Intralocus sexual conflict can resolve the male-female health-survival paradox

C. Ruth Archer, Mario Recker, Eoin Duffy & David J. Hosken

At any given age, men are more likely to die than women, but women have poorer health at older ages. This is referred to as the “male-female, health-survival paradox”, which is not fully understood. Here, we provide a general solution to the paradox that relies on intralocus sexual conflict, where alleles segregating in the population have late-acting positive effects on male fitness, but negative effects on female health. Using an evolutionary modelling framework, we show that male-benefit, female-detriment alleles can spread if they are expressed after female reproduction stops. We provide support for our conflict based solution using experimental Drosophila data. Our results show that selecting for increased late-life male reproductive effort can increase male fitness but have a detrimental effect on female fitness. Furthermore, we show that late-life male fertility is negatively genetically correlated with female health. Our study suggests that intralocus sexual conflict could resolve the health-survival paradox.

1 Science and Engineering Research Support Facility, University of Exeter, Penryn Campus, Penryn, Cornwall TR10 9FE, UK. 2 College of Engineering, Mathematics and Physical Sciences, University of Exeter, Penryn Campus, Penryn, Cornwall TR10 9FE, UK. Correspondence and requests for materials should be addressed to D.J.H. (email: D.J.Hosken@exeter.ac.uk)
While broadly understand why mortality risk rises as fertility and general performance decline with age, it is less clear why the tempo and severity of these changes often differ between the sexes. In humans, survival, fertility, and performance show sex-specific patterns of decline with age. Strikingly, women stop reproducing decades before dying, while men can reproduce throughout their adult lives. Additionally, men are more likely to die than women in most age-classes, but are healthier than women late-in-life. To be clear, this is not just due to the selective loss of low quality males, as female mortality rates are lower than male rates at nearly all ages despite poorer female health. This sex difference has been termed the "male–female, health-survival paradox", and while its causes are not well understood, some resolution of it is needed if we are to ensure healthy aging as human lifespan increases.

The sex-specific mortality element of the health-survival paradox is widespread as women outlive men in most countries studied. There is also abundant evidence that older men are more likely to report health problems than women, but sex differences in mortality are present even in communities where both sexes avoid risky behaviours. Similarly, men and women with similar medical conditions are equally likely to report health problems. While sex differences in behaviour may contribute towards the male–female, health-survival paradox, the consistent observation of lower female mortality but poorer female health at older ages in human populations across the world suggests that this paradox has, at least in part, a genetic basis. To date, most biological mechanisms implicated in the paradox focus on understanding why men die earlier than women (reviewed in refs. 2,11) rather than on why late-life health differs between the sexes. Here we focus on the health aspect of the paradox and suggest that intralocus sexual conflict might explain why women are less healthy than men late-in-life.

Intralocus sexual conflict occurs when the sexes have different optimal values for a shared trait with a common genetic basis. For example, male broad-horned flour beetles develop enlarged mandibles and males with larger mandibles have higher fitness. However, daughters of males with large mandibles have lower fitness because of the masculinisation of the body that occurs with these genotypes. This means that alleles associated with mandibles are subjected to an intersexual tug-of-war over optimal values, with high fitness male genotypes making low fitness females. This type of conflict means that the alleles encoding a high-quality male often produce low quality females and vice versa. A role for sexual conflict in aging, lifespan and sex differences in health has been recognised for years. However, to the best of our knowledge, sexual conflict has not been considered as a driver of the male–female, health-survival paradox. Given the existence of the menopause, which enables selection to bias allelic values towards male-benefit late-in-life, there is enormous potential for sexual conflict to be at the heart of this paradox and by recognising its role, we may better understand what (if anything) we can do about it. The aim of this paper is simply to highlight that as well as explaining sex differences in health and aging in general, intralocus sexual conflict could be central to a long-standing puzzle in medical sciences: why do men die, while women suffer?

Intralocus sexual conflict could cause sex differences in health if there are sexually antagonistic alleles segregating in a population that increase male fitness but reduce female health. Fitness and health are different concepts. Fitness can be thought of as the number of offspring an individual contributes to the next generation, and is often measured as lifetime reproductive success, while health reflects how well individuals maintain homeostasis. Typically, healthier individuals are fitter. For example, more attractive people are likely to have high fitness (because they can attract more mates) and are also less likely to report ill health. However, health and fitness are at least partially decoupled in humans because after the menopause female fitness plummets while health does not to anywhere near the same degree (after menopause direct fitness contributions fall to zero, but health declines are not as severe; not all females experience a complete loss of homeostasis). Furthermore, sexually antagonistic alleles exist in human populations. For example, men that look particularly masculine tend to have more short-term partners, which is likely to increase their fitness, and are more resistant to respiratory conditions. On the other hand, a masculine appearance in women is associated with a greater risk of developing these conditions. Clearly, females that express alleles that are positively associated with male fitness can experience reduced health.

For intralocus sexual conflict to explain the health-survival paradox, male-benefit sexually antagonistic alleles with late-acting effects must accumulate. This is entirely feasible because women experience the menopause. This means that selection against any alleles with costly effects when expressed in females will weaken dramatically once women undergo the menopause and stop reproducing, because these alleles can only have indirect effects on female fitness. However, in men there will be selection for male-benefit alleles over the entire lifespan because men can keep reproducing until advanced ages. This would allow late-acting, male-benefit sexually antagonistic alleles to spread and accumulate in the human genome and reduce female health late-in-life, as females carrying late-acting male-benefit alleles express trait values closer to male than female optimas.

To formally test this hypothesis, we assessed whether a male-benefit, sexually antagonistic allele could spread through a diploid population using an evolutionary modelling framework. We show theoretically that under biologically realistic assumptions of costs and benefits, such antagonistic alleles can accumulate. Using Drosophila model systems, we then assessed whether sexual conflict solutions are feasible by testing whether populations evolving with selection for late-life male reproduction, but with no direct selection on females (as is the case for post-menopausal women), developed late-life costs to females. Finally, we tested whether late-life male fertility and female performance are negatively associated across a range of standardised genotypes (iso-female lines). Our data broadly support the predictions and suggest that intralocus sexual conflict could help explain the male–female, health-survival paradox.
it and it will spread throughout the population. However, a male-benefit allele could affect female fitness by reducing female health, even if it acted after the menopause. This is because older women can care for grandchildren and gain indirect fitness by improving their grandchildren’s survival. If a male-benefit allele reduces female health so severely that it impairs care, it may reduce longevity. In this case, the spread of a male-benefit allele depends on how much it reduces female health. This is clearly shown in Fig. 1a when the fitness costs of females expressing the allele are low, then the allele is likely to spread even if males only receive a modest fitness benefit. As the costs to females rise and the male benefits fall, then the allele will become less prevalent in the population and it will eventually be lost. The balance of costs and benefits that favour the spread of a sexually antagonistic allele depend on the nature of that allele, i.e. whether it is dominant (Fig. 1a) or recessive (Fig. 1d) in both sexes, or has sex-specific patterns of expression (Fig. 1b, c). However, the most important thing to note is that alleles reducing post-menopausal female health exist over a broad range of parameter space regardless of the genetic detail, and under some conditions non-antagonistic alleles are likely to be rare. This strongly suggests that late-acting male-benefit, female-cost alleles are likely to be common and could therefore be responsible for the relatively unhealthy aging of females.

Selecting on late-life male fertility affects female lifespan. Using an insect model (Drosophila simulans), we then tested whether selecting for increased late-life male reproductive success had a negative impact on females. After 12 generations of selection, we found there was a negative relationship between the fitness ranks of the sexes, such that in populations where there was the greatest increase in late-life male fertility, there was the greatest reduction in female longevity (correlation coefficient = −0.929; $t = -4.38$, df = 3, $P = 0.022$) (Fig. 2).

Association between late-life male fitness and female health. We then used another insect model (D. melanogaster) to directly assess female health associations with late-life male fitness. Across isolines (standardised genotypes), there was a negative correlation between male fertility late-in-life and a measure of physiological
health (the vertical distance females climbed in geotaxis assays) (correlation coefficient $= -0.715$; $t = -2.51, df = 6, P = 0.046$). That is, for genotypes where males had large numbers of offspring late-in-life, females performed particularly badly in negative geotaxis assays (one measure of physiological health) (Fig. 3a). However, there was no relationship between male fertility late-in-life and a second measure of female health, recovery time from anaesthesia (correlation coefficient $= 0.257$; $t = 0.596, df = 5, P = 0.577$) (Fig. 3b).

**Discussion**

Human females tend to live longer but are in poorer health than males late-in-life. This is the health-survival paradox, and given that females have higher survival at any given age despite being in poorer health, this cannot be solely due to selective disappearance of low quality males. We propose a very general resolution to this paradox—intralocus sexual conflict—and both theoretical and empirical data support this solution.

The central premise of our proposed solution, that alleles with fitness benefits to one sex can spread in a population despite costs to the other sex, is supported theoretically here and by earlier models. Our model assumes females are the longer-lived sex. This is true for humans and many primates for which we have high-quality demographic data. We also assume there are alleles that improve male fitness but reduce female health. This assumption is supported by the observation that masculine looking men are both likely to have higher reproductive success and suffer less from some health problems, but masculine looking women have a higher risk of these conditions and by the wealth of direct evidence showing that negative intersexual fitness correlations are widespread across the animal kingdom. While alleles with sexually antagonistic effects are common, their effects could be modified by alleles that alter hormone levels. So for example, sex hormones could affect the expression of shared traits in sex-specific ways, relaxing sexual conflict. However, while sex hormones can relieve sexual conflict, in bank voles there can also be pronounced sexual conflict over optimal levels of circulating sex hormones, and these can lead to negative correlations for fitness across the sexes. Thus, there is the potential for sexual conflict in humans despite a role for sex hormones in generating sexual dimorphism. We also rely on the reductions in health not being so severe that they reduce female lifespan. In other words, we assume that lifespan and health can evolve somewhat
independently in humans. This assumption is supported by the existence of the health-survival paradox. Our idea also relies on there being male-benefit alleles that act after the age of menopause and this relies on men expressing these alleles having reproductive success late-in-life. Although men with higher reproductive success tend to live shorter lives, in many societies men can reproduce long after women experience the menopause (even though most male reproduction occurs at ages when women are still reproductively active). In any case, our model shows that as long as males achieve some fitness late-in-life and as long as this does not trade-off with early-life fitness, these alleles will spread through the population.

Additionally, for the conflict hypothesis to be a feasible solution to the health-survival paradox, the male benefits and female fitness costs that promote the spread of sexually antagonistic alleles must be realistic. We assume that the benefits of expressing these alleles range between 1 (i.e. males with and without this allele produce the same numbers of offspring) and 2 (i.e. males with this allele produce twice as many progeny as those without it). In traditional societies, average male reproductive success is around six offspring but in some societies can vary between 0 and 30 children. Therefore, the advantages of expressing male-benefit alleles could be even higher than those we modelled. In females, the costs of expressing late-acting, male-benefit sexually antagonistic alleles are likely to be low. Although impaired health late-in-life could limit how well women provision their grandchildren, and so reduce indirect female fitness, these effects will be modest because grandchildren only share a quarter of their grandmother’s genome. Stated another way, direct effects tend to be more selectively important than indirect effects. On balance therefore, it seems feasible that late-acting, male-benefit alleles will have greater positive effects on male fitness than costs to females, and so they are likely to spread. However, the female fitness costs of expressing male-benefit, female-detriment alleles have never been fully quantified in humans.

Having shown theoretically that the alleles we envisage could underpin the health-survival paradox, we empirically tested these predictions in a general way using Drosophila models. We selected on male fertility late-in-life, and as a correlated response females suffered a reduction in lifespan. This is clearly not a perfect test of the model, if only because our model predicts reduced female health and not survival. Additionally, our control is the founder population from which experimental flies were derived, an approach akin to translocation experiments in ecology and not a “classical” control. Finally, we are testing a model about human health in flies—the nature of the fitness costs of expressing male-benefit alleles will inevitably differ between flies and humans for many reasons, not least because flies do not experience the menopause or receive any indirect benefits from providing parental care. However, it is important to note that the aim of this experiment was simply to see if biasing selection late-in-life towards one sex, can have costly effects on the other. Our data suggest that it can and indeed, this is precisely what an enormous body of evolutionary theory predicts.

We then showed that genotypes with high late-life male fertility produce females that perform less well in negative geotaxis assays, which is a general measure of fly health. Although geotaxis assays are not a perfect health assay, they do suggest that poorly performing females have reduced motor function. This has clear parallels to human data, where elderly women consistently experience greater physical restrictions in their daily lives than men. We did not detect a relationship between male fertility and recovery time from anaesthesia. However, given that human women do not perform worse than men in all individual measures of performance, despite consistently experiencing poorer overall health, it is not surprising that we found differences across our measures of health. A more robust test of our hypothesis would require adopting a similar approach in humans and testing whether men with greater fertility late-in-life have sisters with poor late-life health. If so, this would indicate that females carrying alleles that build a high-quality male suffer more late-in-life. To our knowledge, these data are not available.

In summary, our results indicate that intralocus sexual conflict could have a pivotal role in reducing female health late-in-life and thus provide a general solution to the human male–female, health-survival paradox. To the best of our knowledge, the male–female, health-survival paradox has never been addressed through a conflict lens. However, there are caveats to our experimental evolution data and historical human pedigree data would enable the conflict explanation to be tested more directly.

Robust testing of this idea is important, particularly as the age of reproduction in many societies is being pushed later in life. This could have impacts on sex differences in healthy aging if our hypothesis is correct, although the nature of these impacts would depend on whether sex differences in reproductive success late-in-life become relaxed or exaggerated as a consequence of age-related reproductive shifts.

**Methods**

**The model.** We used a simple population genetic model to simulate and track the prevalence of a sexually antagonistic allele, \( x \), through a diploid population over time. To allow for the allele to have a sex-specific effect and to be either recessive or dominant, we split the populations into males (\( m \)) and females (\( f \)) that are further divided according to the allelic composition of the genetic locus in question. That is, we account for individuals that are homozygous \( xx \) (\( m_{xx} \) and \( f_{xx} \)), homozygous \( yy \) (\( m_{yy} \) and \( f_{yy} \)) and heterozygous (\( m_{xy} \) and \( f_{xy} \)). We assumed mating to be frequency-dependent and for simplicity assumed the population size remains constant over time. The population dynamics can then be described by the following set of differential equations:

\[
\frac{dm_{ij}}{dt} = \mu (1 - \rho) G_{ij} - \mu_{m_{ij}} m_{ij}, i,j \in \{x,y\} \tag{1}
\]

\[
\frac{df_{ij}}{dt} = \mu + \rho G_{ij} - \mu_{f_{ij}} f_{ij} \tag{2}
\]

with

\[
\mu = \sum_{ij} \mu_{m_{ij}} m_{ij} + \mu_{f_{ij}} f_{ij} \tag{3}
\]

\[
G_{xx} = \pi_{xx}^f (\pi_{xx}^m + 0.5\pi_{xx}^m) + \pi_{xx}^f (0.5\pi_{xx}^m + 0.25\pi_{xx}^m) \tag{4}
\]

\[
G_{yy} = \pi_{yy}^f (\pi_{yy}^m + 0.5\pi_{yy}^m) + \pi_{yy}^f (0.5\pi_{yy}^m + 0.25\pi_{yy}^m) \tag{5}
\]

\[
G_{xy} = \pi_{xy}^f (\pi_{xy}^m + 0.5\pi_{xy}^m) + \pi_{xy}^f (\pi_{xy}^m + 0.5\pi_{xy}^m) + \pi_{xy}^f (0.5\pi_{xy}^m + 0.5\pi_{xy}^m + 0.5\pi_{xy}^m) \tag{6}
\]

where

\[
\pi_{ij}^s = \frac{s_{ij} \rho_m m_{ij}}{\sum_{ij} s_{ij} \rho_m m_{ij}} \tag{7}
\]

is the proportion of males or females with locus \( ij \). \( \rho \) denotes the female sex bias (if considered). We further assumed that the cost or benefit associated with carrying allele \( x \) is solely manifested through a decrease or increase in lifetime reproductive success. We cannot model costs to females as reduced health per se; if reductions in health do not reduce female fitness then trivially, any male-benefit alleles will spread. We therefore assume that alleles that severely reduce female health, reduce female fitness by reducing how well women care for their offspring and grand-offspring. Under this assumption, a shortening or lengthening of an individual’s reproductive period is equivalent to a reduction or increase in reproductive fitness. In the model this can be incorporated by changing \( \pi_{ij}^f \) as follows

\[
\pi_{ij} = \frac{b_{ij} \rho_m m_{ij}}{\sum_{ij} b_{ij} m_{ij}} \tag{8}
\]
and

\[ y_i = \sum_{j=1}^{n} c_{ij} x_j \]  

where \( b_x \geq 1 \) denotes the male-benefit and \( c_y \leq 1 \) denotes the cost to females. Considering \( x \) to take an effect only in homoygous form this results in

\[ b_{xy}y_i + b_{xy} + 0 \leq y_i \leq b_{xy}y_i + b_{xy} + 0 \]  

whereas the situation in which \( x \) is dominant can be described as

\[ c_{xy}y_i + c_{xy} + 0 \leq y_i \leq c_{xy}y_i + c_{xy} + 0 \]  

This formulation also allows us to consider dominant and recessive effects in females and males differentially. To obtain equilibrium population frequencies, we solved Eqs. (1) and (2) numerically using the odeint solver from the NumPy Python package. For each combination of male-fitness benefits \( b_x \) and female fitness costs \( c_y \), we simulated the model forward in time until an equilibrium point was reached, using the following initial conditions: \( m_x(0) = 0.05, m_y(0) = 0.3, m_{xy}(0) = 0.1, f_x(0) = 0.1, f_y(0) = 0.35, f_{xy}(0) = 0.1 \).

Selecting on late-life male fertility affects female lifespan. If intrasexual social conflict is responsible for the health-survival paradox, then selecting for late-life male reproduction should result in the accumulation of late-acting male-benefit alleles that reduce female fitness. To broadly test this idea, we selected for late-life male fertility in replicate populations of *Drosophila simulans* but relaxed selection on females. If sexual conflict is responsible for the health-survival paradox, then in populations where males experienced the greatest increase in fertility late-in-life, females should experience the greatest reduction in lifespan (relative to controls). Selecting on male fertility involved establishing five experimental populations and five female-supply populations of *Drosophila* simultaneously. Flies were collected from our large outbreeding, free-mating lab-stock population (100 males and 100 females (all virgins) per experimental and female-supply population). For 28 days, flies food in the experimental populations was changed every five days to ensure that no eggs laid emerged as adults. On day 28, flies were anaesthetised with CO₂, and removed from the cage. One hundred virgin 3- to 5-day-old females were taken from their paired female-supply populations (to ensure experimental populations were independently evolving), and added to the appropriate experimental population. This should reduce selection for age-reproduction in females. Old (28 days) males from each experimental population and young females from paired-female supply were then left for three days to lay eggs. From these, 100 male and 100 female offspring were collected on emergence and these seeded the next generation in each experimental population. This procedure was repeated for 12 generations. Female-supply populations were also fed every five days, but here new food was added on day 15, removed on day 18 and offspring collected from eggs laid between days 15 and 18. This regime is shown in Supplementary Figure 1. To test for male responses to our artificial selection, we collected 30 virgin males from each experimental population and from the lab-stock population. Each male was housed in a 40 mL vial, with 8 mL of medium for two days to mature. Then two virgin lab-stock females were added and these were replaced with young virgin females every week to ensure males had continuous access to young virgins (as is true in selection cages). Any female that died was replaced with a like-age virgin. On day 28, male fertility was assayed. Each focal male was paired with a virgin stock female. Males were removed eight hours later. Females were allowed to lay eggs in three vials over seven days and all offspring that emerged from these vials were counted as our measure of male fertility (full details in Supplementary Figure 2). Males that did not appear to mate (i.e. counts of offspring = zero), were excluded from analyses.

To test for female impacts resulting from selecting on male fertility, 40 females from each experimental population and the lab-stock population were collected as virgins and housed individually in 40 mL vials contain 8 mL of Jazz mix medium. After four days, flies were moved into a new vial. The following morning two stock virgin males (three to five days old) were added and kept with the females for three hours before being removed. This regime was repeated across the entire lifespan of each female fly (full details in Supplementary Figure 3). Pairing females with two males for three hours every five days ensures that females reproduce as normal but lifespan is not reduced due to the direct costs of mating or harassment. Females were checked daily for survival and adult lifespan was calculated.

To assess how late-life male fertility evolved in the experimental populations, mean male fertility in the stock population was subtracted from the mean value for each experimental population. High positive values meant that males from the experimental population performed much better than non-selected males, and a negative value meant that experimental males had worse late-life fertility. Female lifespan was treated in the same way (i.e. average population value—average stock-population value) and a Pearson’s correlation coefficient was calculated in R version 3.4.1.31 to analyse the correlation between male late-life fertility and female lifespan.

Association between late-life male fitness and female health. The above “selection then assay protocol” tests the general principle of the health-survival paradox, but does not directly test for female health declines as late-life male fitness increases. If intrasexual social conflict is responsible for the health-survival paradox, then genotypes that produce high fitness males should produce females with poor health and vice versa. This was tested using *Drosophila melanogaster* Genetic Reference Panel (DGRP) lines and two biomarkers of female physical function to reveal underlying health.

Ten DGRP lines were used (ID = 28, 101, 136, 360, 382, 443, 595, 737, 773, 796). Flies were maintained under a 12:12 L:D cycle at 25°C. Lines were excluded from analyses, if fewer than two animals survived to assay. The final sample sizes for each line and trait are given in Supplementary Table 1. Experimental flies were collected as virgins and aspirated into individual 100 mL vials on their day of hatching with 8 mL of medium. Dahomey tester flies were used as mating partners to assess reproductive performance. Ivanov et al.,32 recorded lifespan in DGRP lines and median lifespan ranged between 21 and 79 days. We therefore conducted tests when adults were 35 days old, approximately the median lifespan for some of the shorter lived genotypes.

Negative geotaxis (vertical climbing in response to shock) was one measure of fly health. It is a measure of motor ability that shows an age-dependent decline in *Drosophila*.33 To assess negative geotaxis, flies were aspirated into 15 mL vials attached to a drop mechanism, which was raised 10 cm and dropped. A camera recorded every trial, to record the distance that flies climbed in the two minutes after dropping. Flies were then given two minutes to recover before the process was repeated. All observations were made blind—flies were labelled with a random number and videos were analysed independently by two different observers, and any values that differed by >3 mm were observed by a third experimenter to reach agreement. Anesthesia was also used as a measure of female health, as this can indicate metabolic performance. Assay flies were transferred into a 15 mL vials where they were exposed to CO₂ (1 L/min) for thirty seconds. Flies were then put onto a piece of white paper and the time until flies stood upright was recorded and used as our measure of performance. All assays were recorded within two hours of lights going (i.e. 9–11 am) and flies assayed in a random order. Measures were made blind; flies were labelled with a random number by one lab member, before vials were passed to the observer.

Reproductive performance of the DGRP males was assessed after matings with virgin females from our wildtype stock animals. Each experimental male was housed with two virgin tester females, aged between 3 and 6 days old, that had been left overnight in 40 mL mating vials containing surplus food. Flies were then left for 48 h, after which, females were transferred to a vial for oviposition for a further 48 h, while males were removed and frozen. Females were then moved to one more vial for a further 48 h, such that their egg laying over 6 days was recorded. Oviposition vials were then incubated at the temperature from which their sire originated and offspring were counted 8 days after the first day of offspring eclosion.

Please note, differences in timings between the two male fertility assays represent species-specific lab protocols. Additionally, males were kept as virgins prior to being assayed in this experiment to allow comparison with females who were also maintained as virgins to ensure that direct physiological damage caused by male harassment did not reduce their physical performance.

To test for effects, we created a single average value, for each line and each trait for analyses (i.e. we conducted derived variable analyses). For male fertility, zero counts were once more excluded as we could not be sure that males had mated if no offspring were produced, but note this is conservative for our hypothesis. For female geotaxis distance, we included zero values but note that the results of analyses are the same irrespective of whether we include all data for both traits or exclude zeros. We then used a Pearson’s correlations, calculated in R, to analyse the associations between male late-life fertility, and female health measures.

Data availability

The data that support the findings of this study are archived in the Dryad Digital Repository (https://doi.org/10.5061/dryad.p888r2v). A Reporting Summary for this article is available as a Supplementary Information file.

Received: 1 February 2018 Accepted: 8 November 2018
Published online: 28 November 2018

References

1. Baudisch, A. & Vaupel, J. W. Getting to the root of aging. *Science* 338, 618–619 (2012).
2. Oksuzyan, A., Juel, K., Vaupe1, J. W. & Christensen, K. Men: good health and high mortality. Sex differences in health and aging. Aging Clin. Exp. Res. 20, 91–102 (2008).

3. Gordon, E. H. et al. Sex differences in frailty: a systematic review and meta-analysis. Exp. Gerontol. 89, 30–40 (2017).

4. Lenart, A. & Vaupel, J. W. Questionable evidence for a limit to human lifespan. Nature 546, E13–E14 (2017).

5. Alberts, S. C. et al. in Sociality, Hierarchy, Health (eds Weinstein, M. & Lane, M. A.) Ch. 15 (National Academies Press, Washington, D.C, 2014).

6. Jagger, C. et al. Inequalities in healthy life years in the 25 countries of the European Union in 2005: a cross-national meta-regression analysis. Lancet 372, 2124–2131 (2008).

7. Oksuzyan, A., Brønnum-Hansen, H. & Jeune, B. Gender gap in health: results from SHARE, ELSA and HRS. Eur. J. Public Health 21, 81–91 (2011).

8. C. & R. et al. The male-female health-survival paradox and sex differences in cohort life expectancy in Utah, Denmark, and Sweden 1850–1910. Ann. Epidemiol. 23, 161–166 (2013).

9. Case, A. & Paxson, C. Sex differences in morbidity and mortality. Demography 42, 189–214 (2005).

10. Austad, S. N. & Fischer, K. E. Sex differences in lifespan. Cell. Metab. 23, 1022–1033 (2016).

11. Harano, T., Okada, K., Nakayama, S., Miyatake, T. & Hosken, D. J. Intralocus sexual conflict unresolved by sex-limited trait expression. Curr. Biol. 20, 2036–2039 (2010).

12. Harano, T., Okada, K., Nakayama, S., Miyatake, T. & Hosken, D. J. Intralocus sexual conflict unresolved by sex-limited trait expression. Curr. Biol. 20, 2036–2039 (2010).

13. Maklakov, A. A. & Lummaa, V. Evolution of sex differences in lifespan and aging: causes and constraints. Bioessays 35, 717–724 (2013).

14. Bonduriansky, R., Maklakov, A., Zajitschek, F. & Brooks, R. Sexual selection, sex differences and ontogeny: the evolution of age and life span. Funct. Ecol. 22, 443–453 (2008).

15. Pekkalna, N., Kotiaho, J. S. & Puurtinen, M. Laboratory relationships between adult lifetime reproductive success and fitness surrogates in a Drosophila littoralis population. PLoS ONE 6, e24561 (2011).

16. Nedelec, J. L. & Beaver, K. M. Physical attractiveness as a phenotypic marker of healthy aging: an assessment using a nationally representative sample of American adults. Evol. Human. Behav. 35, 456–463 (2014).

17. Rhodes, G., Simons, L. W. & Peters, M. Attractiveness and sexual behavior: Does attractiveness enhance mating success? Evol. Human. Behav. 26, 186–201 (2005).

18. Thornhill, R. & Gangestad, S. W. Facial sexual dimorphism, developmental stability, and susceptibility to disease in men and women. Evol. Human. Behav. 27, 131–144 (2006).

19. Tulaupurkar, S. D., Puleston, C. O. & Gurven, M. D. Why men matter: mating patterns drive evolution of human lifespan. PLoS ONE 2, e785 (2007).

20. Shanley, D. P., Sear, R., Mace, R. & Kirkwood, T. B. Testing evolutionary theories of menopause. Proc. Roy. Soc. Lond. B 274, 2943–2949 (2007).

21. Rice, W. R. Sex chromosomes and the evolution of sexual dimorphism. Evolution 38, 735–742 (1984).

22. Bronkowsk1, A. M. et al. Aging in the natural world: comparative data reveal similar mortality patterns across primates. Science 331, 1325–1328 (2011).

23. Chippindale, A. K., Gibson, J. R. & Rice, W. R. Negative genetic correlation for adult fitness between sexes reveals ontogenetic conflict in Drosophila. Proc. Natl. Acad. Sci. USA 98, 1671–1675 (2001).

24. Peterson, M. P. et al. Testosterone affects neural gene expression differently in male and female juncos: a role for hormones in mediating sexual dimorphism and conflict. PLoS ONE 8, e61784 (2013).

25. Mills, S. C., Koskela, E. & Maples, T. Intralocus sexual conflict for fitness: sexually antagonistic alleles for testosterone. Proc. Roy. Soc. B 279, 1889–1895 (2012).

26. Westendorp, R. G. J. & Kirkwood, T. B. L. Human longevity at the cost of reproductive success. Nature 396, 743–746 (1998).

27. Betzig, L. Means, variances, and ranges in reproductive success: comparative evidence. Evol. Human. Behav. 33, 309–317 (2012).

28. Hawkes, K., O’Connell, J. F., Jones, N. B., Alvarez, H. & Charnov, E. L. Grandmothering, menopause, and the evolution of human life histories. Proc. Natl. Acad. Sci. USA 95, 1336–1339 (1998).

29. Oksuzyan, A., Brønnum-Hansen, H. & Jeune, B. Gender gap in health expectancy. Eur. J. Ageing 7, 213–218 (2010).

30. Taylor, M. L., Wigm1ore, C., Hodgson, D. J., Wednell, N. & Hosken, D. J. Multiple mating increases female fitness in Drosophila simulans. Anim. Behav. 76, 963–970 (2008).

31. R Core Development Team. A Language and Environment for Statistical Computing (R Foundation for Statistical Computing, Vienna, Austria, 2017).

32. Ivanov, D. K. et al. Longevity GWAS using the Drosophila genetic reference panel. J. Gerontol. A. Biol. Sci. Med. Sci. 70, 1470–1478 (2015).

33. Rhodesizer, D., Martin, I., Bhandari, P., Pletcher, S. D. & Grotewiel, M. Genetic and environmental factors impact age-related impairment of negative geotaxis in Drosophila by altering age-dependent climbing speed. Exp. Gerontol. 43, 739–748 (2008).

Acknowledgements

Thanks to all members of DJH’s lab group for useful discussions that informed this manuscript and help collecting data or maintaining fly stocks: Cleo Alper, Mathilda Janicott Bale, Matt Carey, Jacqui Glencross, Noah Hosken, Tomo Noda, Sam Oyensku-B1akemore, MD Sharma, Stefan Store, and Andreas Sutter. We thank the National Science Centre (Poland: 2013/09/N/ZS0/03231) and the Leverhulme Trust (UK: RF-2015-01) for funding, which partially supported this work, and the University of Exeter’s Dean’s Fellowship for additional support.

Author contributions

C.R.A. and D.J.H. conceived the study and wrote the manuscript. M.R. devised the mathematical model, C.R.A., D.J.H., and E.D. designed the experiments, C.R.A. and E.D. collected data, all authors helped edit the manuscript.

Additional information

Supplementary Information accompanies this paper at https://doi.org/10.1038/s41467-018-07541-y.

Competing interests: The authors declare no competing interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2018