Evolution of sex ratio through gene loss

Da Yin* and Eric S. Haaga†

*Department of Biology and Biological Sciences Graduate Program, University of Maryland, College Park, MD 20742

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The maintenance of males at intermediate frequencies is an important evolutionary problem. Several species of Caenorhabditis nematodes have evolved a mating system in which selfing hermaphrodites and males coexist. While selfing produces XX hermaphrodites, cross-fertilization produces 50% XO male progeny. Thus, male mating success dictates the sex ratio. Here, we focus on the contribution of the male secreted short (mss) gene family to male mating success, sex ratio, and population growth. The mss family is essential for sperm competitiveness in gonochoristic species, but has been lost in parallel in androdioecious species. Using a transgene to restore mss function to the androdioecious Caenorhabditis briggsae, we examined how mating system and population subdivision influence the fitness of the mss+ genotype. Consistent with theoretical expectations, when mss+ and mss-null (i.e., wild type) genotypes compete, mss+ is positively selected in both mixed-mating and strictly outcrossing situations, though more strongly in the latter. Thus, while sexual mode alone affects the fitness of mss+, it is insufficient to explain its parallel loss. However, in genetically homogenous androdioecious populations, mss+ both increases male frequency and depresses population growth. We propose that the lack of inbreeding depression and the strong subdivision that characterize natural Caenorhabditis populations impose selection on sex ratio that makes loss of mss adaptive after self-fertility evolves.

sex ratio | androecy | nematodes | sperm

Evolutionary transitions in sexual systems are common in many animal and plant taxa (1). One of the most extreme is the adoption of self-fertility. Historically hermaphroditic organisms, such as flowering plants, evolve selfing through loss of mechanisms that prevent union of gametes of the same individual (2–4). However, ancestrally male/female ( dioecious or gonochoristic) organisms can also evolve selfing when the female sex evolves male function to become a neohermaphrodite (5–7). This process creates an androdioecious mating system in which hermaphrodites and males coexist (8–10). Androdioecious populations can have highly skewed sex ratios, as males are optional for reproduction.

In Caenorhabditis nematodes, Caenorhabditis elegans, Caenorhabditis briggsae, and Caenorhabditis tropicalis have evolved self-fertility independently from male/female ancestors (11–13). Selfing increases homozygosis and unMASKS deleterious mutations (14–16). With the right amount of selfing, however, deleterious mutations can be purged from the genome (17, 18). Indeed, selfing Caenorhabditis have healthy, naturally homozygous genotypes and are actually susceptible to outbreeding depression (19, 20). Nevertheless, experimental evolution studies reveal that the frequency of males and outcrossing become elevated with increased mutation rate or novel environmental challenges, such as coevolving parasites (21–23). A similar benefit for outcrossing in facilitating adaptation has also been observed in hermaphroditic snails (24).

In addition to possible fitness differences between selfed and outcross progeny, the frequency of males is strongly influenced by male mating ability and hermaphroditic receptivity, which vary among species and populations (21, 25). To successfully produce outcrossed progeny, a male must locate a hermaphrodite, initiate a complex suite of copulating behaviors, and successfully transfer sperm (26, 27). After insemination, male sperm must compete with the hermaphrodite’s self-sperm for fertilization. The XX–XO system of genetic sex determination of Caenorhabditis means that variation in pre- or postmating male mating success will impact the frequency of males (28). However, the genetic basis of male mating ability and how it links to the difference in male frequency remains unclear.

XO Caenorhabditis males are produced from either outcrossing or from the spontaneous nondisjunction of the X chromosome during meiosis. The frequency of males varies in different species and populations (21, 29, 30), but it is always below 0.5. N2 Bristol, the common C. elegans laboratory strain, has a male frequency of ≤0.002 (8, 30), near the rate of production by nondisjunction (31–33). The low frequency of males in these selfing lineages may often be adaptive. Under benign conditions, hermaphrodites enjoy a population growth rate advantage that allows invasion of male/female populations by eliminating the cost of males (34, 35). In addition, selfing provides reproductive assurance where mates are limited, facilitating colonization of new habitat patches (36–38). Finally, the process of mating with males can physically and physiologically damage the hermaphrodites (39–42).

The simultaneous benefits of selfing and need for at least occasional outcrossing may create an intermediate optimum sex ratio in androdioecious species like C. elegans. The modulation of sex ratio could be implemented at the level of mating success. Previously we identified a Caenorhabditis gene family, mss (for ‘‘male secreted short’’), that can affect male frequency. mss genes are conserved in outcrossing species, but are consistently lost in the androdioecious species (43). Mss proteins are heavily glycosylated surface factors that confer male sperm competitiveness. In the outcrossing Caenorhabditis remanei, mutant males lacking all mss paralogs are fully fertile, yet sire fewer progeny than wild-type (WT) mss+ males in competition. In the selfing C. briggsae, transgenic males in which mss has been restored sire more outcrossed progeny than mss− (i.e., WT) males, and are more effective at suppressing hermaphrodite self-sperm use. The strong advantage conferred by mss thus presents a mystery: Why

Significance

Several species of Caenorhabditis nematodes, including Caenorhabditis elegans, have recently evolved self-fertile hermaphrodites from female/male ancestors. These hermaphrodites can either self-fertilize or mate with males, and the extent of outcrossing determines subsequent male frequency. Using experimental evolution, the authors show that a gene family with a historical role in sperm competition plays a large role in regulating male frequency after self-fertility evolves. By reducing, but not completely eliminating outcrossing, loss of the mss genes contributes to adaptive tuning of the sex ratio in a newly self-fertile species.

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*To whom correspondence may be addressed. Email: ehaag@umd.edu.

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would the males of androdioecious species consistently lose \textit{mss} if it benefits them? One possibility is that the reduction of sexual selection leads to loss of \textit{mss} purely through drift. Here, we consider an alternative, namely that \textit{mss} loss was adaptive in the incipient selfing ancestors of \textit{C. elegans} and \textit{C. briggsae} because it resolved a sexual conflict over mating (44).

Hamilton (45) proposed that in isolated small populations where only mated females disperse, interbreeding selection favors a female-biased sex ratio. The inbreeding associated with this mating system imposes additional selection to minimize male frequency (46). Though these models were produced with fig wasps and other gonochoristic organisms in mind, they are also highly relevant to \textit{Caenorhabditis}, which colonize spatially isolated fruits in small numbers (13, 47, 48). These circumstances resemble Hamilton’s local mate competition (LMC) scenario, but offer the extreme case of zero males as a viable option to selfing hermaphrodites.

We have used simple population models and experimental cultures to simulate natural populations of the sort that may have existed as \textit{C. briggsae} was evolusioning, and adapting to, selfing. We find that the change to partial self-fertility is insufficient to explain the parallel loss of the \textit{mss} genes under a model of global panmixia. We therefore hypothesized that perhaps \textit{C. briggsae} populations without \textit{mss} (which we will refer to as \textit{mss−}) would grow faster than others with \textit{mss+}, owing to a lower male frequency and increased egg output in the former. We provide theoretical and experimental support for the plausibility of this scenario. Overall, our results suggest that mating system and population structure have interacted to promote the adaptive loss of the \textit{mss} family in selfing \textit{Caenorhabditis} species.

**Methods**

\textit{C. briggsae} Strains. AF16, a \textit{C. briggsae} wild isolate originally isolated in India, was obtained from the \textit{Caenorhabditis} Genetics Center. CP161 (nms7[Cni-mss-1(+)] Cni-mss-2(+) Cbr-myo-2::GFP unc-119(+) ish); Cbr-unc-119(nm67) III) and CP162 (nms8[Cni-mss-1(+) Cni-mss-2(+) Cbr-myo-2::GFP unc-119(+) ish); Cbr-unc-119(nm67) III) are independent \textit{mss} chromosomally integrated transgenic strains generated through microparticle bombardment as described previously (43). In addition to \textit{mss} genes from \textit{C. briggsae} nigonii, the construct has a dominant \textit{Cbr-myo-2::GFP} reporter and \textit{Cbr-unc-119(+)}, which serves as a marker for identification of successful transgenic lines. CP164 was generated by crossing \textit{Cbr-myo-2::GFP} reporter and \textit{Cbr-unc-119(+)}, followed by selfing and sibling mating to isolate an outcrossing strain that is colonizing \textit{Caenorhabditis}, which colonize spatially isolated fruits in small numbers (13, 47, 48). These circumstances resemble Hamilton’s local mate competition (LMC) scenario, but offer the extreme case of zero males as a viable option to selfing hermaphrodites.

Measuring Sex Ratios and Population Growth in Genetically Homogeneous Cultures. For each strain (AF16, CP161, and CP162), 50 L4 males and 50 L4 hermaphrodites were set up for crossing for 24 h, after which an individual hermaphrodite (as verified by mating plug) was transferred to a 6-cm NGM plate seeded with \textit{E. coli} strain OP50 (3-cm diameter) and allowed to lay embryos for 40 h before the hermaphrodite mother was removed. After 3 d of incubation at room temperature, the number of F1 males and hermaphrodites (L4 and adult stages) was counted. Worms were washed off of the M9 buffer surface (90 mm; 0.05% Tween; (totally submerged worms sticking to pipette tips) and resuspended in 200 μL in a 2-μL microtube. The entire suspension was transferred onto a plain glass slide (AmScope) as thin streaks of liquid for counting with a stereo microscope. F2 populations were cultured as described above, except the mated P0 hermaphrodite was transferred to a 10-cm NGM plate (OP50 bacterial spot diameter: 6 cm). The larger plates ensured the worms did not run out of food after two generations. After 5 d at room temperature, the worms were washed into a 15-mL Falcon tube. For sex ratio estimation, -100 worms were transferred onto a plain glass slide. For automated counting of entire cultures, a separate set of culture plates was used. Each worm suspension was diluted 10-fold with M9 buffer and aliquoted to 96-well microtiter plates at a density of approximately 10 worms per microliter. The entire population of animals in the 96-well microtiter plate was then counted using the large particle nematode sorter (COPAS Biosort, Union Biometrica). Initially no specific size gating was set, so that worms of all sizes were included in the counts. Subsequently, we realized F2 animals were beginning to lay embryos and these were hatching in L1 larvae. Smith et al. (50) demonstrated extinction (EXT) values below 39 (log EXT -3.669) correspond to L1 larvae and detritus. We therefore used this as a cutoff to distinguish F2 adults from young F3 larva. Only older F2 worms were used in the summary statistics boxplots and calculation of its statistical significance.

**Results**

Fitness of \textit{mss+} Allele in Competitive Context Varies with Mating System. In extant \textit{C. briggsae}, traces of \textit{mss} genes remain as pseudogenes (45), indicating they were lost recently. To assess how the shift to selfing may have impacted \textit{mss} fitness, we first modeled the population dynamics of an \textit{mss}-like modifier of male fertilization success in both the derived self-fertile condition and in the ancestral, obligately outcrossing system. Our approach was similar to previous models of the maintenance of males in androdioecious nematodes (35, 51, 52). In principle, selfing rate, male fertilization success (\(\alpha\)), inbreeding depression, the rate of nondisjunction at the X chromosome, and relative viability difference between males and hermaphrodites all can impact the frequency of males. However, since inbreeding depression is low in long-selfing species (19, 20, 53), X nondisjunction is rare, and male and hermaphrodite viability is equal in the early, reproductive phase of life, a simplified model for the change of male frequency overtime would be \(m'=am/2\), where \(m\) and \(m'\) are male frequency at current and subsequent generation, respectively, and \(\alpha\) is the male fertilization success. \(\alpha\) combines both male mate-finding (premating) and sperm precedence (postmating)
success, and can be anywhere from 0 to 2. \( \alpha m \) is the proportion of hermaphrodite eggs that are outcrossed.

When \( \alpha \) is maximal, \( m^* = m \), and males are maintained. If males additionally start as half the population, all progeny are cross-fathered and the population is a male/female equivalent. Assuming a constant value of \( \alpha \) at all male frequencies, any value of less than two leads to eventual loss of males. There is some indication that \( \alpha \) may vary as a function of male frequency (35), but in common laboratory strains (e.g., \( C. elegans \) N2, \( C. briggsae \) AF16) populations enriched for males steadily lose them (8, 35, 43). However, even in this case, \( mss^+ \) males still have a transient advantage over \( mss^- \) males, and the \( mss^+ \) genotype is expected to increase in frequency whenever males are present in appreciable numbers (Fig. 1A). In the extreme case of obligate outcrossing, \( mss^+ \) should steadily increase in frequency until it is fixed in the population (Fig. 1B), consistent with its consistent presence in gonochoristic \( Caenorhabditis \).

To experimentally assess how the change to selfing impacts the fitness of the \( mss^+ \) genotype, we carried out two experimental evolution studies. One employed self-fertile stocks mixed with males, and the other a self-sterile, obligately outcrossing strain homozygous for a \( she-1 \) (sporeless hermaphrodite) loss-of-function mutation (49). \( she-1 \) mutants reproduce effectively as a male/female strain, similar to \( fog-2 \) mutant \( C. elegans \) (54). The frequency of the \( mss^+ \) transgene, which is marked with a dominant \( myo-2::GFP \) reporter, was tracked in mixed-genotype populations for 25 generations (Fig. 1C). Each replicate began with equal numbers of mated AF16 (\( mss^+/- \), GFP−) and CP161 (\( mss^+/- \), GFP+), where selfing and outcrossing exist, GFP+ (sum of \( mss^+/+ \) and \( mss^-/- \) diploid genotype frequencies is also shown (GFP+, thin solid lines) for comparison with experimental data. (B) Modeling the competition between \( mss^- \) and \( mss^+ \) genotypes in a panmictic, obligately outcrossing (i.e., male/female) population. The sperm competitiveness factor (\( C \) for \( mss^+ \) is set at 1, while that for \( mss^- \) males can vary from 0 (sterility) to 1 (equal to \( mss^+ \)). (C) Experimental evolution scheme for assessing \( mss^+ \) fitness in androdioecious and gonochoristic population. (D) In a mixed population of \( C. briggsae \) WT AF16 (\( mss^-/- \), GFP−) and CP161 (\( mss^+/- \), GFP+), where selfing and outcrossing are allowed, GFP+ (sum of \( mss^+/+ \) and \( mss^-/- \) diploid genotype) frequency increases from 50 to a median of 63.5% over 25 generations. The estimated mean and median \( mss^+ \) frequency at the 25th generation, 0.59 and 0.57, are significantly higher than 0.50 (\( P = 0.0287 \); one-tailed sign test, \( n = 14 \)). (E) Competition in a mixed population of \( C. briggsae \) she-1(v49); \( mss^-/- \), GFP−; \( mss^+/- \), GFP+), where all XX animals lack self-sperm. Median \( mss^+ \) frequency increased to 0.72 at the third generation, and was 0.87 at the 25th generation. The GFP+ frequency distribution at generation 25 is significantly higher than 0.75 (\( P = 0.0002594 \); one-tailed sign test, \( n = 16 \)). Similarly, the mean \( mss^+ \) allele frequency at the 25th generation is 0.67, significantly higher than the initial 0.50. (\( P = 7.629e-06 \); one-tailed sign test, \( n = 16 \)).
mss+; she-1 (GFP+) worms, GFP+ frequency increased to a median 0.72 at the third generation. This initial increase can be fully explained by the formation of the first heterozygotes. However, the frequency of GFP+ genotypes continued to rise, to a median value of 0.87 at the 25th generation (Fig. 1E). This is significantly higher than the 0.75 expected from genetic drift alone. In addition, the mean mss+ allele frequency at the 25th generation is 0.67, significantly higher than the initial 0.50. These results indicate that mss+ also enjoys higher fitness than mss− in she-1 mutants and confirm that obligate outcrossing selects more strongly for mss+ than the mixed-mating case.

**mss+ Increases Male Production and Depresses Growth.** Given the measurable positive selection for mss+ in large mixed-mating populations, we considered an alternative situation, in which small, genetically homogenous subpopulations compete against each other. We previously observed that CP161 and CP162, two transgenic mss+ *C. briggsae* strains, were better at maintaining males compared with the mss− AF16 WT strain (43). Modeling predicts that in populations of initially identical size, those that are mss+ (i.e., whose males have a higher value of α) would produce more male offspring (and thus fewer hermaphrodites) and grow more slowly than those that are mss− (Fig. 2A and B).

To test the first prediction, mated hermaphrodites of the AF16 wild-type, CP161, and CP162 strains were used to found single-genotype populations (Fig. 2C). As expected, the percentage of males in the AF16 (mss−) population was significantly lower than in the two mss+ populations in the second (i.e., F1; Fig. 2D) and third (i.e., F2; Fig. 2E) generations.

Using these results, the proportion of hermaphrodite eggs that are outcrossed (αm) can be estimated for each of the genotypes. From P0 to F1, they are 89.2% (AF16), 100% (CP161), and 100% (CP162), respectively. From F1 to F2, they are 72.6% (AF16), 86% (CP161), and 89.2% (CP162). In the P0 to F1 generation, α for the three strains is estimated to be 1.79 (AF16), 2 (CP161), and 2 (CP162). Because only hermaphrodites that bore copulatory plugs were picked for the P0 founders, these values reflect only postmating sperm competition. From the F1 to F2 generations, where successful sperm transfer by the F1 males was not guaranteed, α was estimated to be 1.62 for AF16, 1.67 for CP161, and 1.71 for CP162. Thus, male fertilization success is consistently higher in the mss+ populations than in the mss− AF16. Nevertheless, by the F2 generation some hermaphrodites were not outcrossed despite the presence of many males.

We next examined the prediction of slower population growth in mss+ cultures. The number of progeny laid by a mated hermaphrodite in 40 h was not significantly different between mss− AF16 and the two mss+ transgenic lines (Fig. 3A). Thus, an intrinsic fertility difference between mss+ *C. briggsae* and mss− *C. briggsae* can be ruled out, validating a key assumption of our model. After the F1s produced F2s, we used an automated worm sorter to precisely count all progeny in these much larger populations. Wild-type AF16 produced a significantly higher number of F2s (n = 16 cultures, median = 16,140) than CP161 (n = 12, median = 12,870) and CP162 (n = 12, median = 14,050) (Fig. 3B and C). These results confirm the expected reduction of population growth incurred by high male mating success. To better understand the dynamics of population growth difference as a function of sex ratio difference, we can model population growth as \( \alpha = (p) \times (h) \times (f) \), where \( p \) is the population size at generation N, \( p \) is the population size at generation N-1, \( h \) is the fraction of hermaphrodites, and \( f \) is the fecundity of each hermaphrodite. The median fractions of hermaphrodites at the first generation (0.552 for AF16 WT, 0.484 for CP161, and 0.479 for CP162) predict that in the second generation AF16 WT will have 1.14 times more progeny than CP161, and 1.15 times more than CP162. These are close to our experimental finding: AF16 has 1.25 times more progeny than CP161 (expected 1.14), and 1.15 times more progeny than CP162 (expected 1.15). We conclude the F2 growth of these experimental cultures can be predicted by F1 sex ratios.

**Discussion**

The repeated evolution of self-fertility suggests that it benefits species that adopt it, likely through reproductive assurance and a large boost in intrinsic growth rate (55, 56). The ongoing role of males, however, is less clear. While *Caenorhabditis* hermaphrodites lay more embryos after crossing (25), total brood size is not likely to be the most significant contributor to the growth of the population. First, the reproductive value of late progeny is discounted compared with the progeny laid earlier (57). Furthermore, *C. elegans* and

![Fig. 2. mss+ *C. briggsae* produce more males. (A) Modeling the impact of varying fertilization success (α) on sex ratio over time in a genetically homogenous population. Males and hermaphrodites are both at 50% frequency in the first generation. At any value of α < 2 male frequency declines and hermaphrodite frequency increases. (B) Expected impact of α on population growth. At generation one, the population is founded with one mated hermaphrodite. At generation two, population size is determined solely by the fecundity of this hermaphrodite (here we assume each produces 100 eggs). The advantage of low male fertilization success is seen in the substantially larger number of grand-offspring in generation three. (C–E) Experimental examination of male frequency. (C) Schematic of experimental design. Replicated populations of the indicated genotype were started with mated hermaphrodites and subsequently scored for male frequency in the first (F1) or second (F2) generation. (D) In F1 populations (progeny of a single mated hermaphrodite), CP161 (mss+) (n = 20) and CP162 (mss+) (n = 22) both have a significantly higher ratio of males compared with AF16 (wild type) (n = 16). The median percentage is 0.448, 0.516, and 0.521 for each strain. (E) In the F2 generation, CP161 (mss+) (n = 12) and CP162 (mss+) (n = 10) again both showed a significantly higher ratio of males compared with AF16 wild type (Kolmogorov–Smirnov test). Individual data points represent the frequency of males in each replicate population.](Image 327x413 to 538x712)
C. briggsae hermaphrodites evolved traits that actually reduce their chance of mating. They secrete less potent sex pheromones (8, 58–60), are less receptive to males that do locate them (27), and can eject the male sperm after transfer (61, 62). While these traits make outcrossing inefficient, they create more hermaphrodite-biased sex ratios and reduce the potential for sterilizing matings with males of sympatric, gonochoristic congeners (41, 63). It appears that outcrossing is minimized in androdioecious species by a suite of changes in both sexes, yet complete male inviability or sterility has not evolved (with exceptions; ref. 64). Our results suggest that loss of mss+ was one of these changes.

**Effects of Mating System on mss Fitness.** We predicted that in both mixed-mating populations with selfing hermaphrodites and males, and in obligately outcrossing populations, the mss+ transgene will increase in frequency (Fig. 1 A and B). This is expected to continue to fixation in the outcrossing competition, but be muted and dependent on the presence of males in the mixed-mating situation. Our experimental populations generally support these conclusions. In the male/female (she-1) mixed population, the mss+ allele had a strong advantage, consistent with the observed retention of mss+ in obligately outcrossing species (43). In the mixed-mating case, we also found an increase in mss+ frequency, and as expected it was less pronounced than in the outcrossing case. However, we did not observe an initial sharp increase in GFP+ frequency as expected (compare Fig. 1 A and D), indicating that mating between mss− and mss+ animals in the first generations was less than the values of α estimated from other experiments would predict.

Given the initially low mating inferred (by GFP+ frequency) in the first few generations of the mixed-mating experiment, it is surprising that male frequency (m) remained as high as it did (median 0.09) at generation 25. Using the empirically calculated value of α for the mss+ CP161 strain in the second generation (1.67) for all subsequent generations, m is predicted to be 0.007 in generation 25, an order of magnitude smaller than observed. A likely explanation is that α (fertilization success) of males varies as a function of male frequency, as suggested by Stewart and Phillips (35). Rare males may encounter virgin hermaphrodites (instead of mated hermaphrodites or other males) more reliably, allowing more matings per male. It is also possible that some assortative mating occurred with regard to mss genotype, though no mechanism to mediate this is apparent.

**Loss of mss Boosts Population Growth.** The parallel loss of mss in three distinct lineages is striking. However, the measurable positive selection for mss+ we observed in partially selfing populations indicates that this loss is unlikely to be via drift alone. Our results are consistent with a model whereby loss of the mss family was a response to LMC-like conditions that selected for minimal outcrossing. To approximate the LMC scenario, we compared male frequencies and population growth rates in populations of pure mss− and mss+. While mss+ and mss− C. briggsae hermaphrodites do not differ in their intrinsic fecundity or viability (Fig. 3A), we found higher frequencies of males in mss+ populations (Fig. 2 D and E). Presumably as a direct consequence of this elevated male frequency, the third generation mss+ populations had lower census sizes (Fig. 3B), as predicted (Fig. 2 A and B). With a transient food supply and in the presence of other competing species, this enhanced growth would produce more dispersive L3-dauer larvae. The loss of the mss+ allele could therefore increase outmigration and the probability of successful colonization of the next patch. In contrast, similar small populations of strict outcrossers would gain nothing from mss loss, because male frequency is a constant 50%.

**Conclusions**

Males are rare in natural populations of androdioecious Caenorhabditis, often rarer than observed in the laboratory conditions (65, 66). Outcrossing is associated with growth-retarding effects of males addressed above, as well as outcrossing depression and other genetic signatures (19, 67–69). Nevertheless, males persist and are generally fertile worldwide. Exposure to the dauer lifestyle, environmental stresses such as heat shock, mutational load, and novel pathogens all can lead to increased outcrossing (22, 70). Thus, the sex ratio of an androdioecious population likely represents a balance between selection for

![Fig. 3. mss+ populations grow slower. (A) The F1 progeny produced by a single mated hermaphrodite of strains AF16 (n = 16), CP161 (n = 20), and CP162 (n = 21) are not significantly (n.s.) different. (B) The F2 strains CP161 (n = 12) and CP162 (n = 12) both produced significantly smaller numbers of F2 progeny than the mss− AF16 (n = 16). P value indicated for each comparison is by the Kolmogorov–Smirnov test. (C) Examples of population size determination by automated worm sorter. Eight replicates with the number of worms closest to the medium value were used to make the scatterplots of extinction (EXT) vs. time-of-flight (TOF) data for individual worms. Data points in gray represent small, translucent embryos, larvae, which are F3 progeny. Data points in red represent animals developed enough to be F2 (or older) after 5 d of growth.](image)
outcrossing under stressful conditions and selection to maximize hermaphrodite selfing under more benign conditions. We propose that this loss represents a mechanism to help tune the sex ratio to a hermaphrodite-biased value while still allowing outcrossing.