INTRODUCTION

It has been known for decades that moderate dietary restriction (DR) extends longevity in a wide variety of experimental animals (Fontana et al., 2010). Many studies have investigated the specific diet components (especially calories, protein or specific amino acids) whose restriction can induce this response (Grandison et al., 2009; Lee, 2015; Lee et al., 2008; Piper & Partridge, 2007; Soultoukis & Partridge, 2016), and the physiological mechanisms involved (Green et al., 2022; Zanco et al., 2021). While DR is seen as a potential means to extend human lifespan and healthspan (Heilbronn & Ravussin, 2003; Maegawa et al., 2017; Pifferi & Aujard, 2019), the life-extending effects of DR are also of considerable interest from an evolutionary perspective. However, the evolutionary interpretation
of these effects remains controversial (Adler & Bonduriansky, 2014; Kirkwood & Rose, 1991; Kirkwood & Shanley, 2005; Moatt et al., 2020; Piper et al., 2023).

An influential evolutionary hypothesis posits that the physiological responses that enhance longevity when nutrient intake is restricted represent an evolved strategy that functions to enable animals to survive periods of famine (Holiday, 1989; Kirkwood & Rose, 1991; Kirkwood & Shanley, 2005). According to this ‘adaptive resource reallocation’ hypothesis, DR induces a reallocation of metabolic resources from reproduction to somatic maintenance, and an up-regulation of cellular recycling and repair processes such as apoptosis, because these responses reduce mortality and thereby increase the probability of surviving until resources become abundant again and reproduction can resume. This hypothesis is supported by evidence that Drosophila melanogaster females are able to resume full reproduction after a period of DR (Sultanova et al., 2021), although increased nutrient abundance following a period of DR can result in elevated mortality that could negate the reproductive gains (McCracken et al., 2020).

Responses to DR have been likened to seasonal diapause, when reproduction ceases and specialized survival mechanisms are activated (Regan et al., 2020). However, a key assumption of the adaptive resource reallocation hypothesis is that the strategy of foregoing reproduction while resources are scarce can pay off in wild insects and other animals during the breeding season. This assumption is problematic in light of the ecology of natural populations and the physiological effects of DR. Although estimates of mortality rates in wild insects are scarce, the available data suggest that reproductively active adult flies experience a mortality rate of ~10% per day (Bonduriansky & Brasili, 2002; Kawasaki et al., 2008; Mautz, Rode, et al., 2019), and estimates from many other insects are similarly high (Zajitschek et al., 2019; Zajitschek & Bonduriansky, 2014). Such a high risk of death means that any delay in reproduction is likely to be very costly for most insects and other small-bodied animals. This problem could be overcome if physiological responses to DR substantially reduce mortality risk in the wild, but there is little evidence to suggest such an effect. Mair (2005) found that DR enhanced the ability of Drosophila melanogaster females (but not males) to resist starvation, but reduced females’ ability to resist desiccation; no effect was observed on resistance to cold, heat or oxidative stress (induced by paraquat). Other studies have reported increased frailty in nutrient-restricted animals subjected to stressful conditions, including a reduced capacity to mount an immune response against pathogens, reduced ability to heal wounds and reduced cold tolerance (reviewed in Adler & Bonduriansky, 2014). For example, in Drosophila ananassae, dietary protein enhanced desiccation and heat shock resistance (Sisodia & Singh, 2012) and, in Drosophila melanogaster, wing clipping reduced longevity of protein-restricted flies but not of fully fed flies (Ghimire & Kim, 2015). The ability to tolerate environmental challenges such as thermal stress and wounding is relatively unimportant in the typically benign conditions of the laboratory, but is likely to have a considerable impact on survival of wild animals living in natural environments.

Furthermore, in insects and other small-bodied animals, many physiological processes are tightly coupled to ambient temperature (Keil et al., 2015; Moloń et al., 2020; Sestini et al., 1991). Consequently, physiological responses (such as reduced metabolic rate) and behavioural strategies (such as reduced activity) that can be deployed to reduce mortality risk during cold-season diapause might be limited or impossible during the breeding season, regardless of nutrition. Moreover, protein restriction induces increased activity levels in D. melanogaster (Ghimire & Kim, 2015; Krittika & Yadav, 2020), and this behavioural response could elevate risk of predation for protein-restricted animals in the wild.

Assessing the adaptive resource reallocation hypothesis therefore requires verifying the key premise that nutrient limitation can increase survival of nondiapausing animals not only under benign conditions but also in challenging, stressful environments. If nutrient limitation fails to reduce (or even increases) mortality risk under stress, then life-extension under DR is likely to be an artefact of laboratory conditions rather than an evolved strategy that enhances fitness in natural populations.

Much of the evidence on the lifespan-extending effects of DR comes from experiments with flies and other insects, and a general finding across many insect species is that lifespan is extended when protein intake is reduced at the adult stage (Adler et al., 2013; Lee, 2015; Maklavok et al., 2008; Regan et al., 2020; Zanco et al., 2021), and perhaps also during development (Runagall-McNaull et al., 2015). In the most important model insect system for DR research, Drosophila melanogaster, it is well-established that protein-restricted flies typically outlive fully fed flies, but protein-restricted females tend to be less fecund (Lee, 2015; Lee et al., 2008; Simpson et al., 2017; Sultanova et al., 2021). However, the vast majority of DR experiments on D. melanogaster and other insects have been carried out under benign laboratory conditions, including a stable ambient temperature to which the flies are well-adapted (25-27°C), few pathogens and parasites, and no predators. Only a few studies have investigated effects of DR on survival in more challenging environments. Burger et al. (2006) found that diluted diets reduced mortality from bacterial infection but also reduced oxidative stress resistance and cold tolerance in D. melanogaster females. Savola et al. (2021) found that mortality from infection with a bacterial pathogen was reduced under moderate protein restriction but increased under severe protein restriction in D. melanogaster females. Savola et al. also found that survival of injured females was not affected by dietary protein. Mautz, Rode, et al. (2019) manipulated access to protein in both captive and wild cohorts of male antler flies (Protopiophila litigata), and found that protein supplementation affected the longevity of adult males differently in the laboratory versus the wild, although this protein × environment interaction was supported in only one of the 2 years of the study. The results of these studies therefore suggest that effects of dietary protein on longevity can be modulated by stress, but also underscore a need for further research.

Here, we asked whether DR (protein restriction) enhances longevity of D. melanogaster females and males not only in benign
environments but also under thermal stress. Temperature fluctuations are experienced by all natural populations, and their effects are likely to pose especially severe challenges to small-bodied, ectothermic animals such as insects. Ambient temperature affects the expression of a range of physiological and life-history traits in D. melanogaster (Mołoń et al., 2020; Sestini et al., 1991), and exposure to both cold and hot temperature extremes can be stressful (Arias et al., 2012; Klepsatel et al., 2016; Mockett & Matsumoto, 2014). The potential for temperature stress to modulate the effects of DR on lifespan is therefore especially relevant to understanding how DR might affect fitness in wild flies and other wild animals. We manipulated adult dietary protein (40%, 100%, and 150% of the normal yeast concentration, with other diet components held constant) across a gradient of ambient temperature, including below-normal temperature that is likely to induce cold stress (21°C and 23°C), normal rearing temperature to which the flies are likely to be well adapted (25°C and 27°C), and above-normal temperature that is likely to induce heat stress (29°C). Given the potential for trade-offs between survival and reproduction in Drosophila (Flatt, 2011), we quantified both longevity and reproductive performance (female fecundity, male mating success) to determine whether negative effects of environmental conditions on longevity might be offset by positive effects on reproduction. Likewise, we examined effects of diet and temperature treatments on performance of the offspring, grand-offspring and great-grand-offspring of exposed flies to determine whether treatment effects on exposed individuals might be offset by effects on their descendants (see Mautz, Lind, et al., 2019).

2 MATERIALS AND METHODS

Experimental flies were sourced from the outbred Dahomey population. This population was started in 1970 from founders caught in the wild in Dahomey (now Benin), West Africa, and has been maintained in population cages, containing several thousand individual males and females, with overlapping generations. In the laboratory, D. melanogaster are usually maintained at a constant temperature of ~25°C (Stocker & Gallant, 2008), but the Dahomey population was probably adapted to somewhat warmer conditions in its natural environment in Benin, where mean temperature ranges from 29 to 36°C during the day and 22 to 25°C at night over the course of the year (https://www.worlddata.info/africa/benin/climate.php). We therefore assumed that 25–27°C is a temperature range to which the flies are adapted. D. melanogaster and other insects generally live longer at cooler temperatures, but also take longer to develop and reproduce (Keil et al., 2015). In the wild, where extrinsic mortality rate is very high for flies, cooler temperatures are therefore likely to result in reduced lifetime reproductive output. High temperature (over 27°C, and especially over 30°C) is generally stressful for Drosophila and other insects (Klepsatel et al., 2016; Mołoń et al., 2020). We therefore chose 21°C and 23°C to represent moderately stressful cold temperatures, 25°C and 27°C to represent ‘benign’ temperatures, and 29°C to represent a moderately stressful hot temperature.

Females in population cages were given the opportunity to lay eggs in vials containing standard sugar-yeast (1.0 SY) diet. Eggs were collected and distributed to 100 new 1.0 SY food vials (10mL food per vial; 50 eggs per vial). Offspring were collected and allowed to mate with offspring of a different vial for the first 48h of adult life (20 males and 20 females from each vial, resulting in 40 adults per combined vial). At age 3 days, males were removed, females were transferred into new vials (20 per vial) and allowed to lay eggs for 5h. Egg number per vial was trimmed to 60–70. Flies were reared for another generation, using the same protocols, and the eclosing adults of the third generation were used as focal experimental subjects. Males and females were given the opportunity to mate for 48h after eclosion, before the sexes were separated and flies were distributed in treatment groups.

We created three different diets by manipulating the protein content and therefore the protein to carbohydrate ratio (protein content in gram yeast per litre diet: 40 = protein restricted, 100 = standard, 150 = protein rich; Table S1), while maintaining the amount of sugar in each diet (at 50 g/litre diet). The effects of each of the three diets were tested at each of five temperatures (21, 23, 25, 27 and 29°C), resulting in 15 treatment groups. The experiment was performed in a controlled temperature and humidity room, at 21°C and 60% relative humidity, with a 12:12h light: dark cycle. Treatment temperatures above 21°C were established with heat mats, regulated by digital thermostats, placed in clear plastic containers (52 × 35 × 28 cm). Vials in the 21°C treatment were placed in an identical container, containing no heat mat. Vials were placed upright in cardboard trays (on plastic frames, 10 cm above heating mats), on top of a piece of cardboard with the same area as the tray (0.5 cm thick in total) to avoid differences in temperature due to potentially localized differences in heat produced by the heating mat (temperatures at 10 cm above heating mats were tested at different locations in each temperature container and not found to be different, prior to the experiment). Vials were randomly allocated within trays every time flies were flipped to new vials. Containers were fitted with lids, leaving a 2 cm wide space open at one short side of the container to allow for air exchange. We monitored temperature inside the boxes with individual temperature loggers, placed at the height of the food surface in experimental vials. Temperatures in the highest temperature treatment never exceeded the set treatment temperature by more than 0.4°C (mean, variance, minimum, maximum for temperature: ambient: 20.58°C, 0.09, 20.20°C, 22.30°C; 21°C container: 20.69°C, 0.06, 20.21°C, 22.15°C; 23°C container: 23.19°C, 0.13, 22.84°C, 23.66°C; 25°C container: 24.83°C, 0.11, 23.88°C, 25.42°C; 27°C container: 27.14°C, 0.02, 26.40°C, 27.70°C; 29°C container: 28.58°C, 0.29, 26.65°C, 29.4°C). Humidity inside the containers was not monitored. To minimize potential interactive effects between treatment temperature and humidity within vials, we used high-density foam Drosophila vial plugs which are specifically designed to reduce evaporation.
For survival estimates, each vial was populated with 10 individuals, with 10 replicate vials per sex per treatment combination (see Table S2 for more details).

Live flies were transferred to new food vials and the number of dead flies per sex was recorded every Monday, Wednesday and Friday. Female fecundity was estimated by collecting and counting eggs laid by females that were used to measure survival, over 18 h, once a week (Wednesday to Thursday), for the first 5 weeks of the experiment (see Table S3 for more details).

2.1 Parental effects and male mating behaviour

To measure male mating behaviour and test for parental temperature and diet treatment effects on offspring, we established additional vials per treatment and sex (six vials with males, three vials with females), containing 20 individuals (male mating behaviour F0 [MMB], paternal effect F0 [PE] and maternal effect F0 [ME]). Males and females experienced the same treatment as the flies used to measure survival, except females were collected as virgins and their fecundity was not measured.

To test for paternal effects, we mated PE males at age 16 days to 5-day-old standard virgin females which originated from the same Dahomey population and were bred for three generations following the same protocol as used for the focal experimental flies (F0 survival, PE and ME flies). For this, the 20 PE males per vial were paired with 20 virgin females for 24 h. Sexes were separated and females were allowed to lay eggs for 24 h on standard 1.0 SY food vials (7 mL food per vial). To test maternal effects, we mated virgin ME females at age 16 days to 5-day-old standard virgin females which originated from the same Dahomey population and were bred for three generations (mothers' age changed from 16 days in F0 to 10 days in F1 and F2 due to logistical reasons). Transgenerational effects can wane over generations (Bonduriansky, 2021). To maximize the potential to detect effects of F0 diet and temperature treatments on F2 and F3 descendants, we therefore maintained F2 and F3 adult flies at a high ambient temperature (29°C) to gauge their capacity to cope with heat stress. F2 and F3 flies were maintained on standard 1.0 SY diet, and in two replicate vials (with 15 individuals each) per sex for each parental F0 sex-specific treatment combination. For practical reasons, F2 and F3 descendants were obtained only from a subset of F0 sex-specific treatment combinations (21°C, 25°C and 29°C, and restricted (0.4 SY) and rich (1.5 SY) diets, resulting in six grandparental treatments and six great-grandparental treatments), and we only tested for effects of F0 diet and temperature treatments on male mating performance.

Male mating performance was tested for 14 males per treatment, seven sourced from each of two replicate vials, in the parental (F0), offspring (F1), grand-offspring (F2) and great-grand-offspring (F3) generations. First, standard 5-day-old virgin females, maintained in groups of 20 females until the behavioural assay, were aspirated into behavioural vials (one female per vial). Vials contained 5 mL of 0.4 SY food and were closed with a white foam plug, leaving a 5 cm high (~20 mL) mating arena. Treatment males were then aspirated and paired one-on-one with females. Pairs were randomly placed in the behaviour observation trays. We recorded time until mating (latency to mate), mating duration and whether a mating took place (mating success). Observations were performed for 3 h, starting at 9 am, in a controlled temperature room at 25°C.

2.2 Statistical analysis

We analysed survival of F0 flies in binomial generalized linear mixed models (GLMMs; function glmer, package lme4, Bates et al., 2015), since the assumption of proportionality of hazard rates in Cox proportional hazards models (function coxph, package survival, Therneau, 2020b) was violated (tested with function cox.zph, package survival and visually inspected; Therneau, 2020b). In two separate model sets for males and for females, we compared five models, starting with the most complex model that included the interaction between diet and the nonlinear term of temperature (modelled as natural splines, function ns, R core package splines), and including all nested models. In all models, individual vial was modelled as a random effect. We used AIC to compare models within sex (function model.sel, package MuMIn, Barton, 2020). If a model that contained diet was supported most, we further tested the effect of diet at each temperature with logrank tests and Benjamini & Hochberg correction for multiple comparisons (function pairwise_survdiff, package survminer, Kassambara & Kosinski, 2019). Sex-specific parental effects on survival on male and female F1 flies were tested with sex-specific mixed effects Cox proportional hazards models (function coxme, package coxme, Therneau, 2020a), since the proportionality of hazards assumption, as tested with function cox.zph (package survival, Therneau, 2020b), was supported.

We estimated total fecundity for each vial as total sum of weekly egg counts in the first 5 weeks after establishing the vial. For F0, we
analysed the square root of fecundity values in robust general linear models (function `glmrob`, package `ROBUSTbase`, Maechler et al., 2020), as the diagnostics of general linear models showed problematic patterns at both sides of the observed range of data. Temperature was modelled as a cubic B-spline (function `bs` in R core package `splines`) and was tested in interaction with diet (`glm` with the same model specifications yielded qualitatively similar results). Diet was included as a numerical variable, corresponding to its protein content, in linear and quadratic form. We tested the effects of model terms by first removing the most complex term, in this case the interaction between temperature and the quadratic term of diet, and comparing it to the full model, using Wald tests. To test effects of F0 parental diet and sex on F1 fecundity, we adjusted our analytical approach, due to lower F1 fecundity sample sizes, compared to F0 fecundity. Variables diet and temperature were categorized or diet experienced by male or female parents (Tables S6 and S7; compared to daughters from fathers on low diet and higher temperatures (Figure 1).)

Survival of offspring (F1) was not affected by either temperature or diet experienced by male or female parents (Tables S6 and S7; model simplification did not change this conclusion).

3.2 | Fecundity

The most complex model of F0 fecundity that includes a cubic B-spline of temperature, diet squared and their interaction provided the best fit (compared to a reduced model without the interaction term `diet^2` × temperature: change in robust quasi-deviance = 40.13, df = 3, \(p < 0.001\)). Fecundity of F0 females on the restricted diet was substantially lower than fecundity of females on the standard and rich diets at all temperatures except 29°C (Table S8; Figure 2). For females on the standard and especially the rich diet, fecundity tended to decline with temperature, reaching levels similar to the restricted diet at 29°C (Table S8; Figure 2).

Fecundity of F1 females in maternal and paternal groups was predicted best by the most complex model (maternal: \(W = 2.754, p = 0.043\); paternal: \(W = 3.993, p = 0.010\)). Differences in fecundity between temperatures within maternal diet treatment groups were only significant in four comparisons in the rich maternal diet group, with fecundity of flies from low temperature mothers higher than from medium temperature mothers (Table S9; Figure 3; estimates of differences (standard error) between 25°C and 21°C: −207.50 (43.86), \(t = −4.066, p = 0.010\); 27°C–21°C: −168.00 (53.67), \(t = −3.466, p = 0.026\); 25°C–23°C: −111.50 (7.91), \(t = −14.104, p < 0.001\); 27°C–23°C: −90.00 (18.39), \(t = −4.895, p = 0.003\)). Daughters from fathers from the low diet and low temperature group had higher fecundity compared to daughters from fathers on low diet and higher temperatures (Table S10; Figure 4; 25°C–21°C: −79.50 (21.51), \(t = −3.697, p = 0.022\); 27°C–21°C: −118.00 (33.53), \(t = −3.520, p = 0.023\); 29°C–21°C: −165.00 (43.86), \(t = −3.762, p = 0.022\); 27°C–23°C: 96.50 (17.10), \(t = 5.642, p = 0.001\)).

3.3 | Male mating behaviour

Latency to mate of males in the parental (F0) generation increased linearly with treatment temperature, independently of diet (Figure 5;
There were no effects of temperature or diet treatments on F0 males’ mating duration or mating success (Tables S12 and S13).

Maternal temperature and diet treatments did not affect sons’ (F1) latency to mate, mating duration or mating success (Tables S14–S16). We found no paternal effect on sons’ (F1) latency to mate, mating duration or mating success.

**Table S11**: Temperature-dependent smooth model, approximate significance of temperature smooth term: edf = 1, F = 14.29, p < 0.001).
latency to mate (Table S17). However, paternal temperature treatment had a nonlinear effect on sons’ (F1) mating duration (Figure 6; Table S18; Temperature-dependent smooth model, approximate significance of temperature smooth term: edf = 2.75, $F = 3.712, p = 0.049$). Paternal diet did not affect sons’ mating performance. The best model of paternal effects on sons’ mating success contained a nonlinear temperature term and no diet effect, but the smooth term itself was not significant (Table S15; edf = 1, $\chi^2 = 0.003, p = 0.954$).

Grandmaternal temperature had a quadratic effect on grandsons’ (F2) latency to mate, independent of grandmaternal diet (Figure 7; Table S19). Males whose grandmothers had been maintained at 25°C took longer, on average, to start mating compared to males whose grandmothers had been maintained at 21°C or 29°C (Figure 7). Males whose grandfathers had been maintained on low diet had a shorter mating duration compared to males whose grandfathers had been maintained on high diet at 25°C but not at 21 or 29°C (Figure 8; Table S20). For all other mating behaviour traits in F2 and F3 males, we found no evidence of grandparental or great-grandparental effects (Tables S21–S30).
4 | DISCUSSION

We found that a protein-restricted adult diet was associated with increased longevity of Drosophila melanogaster at benign ambient temperatures, prolonging life of males at 25°C and of both sexes at 27°C. By contrast, a protein-restricted diet did not increase longevity of either sex under conditions of cold stress (21°C and 23°C) or heat stress (29°C). At stressful temperatures, protein-restricted males had similar lifespans to males on protein-standard and protein-rich diets. Protein-restricted females had substantially shortened lifespans at 21°C and 23°C, but protein restriction had little effect at 29°C. These results are consistent with previous evidence that dietary restriction can increase frailty under some ecologically relevant forms of environmental stress (Adler & Bonduriansky, 2014; Burger et al., 2006; Savola et al., 2021). Our findings suggest that protein-restricted insects are unlikely to achieve extended lifespans in natural environments, where temperature stress is ubiquitous. Our results therefore challenge the idea that the longevity-extending effect of DR seen under benign laboratory conditions represents an adaptive survival strategy.

Treatment effects on reproductive performance did not offset the effects of temperature stress on longevity. In females, fecundity of protein-restricted individuals was lower than that of individuals on standard or high-protein diets at all temperatures except 29°C. Interestingly, at cold temperatures (21–23°C), females maintained on high- and standard-protein diets simultaneously maximized both their longevity and their lifetime fecundity, whereas protein-restricted females had dramatically shortened lives and laid fewer eggs over their lifetimes. In males, mating success was not affected by either diet or temperature treatments but male latency to mate (under standard temperature conditions) increased with treatment temperature. Thus, we found no evidence that the failure of protein restriction to prolong life under cold- or hot-temperature stress was associated with positive effects on reproduction. While the costs of reproduction can result in trade-offs between lifespan and reproductive effort (Flatt, 2011), our results instead suggest that temperature stress imposes physiological costs that reduce both survival and reproduction. If our results can be generalized to natural populations, they suggest that protein-restricted flies experiencing temperature stress might not only fail to achieve greater longevity but also fail to achieve a higher reproductive rate than fully fed flies.

Investment in somatic maintenance or fecundity could trade-off with investment in offspring quality (Fox & Czesak, 2000; Kölliker et al., 2015). However, we found no clear evidence that protein-restricted flies produced higher-quality offspring. Protein-restricted maternal diet reduced daughters’ fecundity at both low and high maternal ambient temperatures. Thus, for females experiencing low ambient temperatures, access to abundant dietary protein enhanced longevity, fecundity and offspring quality. Dietary protein could enhance offspring quality by providing the essential building blocks for yolk synthesis (Mirth et al., 2019), and abundant protein could be especially important at low temperatures, which limit flies’ ability to feed (Klepsatel et al., 2019). Protein-restricted paternal diet increased daughters’ fecundity, but this effect was only apparent at low paternal temperature treatments (21–23°C). Protein restriction did not enhance F0 male mating performance, nor mating performance of sons (F1), grandsons (F2) or great-grandsons (F3). Indeed, the only effect of diet treatment on mating performance of male descendants was a positive effect of grandpaternal dietary protein at 25°C on mating duration of grandsons (F2). Thus, protein restriction did not induce consistent, positive effects on offspring quality that could offset the negative effects on parental longevity and fecundity.

Rather, the temperature- and diet-induced maternal and paternal effects that we observed were complex and likely to reflect a combination of adaptive and deleterious responses. Environment-induced parental effects could reflect adaptive parental strategies that enhance offspring performance in the environmental conditions that offspring are likely to encounter (Bernardo, 1996), or that enhance the performance of offspring produced by parents in high conditions (Bonduriansky, 2021). Our results suggest that D. melanogaster females maximized their condition when ambient temperature was low and dietary protein was abundant, and females maintained under such conditions also appeared to transfer their high condition to their daughters. Alternatively, such effects could occur as deleterious consequences of parental stress (Bell & Hellmann, 2019; Bernardo, 1996; Bonduriansky, 2021). Higher temperature might result in elevated stress for D. melanogaster males, potentially explaining why daughters’ fecundity tended to decrease with the ambient temperature experienced by their fathers.

Our results suggest that thermal stress imposes physiological costs that elevate requirements for dietary protein in homeostasis and somatic maintenance. High ambient temperature accelerates metabolic processes in insects, and this probably results in more rapid deterioration of somatic cells and tissues (Molordi et al., 2020). Dietary protein requirements might therefore increase at higher ambient temperatures because of a greater need to repair and replace damaged cells. Moreover, D. melanogaster and many other animals respond to heat stress by producing protective heat shock proteins (Tower, 2011), and the need to synthesize these proteins could elevate requirements for dietary protein. Because cold temperature slows metabolism in Drosophila, the positive effects of dietary protein on survival at colder ambient temperatures are more intriguing. However, some Drosophila enzymes exhibit increased activity in response to low temperatures (Burnell et al., 1991), and the need to synthesize these enzymes could elevate requirements for protein. Another possibility is that, because cold shock alters concentrations of some amino acids in Drosophila suzukii (Enriquez et al., 2018), the need to maintain homeostasis under cold conditions could increase dietary requirements for particular amino acids. Studies examining the effects of DR on expression of proteins (e.g. Gao et al., 2020), especially if combined with manipulation of ambient temperature, could provide additional clues on how dietary protein requirements for homeostasis and somatic maintenance are altered by thermal stress.
The life-extending effect of DR has been reported in a broad range of animals (Fontana et al., 2010), and involves highly conserved physiological pathways (Kapahi et al., 2017). The DR response has therefore been interpreted as a highly conserved physiological mechanism that enhances fitness in insects, mammals and many other animals by helping individuals to survive periods of famine (Kirkwood & Shanley, 2005). However, this idea is based on the assumption that dietary restriction tends to prolong life not only under benign laboratory conditions but also under the more stressful conditions experienced by wild animals, including small-bodied animals such as insects. This assumption has rarely been tested, and is challenged by evidence that dietary restriction reduces ability to cope with a range of stresses (Adler & Bonduriansky, 2014). Our results show that protein restriction extends lifespan under benign temperature conditions but fails to extend (and can even shorten) life of flies experiencing thermal stress. Our findings therefore suggest that the lifespan-extending effect of DR (specifically, protein restriction) reported in many laboratory experiments is more plausibly interpreted as an artefact of benign laboratory conditions than as a fitness-enhancing strategy that evolved in natural populations.

Our experimental diets varied in protein concentration while keeping carbohydrates (and other diet components) constant. These diets were therefore not isocaloric: the protein-restricted diet was less calorie dense than the standard diet, while the protein-rich diet was more calorie dense than the standard diet. Consequently, we cannot exclude the possibility that caloric content rather than protein content was responsible for diet treatment effects on lifespan and reproduction. Nonetheless, because dietary protein has been shown to affect longevity and reproduction in many studies on Drosophila melanogaster and other animals, variation in dietary protein content in our experimental diets is the most plausible factor responsible for the effects we observed. Note, however, that our broad interpretation would remain unchanged regardless of whether protein or calories were responsible for these effects. If our restricted diet tended to prolong life at benign temperatures but not at stressful temperatures because of an interaction of caloric intake (rather than protein intake) with temperature stress, this would still challenge the adaptive resource reallocation hypothesis by showing that dietary restriction fails to prolong life in stressful environments. However, it is possible that caloric restriction (or restriction of other diet components) would interact differently with temperature stress or other types of stress. Further research is needed to investigate this possibility.

DR’s effects on frailty appear to depend strongly on the type of stress that experimental animals experience (Burger et al., 2006; Mair, 2005; Savola et al., 2021), and could also be taxon specific (Adler & Bonduriansky, 2014). For example, it is possible that DR could promote extended longevity in large-bodied animals that experience relatively low mortality rates in the wild. Further studies are needed to determine how DR affects longevity in different species, and in response to different types of stress, such as pathogens and parasites, toxins, thermal stress and interactions between these different types of stress. Further research is also needed to identify the most important sources of stress and mortality in natural populations of insects and other animals. Such work could help to clarify how DR affects survival and fitness in natural populations, how such effects vary across taxa, and how the life-extending effect of DR evolved.

**AUTHOR CONTRIBUTIONS**

Felix Zajitschek and Russell Bonduriansky designed the study. Felix Zajitschek, Susanne R. K. Zajitschek and Ana C. O. Vasconcelos collected the data. Felix Zajitschek analysed the data. Russell Bonduriansky and Felix Zajitschek wrote the first draft of the manuscript, and all authors contributed substantially to revisions.

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**CONFLICT OF INTEREST STATEMENT**

The authors have no conflict of interest to declare.

**DATA AVAILABILITY STATEMENT**

Data have been deposited in the Dryad Digital Repository http://doi.org/10.5061/dryad.rv15dv4d3 (Zajitschek et al., 2023).

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**SUPPORTING INFORMATION**

Additional supporting information can be found online in the Supporting Information section at the end of this article.

**Table S1.** Diet composition.

**Table S2.** Summary statistics for survival of F0 female and male flies.

**Table S3.** Summary statistics for female fecundity in the F0 generation.

**Table S4.** Cox GAM survival model comparisons for generation F0.

**Table S5.** Pairwise logrank survival model comparisons (corrected for multiple comparisons) for generation F0.

**Table S6.** Cox survival model results for female offspring (generation F1).

**Table S7.** Cox survival model results for male offspring (generation F1).

**Table S8.** Robust GLM model results for female fecundity (generation F0).

**Table S9.** Robust GLM model results for female fecundity (generation F1).

**Table S10.** GAMM male latency to mate model comparison for generation F0. Columns ‘Temperature’ and ‘Diet’ indicate whether these predictors (either as smooths or as linear/intercepts) were part of the specific model.

**Table S11.** GAMM male mating duration model comparison for generation F0. Columns ‘Temperature’ and ‘Diet’ indicate whether these predictors (either as smooths or as linear/intercepts) were part of the specific model.

**Table S12.** GAMM male mating success model comparison for generation F0. Columns ‘Temperature’ and ‘Diet’ indicate whether these predictors (either as smooths or as linear/intercept term) were part of the specific model.

**Table S13.** Maternal effect GAMM male latency to mate model comparison for generation F1. Columns ‘Temperature’ and ‘Diet’ indicate whether these predictors (either as smooths or as linear/intercept term) were part of the specific model.

**Table S14.** Maternal effect GAMM male mating duration model comparison for generation F1. Columns ‘Temperature’ and ‘Diet’ indicate whether these predictors (either as smooths or as linear/intercept term) were part of the specific model.

**Table S15.** Maternal effect GAMM male mating success model comparison for generation F1. Columns ‘Temperature’ and ‘Diet’ indicate whether these predictors (either as smooths or as linear/intercept term) were part of the specific model.

**Table S16.** Paternal effect GAMM male latency to mate model comparison for generation F1. Columns ‘Temperature’ and ‘Diet’ indicate whether these predictors (either as smooths or as linear/intercept term) were part of the specific model.

**Table S17.** Paternal effect GAMM male mating duration model comparison for generation F1. Columns ‘Temperature’ and ‘Diet’ indicate whether these predictors (either as smooths or as linear/intercept term) were part of the specific model.

**Table S18.** Paternal effect GAMM male mating success model comparison for generation F1. Columns ‘Temperature’ and ‘Diet’ indicate whether these predictors (either as smooths or as linear/intercept term) were part of the specific model.

**Table S19.** Final grandmaternal effect LMM of male latency to mate for generation F2.

**Table S20.** Final grandpaternal effect LMM of mating duration of grandsons (generation F2).

**Table S21.** Global grandpaternal effect LMM of male latency to mate for generation F2.

**Table S22.** Global grandmaternal effect LMM of mating duration of grandsons (generation F2).

**Table S23.** Global grandmaternal effect GLM of mating success of grandsons (generation F2).

**Table S24.** Global grandpaternal effect GLM of mating success of grandsons (generation F2).

**Table S25.** Global great-grandmaternal effect LMM of latency to mate of great-grandsons (generation F3).

**Table S26.** Global great-grandpaternal effect LMM of latency to mate of great-grandsons (generation F3).
Table S27. Global great-grandmaternal effect LMM of mating duration of great-grandsons (generation F3).

Table S28. Global great-grandpaternal effect LMM of mating duration of great-grandsons (generation F3).

Table S29. Global great-grandmaternal effect GLM of mating success of great-grandsons (generation F3).

Table S30. Global great-grandpaternal effect GLM of mating success of great-grandsons (generation F3).