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Sexual Conflict and the Tragedy of the Commons

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Abstract

It is widely understood that the costs and benefits of mating can affect the fecundity and survival of individuals. Sexual conflict may have profound consequences for populations, due to the negative effects it causes males and females to have on one another’s fitness. Here we present a model describing the evolution of sexual conflict, in which males inflict a direct cost on female fitness. We show that these costs can drive the entire population to extinction. To males, females are an essential, but finite, resource over which they have to compete. Population extinction owing to sexual conflict can therefore be seen as an evolutionary ‘tragedy of the commons’. Our model shows that a positive feedback between harassment and the operational sex ratio is responsible for the demise of females, and thus for population extinction. We further show that the evolution of female resistance to counter harassment can prevent a tragedy of the commons. Our findings not only demonstrate that sexual conflict can drive a population extinct, but also highlight how simple mechanisms, such as harassment costs to males and females and the coevolution between harassment and resistance, can help avert a tragedy of the commons caused by sexual conflict.

Keywords: adaptive dynamics, population dynamics, coevolution, social evolution, evolutionary suicide
Introduction

Sexual reproduction is widely considered an evolutionary mystery, because it involves a twofold cost. This cost is a result of anisogamy and is due to the production of males halving the population growth rate (West et al. 1999; Agrawal 2006). Under anisogamy, one of the sexes evolves to produce small gametes that contribute virtually no resources to the zygote. The shift from investing in few gametes to producing numerous gametes (which is a specialty of the male sex) represents escalating competition between males. The cost of sex only arises when males shift resource use away from offspring provisioning, and it is thus often remarked that the twofold cost of sex is more appropriately referred to as the cost of male production (Maynard Smith 1978; Jennions and Kokko 2010).

This twofold cost is key to understanding the evolution of sexual reproduction, but it also has other consequences. For males, females form the limiting resource. Numerous male traits, such as sperm competition, have evolved as a consequence of competition among males, which can lead to male gametes vastly outnumbering female gametes, or to investment in weaponry (Emlen 2008). Such scenarios can lead to a “tragedy of the commons”, as behaviors are favored that are advantageous to individuals, but detrimental to the group or population as a whole (Hardin 1968; Rankin et al. 2007a). Sexual conflict over mating is a tragedy of the commons whenever male-male competition reduces the availability of females (or their gametes) for all males, e.g. via elevated female mortality (Le Galliard et al. 2005; Rankin and Kokko 2006). Given the increasing emphasis in the evolutionary literature on understanding the demographic consequences of group-detrimental behaviors using the framework of the tragedy of the commons (e.g. Rankin et al. 2007a; Frank
(2010), it is surprising that sexual conflict has not featured more strongly in such work: so far, we have virtually no theoretical background for understanding the potentially negative population-level consequences of male behaviors towards females.

If male behavior harms female reproduction, the cost of sex can be larger than twofold. Studies of sexual conflict have documented numerous examples of reproductive costs (Chapman et al. 2003; Arnqvist and Rowe 2005). Extreme cases include toxic seminal fluids that appear to harm females as a form of ‘collateral damage’ (Chapman et al. 1995; Wigby and Chapman 2005), as well as male mating behaviors that physically damage females (Le Galliard et al. 2005). Milder cases, which nevertheless can increase the cost of sex, include size dimorphisms that force mothers to invest more energy into male offspring (Rankin and Kokko 2007). Also sexual harassment of females by males has been shown to impose costs on females, in terms of reduced fecundity or survival, in a number of systems (e.g. Réale et al. 1996; Crudgington and Siva-Jothy 2000; Shine et al. 2000; Shine et al. 2001; Maklakov et al. 2006; Rönn et al. 2006; Rönn et al. 2007; Godsen and Svensson 2009). A recent study of water striders demonstrated that, while aggressive males had an advantage in local competition for mates, this had a negative impact on their fitness when considered across a wider metapopulation, due to the harm they imposed on females (Eldakar et al. 2009).

The population-level consequences of sexual conflict, albeit rarely documented, can be severe (Kokko and Brooks 2003; Le Galliard et al. 2005; Valero et al. 2008), but typically depend on both the population density and sex ratio. For example, in the common lizard *Lacerta vivipara*, males harm females by inflicting mating scars and
damaging the skin on the backs of females (Le Galliard et al. 2005). Male-biased populations of these lizards have been shown to exhibit higher female mortality than female-biased populations, owing to a higher level of harassment of females by males (Le Galliard et al. 2005). The harm induced by male lizards was predicted to drive populations extinct within 40 years, if they started out with a male bias (Le Galliard et al. 2005). This highlights that the ‘tragedy’ does not have to stop at a 50% reduction in population growth rate, which would be predicted if the cost of sex were always precisely twofold. Instead, individually selected adaptations can lead to detrimental effects up to the point at which a whole population or species is driven to extinction (Matsuda and Abrams 1994a; Matsuda and Abrams 1994b; Boots and Sasaki 2003; Parvinen 2005; Rankin and López-Sepulcre 2005; Rankin et al. 2010).

Given that sexual conflict among animals is widespread (Arnqvist and Rowe 2005), the population-level cost of sex could often be greater than twofold. If sexual conflict elevates female mortality and becomes more severe when populations are male-biased, what prevents sexual conflict from regularly driving populations extinct? We investigate two potential mechanisms that may avert a tragedy of the commons: (1) costs to males of harassing females (as such direct costs are known to prevent a tragedy of the commons in other situations – Rankin et al. 2007a) and (2) joint evolution between male harassment and female resistance (Arnqvist and Rowe 2002; Rönn et al. 2007). We do this by building a model based on adaptive dynamics theory that separately treats the evolution of phenotypic traits in males and females and does not make any specific assumptions about the underlying genetic system, such as diploidy or additive genetics. To evaluate the robustness of our results, and to see how they might be influenced by explicit genetic assumptions, we further analyze a
corresponding individual-based model. We show that costs of harassment and the underlying population ecology are key determinants of whether populations can be driven extinct by sexual conflict.

**Model and Results**

*Evolving Traits*

We examine the evolution of male harassment of females and of female resistance to male-induced harm, both of which are sex-specific. Males express a harassment level $y$ that improves male mating success, but reduces survival for males as well as females. We denote the corresponding costs by $g$ and $h$ for males and females, respectively. Such costs may be of very unequal magnitude for the harasser and the target of harassment. Females express a level of resistance $x$ ($0 \leq x \leq 1$) that improves their survival, but simultaneously reduces their fecundity by a factor $p(x)$ ($0 \leq p(x) \leq 1$). A description of the variables and parameters used in the model is given in Table 1.

*Population Dynamics*

Since our model links trait evolution with its demographic consequences, we must specify population dynamics explicitly. To do this, we assume that the population dynamics of males and females unfold in continuous time. Male density is denoted by $M$ and female density by $F$. We assume that reproduction takes negligible time, so all males and females can mate at any time, and the operational sex ratio (OSR) is the same as the adult sex ratio. Females have an intrinsic birth rate $b$ and produce males with probability $r$ and females with probability $1-r$ (if $r=1/2$, the primary sex ratio is even).
To capture the assumed demographic consequences of the traits $x$ and $y$ as described above, we consider the following population dynamics of male and female densities:

\[
\frac{dF}{dt} = (1-r)p(x)bF - h(x,y,M,F)F - l(M,F)F ,
\]

\[
\frac{dM}{dt} = rp(x)bF - g(y)M - l(M,F)M .
\]

(1a)

(1b)

In these equations, density-dependent mortality is represented by the function $l(M,F)$. For simplicity, we assume logistic density dependence, such that $l(M,F)$ increases linearly with population density, $l(M,F)=\mu(M+F)$, where the parameter $\mu$ scales the density-dependent mortality. In keeping with adaptive dynamics theory, we assume a homogeneous population such that all males and females (except for the occasional rare mutant) have the same trait values $x$ and $y$, respectively (we later relax this assumption in an individual-based model).

We assume that a male’s investment $y$ in harassment imposes a cost on the male by elevating his mortality in proportion to $y$. We describe this by the function $g(y)=cy$, where $c$ scales these costs to males. Similarly, male harassment increases female mortality, which is described by the function $h(x,y,M,F)$. We derive our examples below using the function $h(x,y,M,F)=(1-dx)yM/F$, which implies that the harassment costs experienced by females decrease with female resistance and increase as males become more common relative to females. The first effect is described by $1-dx$, where $d$ scales the efficiency of female resistance. As $h$ cannot be negative, we assume that $0\leq d\leq 1$ (implying that $0\leq 1-dx\leq 1$). The second effect is described by $kyM/F$, where $k$ scales how strongly harassment increases female mortality. As female resistance $x$ may impose a cost on females, we assume that female fecundity is
reduced by the factor $p(x)=1-ax$, where $a$ scales a female’s cost of investing in resistance ($0\leq a \leq 1$). This fecundity cost can be large, as in the case of bedbugs developing a harder outer shell (Reinhardt et al. 2003), or small, as in the case of females kicking to avoid copulation in the seed beetle *Callosobruchus maculatus* (Edvardsson and Tregenza 2005).

**Extinction Threshold**

There are three equilibria of the dynamics described by equations (1): the first is the extinction equilibrium at $\hat{M}=0$ and $\hat{F}=0$, whereas male and female densities are positive at the other two equilibria. Calculating the Jacobian matrix of equation (1) and determining its eigenvalues, we see that only one of the two non-extinction equilibria is stable, and we therefore focus on this equilibrium in the following analyses. Examining the eigenvalues of the Jacobian matrix at $\hat{M}=0$ and $\hat{F}=0$ yields a threshold condition for the levels of male harassment $y$ that cause extinction,

$$y \geq y_{\text{ext}} = bp(x)\frac{2\sqrt{rk(1-dx)(rk(1-dx) - c(1-r)) + 2rk(1-dx) - c(1-r)}}{c^2}. \quad (2)$$

For $r > c/(c+k(1-dx))$, the extinction threshold $y_{\text{ext}}$ is real and positive, and for $x=0$, it simplifies to $y_{\text{ext}} = b\left(2\sqrt{rk-[rk-c(1-r)]} + 2rk - c(1-r)\right)/c^2$.

We note in passing that, had we assumed density-dependent harassment, $h(x,y,M,F)=(1-dx)kyM$, instead of frequency-dependent harassment, $h(x,y,M,F)=(1-dx)kyM/F$, deterministic extinction would not be possible. This finding is consistent
with analogous results for parasite-mediated extinction, where it has been shown that deterministic extinction can only occur under frequency-dependent transmission and not under density-dependent transmission (e.g. Boots and Sasaki 2003; Rankin et al. 2010).

Evolution of Male Harassment

We now investigate whether evolution drives the harassment level $y$ above the extinction threshold $y_{\text{ext}}$. For our first model, we use adaptive dynamics theory (Metz et al. 1992; Dieckmann and Law 1996; Metz et al. 1996; Dieckmann 1997; Geritz et al. 1998; Meszéna et al. 2001; Dieckmann and Metz 2006) to account for the eco-evolutionary feedback between the population dynamics and the evolution of our two evolving traits. As such, our analysis assumes that ecological and evolutionary timescales are sufficiently separated (so that the densities of males and females rapidly reach equilibrium). We do not distinguish between genotypes and phenotypes, and the terms “mutants” and “residents” refer, respectively, to individuals with rare or prevalent trait values. Since our results in equations (3) to (10) are based on selection gradients, equivalent results could be derived if we had instead followed the quantitative genetics method of Lande (Lande 1976; Iwasa et al. 1991), with uncorrelated traits and small additive genetic variances.

We assume that harassment at level $y$ provides a male with an absolute mating benefit $f(y)$. However, since mating opportunities are limited by the availability of females, a male’s mating success must be evaluated relative to that of all other males: this implies that selection on $y$ is frequency-dependent, as a male’s mating success depends not only on his own $f(y)$, but also on the $f(y)$ of other males. In a first step of
analysis, we assume that females possess no resistance (i.e., \( x=0 \), and thus \( p(x)=1 \)).

The invasion fitness (Metz et al. 1992) of a mutant male with trait value \( y' \) in a population dominated by the resident trait values \( x \) and \( y \) is given by the mutant’s per capita growth rate when rare:

\[
\lambda_m(y', x, y) = rb \frac{\hat{F}}{M} \frac{f(y')}{f(y)} - g(y') - l(M, \hat{F}) .
\]  

(3)

The factor \( \frac{f(y')}{f(y)} \) represents the relative benefits of mating to a male investing \( y' \) in harassment. In a homogeneous population (where \( y'=y \)), this term equals 1. We derive our examples below using the function \( f(y)=y^u \), so the shape of the dependence of the mating benefit \( f(y) \) on the harassment level \( y \) is determined by \( u \). If \( u>1 \), the benefits of harassment are accelerating, while if \( u<1 \) these benefits are decelerating.

The selection gradient for the male harassment level \( y \) is

\[
s_m(x, y) = \frac{\partial \lambda_m(y', x, y)}{\partial y'} \bigg|_{y'=y} = \frac{bru \hat{F}}{y M} - c .
\]  

(4)

The selection pressure on \( y \) vanishes at the evolutionarily singular strategy \( y^* \), which is thus obtained by solving the equation \( s_m(y^*)=0 \),

\[
y^* = bu \frac{c(1-r) - kru}{c^2 (1-u)} .
\]  

(5)
If the singular strategy \( y^* \) satisfies inequality (2), the evolution of harassment causes population extinction through sexual conflict when \( y \geq y_{\text{ext}} \). Such selection-driven extinction is known as evolutionary suicide (Ferrière 2000; Dieckmann and Ferrière 2004; Parvinen 2005; Rankin and López-Sepulcre 2005). Fig. 1 shows an example of the evolutionary dynamics of male harassment, including its effect on male and female densities. The evolutionary dynamics of \( y \) are determined by the equation \( \frac{dy}{dt} = \delta M(y) \), where \( \delta \) is a small number that scales the speed of evolution (Lande 1976; Hofbauer and Sigmund 1990; Iwasa et al. 1991). Had we used the canonical equation of adaptive dynamics, Dieckmann and Law 1996, the right-hand side would also contain \( \hat{F} \).

Fig. 1A and 1B show a case in which the population persists despite the evolutionary aggravation of sexual conflict. Here the number of females exceeds the number of males, but the population is not driven extinct, since the per capita harassment experienced by females remains mild despite the high level of harassment imposed by each male. In contrast, fig. 1C and 1D show a case in which the population is driven extinct. Here the level of male harassment remains relatively low, but since the population contains many more males than females, the positive feedback between the declining fraction of females and the increasing harassment they experience causes extinction.

By replacing the \( y \) in inequality (2) for \( x=0 \) with \( y=y^* \) according to equation (5), we see that harassment evolution causes extinction, if

\[
 u > 1 \quad \text{and} \quad 1 \geq r \geq \frac{c}{c+k}.
\]
Interestingly, this condition is independent of both the intrinsic birth rate $b$ and of the scale $\mu$ of the density-dependent death rate. Inequality (6) shows that accelerating returns to males from investing in harassment ($u>1$) are needed for evolutionary suicide to occur. It also shows that the potential for evolutionary suicide critically depends on the costs of male harassment, both to males from expressing the trait ($c$) and to females from male-induced harm ($k$). If the ratio of male costs to female costs, $c/(c+k)$, is less than the primary sex ratio $r$, then evolutionary suicide occurs, whereas if this ratio exceeds $r$, the population can persist. For an even primary sex ratio, $r=1/2$, the condition $r>c/(c+k)$ simplifies to $k>c$, so the harm inflicted on females must exceed the cost males pay for harassment.

We can understand these results by considering the effect of harassment on the relative densities of males and females. For $u>1$, the evolution of harassment may continue to accelerate, increasingly affecting the population densities of males and females. If $r>c/(c+k)$, females suffer from more harassment-related mortality than males, which leads to greater harassment per female, as there are more males than females. This further increases female mortality, until the population is driven extinct. In contrast, if $r<c/(c+k)$, males suffer from more harassment-related mortality than females, which decreases male density, and thus further reduces the per capita level of harassment females experience.

These findings are confirmed by fig. 2, which shows how the evolutionarily singular level of harassment (fig. 2A) and the male and female density (fig. 2B) vary with the costs of harassment to males. We see that the population is driven to extinction for $c<k$. Low costs to males result in high levels of harassment, and thus extinction. High
costs to males enable population persistence, but decrease the density of males through their effect on male fitness. It is also interesting to observe that the greater the cost of harassment to males, the greater the density of females. This effect arises through a feedback between male density and the costs to males: if harassment increases male mortality, the lower resulting density of males creates conditions that are more favorable to females. Under an even primary sex ratio, the more females reproduce, the more even the OSR will be, and this, in turn, further reduces the harm inflicted on females.

**Evolution of Female Resistance**

In a second step of analysis, we can now look at the evolution of female resistance $x$ ($0 \leq x \leq 1$) to male harassment. Following the assumptions described previously, the invasion fitness of a mutant female investing $x'$ in a resident population investing $x$ is

$$
\lambda_r(x', x, y) = (1-r)p(x')b - h(x', y, \hat{M}, \hat{F}) - l(\hat{M}, \hat{F}) .
$$

(7)

From equation (7), the selection gradient for $x$ is thus given by

$$
\frac{\partial \lambda_r(x', x, y)}{\partial x'} \bigg|_{x' = x} = dyx' \frac{\hat{M}}{\hat{F}} - ab(1-r) .
$$

(8)

We can study the coevolution of male harassment and female resistance by examining the male and female selection gradients in equations (4) and (8). An example of the joint evolutionary dynamics of male harassment and female resistance,
\[
\frac{d(x, y)}{dt} = \delta(s_F(x, y), s_M(x, y)),
\]
is given in fig. 3. Solving this for \( \frac{d(x^*, y^*)}{dt} = 0 \), gives us the evolutionarily singular levels of female resistance \( x^* \) and male harassment \( y^* \),

\[
x^* = \frac{1}{a} - \frac{c(1-r)}{dkru}, \quad (9a)
\]

\[
y^* = \frac{bu(d-a)(1-r)}{cd(1-u)}. \quad (9b)
\]

We can now use the threshold in inequality (2) to determine where evolution toward \( x^* \) and \( y^* \) imply population extinction. If we assume that the benefits of resistance are greater than the costs (i.e., \( a > d \)), sexual conflict drives the population extinct if

\[
u > 1 \text{ and } 1 \geq r > \frac{ac(1-u)}{ac + du - au(c + k)}. \quad (10)
\]

In other words, greater values of \( a \) or \( k \), or smaller values of \( c \), will result in evolutionary suicide. As for the case of male harassment evolving in the absence of female resistance, more male-biased primary sex ratios (greater values of \( r \)) or more accelerating harassment benefits (greater values of \( u \)) will increase the extinction risk resulting from sexual conflict.

Our results make intuitive sense as, if \( u > 1 \), runaway selection for greater levels of sexual conflict may occur. If harassment is costly to males, this can prevent the OSR from becoming more biased: these costs prevent harassment from evolving toward the extinction threshold, not only because they directly hamper the evolution of strong (and thus costly) harassment, but also because the costs to males manifest themselves
as increases in male mortality. The reduction in male density resulting from the
greater costs to males obviously limits the dangerous increase in the OSR. The
opposite is true if the male-induced harm to females increases with $k$, as this
magnifies the OSR bias.

Fig. 4 shows how the evolution of female resistance depends on the cost to males of
investing in harassment. We see that female resistance is favored at intermediate
levels of the cost to males. If this cost is small, males evolve to be very harmful to
females, so it does not pay to females to invest in resistance (Lee and Hays 2004). At
the other extreme, high costs to males disfavor male harassment, and thus reduce the
need for resistance. It is in the intermediate range that female resistance is
evolutionarily favored. Fig. 4 shows that the evolution of female resistance can rescue
the population from extinction in a significant part of the range $0 \leq c \leq k$ we had
previously shown to lead to population extinction in the absence of female resistance.

*Individual-based Model*

To check the robustness of results and relax the simplifying assumptions made in the
analyses above, we now examine a genetically explicit individual-based model with
diploid inheritance of harassment and resistance alleles. We consider one locus for
harassment and one locus for resistance, and assume free recombination between
these loci. We also assume additive genetics and the absence of environmental
variation, so an offspring’s phenotype is the mid-parental value of the alleles inherited
from the mother and the father. For example, the offspring’s resistance is
$x = (x_F + x_M)/2$, where $x_F$ denotes the allelic value inherited from the mother and $x_M$ the
allelic value inherited from the father. Sexual reproduction involves segregation and
recombination, and one of the parent’s two homologous alleles is chosen at random to be inherited. Each individual thus possesses two harassment alleles and two resistance alleles. The harassment alleles are only expressed in males, and the resistance alleles only in females. All other assumptions are as described above; in particular, we use the same functions to link phenotypes to per capita birth and death rates. Since populations in the individual-based model are finite, population dynamics are stochastic and evolution is subject to genetic drift. Full details of the individual-based model are given in Appendix 1.

For \( c < k \), and when harassment is not resisted by females, the population is driven extinct due to harassment evolution (fig. 5A and 5B). Our assumption of an even primary sex ratio results in a vicious circle in which males become more abundant as harassment shortens female lifespan, diminishing female density, and ultimately driving the population extinct. Allowing female resistance and male harassment to coevolve (fig. 5C and 5D) can prevent extinction. The two examples in fig. 5 reflect a general pattern. When examining 25 model runs when costs to males were low (\( c=0.8 \)) and female resistance was not favored (\( a=1 \) and \( d=0 \)), all 25 resulted in evolutionary suicide. When the costs to males were higher (\( c=1.2 \)), the population persisted in 23 out of the 25 considered replicate model runs, with an average final population size of \( 971.52 \pm 82.7 \) (mean ± S.E.) individuals. Allowing females to evolve resistance (for \( c=0.8 \), \( a=0.5 \), and \( d=1 \)), persistence occurred in all 25 considered replicate model runs, with an average final population size of \( 1770.6 \pm 17.5 \) (mean ± S.E.). All of these results are in good qualitative agreement with our analytical model.
Discussion

Here we have shown that the evolution of male harassment can result in the extinction of an entire population. In our model, the evolution of male harassment can lead to a vicious circle in which fewer females survive while the mortality of the remaining females increases due to a high male-to-female ratio, until the population collapses. Our findings therefore confirm the results of Le Galliard et al. (2005). In particular, we have shown that an evolutionary ‘tragedy of the commons’ can occur both in a simplified analytical model and in a genetically explicit individual-based model. While earlier studies based on simpler models have speculated that sexual conflict may cause population extinction (for a review, see Kokko and Brooks 2003), our model is, to our knowledge, the first to confirm this effect whilst being explicit about the underlying population dynamics.

Our analytical model highlights the importance of female-biased primary sex ratios as a mechanism that can limit the harm inflicted on females. However, a more important factor influencing extinction in our model is the ratio of male costs of harassment to male-induced harm to females. Under an even primary sex ratio, if investing in harassment is cheap for males and such harassment inflicts a large mortality cost on females, selection favors ever-greater levels of harassment. Elevated female mortality creates a male-biased operational sex ratio (OSR), which intensifies harassment and feeds back to increase female mortality even further, until the population is driven extinct. This is a fully expressed tragedy of the commons, as selection at no point turns to favor less harassing males as the shared resource (here the density of remaining females) diminishes.
The simplest way in which a tragedy of the commons can be averted is to make it costly for the exploiter (males) to exploit the resource (females), so harassing should be costly for males. Harassment is costly for males when competition among males and/or the pursuit of females for copulation implies higher mortality, increased predation risk, elevated energy requirements, or reduced time available for performing other functions. Thus, if a strong trade-off exists between the benefits males get from harassing, in terms of extra matings, and the costs of obtaining those matings, it will be advantageous for males to reduce harassment, which is likely to prevent extinction. The extent to which harassment is costly will depend very much on the mating system, but our model predicts that a tragedy of the commons through sexual conflict is likely if harassment is cheap for males, but costly to females.

An interesting result of our analysis is that if male costs of harassment exceed those inflicted on females, more male-induced harm actually increases population density, despite causing the sexual conflict to intensify. This occurs because conflict in this case elevates male mortality more than female mortality, and removing males can increase population growth via density dependence when males and females utilize the same resources (e.g. Wedekind 2002; Kokko and Brooks 2003; Rankin and Kokko 2007). In addition, the shape of the relationship between the benefits and costs of harassment to males could play a large role in promoting or inhibiting extinction. For example, decelerating costs to males might select for runaway evolution to greater levels of male harassment, thus favoring extinction, while accelerating costs are expected to have the opposite effect.
Coevolutionary dynamics occur under a wide range of circumstances, such as between predators and their prey and parasites and their hosts (Dieckmann et al. 1995; van der Laan and Hogeweg 1995; Dieckmann and Law 1996; Gilchrist and Sasaki 2002), as well as between male harassment and female resistance (Arnqvist and Rowe 2002; Wigby and Chapman 2004; Rönn et al. 2007; Godsen and Svensson 2009). We have established conditions under which the coevolution of female resistance with male harassment can prevent extinction. Adaptations in females that counter harm caused by males results in less damage to females, thus increasing overall female fitness and a greater population density (fig. 4). Coevolutionary dynamics are influenced by the speed at which different traits can evolutionarily respond to changes in other traits (e.g., host-parasite dynamics are strongly influenced by the typically faster life cycles of the parasite). If the harming sex evolves faster than the harmed sex, extinction could occur under a wider range of circumstances than observed in our model. Conversely, if females can resist males by plastically reducing their mating rate (Lessells 2005), the harm to females is reduced, potentially preventing extinction.

Our models obviously do not consider all mechanisms that may conceivably prevent extinction through sexual conflict. For instance, group structure can disfavor male harassment (Eldakar et al. 2009; Rankin 2011). Although we considered the feedback between sex-specific mortality and the per capita harassment experienced by females, our model has not explored all potential forms of feedback between demography and conflict. There are many other feedbacks that make selection a function of population characteristics such as density (Dieckmann and Metz 2006; Kokko and López-Sepulcre 2007), and such feedbacks can potentially resolve conflicts (Kokko and Rankin 2006; Rankin 2007; Knell 2009). However, since sexual selection on males
can either intensify or relax with increasing density (Kokko and Rankin 2006; Klug et al. 2010), we based our model on the assumption of no change (the null case in the absence of any specific information) and instead focused on density-dependent shifts in the OSR. In nature, predicting even just the direction of such shifts is far from obvious. If conflict intensified at lower population densities (Kokko and Rankin 2006), this would increase the risk of evolutionary suicide. Any mechanism that favors male over female survival in low-density conditions (e.g., if males dominate females behaviorally during and after a population crash) could have this effect. On the other hand, there are conditions in which the mechanism we envisage to cause vicious circles will not apply as efficiently as we have assumed. OSRs can differ from primary sex ratios whether or not breeding is continuous, but any effects of male harassment are probably weaker with discrete generations than with overlapping generations (as modeled here), because in the former case each generation starts again with a roughly even primary sex ratio.

Conclusions

Male harassment can increase female mortality (Arnqvist and Rowe 2005) and female mortality often is a strong determinant of equilibrium population densities (Rankin and Kokko 2007; Candolin and Heuschele 2008). Our model shows that, through the vulnerability implied by this relationship, male harassment can cause a tragedy of the commons, and even result in a population being driven extinct. We have shown how this tragedy can be averted through the costs to males of investing in harassment, and the coevolution of male harassment and female resistance. In addition, any individually selected trait that can result in local population extinction is subject to contravening selection at higher evolutionary levels (Rankin et al. 2007b). If sexual
conflict frequently impairs a population’s reproduction and viability, such higher-level selection may be an important factor limiting the intensities of male harassment we observe in nature.

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**Appendix 1: Individual-based model**

To test the robustness of our analytical model, we built an individual-based model to examine how our results are affected by introducing demographic stochasticity and an explicit genetic system, namely diploid, additive genetics. In our individual-based model, the population can have any distribution of trait values, and there is no assumption that all but one trait value should be rare. The model assumes diploid inheritance of harassment and resistance alleles at two freely recombining loci. An individual thus possesses two harassment alleles, one inherited from the father, the other from the mother. We further assume additive genetics (i.e., no dominance or epistasis), so the harassment trait is given by the arithmetic mean of the two
harassment alleles. The harassment trait is only expressed in males; analogous rules apply to the resistance trait only expressed in females.

The individual-based model is initialized with a population of 400 individuals, randomly designated as either male or female. Each allele at each locus is drawn at random. To ensure sufficient variation in the traits, male harassment alleles (which can take any positive value) are drawn from a random exponential distribution with a mean of 1, while female resistance alleles (which can take values between 0 and 1) are drawn from a random uniform distribution between 0 and 1. Males express the harassment trait \( y = (y_M + y_F)/2 \), where \( y_M \) denotes the harassment allele inherited from the male and \( y_F \) the harassment allele inherited from the female. Females express the resistance trait \( x = (x_M + x_F)/2 \), where \( x_M \) denotes the resistance allele inherited from the male and \( x_F \) the resistance allele inherited from the female. For each trait, one of each parent’s alleles is chosen at random to be inherited to the diploid offspring. We use the same functions to describe per capita birth and death rates as in the analytic model. The birth rate of a given female is thus given by \( b(1 - ax) \), and the death rate of a given female by \( (1 - dx)(k \overline{y} M/F) + \mu (M+F) \). We assume that all individuals mate sufficiently frequently, so females experience the same level of male harassment \( \overline{y} \), given by the population average of all harassment traits \( y \) expressed by males. The death rate of a given male is given by \( cy + \mu (M+F) \). We assume that the relative mating success of males is proportional to their mating ability \( y^\mu \).

The model keeps track of birth and death events, which occur in continuous time. Each individual female has a unique birth rate and death rate, while males express
death rates only. The next event that occurs in the population is chosen with a probability proportional to its rate. If the event is a death, the chosen individual is removed from the population. If it is a birth, a father is chosen with a probability proportional to his mating ability $y^u$ and is mated with the chosen mother. The resultant offspring inherits one randomly chosen resistance allele and harassment allele from each parent. With probability $r$ the offspring is male, while with probability $1-r$ it is female. The model is iterated for 500,000 events, or until the population is driven extinct. The reported examples use parameter values $r=1/2$, $b=20$, $\mu=0.001$, $k=1$, and $u=1.01$, and we change the cost $c$ of male harassment, and the cost $a$ and benefit $d$ of female resistance.
References


Figure and Table Legends

Table 1. Model variables and parameters.

Figure 1: Evolution of male harassment (A and C) and implications for the population densities of males and females (B and D) in the analytical model. Panels A and B show a case without selection-driven extinction (c=1.2), whereas panels C and D show a case with evolutionary suicide (c=0.8). Other parameters: r=1/2, b=50, µ=1, k=1, u=1.01, a=1, d=0, x=0, and δ=0.01.

Figure 2: Influence of harassment cost to males on the evolutionary equilibrium of male harassment (A) and the resultant equilibrium densities of males and females (B) in the analytical model. The dashed line shows the equilibrium density of males and females in the absence of male harassment (y=0) and female resistance (x=0). Parameters: r=1/2, b=50, µ=1, k=1, u=1.01, a=1, d=0, and x=0.

Figure 3: Coevolution of male harassment and female resistance (A) and implications for the population densities of males and females (B) in the analytical model. Parameters: r=1/2, b=50, µ=1, u=1.01, k=1, a=1/2, d=1, c=0.8, and δ=0.01.

Figure 4: Influence of harassment cost to males on the evolutionary equilibrium of male harassment and female resistance (A) and the resultant equilibrium densities of males and females (B) in the analytical model. The dashed line shows the equilibrium density of males and females in the absence of male harassment (y=0) and female resistance (x=0). Parameters: r=1/2, b=50, µ=1, k=1, u=1.01, a=1/2, and d=1.
**Figure 5:** Evolution of male harassment (A and C) and implications for the population sizes of males and females (B and D) in two model runs of the individual-based model. In panels A and B female resistance could not evolve ($x=0$), whereas in panels C and D it could coevolve together with male harassment ($a=0.5$ and $d=1$).

Other parameters: $r=1/2$, $b=20$, $\mu=0.001$, $k=1$, $u=1.01$, and $c=0.8$. 
Figure 1
Figure 3

(A) Graph showing the relationship between Time and Male Harassment and Female Resistance. Male Harassment increases significantly over time, while Female Resistance shows a more gradual increase.

(B) Graph showing the relationship between Time and Population Density. Both Male and Female Density decrease over time, with Male Density decreasing more rapidly than Female Density.
**Figure 4**

(A) Graph showing the relationship between the cost of male harassment and male harassment, as well as the cost of male harassment and female resistance.

(B) Graph showing the relationship between the cost of male harassment and population density, with separate curves for female density and male density.
Figure 5

A

Male harassment

Events

B

Number of males

Number of females

Events

C

Male harassment

Female resistance

Events

D

Number of females

Number of males

Events
<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$M$</td>
<td>Density of males</td>
</tr>
<tr>
<td>$F$</td>
<td>Density of females</td>
</tr>
<tr>
<td>$y'$ and $y$</td>
<td>Level of male harassment for a mutant ($y'$) and a resident ($y$)</td>
</tr>
<tr>
<td>$x'$ and $x$</td>
<td>Level of female resistance for a mutant ($x'$) and a resident ($x$)</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
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<tbody>
<tr>
<td>$r$</td>
<td>Primary sex ratio (proportion of males)</td>
</tr>
<tr>
<td>$b$</td>
<td>Intrinsic birth rate</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Scale of density-dependent death rate</td>
</tr>
<tr>
<td>$c$</td>
<td>Cost to males of investing into harassment</td>
</tr>
<tr>
<td>$k$</td>
<td>Cost to females of experiencing harassment</td>
</tr>
<tr>
<td>$u$</td>
<td>Shape of returns to males from investing into harassment</td>
</tr>
<tr>
<td>$a$</td>
<td>Cost to females of resisting harassment</td>
</tr>
<tr>
<td>$d$</td>
<td>Efficiency of female resistance in reducing the costs of male harassment</td>
</tr>
</tbody>
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