

Basal resistance enhances warming tolerance of alien over indigenous species across latitude

Charlene Janion-Scheepers^{a,1}, Laura Phillips^a, Carla M. Sgrò^a, Grant A. Duffy^a, Rebecca Hallas^a, and Steven L. Chown^a

^aSchool of Biological Sciences, Monash University, Melbourne, VIC 3800, Australia

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Soil systems are being increasingly exposed to the interactive effects of biological invasions and climate change, with rising temperatures expected to benefit alien over indigenous species. We assessed this expectation for an important soil-dwelling group, the springtails, by determining whether alien species show broader thermal tolerance limits and greater tolerance to climate warming than their indigenous counterparts. We found that, from the tropics to the sub-Antarctic, alien species have the broadest thermal tolerances and greatest tolerance to environmental warming. Both groups of species show little phenotypic plasticity or potential for evolutionary change in tolerance to high temperature. These trait differences between alien and indigenous species suggest that biological invasions will exacerbate the impacts of climate change on soil systems, with profound implications for terrestrial ecosystem functioning.

climate change | introduced species | soil fauna | thermal limits | laboratory evolution

The impacts of climate change and biological invasions continue to increase in magnitude (1, 2), yet investigation of their interactions remains fragmented and incomplete (3). The general expectation is that such interactions will benefit alien over indigenous species, and thus have negative consequences for native biodiversity (4). Soil systems are globally important reservoirs of diversity, influence aboveground ecological dynamics, and are critical for food security (5). They are also highly susceptible to climate change and to biological invasions, which both have major effects on soil ecosystem functioning and are expected to interact to the detriment of these systems (6, 7). Understanding how climate change–invasion interactions will play out in soil systems is, therefore, pressing (8). For the soil biota, increasing temperature is one of the most significant components of climate change (9). Differential fitness effects of rising temperatures on indigenous compared with alien species must necessarily be mediated by consistent thermal trait differences between these two groups of species (10). Moreover, if such effects are general, these trait differences between alien and indigenous species should be maintained across a broad latitudinal—and hence climatic—range. No study has yet evaluated for any group of soil macrobiota whether such latitudinally extensive differences in thermal traits exist that advantage alien species under warming conditions, and thus whether forecast impacts are likely to be realized.

To assess this expectation, we investigated the thermal limits to activity in 16 alien and 14 indigenous species of springtails (Collembola) spanning a climatic gradient from the Australian tropics (16°S) to south of the Antarctic Polar Front (54°S) (Fig. 1). Springtails are a globally distributed, functionally significant component of the soil biota (5, 11). The group includes many alien species that have invaded soil systems from the polar regions to the tropics (6, 12). Critical thermal limits to activity form a useful proxy for adult fitness because organisms are incapable of movement outside of these limits, thus reducing fitness (13, 14), with demographic effects reflected in associations between critical thermal limits and geographic ranges in several taxa (15, 16). Upper critical thermal limits are also now widely used to assess tolerance to climate change-associated warming in a wide range of organisms (17, 18). Using a single consistent method [in contrast with multiple methods represented in and which complicate interpretations of

meta-analyses (19)], the F2 generation of populations of each species to overcome parental effects and the potential for laboratory adaptation (20), and taking account of the influence of phylogenetic relatedness (21), we investigated both basal thermal tolerance and the extent of phenotypic plasticity in the upper critical thermal limits.

Although phenotypic plasticity of thermal tolerance traits may only play a limited role in reducing exposure to climate change in some ectotherms (22), it has been identified as a potentially significant mediator of responses to thermal challenge (23), including in springtails (9, 24). Phenotypic plasticity is also thought to be a major contributor to the success of invasive species (25, 26). In addition, total thermal tolerance range and lower critical thermal limits play a role in determining range shifts in response to environmental change (16). Thus, we not only assessed basal tolerance, but also phenotypic plasticity for both upper and lower critical thermal limits. We evaluated whether climate-related variation in upper and lower critical thermal limits, thermal tolerance range, and warming tolerance—the difference between species' upper critical thermal limits and maximum environmental temperatures (17)—differ systematically between the indigenous and alien springtail species. Tests were conducted using phylogenetic generalized least squares (PGLS) (21), and a phylogenetic tree constructed from molecular markers to take into account phylogenetic signal in the traits measured.

Evolutionary responses have the potential to reduce substantially the impacts of climate change on populations (27, 28). Therefore, we also examined the extent to which thermal traits might evolve in springtails and whether this potential differs between indigenous and alien species. We used laboratory natural selection (29) to assess the extent of evolutionary potential in the critical thermal limits

Significance

How climate change and biological invasions interact to affect biodiversity is of major concern to conservation. Quantitative evidence for the nature of climate change–invasion interactions is, however, limited. For the soil ecosystem fauna, such evidence is nonexistent. Yet across the globe, soil-dwelling animals regulate belowground functioning and have pronounced influences on aboveground dynamics. Using springtails as an exemplar taxon, widely known to have species-specific effects on below- and aboveground dynamics, we show that across a wide latitudinal span (16–54°S), alien species have greater ability to tolerate climate change-associated warming than do their indigenous counterparts. The consequences of such consistent differences are profound given globally significant invasions of soil systems by springtails.

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¹To whom correspondence should be addressed. Email: charlene.janionscheepers@monash.edu.

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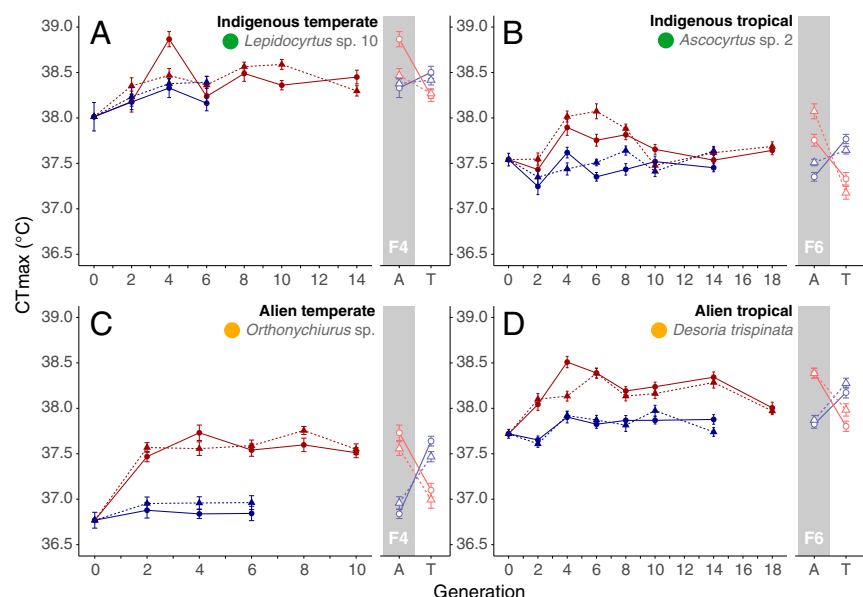


Fig. 3. The response of critical thermal maximum to laboratory natural selection in four species of springtails. Mean CT_{max} ($\pm$ SE) for the temperate indigenous *Lepidocyrtus* sp. 10 (A), tropical indigenous species *Ascocyrtus* sp. 2 (B), temperate alien *Orthonychiurus* sp. (C), and tropical alien *Desoria trispinata* (D) for each of two lines (filled circles/triangles and solid/dashed lines) evolved under an elevated temperature (red: tropical = 27 °C, temperate = 25 °C) and control lines (blue: tropical = 20 °C, temperate = 15 °C). Open symbols to the right indicate the outcomes of a reciprocal transplant experiment at generations four or six, determining the contribution of developmental plasticity to CT_{max} . Here, springtails from selection and control groups were reared under either their standard acclimation temperature (A) or transplanted to the thermal environment of the opposing group (T) and reared for one generation. Inter-generational change in CT_{max} is typically within the scope of developmental plasticity. Effect sizes are typically less than 1 °C (see *SI Appendix*, Tables S2 and S4 for statistics).

evolutionary change in CT_{max} in the springtail species assessed (and in contrast to substantial phenotypic plasticity and potential for evolutionary change in CT_{min}) are in keeping with what has been found for other animal groups (22, 34). They indicate that thermal tolerance differences between the indigenous and alien groups are likely to persist through time. Nonetheless, thermal conditions different to those adopted in the laboratory selection experiment, such as occasional extreme events (39), might be more effective in driving evolutionary change in CT_{max} (28). Indeed, past evolutionary responses to divergent thermal conditions, in conjunction with the ecological conditions that promote the success of alien species (7), may explain the observed trait differences between the two groups of species in the first place. Such a mechanistic macro-physiology of biological invasion remains to be developed.

Given that trait differences between the alien and indigenous groups will likely endure, springtail assemblages should, with ongoing climate change, be more prone to dominance by alien species, irrespective of whether these assemblages are tropical, temperate, or more polar. Indeed, evidence from some areas suggests that such a process is already underway (6, 12, 32). Clearly, interspecific variation exists in thermal traits and warming tolerance among the springtail species investigated here (Fig. 1), and other life-history traits might also play a role (15), suggesting that specific predictions may be complicated. Nonetheless, the general trends we demonstrate support growing concerns that soil systems will be significantly impacted by interactions among climate change and biological invasion (7, 8).

The broader influence of such climate change–invasion interactions is likely to be realized through changes to soil-system dynamics. Springtails are widely known to have considerable effects on soil system functioning, which can result in aboveground changes to system properties and functioning, influencing higher trophic levels (40, 41). For example, soil-dwelling springtails alter aboveground plant biomass and either lower or increase herbivore reproduction depending on the specific host plant (42). Moreover, these effects differ substantially between different springtail species, and influence belowground faunal interactions, such as between earthworms and springtails, again affecting the aboveground component of systems (11, 42). These findings, widespread invasion of soil systems by springtails (6), warming-related dominance of springtail assemblages (12), and different functional effects between alien and indigenous springtail species (43), indicate that interactions between warming and soil invasions will result in substantial changes to system functioning. Thus, differential success of alien over indigenous species as climates change is likely not only to have an

effect on soil biodiversity and belowground ecological functioning, but also on aboveground ecosystems. In consequence, our work suggests that biological invasions stand to increase substantially the already pronounced impacts of climate change on terrestrial ecosystems across the planet.

Materials and Methods

Extended protocol descriptions are provided in the *SI Appendix*.

Collection, Identification, and Alien Species Assignment. The 30 springtail species were collected typically from across Australia (*SI Appendix*, Table S8) between 2013 and 2016. The focus was on hemiedaphic (litter-dwelling) species. Individuals were initially assigned to species in the field and at least 200 individuals collected per species. Collections were returned to the laboratory typically within 1 wk of collection. Species were identified to genus, and where species had been described, to species level using available keys (e.g., refs. 44 and 45). DNA barcoding was used to confirm species identifications. Mitochondrial DNA extraction and sequencing of the cytochrome c oxidase subunit I gene was undertaken by the Biodiversity Institute of Ontario, University of Guelph, Canada, following standard protocols developed for springtails (46). Sequences of 74 specimens from 23 species were compared with the >75,000 springtail sequences available through the Barcode of Life Data Systems (BOLD; www.barcodinglife.org) (*SI Appendix*, Table S9). Individuals that could not be identified using available keys and which were not represented in BOLD were examined by one of the authors (C.J.-S.), in discussion with other systematic experts, and assigned to uniquely identifiable species based on morphological characteristics or a barcoding gap of at least 2.5% (47). Sequences are available on BOLD (www.boldsystems.org) as part of Project COLMU (Collembola of Monash University). Species that were clearly identified as indigenous to Australia or to the sub-Antarctic islands in faunal treatments and those similarly identified as alien to either of these regions were retained as such in the classification of species as alien or indigenous. Undescribed species not represented in BOLD previously, or represented only from individuals already collected across Australia, New Zealand, or south of the Wallace line were considered indigenous. Following previous authors (6), undescribed species that had sequences present in BOLD from other distant tropical regions (such as the Neotropics) or from the Holarctic (typically Europe) were considered alien species (*SI Appendix*, Tables S8 and S9). Given the extensive nature of the BOLD information on springtails (>75,000 sequences, representing several thousand species), and the systematic expertise we consulted, we have high confidence in the species assignments to alien or indigenous species.

Site Microclimate Characteristics. The soil microclimate characteristics of each site were calculated using remote-sensed daytime land-surface temperature data from the MODIS satellite network (MOD11C2 v006; 30 arcseconds spatial resolution; 8-d temporal resolution from January 2001 to December 2015; doi:10.5067/MODIS/MOD11C2.006), which were linearly transformed to account for the diffusion of heat from the land surface to 2.5-cm below the soil surface. The slope of this linear

transformation was derived from the microclim dataset (48), which contains validated estimates of soil temperature for each hour in a 24-h cycle of an average day in each month of an average year under varying shade conditions. The median (MODIS soil median), 99% quantile (MODIS soil99), and maximum soil temperature (MODIS soil max; i.e., the warmest 8-d mean) of each site were calculated from our linearly transformed MODIS time series.

Colony Maintenance. Species were reared at constant rearing temperatures that typically reflect the average soil temperatures of the sites at which they were collected (*SI Appendix, Table S8*) in controlled-temperature incubators (MIR-154; SANYO Electric) on a 12-h light:12-h dark light cycle. Incubator temperatures were monitored using Hygrochron iButtons (DS 1923-F5; Maxim Integrated) (*SI Appendix, Table S8*). The F2 generation was the focus of this work (*SI Appendix, Fig. S4*) to minimize any carryover effects from the environment of origin, including parental effects, and to reduce the possibility of adaptation to laboratory conditions (20). Between 50 and 200 adults from the collected (F0) individuals were randomly assigned to two to four 60-mL pots lined with moistened Plaster-of-Paris: Charcoal powder (9:1) substrates. De-ionized water was added once to twice a week to maintain high humidity and, depending on the species, individuals were fed two to three times a week with algae from the bark of *Platanus* sp. or on slime mold *ad libitum* (49), enabling individuals to select nutrients optimally. Adults from the F2 generation [average egg to adult development time among the species is 74.16 ± 40.56 (SD)] were used for most experiments (*SI Appendix, Fig. S4*).

Acclimation to Assess Phenotypic Plasticity. Before the experimental trials, all species were subject to temperature treatments (referred to “acclimation” hereafter), undertaken in controlled-temperature incubators (MIR-154; SANYO), with temperatures verified using Hygrochron iButtons and under 12-h light:12-h dark conditions (*SI Appendix, Table S8*). Acclimation treatments lasted 7 d, given that complete responses usually occur within less time in terrestrial arthropods (50). Low, medium, and high acclimation temperatures were set 5 °C below, and 5 and 10 °C above standard rearing temperatures, respectively (*SI Appendix, Fig. S4*). For control temperatures, individuals were subject to the same manipulation as those in the acclimation treatments.

Critical Thermal Limit and Warming Tolerance Determinations. Critical thermal limits (CT_{max} and CT_{min}) were determined following standard methods (24, 51). Assays began at the control (rearing) temperature for each species to ensure that comparisons between the control and acclimated groups could be made. Heating (for CT_{max}) and cooling (for CT_{min}) rates were set at 0.05 °C/min, rates that are within the range and close to the mean (for increasing temperatures) of those recorded for tropical to temperate microhabitats, and which empirical data and modeling show provide comparable acclimation and warming tolerance estimates across species and environments (24). Temperature of the substrate was recorded with a type K thermocouple, using a digital thermometer (Model RDXL 125D; Omega Engineering). CT_{min} and CT_{max} were defined as temperature at which a loss of righting response occurred (24, 51). Because body mass may contribute to variation in critical thermal limits (e.g., refs. 9 and 52), species' mean body mass (mg) was determined from a randomly selected, separate sample of 40 adult individuals for each species using a high-resolution (0.1 µg) microbalance (Mettler-Toledo XP2U) (*SI Appendix, Table S8*). For each species, basal thermal tolerance was calculated as the mean CT_{min} for the lower critical thermal limit and mean CT_{max} for the upper critical thermal limit, obtained from individuals reared under control conditions and subjected to the same temperature for acclimation. Basal thermal tolerance range was calculated as the difference between these two mean values. To determine the extent of phenotypic plasticity, the ARR (22) was calculated for each CT_{min} and CT_{max} for each species. Here, the ARR was calculated as the largest difference between mean CT_{min} (or CT_{max}) across any of the acclimation treatments, divided by the maximum temperature range represented by those treatments. Warming tolerance was calculated as the difference between mean basal CT_{max} (individuals reared under control conditions) of each species and the MODIS 99% quantile (MODIS soil99).

Selection Experiment. Laboratory natural selection (29) was used to investigate the ways in which critical thermal limits respond to elevated rearing temperatures. Four species were selected for this experiment, representing a tropical alien (*Desoria trispinata*), a tropical indigenous (*Ascochyrtus* sp. 2), a temperate alien (*Orthonychiurus* sp.), and a temperate indigenous species (*Lepidocyrtus* sp. 10). Laboratory natural selection rather than artificial selection was chosen to minimize the risk of sterilization and other cellular damage that can be associated with exposure to extreme temperatures (53–55). Selection and control groups were initiated for each species from the F2 generation of field-caught individuals. F2 individuals were used to minimize any carryover effects from the environment of origin, including parental effects, and to reduce the possibility of adaptation to laboratory conditions (20, 53). Each group

contained two independent replicate lines starting with 150 individuals divided into two separate vials per replicate line. Control lines were maintained under the original rearing temperatures (temperate = 15 °C, tropical = 20 °C), while selection lines were maintained under warmer temperatures (temperate = 25 °C, tropical = 27 °C). Temperatures for the selection treatment were based on results from a pilot study, which indicated that 25 °C and 27 °C were the highest temperatures at which the temperate and tropical species could still reproduce, respectively (*SI Appendix, Table S10*). Throughout the experiment, generations remained discrete, and eggs from replicate vials within each replicate selected and control line were randomly combined within generations to maintain genetic diversity. Population size in each line (control and selection lines) was on average 775 individuals. Critical thermal limits were assessed, as for the interspecific comparisons, for adults of each species before selection commencing (at generation 0), then every second generation for individuals in the selection and control groups up to generation 10, and every fourth generation thereafter. Approximately 45 individuals were assessed per replicate line per treatment at each sampling period. The degree of plasticity associated with any phenotypic changes observed during the selection experiment was assessed using a reciprocal transplant experiment, investigating developmental plasticity. This involved switching individuals from the selection conditions to the control conditions and vice versa. Individuals were switched within 1 d of hatching at generation four for the temperate species and generation six for the tropical species. These different generations were used because of the slower development time of the temperate species. Thermal limits were assessed when the switched individuals had reached adulthood.

Statistical Analyses. All analyses were conducted in R v3.3.1 (56). PGLS (57), as implemented in the caper v0.5.2 (58) and APE (59) packages, was used to investigate relationships between species mean critical thermal limits (either CT_{min} , CT_{max} , or thermal tolerance range), environmental characteristics (MODIS soil median), springtail species mean mass, and species status (alien or indigenous).

A phylogeny for the species was constructed based on joint considerations of two recent molecular phylogenies for the group (60, 61), with species relative positions based on the cytochrome c oxidase subunit I gene phylogeny, or in a few cases on morphological similarity adjudicated by one of us (C.J.-S.). The barcoding placements were obtained from a neighbor-joining tree using standard methods. For the final tree, branch lengths were assigned using Grafen's method (62), and the tree (*SI Appendix, Fig. S5*) is available as a Newick file. Initially, two covariance matrices were constructed following either Brownian motion (BM) or Ornstein–Uhlenbeck (OU) models of evolution (63). Akaike Information Criterion values of BM and OU models were compared to identify which model of evolution provided the best fit to observed data using the APE (59) and nlme v3.1-131 (64) packages. Phylogenetically corrected models based on BM covariance matrices were a consistently better fit than those based on other evolutionary assumptions (*SI Appendix, Table S11*), thus the outcomes of these models are reported primarily, although for comparative purposes we provide the OU outcomes as well (*SI Appendix, Table S12*). For the BM models, a maximum-likelihood approach provided Pagel's λ (21) to indicate the degree of phylogenetic correlation in the data. The PGLS approach was used to investigate relationships between warming tolerance, maximum soil temperature (MODIS soil maximum), species mass, and species status, and to investigate relationships between CT_{min} and CT_{max} . Analyses were repeated using ordinary least-squares approaches as implemented in the linear model function of R v3.3.1, and coefficients were typically similar to those found in the PGLS models (*SI Appendix, Table S13*). Throughout, mass did not appear as a significant term in the models, and in no cases did slopes of the relationships between critical thermal limit traits and environmental features differ between the alien and indigenous species groups. For investigations of the ARR and its relationship with mean trait values, ordinary least-squares methods indicated no significant relationships and PGLS bore out these conclusions.

For the selection experiment, to analyze differences in critical thermal limits between selection and control lines, nested mixed-effect model analyses were conducted using the lmer function in the lme4 package (v1.1-13) (65) in R (v3.3.1). “Treatment” (control or selection) and “generation number” were treated as fixed effects, and “replicate line” was nested within treatment as a random effect (66). Nested mixed-effect analyses were also undertaken to analyze data from the reciprocal transplant experiment examining developmental plasticity. This involved comparing the critical thermal limits of the selection, control, and reciprocally transplanted lines at the respective generation of the reciprocal transplant experiment, with “replicate line” nested within treatment (control, selected, reciprocally transplanted) as a random effect. Separate analyses were performed for each of the four species.

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