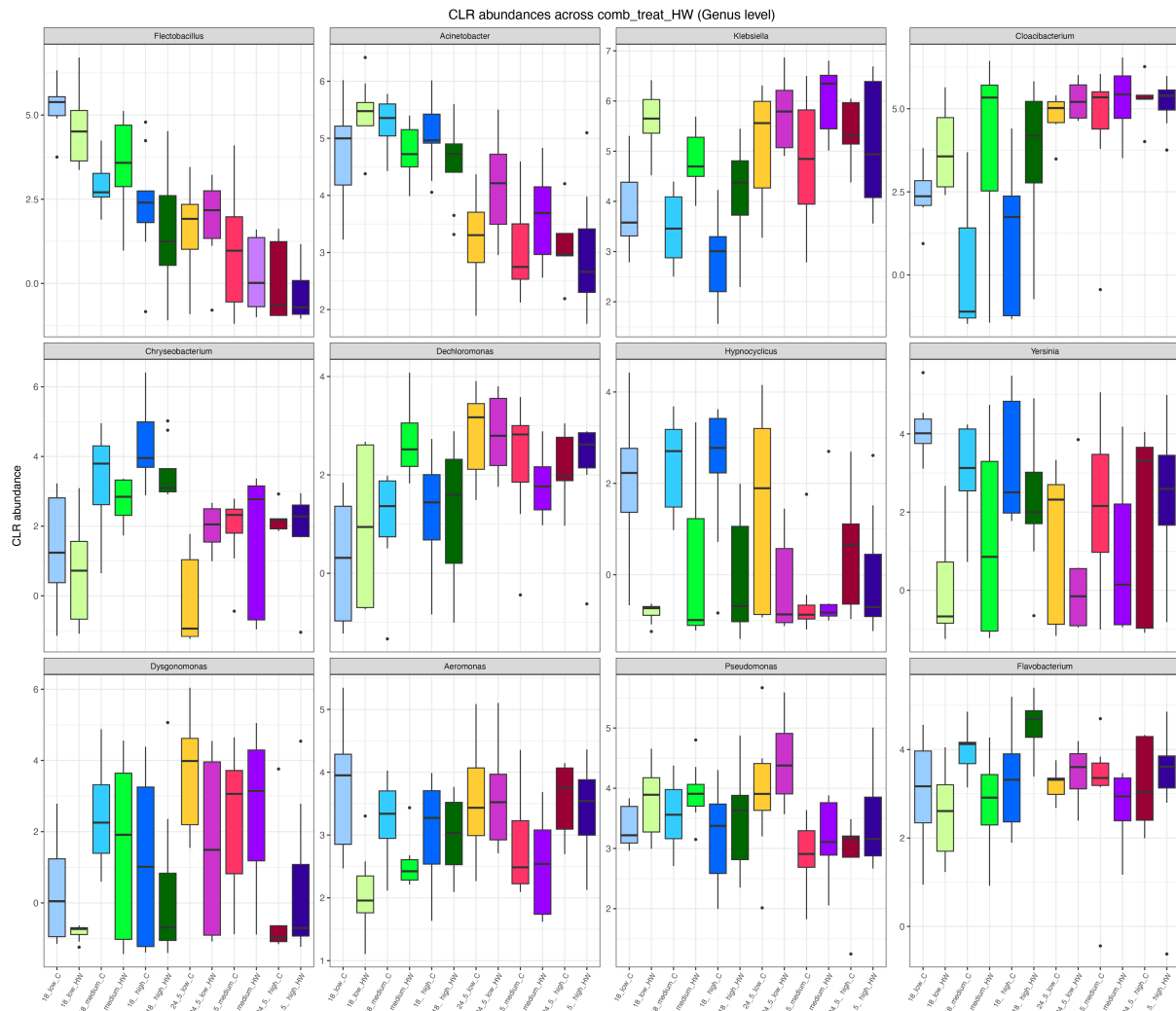
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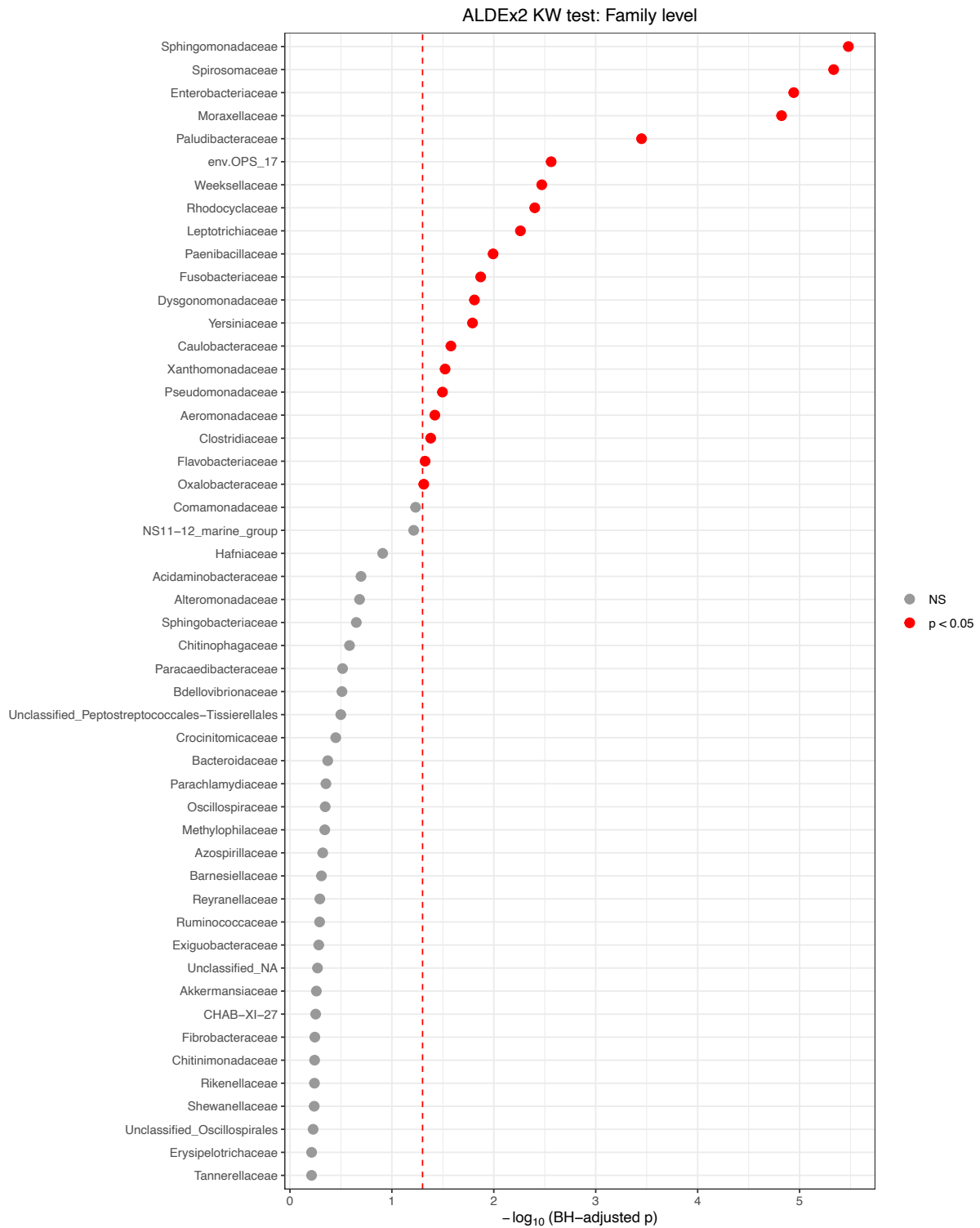
1513 Fig. S25. Genera of bacteria identified (using ALDEx2) as significantly different among
 1514 treatments comparing gut microbiomes from *Rana temporaria* larvae fed three diets with
 1515 increasing levels of protein, fat, and animal components (considered as low-, medium- and
 1516 high-quality), reared at either 18 °C or 24.5 °C and exposed or not to a heatwave, in a crossed
 1517 experimental design.



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1519 Fig. S26. Comparison of genera of bacteria in gut microbiomes from *Rana temporaria* larvae
 1520 fed three diets with increasing levels of protein, fat, and animal components (considered as
 1521 low-, medium- and high-quality), reared at either 18 °C or 24.5 °C and exposed or not (control
 1522 = C) to a heatwave (HW), in a crossed experimental design.

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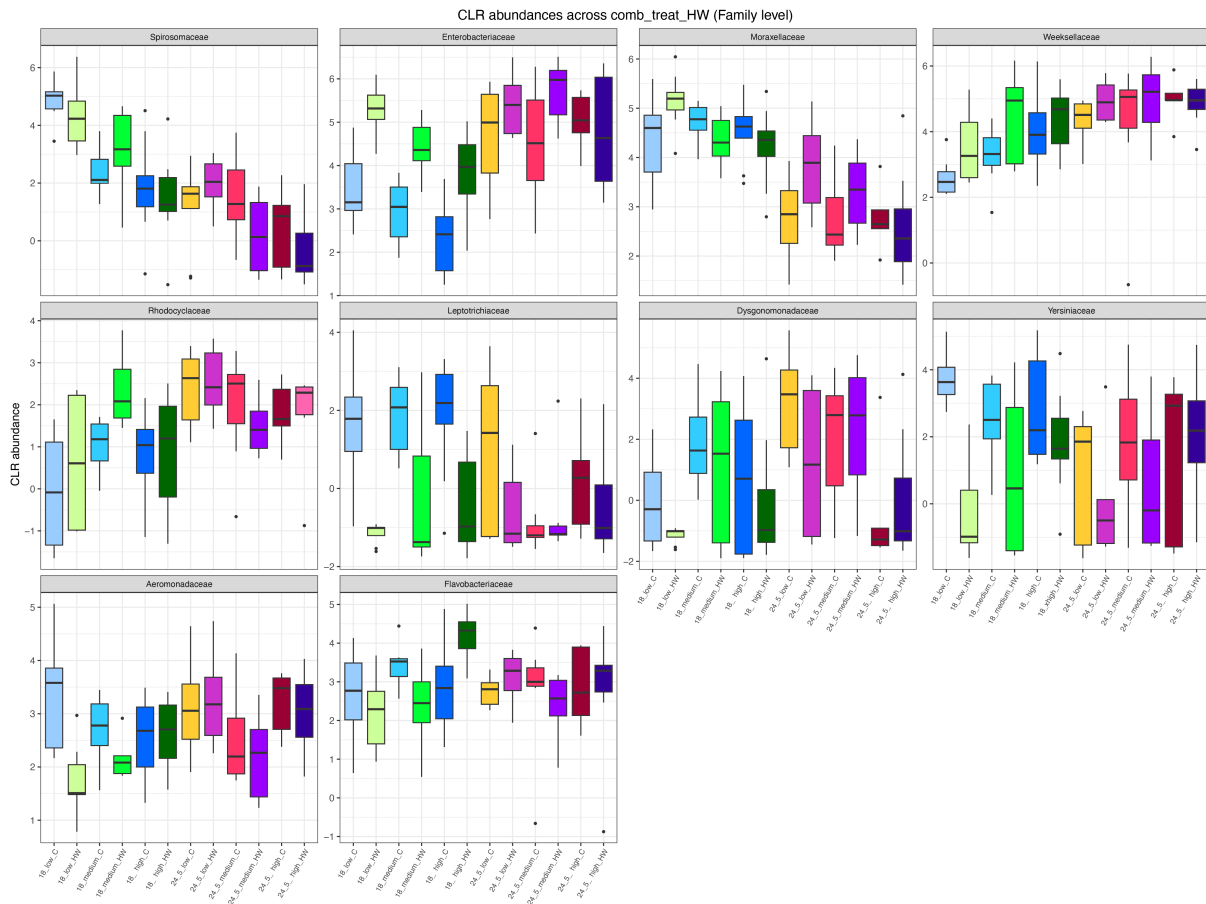


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1525 Fig. S27. Families of bacteria identified (using ALDEx2) as significantly different among
 1526 treatments comparing gut microbiomes from *Rana temporaria* larvae fed three diets with
 1527 increasing levels of protein, fat, and animal components (considered as low-, medium- and
 1528 high-quality), reared at either 18 °C or 24.5 °C and exposed or not to a heatwave, in a crossed
 1529 experimental design.

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1533 Fig. S28. Comparison of families of bacteria in gut microbiomes from *Rana temporaria*
 1534 larvae fed three diets with increasing levels of protein, fat, and animal components
 1535 (considered as low-, medium- and high-quality), reared at either 18 °C or 24.5 °C and exposed
 1536 or not (control = C) to a heatwave (HW), in a crossed experimental design.

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Table S1. Permanova pairwise comparisons among treatments applied to *Rana temporaria* larvae based on unweighted unfrac distances. Treatments corresponded to three diets with increasing levels of protein, fat, and animal components (considered as low-, medium- and high-quality), two rearing temperatures (18 °C or 24.5 °C), and exposed or not (C = control) to a heatwave (HW).

Group 1	Group 2	Sample size	Permutations	pseudo-F	p-value	q-value
18C_highC	18C_highHW	17	999	3.930	0.002	0.003
18C_lowC		17	999	2.343	0.023	0.026
18C_lowHW		18	999	7.540	0.001	0.002
18C_mediumC		16	999	1.173	0.278	0.286
18C_mediumHW		17	999	3.086	0.001	0.002
24.5C_highC		14	999	4.626	0.001	0.002
24.5C_highHW		16	999	4.023	0.001	0.002
24.5C_lowC		18	999	6.196	0.001	0.002
24.5C_lowHW		15	999	4.923	0.001	0.002
24.5C_mediumC		17	999	4.745	0.001	0.002
24.5C_mediumHW		17	999	5.242	0.001	0.002
18C_highHW	18C_lowC	16	999	4.995	0.002	0.003
18C_lowHW		17	999	4.244	0.001	0.002
18C_mediumC		15	999	5.374	0.001	0.002
18C_mediumHW		16	999	2.238	0.021	0.025
24.5C_highC		13	999	3.895	0.007	0.009
24.5C_highHW		15	999	2.699	0.012	0.015
24.5C_lowC		17	999	6.442	0.001	0.002
24.5C_lowHW		14	999	3.416	0.003	0.004
24.5C_mediumC		16	999	2.896	0.003	0.004
24.5C_mediumHW		16	999	4.640	0.002	0.003
18C_lowC	18C_lowHW	17	999	6.007	0.001	0.002
18C_mediumC		15	999	2.496	0.031	0.034
18C_mediumHW		16	999	2.567	0.010	0.013
24.5C_highC		13	999	6.103	0.001	0.002
24.5C_highHW		15	999	4.878	0.001	0.002
24.5C_lowC		17	999	6.651	0.001	0.002
24.5C_lowHW		14	999	4.584	0.003	0.004
24.5C_mediumC		16	999	5.013	0.001	0.002
24.5C_mediumHW		16	999	5.295	0.001	0.002
18C_lowHW	18C_mediumC	16	999	8.262	0.001	0.002
18C_mediumHW		17	999	3.289	0.002	0.003
24.5C_highC		14	999	5.691	0.002	0.003
24.5C_highHW		16	999	3.821	0.001	0.002
24.5C_lowC		18	999	8.517	0.001	0.002
24.5C_lowHW		15	999	2.451	0.027	0.030
24.5C_mediumC		17	999	3.688	0.002	0.003
24.5C_mediumHW		17	999	4.716	0.001	0.002

18C_mediumC	18C_mediumHW	15	999	3.181	0.002	0.003
24.5C_highC		12	999	6.624	0.002	0.003
24.5C_highHW		14	999	4.186	0.001	0.002
24.5C_lowC		16	999	8.238	0.001	0.002
24.5C_lowHW		13	999	5.406	0.001	0.002
24.5C_mediumC		15	999	4.792	0.002	0.003
24.5C_mediumHW		15	999	4.985	0.003	0.004
18C_mediumHW	24.5C_highC	13	999	3.343	0.001	0.002
24.5C_highHW		15	999	2.502	0.003	0.004
24.5C_lowC		17	999	3.319	0.004	0.005
24.5C_lowHW		14	999	1.998	0.022	0.025
24.5C_mediumC		16	999	2.708	0.002	0.003
24.5C_mediumHW		16	999	3.227	0.001	0.002
24.5C_highC	24.5C_highHW	12	999	0.817	0.644	0.644
24.5C_lowC		14	999	4.226	0.001	0.002
24.5C_lowHW		11	999	3.547	0.005	0.006
24.5C_mediumC		13	999	2.029	0.037	0.040
24.5C_mediumHW		13	999	2.342	0.017	0.021
24.5C_highHW	24.5C_lowC	16	999	5.245	0.002	0.003
24.5C_lowHW		13	999	2.052	0.015	0.019
24.5C_mediumC		15	999	1.557	0.121	0.128
24.5C_mediumHW		15	999	1.003	0.423	0.429
24.5C_lowC	24.5C_lowHW	15	999	4.575	0.001	0.002
24.5C_mediumC		17	999	4.276	0.001	0.002
24.5C_mediumHW		17	999	6.036	0.001	0.002
24.5C_lowHW	24.5C_mediumC	14	999	1.933	0.037	0.040
24.5C_mediumHW		14	999	2.296	0.022	0.025
24.5C_mediumC	24.5C_mediumHW	16	999	1.612	0.144	0.150

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