1 Vulnerability of amphibians to global warming

- 2 Patrice Pottier^{1,2*}, Michael R. Kearney³, Nicholas C. Wu⁴, Alex R. Gunderson⁵, Julie E. Rej⁵, A.
- 3 Nayelli Rivera-Villanueva^{6,7}, Pietro Pollo¹, Samantha Burke¹, Szymon M. Drobniak^{1,8+}, and
- 4 Shinichi Nakagawa^{1,9+}
- 5
- ¹ Evolution & Ecology Research Centre, School of Biological, Earth and Environmental
 Sciences, University of New South Wales, Sydney, New South Wales, Australia.
- ² Division of Ecology and Evolution, Research School of Biology, The Australian National
 University, Canberra, Australian Capital Territory, Australia
- 10 ³ School of BioSciences, The University of Melbourne, Melbourne, Victoria, Australia
- ⁴ Hawkesbury Institute for the Environment, Western Sydney University, Richmond, New South
 Wales, Australia
- ⁵ Department of Ecology and Evolutionary Biology, Tulane University, New Orleans, Louisiana,
 USA
- ⁶ Centro Interdisciplinario de Investigación para el Desarrollo Integral Regional Unidad Durango
 (CIIDIR), Instituto Politécnico Nacional, Durango, México
- ⁷ Laboratorio de Biología de la Conservación y Desarrollo Sostenible de la Facultad de Ciencias
 Biológicas, Universidad Autónoma de Nuevo León, Monterrey, México
- ⁸ Institute of Environmental Sciences, Jagiellonian University, Kraków, Poland.
- ⁹ Department of Biological Sciences, University of Alberta, Edmonton, Alberta, Canada.
- 21 ^{*}Corresponding author
- ²² ⁺These authors supervised the work equally
- 23 Corresponding author: Patrice Pottier (<u>p.pottier@unsw.edu.au</u>)

24 **ORCID**

- 25 Patrice Pottier https://orcid.org/0000-0003-2106-6597
- 26 Michael R. Kearney https://orcid.org/0000-0002-3349-8744
- 27 Nicholas C. Wu https://orcid.org/0000-0002-7130-1279
- 28 Alex R. Gunderson <u>https://orcid.org/0000-0002-0120-4246</u>
- 29 Julie E. Rej <u>https://orcid.org/0000-0002-3670-067X</u>
- 30 A. Nayelli Rivera-Villanueva https://orcid.org/0000-0002-9190-4317
- 31 Pietro Pollo <u>https://orcid.org/0000-0001-6555-5400</u>
- 32 Samantha Burke https://orcid.org/0000-0001-6902-974X
- 33 Szymon M. Drobniak <u>https://orcid.org/0000-0001-8101-6247</u>
- 34 Shinichi Nakagawa https://orcid.org/0000-0002-7765-5182

35 Summary

36 Amphibians are the most threatened vertebrates, yet their resilience to rising temperatures remains poorly understood^{1,2}. This is primarily because knowledge of thermal tolerance is 37 taxonomically and geographically biased³, compromising global climate vulnerability 38 39 assessments. Here, we employed a phylogenetically-informed data imputation approach to 40 predict the heat tolerance of 60% of amphibian species and assessed their vulnerability to daily temperature variation in thermal refugia. We found that 104 out of 5203 species (2%) are currently 41 42 exposed to overheating events in shaded terrestrial conditions. Despite accounting for heat tolerance plasticity, a 4°C global temperature increase would create a step-change in impact 43 44 severity, pushing 7.5% of species beyond their physiological limits. In the Southern Hemisphere, tropical species encounter disproportionally more overheating events, while non-tropical species 45 46 are more susceptible in the Northern Hemisphere. These findings challenge evidence for a general latitudinal gradient in overheating risk^{4–6} and underscore the importance of considering 47 48 climatic variability in vulnerability assessments. We provide conservative estimates assuming 49 access to cool shaded microenvironments. Therefore, the impacts of global warming will likely exceed our projections. Our microclimate-explicit analyses demonstrate that vegetation and water 50 bodies are critical in buffering amphibians during heat waves. Immediate action is needed to 51 52 preserve and manage these microhabitat features.

53 Keywords

54 Anura, Caudata, critical thermal maximum, behavioral thermoregulation, behavioural

55 thermoregulation, microclimate selection, biophysical modelling, global analysis, thermal safety

56 margin, warming tolerance, extreme heat events, climate change.

58 Main text

59 Climate change has pervasive impacts on biodiversity, yet the extent and consequences of this 60 environmental crisis vary spatially and taxonomically^{7,8}. For ectothermic species, such as 61 amphibians, the link between climate warming and body temperature is clear, with immediate 62 effects on physiological processes⁹. Over 40% of amphibian species are currently listed as 63 threatened, and additional pressures due to escalating thermal extremes may further increase 64 their extinction risk^{2,10}. Therefore, it is vital to assess the resilience of amphibians to climate 65 change to prioritise where and how conservation actions are taken.

66 Accurate assessments of resilience to climate change require adequate data on thermal tolerance and environmental exposure^{5,6,11}. However, the most exhaustive dataset on 67 68 amphibian heat tolerance limits only covers 7.5% of known species and is geographically biased towards temperate regions³ (Fig. 1). This discrepancy is problematic, considering the 69 70 high species richness in the tropics and the mounting evidence that tropical ectotherms are most susceptible to rising temperatures^{4–6,12,13}. Such sampling biases call into question the 71 reliability of inferences in under-sampled areas and have implications for conservation 72 73 strategies. Given the rapid pace of climate change and the finite resources available for 74 research, acquiring sufficient empirical data to fill these knowledge gaps within a realistic timeframe is increasingly untenable^{14,15}. Therefore, alternative methods to identify the 75 populations and areas most susceptible to thermal stress are critically needed in a rapidly 76 77 warming climate.

Climate vulnerability assessments also require environmental data with high spatial and temporal resolution, particularly because extreme heat is more likely to trigger overheating events than increased mean temperatures^{16–18}. When heat tolerance limits are known, cuttingedge approaches in biophysical ecology allow fine-scale vulnerability assessments that account for morphology, behaviour, and microhabitat setting in both historical and future climate projections^{19,20}. While broadly applicable, biophysically informed analyses are particularly relevant for amphibians, whose body temperatures depend on evaporative heat loss and whose

microhabitat use span terrestrial, aquatic, and arboreal environments. Because
microenvironmental features are essential for behavioural thermoregulation^{21,22}, modelling
microhabitats allow assessments of the effectiveness of different thermal refugia in buffering the
impacts of extreme heat events.

Here, we assess the global vulnerability of amphibians to extreme heat events in
different climatic scenarios and thermal refugia. By integrating predicted thermal limits for 60%
of amphibian species with daily operative body temperatures, our study offers the first
comprehensive evaluation of the impact of heat extremes on the physiological viability of
amphibians in nature.

94 Thermal limits and environmental exposure

95 We first developed an approach to predict standardised thermal limits for 5,203 amphibian 96 species using data imputation based on phylogenetic niche clustering (Pagel's λ = 0.95 [0.91 – 0.98]) and known correlations between critical thermal limits (CT_{max}) and other variables (n = 97 2,661 estimates measured in 524 species; Fig. S2; Methods). Our phylogenetic model-based 98 99 imputation approach has expanded our understanding of amphibian thermal tolerance by 100 generating testable predictions for 4,679 unstudied species, particularly in biodiversity hotspots 101 (Fig. 1-2). We confirmed our imputation approach was accurate and unbiased by demonstrating 102 a strong congruence between experimental and imputed data in cross-validations (experimental 103 mean \pm standard deviation = 36.19 \pm 2.67; imputed mean = 35.93 \pm 2.54; n = 375; r = 0.86; 104 Extended Data Fig. 2a,b), though, as expected, the uncertainty in imputed predictions was 105 higher in understudied clades (Extended Data Fig. 2c).

We then integrated predicted thermal limits with daily maximum operative body temperature fluctuations estimated from biophysical models to evaluate the sensitivity of amphibians to extreme heat events in terrestrial, aquatic, and arboreal microhabitats (Extended Data Fig. 1; Methods). Operative body temperatures are the steady-state body temperatures that organisms would achieve in a given microenvironment, which can diverge significantly from ambient air temperatures due to, for example, radiative and evaporative heat exchange

processes^{19,20}. For each microhabitat, we modelled daily operative body temperatures during 112 the warmest quarters of 2006-2015 and across the distribution range of each species 113 114 (Methods). We also used projected future climate data from TerraClimate²³ to generate 115 projections assuming 2°C or 4°C of global warming above pre-industrial levels. These 116 temperatures are within the range projected by the end of the century under low and intermediate/high greenhouse gas emission scenarios, respectively²⁴. Notably, recent historical 117 CO₂ emissions most closely align with high warming scenarios²⁵ (i.e., 4.3°C of predicted 118 119 warming by 2100). All microenvironmental projections assumed access to 85% of shade and 120 sufficient humidity to maintain wet skin to simulate amphibians in thermal refugia (Methods).

121 We estimated the vulnerability of amphibians by estimating daily differences between 122 predicted thermal limits and maximum hourly operative body temperatures (Extended Data Fig. 123 1; Methods). We also adjusted daily thermal limits to assume that species were, on any given 124 day, acclimated to local mean weekly operative body temperatures, effectively accounting for 125 plasticity throughout species' distribution ranges (Methods). In total, we predicted vulnerability metrics for 203,853 local species occurrences (individual species in 1° x 1° grid cells) in 126 127 terrestrial conditions (5,177 species), 204,808 local species occurrences in water bodies (5,203 128 species); and 56,210 local species occurrences (1,771 species) in above-ground vegetation, for 129 each warming scenario. The number of species examined in arboreal conditions was lower to 130 reflect morphological adaptations required for climbing in above-ground vegetation. These 131 estimates were then grouped into assemblages (all species occurring in 1° x 1° grid cells), 132 tallying 14,090 and 14,091 assemblages for terrestrial and aquatic species and 6,614 133 assemblages for arboreal species, respectively.

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135

- 136 Fig. 1 | Contrast between the geographical locations at which experimental data were
- 137 **collected and patterns in species richness.** Pink points denote experimental data (587
- species), while the colour gradients refer to species richness calculated in 1 x 1 ° grid cells in
- the imputed data (5,203 species). Density plots on the right panel represent the distribution of
 experimental data (pink) and the number of species inhabiting these areas (blue) across
- 141 latitudes. Dashed lines represent the equator and tropics.
- 142



144

145 Fig. 2 | Phylogenetic coverage and taxonomic variation in climate vulnerability. Heat

146 maps show heat tolerance limits (CT_{max}) and thermal safety margins (TSM), while histograms

show the number of overheating events (days) averaged across each species' distribution

range (n = 5,177 species). Pink bars refer to species with prior knowledge (n = 521), while grey

bars refer to entirely imputed species (n = 4,656). This figure was constructed assuming

150 ground-level microclimates occurring under 4°C of global warming above pre-industrial levels.

Phylogeny is based on the consensus of 10,000 trees sampled from a posterior distribution (see for details). Highlighted species starting from the right side, anti-clockwise: *Neurergus kaiseri*,

153 Plethodon kiamichi, Bolitoglossa altamazonica, Cophixalus aenigma, Tomaptera cryptotis,

154 Lithobates palustris, Allobates subfolionidificans, Phyzelaphryne miriamae, Barycholos ternetzi,

155 Pristimantis carvalhoi, Pristimantis ockendeni, Boana curupi, Teratohyla adenocheira, Atelopus

- 156 spumarius.
- 157

159 Vulnerability to historical and future heat

We first calculated thermal safety margins (TSM, sensu⁶) as the weighted mean difference 160 161 between heat tolerance limits (CT_{max}) and the maximum daily body temperatures of the warmest 162 quarters of 2006-2015 for each local species occurrence. Thermal safety margins averaged 163 from long-term climatology are routinely used in climate vulnerability analyses^{27–29}. We found evidence for a decline in TSM towards mid to low latitudes in all microhabitats, a pattern 164 165 maintained across warming scenarios (Fig. 3, Extended Data Fig. 3). However, warming substantially reduce TSM at all latitudes (Fig. 3), likely reflecting the contrast between weak 166 plastic responses in CT_{max} across latitudes^{11,15} (Extended Data Fig. 3; Fig. S3) and large 167 variation in environmental temperatures (Extended Data Fig. 3). Across all conditions simulated, 168 TSM is always positive, even in the highest warming scenario (Fig. 3, Extended Data Fig. 3). 169 170 The mean TSM is lower for terrestrial (mean [95% confidence intervals]; current = 11.69 [8.86 – 171 14.43; $+4^{\circ}C = 9.41$ [6.53 - 12.09]) and arboreal conditions (current = 12.23 [9.40 - 14.96]; $+4^{\circ}C = 10.07 [7.23 - 12.80]$) than for water bodies (current = 13.60 [10.71 - 16.28]; $+4^{\circ}C =$ 172 11.68 [8.80 – 14.36]; Fig. 3; Extended Data Table 1). 173



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176 Fig. 3 | Assemblage-level patterns in thermal safety margin for amphibians in terrestrial (a), aquatic (b) or arboreal (c) microhabitats. Thermal safety margins (TSM) were calculated 177 178 as the weighted mean difference between CT_{max} and the predicted operative body temperature 179 in full shade during the warmest quarters of 2006-2015 in each assemblage (1-degree grid cell). Black colour depicts areas with no data. The right panel depicts latitudinal patterns in TSM in 180 current climates (blue) or assuming 4°C of global warming above pre-industrial levels (pink), as 181 predicted from generalised additive mixed models. Point estimates are scaled by precision 182 183 (1/s.e.). Dashed lines represent the equator and tropics.

Because extreme heat events are more likely to trigger overheating events than mean 184 temperatures^{5,6,11}, we also calculated the binary probability (0/1) that operative body 185 temperatures exceeded CT_{max} for at least one day across the warmest guarters of 2006-2015 186 187 (i.e., overheating risk). Overall, overheating risk is low, although numerous species are predicted to face overheating events locally (Fig. 4, Extended Data Table 2). In terrestrial 188 189 conditions, we predict that 104 species (836 local species occurrences from 253 assemblages) 190 are likely to experience overheating events in current microclimates (Fig. 4-5). However, under 191 4°C of warming, 391 species (4,248 local species occurrences from 1,328 assemblages) are 192 expected to overheat, which represents nearly a four-fold increase relative to current conditions (Fig. 4-5; Extended Data Table 2-3). The number of species predicted to overheat in each grid 193 194 cell also increases with warming; each assemblage comprises up to 18 vulnerable species in 195 current climates (mean [95% confidence intervals] = 3.19 [0.60 - 6.88] species) and up to 37 vulnerable species with 4°C of global warming (3.08 [0.62 – 6.56]; Fig. 4; Extended Data Table 196

3). In addition, the proportion of species predicted to experience overheating events in each
assemblage varies geographically and between warming scenarios (Extended Data Fig. 5;
Extended Data Table 4). The proportion of species at risk is high in some areas with high
species richness (e.g., Northern Australia, Southeastern United States) and not linearly
predicted by latitude (Extended Data Fig. 5).

202 In current conditions for species that can shelter in trees (arboreal), 74 assemblages 203 (comprising 1-6 species; 1.93 [0.05 – 5.05] species) are predicted to overheat, while 285 assemblages (comprising 1-11 species; 2.51 [0.31 – 5.69] species) are predicted to overheat 204 205 assuming 4°C of global warming (Fig. 4; Extended Data Table 3). While the overheating risk is 206 lower in arboreal conditions, considerably fewer species were examined than in terrestrial 207 conditions (1,771 vs. 5,177 species). In fact, comparing the responses of arboreal species in 208 different microhabitats revealed that occupying above-ground vegetation is only partially 209 beneficial (Extended Data Fig. 4). In current climates, up to 15 arboreal species (320 local 210 species occurrences) are predicted to experience overheating events in terrestrial conditions, 211 whereas 13 arboreal species (152 local species occurrences) are predicted to overheat in 212 above-ground vegetation (Extended Data Fig. 4). Furthermore, under 4°C of warming, 83 213 arboreal species (1,137 local species occurrences) are predicted to overheat in terrestrial 214 conditions, while retreating to above-ground vegetation only reduces the number of species 215 exposed to overheating events by 32.5% (56 species, 748 local species occurrences) 216 (Extended Data Fig. 4). Contrary to terrestrial and arboreal conditions, no amphibian 217 populations are predicted to overheat in water bodies in current or intermediate climate warming scenarios due to the thermal buffering properties of water. However, assuming 4°C of climate 218 219 warming, we predict that 11 species (56 local species occurrences from 48 assemblages) will 220 exceed their physiological limits in aquatic microhabitats (Fig. 4).

221



223 Fig. 4 | Number of species predicted to experience overheating events in terrestrial (a), aquatic (b), and arboreal (c) microhabitats. The number of species overheating was 224 assessed as the sum of species overheating for at least one day in the period surveyed 225 226 (warmest quarters of 2006-2015) in each assemblage (1-degree grid cell). Black colour depicts 227 areas with no data, and grey colour assemblage without species at risk of overheating. The right 228 panel depicts latitudinal patterns in the number of species predicted to overheat in current 229 climates (blue) or assuming 4°C of global warming above pre-industrial levels (pink). Dashed 230 lines represent the equator and tropics.

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231 Finally, we quantified the number of days (out of 910 simulated days across the warmest 232 quarters of 2006-2015) each species was predicted to locally exceed their plasticity-adjusted 233 heat tolerance limits. This metric fully integrates the frequency at which amphibians are predicted to experience temperatures beyond their thermal limits. For current climates, we 234 235 found that species rarely experience overheating events in shaded terrestrial conditions (overall mean overheating days [95% confidence intervals] = 0.01 [0.01 - 0.08]; mean among 236 237 overheating species = 2.15 [0.24 - 5.26] days); but these figures increase considerably with global warming (Fig. 5; Extended Data Table 2). Under 4°C of warming, species are predicted 238 239 to overheat on as many as 207.18 [182.39 – 231.97] days, representing up to 22.8% of the 240 warmest days of the year (overall mean = 0.15 [0.05 - 0.46] days; mean among overheating species = 6.75 [3.14 – 11.38] days; Fig. 5; Extended Data Table 2). This is noticeably more than 241 what is predicted under 2° C of warming (overall mean = 0.02 [0.01 - 0.13] days; mean among 242 243 overheating species = 2.58 [0.41 – 5.86] days; Fig. 5; Extended Data Table 2). In above-ground 11

244 vegetation, the frequency of overheating events is lower, as expected. Under current climates, 245 arboreal species are predicted to overheat on up to 5.65 [1.00 – 10.29] days in total (overall 246 mean = 0.01 [0.01 - 0.04] days; mean among overheating species = 1.62 [0.03 - 4.43] days; 247 Fig. 5; Tab. Extended Data Table 2). Under 4 degrees of warming, arboreal species are 248 predicted to overheat on up to 76.17 [59.79 – 92.54] days (overall mean = 0.08 [0.01 – 0.23] 249 days; mean among overheating species = 5.08 [1.81 – 9.39] days; Fig. 5; Extended Data Table 250 2). Arboreal species retreating to above-ground vegetation are predicted to experience fewer 251 overheating events than those in terrestrial conditions (Extended Data Fig. 4). Interestingly, we 252 found that species predicted to overheat locally have TSMs well above zero, although some are 253 living particularly close to their heat tolerance limits during the warmest months in both 254 terrestrial (mean [95% confidence intervals]; current = 8.20 [6.91 - 9.98], range: 3.02 - 12.19; +4°C = 6.30 [5.02 – 8.09], range: 0.97 – 11.27) and above-ground conditions (current = 8.71 255 256 [7.20 – 10.28], range: 3.70 – 9.76; +4°C = 6.73 [5.44 – 8.48], range: 1.75 – 8.70; Fig. 5c,d). 257 Finally, we found a strong non-linear negative association between the number of overheating 258 events and the thermal safety margin, with stark contrasts between warming scenarios (Fig. 259 5c,d; Extended Data Table 5). In particular, overheating days increase rapidly as thermal safety 260 margins fall below 5°C (Figure 5c,d).





264 Fig. 5 | Latitudinal variation in the number of overheating events in terrestrial (a,c) and 265 arboreal (b,d) microhabitats as a function of latitude (a,b) and thermal safety margin (c,d). 266 The number of overheating events (days) were calculated based on the mean probability that 267 daily maximum temperatures exceeded CT_{max} during the warmest quarters of 2006-2015 for each species in each grid cell. Blue points depict the number of overheating events in current 268 microclimates, while orange and pink points depict the number of overheating events assuming 269 2°C and 4°C of global warming above pre-industrial levels, respectively. For clarity, only the 270 271 species predicted to experience at least one overheating event are depicted across latitudes 272 (a,b).

273

274 The mounting impacts of global warming

275 Quantifying the resilience of biodiversity to a changing climate is one of the most pressing

- challenges for contemporary science^{7,8}. Here, we show that over a hundred species may
- 277 already experience hourly temperatures that would likely result in death over minutes or hours
- of exposure in thermal refugia. This pattern is only predicted to worsen (Fig. 4-5). Assuming 4°C
- of global warming, the number of species and assemblages exposed to overheating events
- would be four to five times higher than currently, totalling 391 out of 5,203 species studied
- 281 (7.5%; Fig. 4-5).

282 We also found striking disparities in overheating risk between the 2°C and 4°C warming 283 projections (Fig. 5; Extended Data Table 1), which are anticipated by the end of the century 284 under low and high greenhouse gas emission scenarios, respectively²⁴. The more extreme 285 warming scenario considerably increased the number overheating events experienced by 286 amphibian populations (Fig. 5), highlighting the escalating and abrupt impacts of global warming^{7,30}. Such an increase is attributable to the contrast between the rapid pace at which 287 288 temperatures are increasing and the low ability of amphibians to acclimate to new thermal 289 environments via plasticity (Extended Data Fig. 3; Fig. S3). Our study clearly demonstrates, as others have suggested^{18,28,31,32}, that physiological plasticity is not a sufficient mechanism to 290 buffer many populations from the impacts of rapidly rising temperatures. 291

292 Extreme heat events drive vulnerability

293 We found large spatial heterogeneity in the vulnerability of amphibians. In tropical areas, most 294 vulnerable species are concentrated in South America and Australia, whereas fewer species 295 are impacted in the African and Asian tropics (Fig. 4). Tropical species also experience 296 disproportionately more overheating events in the Southern Hemisphere, while non-tropical 297 species are more susceptible in the Northern Hemisphere (Fig. 5). Furthermore, the proportion 298 of species experiencing overheating events in each assemblage was not predicted by latitude 299 (Extended Data Fig. 5). Therefore, our findings are inconsistent with the expectation of a 300 general latitudinal gradient in overheating risk based on thermal safety margins^{4–6,13}. In fact, the 301 overheating risk does not increase linearly with TSM (Fig. 5c,d), and species with seemingly 302 comparable TSMs can have markedly different probabilities of overheating due to varying 303 exposure to daily temperature fluctuations (Fig. 5c,d). Therefore, TSMs alone hide critical 304 tipping points for thermal stress (Fig. 5c,d).

305 Our study questions the reliability of thermal safety margins and other climate 306 vulnerability metrics when averaged across large time scales (e.g., using the maximum 307 temperature of the warmest quarter) for detecting species most vulnerable to thermal extremes. 308 It also challenges the general notion that low-latitude species are uniformly most vulnerable to warming^{4-6,13}, revealing a far more nuanced pattern of climate vulnerability across latitudes.
While the reliability of TSM-based assessments has been questioned in previous studies¹¹, our
work further emphasises the need to consider natural climatic variability and extreme hourly
temperatures^{4,16-18} when evaluating the vulnerability of ectotherms to global warming.
Considering alternative metrics, such as the number of predicted overheating events, may
prove particularly useful in identifying the most vulnerable species and populations.

315 The vital yet limited role of thermal retreats

316 Our study highlights the critical yet sometimes insufficient role that thermal retreats play in 317 buffering the impacts of warming on amphibians. Most amphibian species are predicted not to 318 experience overheating events in full shade (Fig. 4), and the availability of water bodies allows 319 nearly all amphibians to maintain their body temperatures below critical levels, apart from 11 320 species in the most extreme warming scenario investigated. This is attributable to the higher 321 specific heat capacity of water relative to air, delaying rapid temperature rises and affording a more stable environment during heat waves³³. Our findings add to the growing evidence that 322 323 finding access to cooler microhabitats is the main strategy amphibians and other ectotherms can use to maintain sub-lethal body temperatures^{6,21,34}. 324

325 However, it is crucial to emphasise that vegetated terrestrial conditions in full shade offer 326 inadequate protection to 7.5% of species, and many arboreal species predicted to overheat at 327 ground level face similar risks in above-ground vegetation (Fig. 4-5, Extended Data Fig. 4). In 328 fact, although reducing the frequency of overheating events (Extended Data Fig. 4), access to 329 shaded above-ground vegetation only reduces the number of vulnerable species by 32.5%. 330 Moreover, although burrows offer cooler microclimates (see Fig. S9), the ability to use 331 underground spaces is not universal among amphibians and can greatly restrict activity. 332 reproduction, and foraging opportunities.

333 Warming impacts may exceed projections

334 Our predictions are largely conservative, and likely overestimate the resilience of amphibians to 335 global warming in two main ways. First, we assume that microhabitats such as shaded ground-336 level substrates, above-ground vegetation, and water bodies are available throughout a species' 337 range, and that amphibians can maintain wet skin. These assumptions will often be violated as habitats are degraded. Deforestation and urbanization are diminishing vital shaded areas^{35,36}, 338 339 while increased frequencies of droughts will cause water bodies to evaporate^{37,38}. These 340 changes compromise not only habitat integrity but also local humidity levels - key for effective thermoregulation^{39,40}. Consequently, amphibians will likely experience higher body temperatures 341 and desiccation stress events than our models predict due to inconsistent access to cooler 342 microhabitats⁴¹, particularly in degraded systems. 343

Second, ectotherms can experience deleterious effects from heat stress before reaching their heat tolerance limits. Prolonged exposure to sub-lethal temperatures can lead to altered activity windows^{42,43}, disruptions to phenology^{44,45}, reduced reproductive fitness (fertility and fecundity)^{29,46,47}, and death^{48,49}. Although comprehensive data on thermal incapacitation times and fertility impacts are sparse in amphibians, integrating both the duration and intensity of thermal stress^{49–51} will likely point to more extreme vulnerability estimates. This represents a vital avenue for future research, albeit one requiring a large collection of empirical data.

Alternatively, species that can retreat underground during heat events are likely to experience fewer overheating events than our models predict (see Fig. S9), and prolonged exposure to high temperatures in the permissive range (*sensu* ⁴⁸) can enhance performance and fitness, thereby reducing the impacts of extreme heat on natural populations. In addition, some species may adapt to changing temperatures. However, evidence for slow rates of evolution and physiological constraints on thermal tolerance^{52,53} challenges the likelihood of local adaptation to occur in rapidly warming climates.

358 The power of data imputation

359 Our imputation approach has generated testable predictions of the thermal limits of 5.203 species, expanding the scope of previous research³ (Fig. 2). We also addressed geographical 360 361 biases by generating predictions in under-sampled but ecologically critical regions of Africa, 362 Asia, and South America (Fig. 2). We found that these understudied regions frequently harbor 363 species exhibiting the highest susceptibility to extreme heat events (Fig. 1.4-5), with 74% (288) 364 out of 391) of vulnerable species remaining unstudied. Targeted research efforts in these 365 vulnerability hotspots are instrumental in validating our model predictions and advancing our 366 understanding of amphibian thermal physiology to inform their conservation. Though underlable 367 logistical and financial challenges exist in accessing some of these remote locations. 368 collaboration with local scientists could expedite data collection and result in timely conservation measures. Exemplary initiatives to sample numerous species in South America (e.g., ^{22,54,55}) are 369 370 promising steps in this direction, and we hope our findings will catalyse research activity in 371 these regions.

372 Amphibian biodiversity in a warming world

373 Our study highlights the dire consequences of global warming on amphibians. Yet it is crucial to 374 differentiate between global extinction and local extirpations – the latter being confined 375 extinctions within specific geographic areas. Most species will not experience overheating 376 events throughout their entire range, and these overheating events may not occur 377 simultaneously. Hence, most species are likely to only experience local extirpation due to 378 overheating, according to our models. Nevertheless, local extirpations carry their own sets of 379 ecological repercussions, such as reshuffling community compositions and eroding genetic and 380 ecological diversity^{56,57}.

381 Some amphibian populations may also undergo range shifts, permanently or transiently 382 relocating to habitats with more hospitable weather patterns⁵⁸. However, this is only possible if 383 suitable habitats are available for establishment. Given the low dispersal rates of some 384 amphibians and their common reliance on water bodies for reproduction and thermoregulation, opportunities for range shifts are likely to be rare for many species. Identifying which species at
high risk of overheating are simultaneously predicted to have limited ability to extend their range
is an interesting avenue for research. In addition, we stress that amphibians living close to their
physiological limits for extended times at the warm edge of their distribution are likely to
experience heat stress that could hamper activity, foraging opportunities, and reproductive
success, adding layers of complexity to their survival challenges and potentially leading to
population declines^{42,48}.

392 Overall, our study contributes to the evidence that climate change is a mounting threat to 393 amphibians^{2,10} and emphasises the importance of limiting global temperature rises below 2°C to 394 minimise the risk of overheating to amphibian populations. A 4°C temperature rise would not 395 just increase these risks but create a step-change in impact severity (e.g., Fig. 5c). The 396 mechanistic basis of our species- and habitat-specific predictions also leads to clear 397 management priorities. Particularly, our analyses revealed the critical importance of preserving 398 dense vegetation cover and water bodies. These microhabitats provide conditions with cooler 399 and more stable temperatures and increase the potential for amphibians and other ectothermic 400 species to disperse to more suitable microhabitats. Establishing protected areas and 401 undertaking habitat restoration initiatives may support amphibians in a changing climate and 402 buffer additional anthropogenic threats, in turn mitigating amphibian population declines^{2,10,59}. These actions are critical for the amphibians at risk and the ecosystems they support⁶⁰ in a 403 404 planet undergoing perilous climatic changes.

405

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545 Methods

546 *Reporting*

547 We report author contributions using the CRediT (Contributor Roles Taxonomy) statement⁶¹ and 548 MeRIT (Method Reporting with Initials for Transparency) guidelines⁶². We also crafted the study 549 title, abstract and keywords to maximise indexing in search engines and databases⁶³. All 550 analyses were performed using R statistical software⁶⁴ (v. 4.3.0), and most computations used 551 the computational cluster Katana supported by Research Technology Services at UNSW 552 Sydney.

553 Amphibian heat tolerance limits

554 We leveraged the most comprehensive compilation of amphibian heat tolerance limits³ for our 555 analyses (Extended Data Fig. 1). Briefly, these data were collated by systematically reviewing 556 the literature in five databases and seven languages, comprising 3,095 heat tolerance limits 557 from 616 amphibian species. To facilitate the comparability and analysis of heat tolerance limits, 558 we only included data matching four specific criteria. First, we only included heat tolerance limits 559 measured using a dynamic methodology (i.e., temperature at which animals lose their motor coordination when exposed to ramping temperatures, critical thermal maximum CT_{max}^{65}) 560 561 because it was the most used and comparable metric. Second, we only selected data for which 562 the laboratory acclimation temperature, or the field temperature during the month of capture, was recorded. Third, we only included data from species listed in the phylogeny from ²⁶. Fourth, 563 we only included species for which their geographical range was reported in the International 564 Union for the Conservation of Nature red list⁶⁶ (accessed in January 2023). 565

These criteria were chosen to perform phylogenetically, climatically, and spatially informed
analyses. In total, we selected 2,661 heat tolerance limits estimates with metadata for 524
amphibian species (mean = 5.08; range = 1 - 146 estimates per species; 287 species with more
than one estimate). We also complemented this dataset with ecotypic data for each species.
Amphibians were grouped into six major ecotypes according to ⁴¹: ground-dwelling, fossorial,

aquatic, semi-aquatic, stream-dwelling and arboreal. Cave specialists were excluded becausethey experience unique microclimatic conditions.

573 Data-deficient species

574 Our objective was to assess the thermal tolerance of amphibians globally. However, the data 575 compiled in ³ are geographically and taxonomically biased. Therefore, we employed a data 576 imputation procedure to infer the thermal tolerance of data-deficient species, totalling 5,203 577 species at a broad geographical coverage (524 species + 4,679 data-deficient species; ~60% of 578 all described amphibian species, amphibiaweb.org; accessed in December 2023). We selected data-deficient species from a species list that matched the phylogeny from ²⁶ (7,238 species), 579 was listed in the IUCN red list⁶⁶ along with geographic distribution data (5,792 species), and for 580 which ecotypes were known (6,245 species). We did not consider Caecilians (order 581 582 Gymnophiona) because, to our knowledge, heat tolerance limits are unknown for all Caecilian 583 species³. Of the 5,792 species for which we had distribution and phylogenetic data, 5,268 were 584 data-deficient for CT_{max}, of which 4,822 had a known ecotype. After removing Caecilians, we 585 were left with 4,679 species to impute. We also supplemented our dataset with published body 586 mass data retrieved from literature sources or estimated based on length-mass allometries^{41,67,68}. We then estimated the geographical coordinates at which all extant species 587 588 occurred in their IUCN distribution range at a 1° x 1° resolution to use for biophysical modelling 589 (Extended Data Fig. 1).

590 Data imputation

We developed a phylogenetic imputation procedure, here named Bayesian Augmentation with Chained Equations (BACE). The BACE procedure combines the powers of Bayesian data augmentation and multiple imputation with chain equations (MICE⁶⁹). Briefly, we ran multiple iterative models using *MCMCgImm*⁷⁰ (v. 2.34) and supporting functions from the *hmi* package⁷¹. In the first cycle, missing data was either taken as the arithmetic mean for continuous predictors, or randomly sampled from existing values for (semi)categorical predictors. Predicted

597 (augmented) values from the models were then extracted from the response variables and used 598 as predictor variables in the next models to predict other response variables. Ultimately, heat 599 tolerance limits were predicted using augmented data from all predictors. We ran 5 cycles 600 where the data from one cycle was iteratively used in the next cycle, and estimations converged 601 after the first cycle (Fig. S1). Although the proportion of missing data was large (89.9%), imputations based on large amounts of missing data are common^{13,72}, and although estimate 602 603 uncertainty increases with the proportion of missing data, as expected, simulation studies have 604 shown estimations remain unbiased^{73,74}. Note, however, that although our approach took the 605 uncertainty of missing data in the response or variable of interest (CT_{max}) into account, we used 606 the most likely values for the predictors. While such an approach could underestimate the 607 uncertainty in the response, point estimates should not be biased. In fact, our cross-validation 608 approach demonstrated the ability of our models to predict back known experimental estimates 609 with reasonable error (experimental mean \pm standard deviation = 36.19 \pm 2.67; imputed mean = 610 35.93 ± 2.54 ; r = 0.86; Extended Data Fig. 2).

611 Heat tolerance limits were imputed based on the species' acclimation temperatures, the 612 duration of acclimation, the ramping rate and endpoint used in assays, the medium used for 613 measuring heat tolerance limits (i.e., ambient temperatures, water/body temperatures), and the 614 life stage of the animals (adults or larvae), and their ecotype. These variables were correlated 615 with amphibian heat tolerance limits (Fig. S2) and were fitted as covariates in Bayesian linear 616 mixed models. We also weighted heat tolerance estimates based on the inverse of their 617 sampling variance, accounted for phylogenetic non-independence using a correlation matrix of phylogenetic relatedness, and fitted random intercepts for species-specific effects and 618 619 phylogenetic effects, as well as their correlation with acclimation temperatures (i.e., random 620 slopes). In other words, we modelled species-specific slopes (plasticity; see Fig. S2) and 621 partitioned the variance among phylogenetic and non-phylogenetic effects. We imputed data for adult amphibians assuming they were acclimated to the median, 5th, or 95th percentile operative 622 623 body temperatures experienced across their geographical range (see *Microenvironmental data*

624 and biophysical modelling) for a duration of 10 days, tested using a ramping rate of 1°C/min in a 625 container filled with water, and for which thermal tolerance endpoint was recorded as the onset 626 of spasms. These methodological parameters were the median values in the experimental 627 dataset, or the most common values (mode). This allowed standardization of heat tolerance limits for the comparative analysis^{75–77}. In amphibians, the onset of spasms usually occurs after 628 the loss of righting response⁷⁸, meaning that our estimates are conservative. While we did 629 630 include data from larvae in the training data, we only imputed data for adults to increase the 631 comparability of our estimates.

For both known species and data-deficient species, we generated three ecologically relevant and standardised heat tolerance estimates, and all analyses were built upon these standardised imputed estimates. In total, we generated data for 5,203 species of amphibians (Extended Data Fig. 1-2). Notably, our imputed estimates are accompanied by standard errors, which provide estimates of uncertainty in the imputation, and errors were propagated throughout our analyses (see *Climate vulnerability analysis*).

638 Microenvironmental data and biophysical modelling

We used the package *NicheMapR*^{79,80} (v. 3.2.1) to estimate microenvironmental temperatures and hourly operative body temperatures in current (2006-2015) and projected climatic conditions (2°C or 4°C of global warming above pre-industrial levels). Operative body temperatures are the steady-state body temperatures that organisms would achieve in a given microenvironment, which can diverge significantly from ambient air temperatures due to, for example, radiative and evaporative heat exchange processes^{19,20,81–86}.

For each geographic location, we generated microclimatic temperatures experienced by amphibians on i) a vegetated ground-level substrate (i.e., terrestrial), ii) in above-ground vegetation (i.e., arboreal), or iii) in a water body (i.e., aquatic) (Extended Data Fig, 1). For terrestrial and aquatic species, we simulated microenvironmental temperatures 1 cm above the surface. For arboreal species, we simulated microenvironmental temperatures 2 meters above 650 ground, applied a reduction of 80% in windspeed to account for reduced wind due to vegetation⁸⁷, and assumed that 90% of the solar radiation was diffused due to canopy cover⁸⁸. 651 652 All microenvironmental projections were made using 85% shade to simulate animals in thermal 653 refugia, i.e., the microhabitats in which animals would retreat during the hottest times of the day. 654 We did not model temperatures in the sun because ectothermic species most likely behaviourally thermoregulate by retreating to thermal refugia during extreme heat events²¹. Our 655 656 calculations thus represent conservative estimates of the vulnerability of amphibians to extreme 657 temperature events.

658 For microclimatic temperature estimates, we used the *micro_ncep* function from *NicheMapR*⁷⁹ 659 (v. 3.2.1), which integrates 6-hourly macroclimatic data from the National Center for Environmental Predictions (NCEP). This function also inputs from the *microclima* package⁸⁹ (v. 660 661 0.1.0) to predict microclimatic temperatures after accounting for variation in radiation, wind 662 speed, altitude, albedo, vegetation, and topography. These data are downscaled to an hourly 663 resolution, producing high-resolution microclimatic data. We used projected future monthly climate data from TerraClimate²³ to generate hourly projections assuming 2°C or 4°C of global 664 665 warming above pre-industrial levels. These temperatures are within the range projected by the 666 end of the century under low (Shared Socioeconomic Pathway SSP 1-2.6 to SSP 2-4.5) and high (SSP 3-7.0 to SSP 5-8.5) greenhouse gas emission scenarios, respectively²⁴. TerraClimate 667 668 projections use monthly data on precipitation, minimum temperature, maximum temperature, 669 wind speed, vapor pressure deficit, soil moisture, and downward surface shortwave radiation. 670 These projections impose monthly climate projections from 23 CMIP5 global circulation models, as described in ⁹⁰. The *micro_ncep* function then downscales monthly TerraClimate inputs to 671 672 hourly by imposing a diurnal cycle to the data and imposes TerraClimate offsets onto the 673 climatic data from NCEP. Because the TerraClimate data is already bias-corrected, adding 674 future climate projections onto the NCEP data did not require further bias correction. We ran all microclimatic estimations between 2005 and 2015 to match the range of pseudo-years available 675 676 for TerraClimate future climate projections. We did not use a larger range of historical records

and only used climate projections available in TerraClimate (i.e., 2°C and 4°C) to reduce
computational demands.

679 We then used microclimate estimates to generate hourly operative body temperatures using the ectotherm function in NicheMapR⁸⁰. This modelling system has been extensively validated with 680 field observations^{91–93} (see also Fig. S12). We modelled an adult amphibian in the shape of the 681 leopard frog Lithobates pipiens, positioned 1 cm above ground (or 2 m for arboreal species), 682 683 and assumed that 80% of the skin acted as a free water surface (wet skin). Estimating body 684 mass-specific operative body temperatures for each grid cell, species, and microhabitat was too 685 computationally extensive, given the geographic and taxonomic scale of our study (464,871 686 local species occurrences). Therefore, we ran the ectotherm models using the median body 687 mass of the species assemblage in each geographical coordinate. When body mass was 688 unknown, we ran models assuming a body mass of 8.4 grams, the median assemblage-level 689 body mass. Given that most amphibians in our dataset are small (median = 1.4 g, mean = 27.5 690 g), body temperatures equilibrate quickly with the environment, and operative body 691 temperatures are likely representative of core body temperatures.

692 To model operative body temperatures in water bodies (e.g., ponds or wetlands), we used the 693 container model from *NicheMapR*. Unlike previously mentioned calculations predicting steady-694 state temperatures, this approach accounts for transient temperature changes, capturing lags due to thermal inertia (i.e., transient heat budget model^{94,95}). For pond simulations, we modelled 695 696 a container permanently filled with water (12 m width and 1.5 m-depth) and decreased direct 697 solar radiation to zero to simulate full shade. This modelling approach serves as a proxy for 698 estimating the body temperature of ectotherms submerged in water bodies such as ponds or wetlands, which was validated with field measurements (e.g., ^{40,92}). Ground-level and water 699 temperatures were modelled for all species regardless of their ecotype (apart from 700 701 paedomorphic salamanders that were only assessed in aquatic environments) because 702 arboreal and terrestrial species may retreat on land or in water occasionally. Temperatures in 703 above-ground vegetation were only estimated for arboreal and semi-arboreal species as

704 reaching 2 meters height in vegetation requires a morphology adapted to climbing. Our 705 biophysical models assume that shaded microhabitats are available to species throughout their 706 range. While this may not hold true, fine-scaled distribution of these microenvironments are not 707 available at global scales. Moreover, assuming that these microenvironments are available 708 serves a functional role; it provides a best-case scenario that is useful for comparative analyses 709 and offers actionable insights for conservation. For instance, reduced exposure to overheating 710 events in aquatic relative to terrestrial environments would suggest that preserving ponds and 711 wetlands may be critical in buffering the impacts of climate change on amphibians.

712 We then estimated, for each geographical coordinate, the maximum daily body temperature and 713 the mean and maximum weekly maximum body temperature experienced in the 7 days prior to 714 each given day to account for acclimation responses and to assess climate vulnerability 715 metrics¹⁸ (see *Climate vulnerability analyses*). We only used data for the 91 warmest days (i.e., 716 warmest guarter) of each year, as we were interested in the responses of amphibians to extreme heat events¹⁸. Note that data from the year 2005 was excluded *a posteriori* as a burn-in 717 718 to remove the effects of initial conditions on soil temperature, soil moisture, and pond 719 calculations. Therefore, our analyses are based on 910 days (91 days per year in the range 720 2006-2015) for each climatic scenario (current climate, 2°C above pre-industrial levels, 4°C 721 above pre-industrial levels).

We also used maximum daily body temperatures on terrestrial conditions to calculate the median, 5th percentile and 95th percentile maximum body temperature experienced by each species across their range of distribution. These values were used as acclimation temperatures in the training data to calibrate the data imputation with ecologically-relevant environmental temperatures (see *Data imputation*); while maximizing the range of temperatures used to infer the plasticity of heat tolerance limits (see *Climate vulnerability analysis*).

728 Climate vulnerability analysis

729 Using the imputed data, we fitted an individual meta-analytic model for each species to estimate 730 the plasticity of imputed heat tolerance limits (CT_{max}) to changes in operative body temperatures 731 using the *metafor* package⁹⁶ (v. 4.2-0). CT_{max} was used as the response variable, acclimation 732 temperature (i.e., median, 5th percentile, or 95th percentile daily maximum body temperature 733 experienced by a species across its distribution range) was used as the predictor variable, and 734 imputed estimates were weighted based on their standard error. From these models, we used 735 out-of-sample model predictions (using the *predict* function) to estimate the CT_{max} of each 736 species in each 1° x 1° grid cell across their distribution range in different warming scenarios, 737 based on predicted mean weekly body temperatures. Specifically, we assumed that species 738 were, on any given day, acclimated to the mean daily body temperature experienced in the 7 739 days prior¹⁸. Therefore, CT_{max} was simulated as a plastic trait, which varied daily, as animals 740 acclimate to new environmental conditions (Extended Data Fig. 1). While evidence in small amphibians suggests the full acclimation potential is reached within 3-4 days^{97–99}, other 741 evidence points to some variation after longer periods¹⁰⁰. Therefore, we chose 7 days to reflect 742 743 that some amphibians may require longer to acclimate. Because we used out-of-sample model 744 predictions, we propagated errors from the imputation when estimating the predicted CT_{max} across geographical coordinates. Predicted CT_{max} values and their associated standard errors 745 746 thus reflect variation in both the imputation procedure and the estimation of plastic responses. 747 Our approach to accounting for plasticity assumes that plasticity is homogeneous within species 748 and ignores the possible influence of local adaptation. However, given the low variability in plasticity among species (Fig. S2-3), lack of evidence for latitudinal variation in plasticity 749 $(^{28,31,101})$, high phylogenetic signal in thermal tolerance (Pagel's $\lambda^{102} = 0.95$ [0.91 – 0.98]; see 750 751 Sensitivity Analyses), and evidence for slow rates of evolution and physiological constraints on 752 CT_{max}^{52,53}, geographic variation in thermal tolerance and plasticity is unlikely to have a major 753 influence on our results.

754 We then estimated the vulnerability of amphibians to global warming using three metrics 755 (Extended Data Fig. 1). First, we calculated the difference between CT_{max} and the maximum 756 daily body temperature, i.e., the thermal safety margin (i.e., TSM, sensu⁶). We calculated weighted means and standard errors (sensu¹⁰³) of thermal safety margins across years to 757 758 estimate the mean difference between CT_{max} and the maximum temperature during the warmest 759 quarters. Using TSM averaged from the maximum temperature of the warmest quarter is 760 common in the literature²⁷⁻²⁹. Second, we calculated the number of days the maximum daily 761 operative body temperature exceeded CT_{max} across the warmest guarters of 2006-2015, i.e., 762 the number of overheating events. To propagate the uncertainty, we calculated the mean 763 probability that daily operative body temperatures exceeded the predicted distribution of CT_{max} 764 (using the *dnorm* function). Note that the standard error (standard deviation of estimates) of 765 simulated CT_{max} distributions were restricted to one (i.e., simulating distributions within ~3°C of 766 the mean) to avoid inflating overheating probabilities due to large imputation uncertainty (cf⁷²: 767 see also Sensitivity analyses; Fig. S8). We then multiplied the mean overheating probability by 768 the total number of simulated days (910) to estimate the number of overheating events and their 769 associated standard error using properties of the binomial distribution. Third, we calculated the 770 binary probability (0/1) that species overheat for at least one day across the 910 days surveyed 771 (warmest guarters of 2006-2015). The latter two metrics provide a finer resolution than TSMs, 772 as they capture daily temperature fluctuations and potential overheating events¹⁸.

773 Macroecological patterns

The objective of this study was to characterise the vulnerability of amphibians to global warming. We investigated patterns at the level of local species occurrences (presence of a given species in a 1° x 1° grid cell based on IUCN data), allowing one to identify specific populations and species that may be more susceptible to heat stress and direct targeted research efforts. We also analysed data at the assemblage level, the species composition within a grid cell. In such case, we calculated the weighted mean and standard error of TSM (*sensu* ¹⁰³) across species in each grid cell. Assemblage-level analyses allow one to identify areas

containing a higher number of vulnerable species, offering actionable insights for broader-scale
 conservation initiatives.

We used the gamm4 package¹⁰⁴ to fit generalised additive mixed models (GAMM) against 783 784 latitude. For local species occurrences, we fitted latitude as a fixed factor, and nested genus 785 and species identity as random terms to account for phylogenetic non-independence. Note that 786 we did not include family as a random term because models failed at estimating higher 787 taxonomic variation. While better methods exist to model phylogenetic patterns, generalised 788 additive linear models do not allow for phylogenetic correlation matrices, and other functions such as *brms*¹⁰⁵ surpassed our computational time and memory limits. Nevertheless, imputed 789 790 estimates already reflect variation due to phylogeny (see Data imputation), and phylogeny was 791 further modelled when deriving mean estimates in each microhabitat and climatic scenario (see 792 below). We fitted models using the three metrics as response variables independently: the 793 thermal safety margin, overheating risk, and number of overheating events. The former was 794 modelled using a Gaussian distribution of residuals, overheating risk was modelled using a 795 binomial error structure, and the latter using a Poisson error structure. Note that overheating 796 risks were rounded to integer values to fit a Poisson distribution. Thermal safety margin 797 estimates were weighted by the inverse of their sampling variance to account for the uncertainty 798 in the imputation and predictions across geographical coordinates. We fitted separate models 799 for each climatic scenario (current climate, 2°C above preindustrial levels, 4°C above 800 preindustrial levels) and microhabitat (terrestrial, aquatic, arboreal).

To investigate the mean TSM in each microhabitat and climatic scenario, we fitted models with the interaction between microhabitat and climatic scenario as a fixed effect using *MCMCglmm*⁷⁰ (v. 2.34) and flat, parameter-expanded priors. In these models, we weighted estimates based on the inverse of their sampling variance, species identity was fitted as a random effect, and we accounted for phylogenetic non-independence using a variance-covariance matrix of phylogenetic relatedness (calculated from the consensus tree of ²⁶). To investigate the overall overheating risk and number of overheating events in each condition, we attempted to fit

models in *MCMCgImm* but these models failed to converge. Therefore, we fitted Poisson and binomial models using $Ime4^{106}$ (v. 1.1-33) and nested genus, species, and observation as random terms. We used similar Poisson models to investigate the relationship between the number of overheating events and thermal safety margins. While the mean estimates from these simpler models should be unbiased, estimate uncertainty is likely underestimated¹⁰⁷.

813 We also investigated patterns of climate vulnerability at the assemblage level. We calculated 814 the weighted average of TSM and overheating risk in each 1-degree grid cell (14,091; 14,090; 815 or 6,614 grid cells for terrestrial, aquatic, and arboreal species, respectively), and mapped 816 patterns geographically. Averaging overheating risk effectively returned the proportion of 817 species overheating in each coordinate; and we also calculated the number of species 818 overheating in each grid cell. For assemblage-level models, we fitted Gaussian, binomial or 819 Poisson models as described above, but without taxon-level random effects because these 820 cannot be modelled at the assemblage level. All models were fitted without a contrast structure 821 to estimate mean effects in each microhabitat and climatic scenario, and with two-sided 822 contrasts to draw comparisons with current terrestrial conditions.

823 Cross-validation and sensitivity analyses

824 We assessed the accuracy of the data imputation procedure using a cross-validation approach. 825 Specifically, we removed heat tolerance estimates for 5% of the species in the experimental 826 data and 5% of the data-deficient species (maintaining the same proportion of missing data) 827 and assessed how well experimental values could be predicted from the models. Of relevance, 828 we only removed data that were comparable to the data that were imputed. That is, data from 829 adult animals tested using a ramping rate of 1°C/min, and where thermal limits were recorded 830 as the onset of spasms. While we could have trimmed any data entry in the experimental data, 831 validation of the imputation performance can only be achieved by comparing comparable 832 entries, and imputing data from species tested in unusual settings would naturally result in large 833 errors. In total, we cross-validated experimental estimates for 77 species.

834 We investigated alternative ways to i) calculate thermal safety margins, ii) account for 835 acclimation responses, and iii) control for prediction uncertainty (see Supplementary methods; 836 Fig. S6-8) and investigated the influence of different parameters of our biophysical models (i.e., 837 shade and burrow availability, plant height, solar radiation, wind speed, pond depth) on 838 predicted vulnerability risks (see Supplementary methods; Fig. S9-11). Our results were 839 generally robust to changes in model parameters, although amphibians are likely to experience more overheating events in open habitats^{6,42} (Fig. S9) and shallow ponds (Fig. S10), and lower 840 841 risks in underground conditions¹⁰⁸ (Fig. S9). We also confirmed that predicted operative body 842 temperatures were comparable to field body temperatures measured in some wild frogs (see 843 Supplementary methods; Fig. S12).

844 Finally, we confirmed the presence of a phylogenetic signal in the experimental dataset by fitting 845 a Bayesian linear mixed model using all complete (no missing data) predictors (i.e., acclimation 846 temperature, endpoint, acclimation status, life stage, and ecotype) in MCMCalmm. We 847 accounted for phylogenetic non-independence using a correlation matrix of phylogenetic 848 relatedness and fitted random intercepts for non-phylogenetic species effects. The phylogenetic signal (Pagel's λ^{102} , which is equivalent to phylogenetic heritability^{109,110}) was calculated as the 849 850 proportion of variance explained by phylogenetic effects relative to the total non-residual 851 variance.

852 Results from all statistical models and additional data visualizations are available at https://p-

853 pottier.github.io/Vulnerability_amphibians_global_warming/.

854 Data availability

- 855 Raw and processed data are available at <u>https://github.com/p-</u>
- 856 <u>pottier/Vulnerability_amphibians_global_warming</u>. Note, however, that some intermediate data
- files were too large to be shared online. These files are available upon request and will be
- uploaded to a permanent repository upon acceptance. TerraClimate data is available from
- 859 <u>https://www.climatologylab.org/terraclimate.html</u> and NCEP data is available from
- 860 https://psl.noaa.gov/thredds/catalog/Datasets/ncep.reanalysis2/gaussian_grid/catalog.html.

861 Code availability

- 862 All code needed to reproduce the analyses is available at <u>https://github.com/p-</u>
- 863 <u>pottier/Vulnerability_amphibians_global_warming</u>.

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989 Authors' contributions

990 This study was conceptualized by PPottier, MRK, SB, SMD, and SN. All data manipulation and 991 analyses were performed by PPottier (with conceptual and technical input from SMD and SN for 992 the imputation methods and statistical analyses, MRK, ARG, JER, and NCW for the biophysical 993 modelling and climate vulnerability analyses). All code was reviewed by NCW, ARG, and JER 994 following the recommendations of ¹¹¹. Ecotype information was collected by NCW, PPollo, and 995 ANRV. PPottier, NCW, and SMD contributed to data visualization. PPottier wrote the initial draft, 996 and all authors were involved in the review and editing. PPottier oversaw the project 997 administration, while SMD and SN were in charge of the supervision.

998 Inclusion & ethics statement

- 999 This study did not involve researchers who collected the original data. All data used for the
- analyses were taken from a previous data compilation, and original references are listed in
- 1001 Supplementary Information (*Data sources*).

1002 **Competing interest declaration**

1003 The authors declare no conflict or competing interests.

1005 Additional information

- 1006 Supplementary Information is available for this paper.
- 1007 Correspondence and requests for materials should be addressed to Patrice Pottier.
- 1008 Reprints and permissions information is available at www.nature.com/reprints.



1012 Extended Data Fig. 1 | Conceptual overview of the methods employed to assess the

1013 vulnerability of amphibians to global warming.





1018Extended Data Fig. 2 | Accuracy of the data imputation procedure. a) Probability density1019distributions (n = 375 observations, 77 species) of experimental CT_{max} (blue) and CT_{max} cross-1020validated using our data imputation procedure (pink). b) Correlation between experimental and1021imputed CT_{max} values. c) Variation in the uncertainty (standard error, SE) of imputed CT_{max} 1022predictions (outer heat map) across studied (blue; n = 524) and imputed (grey; n = 4,679)1023species.





Extended Data Fig. 3 | Thermal safety margin, critical thermal maximum, and operative body temperatures in different microhabitats and climatic scenarios. Weighted mean thermal safety margins (TSM; a-c), critical thermal maximum (CT_{max}; d-f) and operative body temperatures (g-i) in terrestrial (a,d,g), aquatic (b,e,h) and arboreal (c,f,i) microhabitats are depicted in current microclimates (blue data points), or assuming 2°C and 4°C of global warming above pre-industrial levels (orange, and pink data points, respectively) across latitudes. Lines represent 95% confidence intervals of model predictions from generalised additive mixed models. CT_{max} and TSM estimates are scaled by precision (1/s.e.). Each point represents a species in a given grid cell.



Extended Data Fig. 4 | Vulnerability of arboreal amphibians in terrestrial and arboreal 1040 microhabitats. Depicted are the number of overheating events experienced by arboreal 1041 1042 species across latitudes (a-b) and in relation to thermal safety margins (c-d) in terrestrial (a-c) 1043 and arboreal microhabitats (b-d). The number of overheating events were calculated based on the mean probability that daily maximum temperatures exceeded CT_{max} during the warmest 1044 1045 quarters of 2006-2015 for each species in each grid cell. Blue points depict the number of overheating events in historical microclimates, while orange and pink points depict the number 1046 of overheating events assuming 2°C and 4°C of global warming above pre-industrial levels, 1047 1048 respectively. In panel a) and b), only the species predicted to overheat for at least one day are displayed. The number of arboreal species predicted to experience overheating events in 1049 1050 terrestrial (e) and arboreal (f) microhabitats in each assemblage is also depicted. The number of species overheating was assessed as the sum of species overheating for at least one day in the 1051 period surveyed (warmest quarters of 2006-2015) in each assemblage (1-degree grid cell). 1052 Black colour depicts areas with no data, and grey colour assemblages without species at risk. 1053 1054 The right panel depicts latitudinal patterns in the number of species predicted to overheat in current climates (blue) or assuming 4°C of global warming above pre-industrial levels (pink). 1055 Dashed lines represent the equator and tropics. Few species (n = 11) were predicted to 1056 1057 experience overheating events in water bodies, and hence are not displayed. 1058



1060

Extended Data Fig. 5 | Proportion of species predicted to experience overheating events 1061 1062 in terrestrial (a), aquatic (b), and arboreal (c) microhabitats. The proportion of species 1063 overheating was assessed as the sum of species overheating for at least one day in the period 1064 surveyed (warmest quarters of 2006-2015) divided by the number of species in each 1065 assemblage (1-degree grid cell). Black colour depicts areas with no data, and grey colour assemblages without species at risk. The right panel depicts latitudinal patterns in the proportion 1066 of species predicted to overheat in current climates (blue) or assuming 4°C of global warming 1067 1068 above pre-industrial levels (pink). Dashed lines represent the equator and tropics.

Extended Data Table 1 | Statistical model estimates for thermal safety margins calculated 1071 for local species occurrences and assemblages Model estimates for each microhabitat 1072 (terrestrial, arboreal, aquatic) and each climatic scenario (current, +2°C, or +4°C of global 1073 1074 warming above pre-industrial levels) are depicted. mean: mean model estimate; Cl.lb: lower bound of the 95% confidence interval; Cl.ub: upper bound of the 95% confidence interval; k_{sp}: 1075 number of species; kobs: number of observations; Varsp: variance explained by differences 1076 1077 between species; Var_{phy}: variance explained by shared evolutionary history; Var_{obs}: residual 1078 variance.

Local species patterns in thermal safety margin								
	mean	CI.lb	Cl.ub	k _{sp}	kobs	Var _{sp}	Var _{phy}	Varobs
Terrestrial (current)	11.694	8.856	14.428	5177	203853			
Terrestrial (+2°C)	10.914	8.025	13.594	5177	203853			
Terrestrial (+4°C)	9.409	6.530	12.090	5177	203853			
Arboreal (current)	12.235	9.402	14.960	1771	56210			
Arboreal (+2°C)	11.517	8.660	14.236	1771	56210	1.295	11.960	1.828
Arboreal (+4°C)	10.073	7.229	12.797	1771	56210			
Aquatic (current)	13.598	10.708	16.276	5203	204808			
Aquatic (+2°C)	12.827	8.796	14.361	5203	204808			
Aquatic (+4°C)	11.682	8.796	14.361	5203	204808			

Assemblage-level patterns in thermal safety margin

	mean	CI.lb	Cl.ub	Kobs	Varobs
Terrestrial (current)	15.279	15.208	15.330	14090	
Terrestrial (+2°C)	14.328	14.279	14.396	14090	
Terrestrial (+4°C)	12.602	12.542	12.657	14090	
Arboreal (current)	14.279	14.191	14.381	6614	11.06
Arboreal (+2°C)	13.393	13.298	13.478	6614	
Arboreal (+4°C)	11.746	11.666	11.830	6614	
Aquatic (current)	17.408	17.352	17.471	14091	
Aquatic (+2°C)	16.528	16.468	16.581	14091	
Aquatic (+4°C)	15.287	15.225	15.346	14091	

1079

1082 Extended Data Table 2 | Statistical model estimates for overheating risk and the number 1083 of overheating events. Model estimates for each microhabitat (terrestrial, arboreal) and each 1084 climatic scenario (current, +2°C, or +4°C of global warming above pre-industrial levels) are depicted. The estimated number of overheating events in species predicted to experience at 1085 least one overheating event (i.e., overheating species) are also depicted. Model estimates for 1086 1087 aquatic microhabitats are not displayed because no species was predicted to experience 1088 overheating events in this microhabitat. mean: mean model estimate; CI.Ib: lower bound of the 95% confidence interval; Cl.ub: upper bound of the 95% confidence interval; k_{sp}: number of 1089 1090 genera; k_{sp}: number of species; k_{obs}: number of observations; Var_{genus}: variance explained by differences between genera; Var_{sp}: variance explained by differences between species; Var_{obs}: 1091 1092 residual variance.

Overheating risk										
	mean	CI.lb	Cl.ub	k genus	k sp	k obs	Vargenus	Var _{sp}		
Terrestrial (current)	9.98 x 10 ⁻⁷	5.60 x 10 ⁻⁷	1.78 x 10 ⁻⁶	464	5177	203853				
Terrestrial (+2°C)	1.93 x 10 ⁻⁶	1.09 x 10 ⁻⁶	3.43 x 10 ⁻⁶	464	5177	203853				
Terrestrial (+4°C)	9.09 x 10 ⁻⁶	5.13 x 10 ⁻⁶	1.61 x 10 ⁻⁵	464	5177	203853	0 306	69.653		
Arboreal (current)	4.77 x 10 ⁻⁷	2.58 x 10 ⁻⁷	8.80 x 10 ⁻⁷	174	1771	56210	0.000			
Arboreal (+2°C)	9.78 x 10 ⁻⁷	5.45 x 10 ⁻⁷	1.75 x 10 ⁻⁶	174	1771	56210				
Arboreal (+4°C)	3.72 x 10 ⁻⁶	2.08 x 10 ⁻⁶	6.67 x 10 ⁻⁶	174	1771	56210				

Number of overheating events (all species)

-	mean	CI.lb	Cl.ub	k genus	k _{sp}	kobs	Var _{genus}	Var _{sp}	
Terrestrial (current)	0.014	0.001	0.080	464	5177	203853			
Terrestrial (+2°C)	0.025	0.002	0.127	464	5177	203853		52.500	
Terrestrial (+4°C)	0.153	0.046	0.460	464	5177	203853	0 1 1 0		
Arboreal (current)	0.008	0.001	0.043	174	1771	56210	0.110		
Arboreal (+2°C)	0.015	0.001	0.083	174	1771	56210			
Arboreal (+4°C)	0.076	0.012	0.230	174	1771	56210			

Number of overheating events (among overheating species)

	<u> </u>	<u> </u>							
mean	CI.Ib	Cl.ub	Kgenus	Ksp	Kobs	Vargenus	Var _{sp}	Var _{obs}	
2.155	0.239	5.264	38	104	836				
2.576	0.410	5.857	61	168	1424		0.187		
6.747	3.136	11.385	118	391	4248	0 253		0.310	
1.621	0.026	4.429	4	13	152	0.200			
1.956	0.113	4.973	5	16	283				
5.084	1.806	9.387	17	56	748				
	mean 2.155 2.576 6.747 1.621 1.956 5.084	mean Cl.lb 2.155 0.239 2.576 0.410 6.747 3.136 1.621 0.026 1.956 0.113 5.084 1.806	meanCl.lbCl.ub2.1550.2395.2642.5760.4105.8576.7473.13611.3851.6210.0264.4291.9560.1134.9735.0841.8069.387	meanCl.lbCl.ubkgenus2.1550.2395.264382.5760.4105.857616.7473.13611.3851181.6210.0264.42941.9560.1134.97355.0841.8069.38717	meanCl.lbCl.ubkgenusksp2.1550.2395.264381042.5760.4105.857611686.7473.13611.3851183911.6210.0264.4294131.9560.1134.9735165.0841.8069.3871756	meanCl.lbCl.ubkgenuskspkobs2.1550.2395.264381048362.5760.4105.8576116814246.7473.13611.38511839142481.6210.0264.4294131521.9560.1134.9735162835.0841.8069.3871756748	meanCl.lbCl.ubkgenuskspkobsVargenus2.1550.2395.264381048362.5760.4105.8576116814246.7473.13611.38511839142481.6210.0264.4294131521.9560.1134.9735162835.0841.8069.3871756748	meanCl.ubkgenuskspkobsVargenusVarsp2.1550.2395.264381048362.5760.4105.8576116814246.7473.13611.38511839142481.6210.0264.4294131521.9560.1134.9735162835.0841.8069.3871756748	

1094 Extended Data Table 3 | Statistical model estimates for the number of species predicted to experience overheating events. Model estimates for each microhabitat (terrestrial, 1095 arboreal) and each climatic scenario (current, +2°C, or +4°C of global warming above pre-1096 industrial levels) are depicted. The estimated number of species overheating in assemblages 1097 containing at least one species predicted to experience at least one overheating event (i.e., 1098 1099 overheating assemblages) are also depicted. Model estimates for aquatic microhabitats are not displayed because no species was predicted to experience overheating events in this 1100 microhabitat. mean: mean model estimate; CI.lb: lower bound of the 95% confidence interval; 1101 1102 Cl.ub: upper bound of the 95% confidence interval; kobs: number of observations; Varobs: 1103 residual variance.

Number of species overheating (all assemblages)							
_	mean	CI.lb	Cl.ub	k _{obs}	Varobs		
Terrestrial (current)	0.056	0.016	0.118	14090			
Terrestrial (+2°C)	0.096	0.029	0.199	14090			
Terrestrial (+4°C)	0.288	0.083	0.604	14090	55 17		
Arboreal (current)	0.021	0.002	0.054	6614	55.47		
Arboreal (+2°C)	0.040	0.006	0.094	6614			
Arboreal (+4°C)	0.107	0.021	0.243	6614			

Number of species overheating (among overheating assemblages)

mean	Cl.lb	Cl.ub	kobs	Var _{obs}
3.185	0.601	6.883	253	
3.228	0.678	6.810	426	
3.084	0.617	6.557	1328	0.601
1.930	0.054	5.054	74	0.001
2.445	0.189	5.649	111	
2.509	0.312	5.692	285	
	mean 3.185 3.228 3.084 1.930 2.445 2.509	mean Cl.lb 3.185 0.601 3.228 0.678 3.084 0.617 1.930 0.054 2.445 0.189 2.509 0.312	meanCl.lbCl.ub3.1850.6016.8833.2280.6786.8103.0840.6176.5571.9300.0545.0542.4450.1895.6492.5090.3125.692	meanCl.lbCl.ubkobs3.1850.6016.8832533.2280.6786.8104263.0840.6176.55713281.9300.0545.054742.4450.1895.6491112.5090.3125.692285

1104

1107 Extended Data Table 4 | Statistical model estimates for the proportion of species predicted to experience overheating events. Model estimates for each microhabitat 1108 (terrestrial, arboreal) and each climatic scenario (current, +2°C, or +4°C of global warming 1109 above pre-industrial levels) are depicted. The estimated proportion of species overheating in 1110 assemblages containing at least one species predicted to experience at least one overheating 1111 event (i.e., overheating assemblages) are also depicted. Model estimates for aquatic 1112 microhabitats are not displayed because no species was predicted to experience overheating 1113 events in this microhabitat. mean: mean model estimate; CI.lb: lower bound of the 95% 1114 1115 confidence interval; Cl.ub: upper bound of the 95% confidence interval; kobs: number of observations; Varobs: residual variance. 1116

Proportion of species overheating (all assemblages)							
	mean	CI.lb	Cl.ub	k obs	Var _{obs}		
Terrestrial (current)	1.22 x 10⁻⁵	8.96 x 10 ⁻⁶	1.66 x 10 ⁻⁵	14090			
Terrestrial (+2°C)	2.09 x 10 ⁻⁵	1.60 x 10 ⁻⁵	2.72 x 10 ⁻⁵	14090			
Terrestrial (+4°C)	8.13 x 10⁻⁵	6.60 x 10 ⁻⁵	1.00 x 10 ⁻⁴	14090	12.26		
Arboreal (current)	1.19 x 10 ⁻⁵	7.07 x 10 ⁻⁶	2.02 x 10 ⁻⁵	6614	42.20		
Arboreal (+2°C)	1.86 x 10 ⁻⁵	1.19 x 10 ⁻⁵	2.89 x 10 ⁻⁵	6614			
Arboreal (+4°C)	4.99 x 10 ⁻⁵	3.62 x 10⁻⁵	6.87 x 10 ⁻⁵	6614			

Proportion of species overheating (among overheating assemblages)

	mean	Cl.lb	Cl.ub	k obs	Var _{obs}
Terrestrial (current)	0.053	0.046	0.061	253	
Terrestrial (+2°C)	0.058	0.052	0.065	426	
Terrestrial (+4°C)	0.094	0.088	0.100	1328	1 010
Arboreal (current)	0.038	0.029	0.050	74	1.019
Arboreal (+2°C)	0.054	0.043	0.067	111	
Arboreal (+4°C)	0.061	0.053	0.070	285	

1117

1120 Extended Data Table 5 | Statistical model estimates for the association between the number of overheating events and thermal safety margins. Model estimates for each 1121 microhabitat (terrestrial, arboreal) and each climatic scenario (current, +2°C, or +4°C of global 1122 warming above pre-industrial levels) are depicted. Model estimates for aquatic microhabitats 1123 are not displayed because no species was predicted to experience overheating events in this 1124 microhabitat. All model estimates are on the log scale. mean: mean model estimate; se: 1125 standard error; k_{sp}: number of genera; k_{sp}: number of species; k_{obs}: number of observations; 1126 Vargenus: variance explained by differences between genera; Varsp: variance explained by 1127

	mean	se	р	k genus	k _{sp}	k obs	Var _{genus}	Var _{sp}	Var _{obs}
Terrestrial (current)									
Intercept	3.723	0.390	<0.001	464	5177	203853	5.850	3.346	0.116
Slope (TSM)	-1.201	0.031	<0.001						
Terrestrial (+2°C)									
Intercept	6.318	0.310	<0.001	464	5177	203853	5.272	2.380	0.078
Slope (TSM)	-1.452	0.027	<0.001						
Terrestrial (+4°C)									
Intercept	7.611	0.171	<0.001	464	5177	203853	2.954	1.025	0.248
Slope (TSM)	-1.616	0.015	<0.001						
Arboreal (current)									
Intercept	4.929	1.091	<0.001	174	1771	56210	0.001	15.190	0.001
Slope (TSM)	-1.511	0.094	<0.001						
Arboreal (+2°C)									
Intercept	7.836	0.836	<0.001	174	1771	56210	4.359	2.358	0.001
Slope (TSM)	-1.739	0.080	<0.001						
Arboreal (+4°C)									
Intercept	10.093	0.587	<0.001	174	1771	56210	8.789	0.917	0.001
Slope (TSM)	-2.085	0.039	<0.001						

1128 differences between species; Varobs: residual variance.

1129

Vulnerability of amphibians to global warming

- 2 Patrice Pottier^{1,2*}, Michael R. Kearney³, Nicholas C. Wu⁴, Alex R. Gunderson⁵, Julie E. Rej⁵,
- A. Nayelli Rivera-Villanueva^{6,7}, Pietro Pollo¹, Samantha Burke¹, Szymon M. Drobniak^{1,8+}, and
- 4 Shinichi Nakagawa^{1,9+}
- 5

- ¹ Evolution & Ecology Research Centre, School of Biological, Earth and Environmental
 Sciences, University of New South Wales, Sydney, New South Wales, Australia.
- ² Division of Ecology and Evolution, Research School of Biology, The Australian National
 University, Canberra, Australian Capital Territory, Australia
- ³ School of BioSciences, The University of Melbourne, Melbourne, Victoria, Australia
- ⁴ Hawkesbury Institute for the Environment, Western Sydney University, Richmond, New
 South Wales, Australia
- ⁵ Department of Ecology and Evolutionary Biology, Tulane University, New Orleans,
 Louisiana, USA
- ⁶ Centro Interdisciplinario de Investigación para el Desarrollo Integral Regional Unidad
 Durango (CIIDIR), Instituto Politécnico Nacional, Durango, México
- ⁷ Laboratorio de Biología de la Conservación y Desarrollo Sostenible de la Facultad de
 Ciencias Biológicas, Universidad Autónoma de Nuevo León, Monterrey, México
- ⁸ Institute of Environmental Sciences, Jagiellonian University, Kraków, Poland.
- ⁹ Department of Biological Sciences, University of Alberta, Edmonton, Alberta, Canada.
- 21 ^{*}Corresponding author
- 22 ⁺These authors supervised the work equally
- 23
- 24 Corresponding author: Patrice Pottier (<u>p.pottier@unsw.edu.au</u>)
- 25

Supplementary Information for Pottier et al. 2024. Vulnerability of amphibians to global warming.

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48 Sensitivity analyses

In this study, we projected CT_{max} estimates assuming animals were acclimated to the mean weekly temperature experienced prior to each day. We also assessed the climate vulnerability of amphibians assuming they were acclimated to weekly maximum body temperatures (*cf.* ¹), which reflects more conservative estimates (Fig. S7).

53 We also calculated thermal safety margins as the difference between the maximum (or 95th percentile, cf.²) hourly body temperature experienced by each population and their predicted 54 CT_{max} to investigate the consequences of averaging temperatures when calculating TSMs 55 (Fig. S6). To increase the comparability of our estimations with similar studies (e.g., 2), we also 56 57 calculated climate vulnerability metrics more conservatively. Specifically, we excluded 58 temperature data falling below the 5th percentile and above the 95th percentile body temperature for each population to mitigate the impact of outliers (Fig. S6). However, extreme 59 weather events, which are typically captured by these outlier values, are the very phenomena 60 most likely to precipitate mortality events^{3,4}. Omitting these outliers could therefore obscure 61 62 the ecological significance of extreme temperatures, thereby underestimating true overheating 63 risks.

To estimate overheating probabilities, we calculated the mean daily probability that operative 64 body temperatures exceeded the predicted distribution of CT_{max} and restricted the standard 65 66 deviation of simulated distributions to one (i.e., within ~3°C of the mean) to avoid inflating overheating probability for observations with large uncertainty. We also provided alternative 67 results (Fig. S8) where the standard deviation of CT_{max} was restricted to the "biological range", 68 i.e., the standard deviation of the distribution of all CT_{max} estimates across species (range = 69 1.84 - 2.17). We also provide a sensitivity analysis where overheating risk was positive only 70 when the 95% confidence intervals of predicted overheating days did not overlap with zero 71 (Fig. S8). 72

We also investigated the influence of different parameters of our biophysical models (i.e., shade and burrow availability, height in above-ground vegetation, solar radiation, wind speed, pond depth) on predicted vulnerability risks (Fig. S9-11). Specifically, we modelled the responses of the species at highest risk in terrestrial and aquatic conditions, *Noblella myrmecoides*, in its most vulnerable location (latitude, longitude = -9.5, -69.5). For terrestrial conditions, we modelled the response of amphibians with different body sizes (0.5, 4.28, or 50 grams), and with different levels of exposure to open habitat conditions. Specifically, we

80 modelled an amphibian exposed to 50% of shade to simulate an open habitat lightly covered 81 by vegetation, and inferred temperatures at different soil depths (2.5, 5, 10, 15, or 20 cm 82 underground). For aquatic conditions, we adjusted pond depths to simulate a very shallow pond (50 cm) and compared it to deeper ponds (1.5- or 3-meters depth). For arboreal 83 conditions, we modelled the responses of Pristimantis ockendeni, in its most vulnerable 84 location (-4.5, -71.5), and adjusted the height in above-ground vegetation (0.5, 2, or 5 meters), 85 the percentage of radiation diffused by vegetation (50%, 75%, or 90% of radiation diffused), 86 and the percentage of wind speed reduced by vegetation (0%. 50%, or 80% of wind speed 87 reduced by vegetation). We did not estimate the influence of these parameters on all species 88 and at all locations because of the scale of our study, but these results should provide insight 89 into how varying microenvironmental features and biological characteristics may impact our 90 91 general conclusions.

Finally, we compared our predictions of operative body temperatures with field body 92 temperature measurements. We extracted night-time (18:00 – 00:30) field body temperatures 93 measured for 11 species of frogs in Mexico (21.48° N, -104.85° W; and 21.45° N, -105.03° W) 94 95 between June and October of 2013 and 2015 from Table 1 of ⁵. We chose this study because it provided the data and location of body temperature measurements, covered multiple species 96 97 from different sites, and matched our study timeframe (2006-2015). We then compare these estimates with hourly operative body temperatures predicted in shaded terrestrial conditions 98 at the same dates and time windows (Fig. S12). 99

Results from all statistical models and additional data visualizations are available at https://p-
 pottier.github.io/Vulnerability_amphibians_global_warming/.

- 102 Original studies on which our analyses are built upon are listed in *Data sources*^{6–218}.
- 103

105 Supplementary figures



106

107 Fig. S1 | Predicted critical thermal maximum (CT_{max}) across imputation cycles.

108 Boxplots depict median (horizontal line), interquartile ranges (boxes), and whiskers 109 extend to 1.5 times the interquartile range.



111

Ecotype

Fig. S2 | Correlations between critical thermal maximum (CT_{max}) and predictors 112 used for the imputation. CT_{max} from the experimental dataset was plotted against 113 acclimation temperature (a), acclimation time (b, log scale), ramping rate (c). Colours 114 are proportional to the values of the continuous predictors and the line refers to 115 predictions from a simple linear regression between CT_{max} and the predictors. 116 Individual slopes for each species are depicted for species when CT_{max} was estimated 117 at different acclimation temperatures (a). Depicted is also the variation in CT_{max} with 118 different endpoints (d), media used to infer body temperature (e), life stages (f), and 119 ecotypes (g). Boxplots depict median (horizontal line), interquartile ranges (boxes), 120 and whiskers extend to 1.5 times the interguartile range. LRR: loss of righting 121 response. OS: onset of spasms. 122



Acclimation Response Ratio (ARR)

Fig. S3 | Variation in plastic responses across species. The acclimation response
 ratio (ARR) represents the magnitude change in heat tolerance limits for each degree

126 change in environmental temperature. We found limited variation in ARR (mean \pm

standard deviation = 0.134 ± 0.008 ; range = 0.049 - 0.216; n = 5203).

128





Fig. S4 | Assemblage-level patterns in thermal safety margin for amphibians on 132 terrestrial (a), aquatic (b), or arboreal (c) microhabitats. Thermal safety margins 133 (TSM) were calculated as the mean difference between CT_{max} and the predicted 134 operative body temperature in full shade during the warmest guarters of 2006-2015 in 135 each assemblage (1-degree grid cell). Black colour depicts areas with no data. The 136 right panel depicts latitudinal patterns in TSM in current climates (blue) or assuming 137 2°C (orange) or 4°C of global warming above pre-industrial levels (pink), as predicted 138 from generalized additive mixed models. Point estimates are scaled by precision 139 (1/s.e.). Dashed lines represent the equator and tropics. 140





Fig. S5 | Number of species predicted to experience overheating events in 144 terrestrial (a), aquatic (b), and arboreal (c) microhabitats. The number of species 145 overheating was assessed as the sum of species overheating for least one day in the 146 period surveyed (warmest quarters of 2006-2015) in each assemblage (1-degree grid 147 cell). Black colour depicts areas with no data and grey colour assemblages without 148 species at risk. The right panel depicts latitudinal patterns in the number of species 149 predicted to overheat in current climates (blue) or assuming 2°C (orange) or 4°C of 150 global warming above pre-industrial levels (pink). Dashed lines represent the equator 151 and tropics. No species were predicted to experience overheating events in water 152 bodies, and hence are not displayed. 153





Fig. S6 | Variation in thermal safety margins calculated using different 156 assumptions. Thermal safety margins (TSM) were calculated as the mean difference 157 between CT_{max} and the predicted operative body temperature in full shade during the 158 159 warmest quarters of 2006-2015 (grey), as the mean difference between CT_{max} and the predicted operative body temperature in full shade during the warmest quarters of 160 2006-2015 excluding body temperatures falling outside the 5% and 95% percentile 161 162 temperatures (blue), as the difference between the 95% percentile operative body temperature and the corresponding CT_{max} (yellow), or as the difference between the 163 maximum operative body temperature and the corresponding CT_{max} (red). Lines 164 represented 95% confidence interval ranges predicted from generalized additive 165 mixed models. This figure was constructed assuming ground-level microclimates 166 occurring under 4°C of global warming above pre-industrial levels. 167



169

Fig. S7 | Latitudinal variation in the number of overheating events when animals 171 are acclimated to the mean (a,b) or maximum (c,d) weekly body temperature 172 experienced in the seven days prior in terrestrial (a,c) and arboreal (b,d) 173 microhabitats. The number of overheating events (days) were calculated based on 174 the mean probability that daily maximum temperatures exceeded CT_{max} during the 175 warmest guarters of 2006-2015 for each species in each grid cell. Blue points depict 176 the number of overheating events in historical microclimates, while orange and pink 177 points depict the number of overheating events assuming 2°C and 4°C of global 178 warming above pre-industrial levels, respectively. For clarity, only the species 179 predicted to experience overheating events across latitudes are depicted. 180

181

182





Fig. S8 | Latitudinal variation in the number of overheating events using regular 185 (a,b,c), uncertain (d,e,f), or conservative estimates (g,h,i) in terrestrial (a,d,g), 186 aquatic (b,e,h) and arboreal (c,f,i) microhabitats. The number of overheating 187 events (days) were calculated based on the mean probability that daily maximum 188 temperatures exceeded CT_{max} during the warmest guarters of 2006-2015 for each 189 species in each grid cell. Uncertain estimates are those where daily overheating 190 probabilities were calculated based on broad predicted distributions of CT_{max} (i.e., 191 simulated over the whole "biological range"), likely inflating overheating probabilities 192 for observations with large uncertainty. Conservative estimates are those when 193 194 overheating risk was considered only when the 95% confidence intervals of the predicted number of overheating events did not overlap with zero (e,f). Blue points 195 depict the number of overheating events in historical microclimates, while orange and 196 pink points depict the number of overheating events assuming 2°C and 4°C of global 197 warming above pre-industrial levels, respectively. For clarity, only the species 198 predicted to experience overheating events across latitudes are depicted. 199



202

Fig. S9 | Influence of biophysical model parameters on the estimation of 203 terrestrial thermal safety margins. Depicted is the variation in daily thermal safety 204 margins (TSM) as density distributions according to body mass (a), shade availability 205 and soil depth (b). All simulations were performed assuming 4°C of global warming 206 above pre-industrial levels in a specific grid cell (latitude, longitude = -9.5, -69.5; where 207 208 the highest number of overheating events was predicted), for the most vulnerable species (Noblella myrmecoides). Negative daily TSMs were recorded as overheating 209 events, and conditions depicted in dark grey reflect the results presented in the 210 211 manuscript. The number of predicted overheating events is indicated in brackets for each condition. 212

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Fig. S10 | Influence of pond depth on the estimation of aquatic thermal safety 216 margins. All simulations were performed assuming 4°C of global warming above pre-217 industrial levels in a specific grid cell (latitude, longitude = -9.5, -69.5; where the 218 highest number of overheating events was predicted), for the most vulnerable species 219 (Noblella myrmecoides). Depicted is the variation in daily thermal safety margins 220 (TSM) as density distributions. Negative daily TSMs were recorded as overheating 221 events, and conditions depicted in dark grey reflect the results presented in the 222 manuscript. The number of predicted overheating events is indicated in brackets for 223 each condition. 224



Fig. S11 | Influence of biophysical parameters on the estimation of aquatic 227 arboreal safety margins. All simulations were performed assuming 4°C of global 228 warming above pre-industrial levels in a specific grid cell (latitude, longitude = -9.5, -229 69.5; where the highest number of overheating events was predicted), for the most 230 vulnerable arboreal species (Pristimantis ockendeni). Depicted is the variation in daily 231 thermal safety margins (TSM) as density distributions according to height of the animal 232 in above-ground vegetation (a), the percentage of solar radiation diffused by 233 vegetation (b) and the percentage of wind reduced by vegetation (c). Negative daily 234 TSMs were recorded as overheating events, and conditions depicted in dark grey 235 reflect the results presented in the manuscript. The number of predicted overheating 236 events is indicated in brackets for each condition. 237





Fig. S12 | Validation of operative body temperature estimations. Terrestrial 240 operative body temperatures estimated from biophysical models were compared to 241 field body temperatures recorded around Tepic (21.48° N, -104.85° W; panel a) and 242 El Cuarenteño (21.45° N, -105.03° W; panel b) between June and October of 243 2013/2015, for 11 species of frogs⁵. The mean hourly operative body temperatures 244 predicted from our models for the same date and time windows (18:00 - 01:00) are 245 represented by the black horizontal line, along with their standard deviation (dark grey 246 box), and range (light grey box). The mean (point) and range (bars) of field body 247 temperatures recorded for each species are presented in colour. Note that our 248 analyses were based on the maximum daily temperature recorded at each site during 249 the warmest quarters of 2006-2015, which may not match the times and dates at which 250 field body temperatures were recorded. Nevertheless, congruence between night-time 251 predicted and field body temperatures suggests our models are likely to capture true 252 biological variation in operative body temperatures throughout the day. 253

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