Non-Mendelian assortment of homologous autosomes of different sizes in males is the ancestral state in the *Caenorhabditis* lineage

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Organisinal genome sizes vary by six orders of magnitude and appear positively correlated with organismal size and complexity. Neutral models have been proposed to explain the broad patterns of genome size variation based on organism population sizes. In the *Caenorhabditis* genus, hermaphrodite genomes are smaller than those of gonochoristic species. One possible driving force for this genome size difference could be non-random chromosome segregation. In *Caenorhabditis elegans*, chromosome assortment is non-independent and violates Mendel’s second law. In males, the shorter homologue of a heterozygous autosome pair preferentially co-segregates with the X chromosome while the longer one preferentially co-segregates with the nullo-X (O) chromosome in a process we call “skew”. Since hermaphrodites preferentially receive the shorter chromosomes and can start populations independently, their genome size would be predicted to decrease over evolutionary time. If skew is an important driver for genome size reduction in hermaphroditic *Caenorhabditis* species, then it should be present in all congeneric species. In this study, we tested this hypothesis and found that skew is present in all eight examined species. Our results suggest that skew is likely the ancestral state in this genus. More speculatively, skew may drive genome size patterns in hermaphroditic species in other nematodes.

Why is there variation in genome size amongst organisms? Organisinal genome sizes span six orders of magnitude2,3 from ~160 kb (bacterial endosymbiont *Carsonella ruddii*) to ~150 Gb (angiosperm, *Paris japonica*). Genome size is a complex multivariate trait. Nevertheless, for viruses, prokaryotes and many unicellular eukaryotes, genome size can be explained largely by the gene number2. In addition to gene number, population genetic principles have been proposed to explain genome size and complexity2,3. The basic model is that organisms with smaller effective population sizes are less efficient at removing mildly deleterious mutations, in particular transposons and introns5,6,7. Thus, lineages with small population sizes will drift towards larger genomes. While the magnitude of the effect of population size on genome size variation is under debate, ranging from ~8–64%5, the remaining variation in genome size is still largely unexplained.

In the Elegans subgroup of the *Caenorhabditis* genus, all 3 cases of the evolution of hermaphroditism have been associated with a convergent reduction in genome size relative to gonochoristic (male/female) species8–13. However, based on the effective population sizes in *Caenorhabditis* species, the genome sizes are opposite of the simple population size predictions. The gonochoristic species (*C. remanei* and *C. brenneri*) have larger population sizes than hermaphroditic species (*C. briggsae* and *C. tropicalis*), consistent with the expectations for outcrossing and selling species8–10. However, gonochoristic species (*C. remanei, C. brenneri, C. japonica, C. sinica* and *C. mignon*) have larger genome sizes than hermaphroditic species (*C. elegans, C. briggsae* and *C. tropicalis*)8–10.

Non-Mendelian chromosome assortment may also be associated with genome size differences in the *Caenorhabditis* genus. In males, there is preferential segregation of the longer chromosome of a heterozygous homologous pair away from the X chromosome and of the shorter chromosome with the X17. We call this phenomenon “skew”. Given that hermaphrodites tend to inherit the shorter chromosome and because new

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populations of C. elegans are frequently initiated by hermaphrodites\textsuperscript{18–21}, skew may partially explain genome size reduction relative to the ancestral gonochoristic species.

If skew plays a role in genome size reduction within the clade, then skew should be present in other Caenorhabditis species. In this study, we tested this hypothesis by examining transgenic strains of multiple species across the Caenorhabditis genus and then observing chromosome inheritance patterns. We examined both integrated and extrachromosomal transgenes as they are compositionally similar in Caenorhabditis nematodes.

Finally, we wanted to determine whether skew was male germline specific or general. We found that all eight of the tested species exhibited skew, suggesting that skew was probably present in the ancestral Caenorhabditis species.

Results
Integrated transgenes are transmitted preferentially from fathers to sons in both hermaphroditic and gonochoristic species. A previous study demonstrated skew in C. elegans\textsuperscript{17}. For comparison in this study, we retested an outcrossed mIs10 strain by crossing heterozygous mIs10/+ males to Unc hermaphrodites. Counts of their nonUnc cross progeny confirmed skew with a transmission bias ratio (TBR; “preferred”/“anti” gamete combinations, see Methods) of 3.86. This value is smaller than previously observed for mIs10 (TBR = 6.55\textsuperscript{17}). The difference is likely because of one or more skew modifier mutations being removed or introduced during our outcrossing of mIs10 (see Supplementary Text S1).

Next, we tested the two other hermaphroditic species, C. briggsae and C. tropicalis, in a similar fashion. In C. briggsae, examination of three heterozygous integrated transgenes (mfIs42, syIs803, and syIs807) revealed skew in all cases with the TBRs ranging from 1.57 (syIs803) to 4.08 (mfIs42) (Fig. 1, all \( P < 0.001, \chi^2\)-test, df = 1). Likewise, C. tropicalis exhibited skew (ants1, TBR 2.08; \( P < 0.001, \chi^2\)-test, df = 1).

To determine whether skew was present in gonochoristic species, we tested C. brenneri (ants2), C. portoensis (ants7), and C. remanei (mfIs83\textsuperscript{22}). In all three species, we observed skew (Fig. 1, TBR’s from 1.19 to 1.34, all \( P < 0.001, \chi^2\)-test, df = 1). Thus for all six species tested, we observed skew, regardless of the mode of reproduction employed, i.e., hermaphroditism or gonochorism.

TBR varied across the lines and species. In C. elegans, TBR is positively correlated with insertion size\textsuperscript{17}. We tested for a similar correlation across the strains (four species) but found none (\( P = 0.41, \) Spearman’s rank correlation test) indicating that species-specific factors likely modify TBR. However, three C. briggsae strains show a suggestive pattern consistent with a positive correlation (Supplementary Figure S1). Examination of additional insertion strains within each species would clarify the magnitude of TBR explained by insertion length.

Extrachromosomal transgenes were also transmitted preferentially from fathers to sons in hermaphroditic and gonochoristic species. Given the similarity of DNA composition between extrachromosomal and integrated arrays, we examined if the former also shows biased transmission from fathers to sons. We first tested two hermaphroditic species, C. elegans and C. tropicalis, by crossing males carrying extrachromosomal arrays with GFP or mCherry to tester hermaphrodites. These crosses revealed that extrachromosomal arrays were also preferentially transmitted to sons in both hermaphroditic species (both \( P < 0.008, \chi^2\)-test, df = 1, Fig. 2; and Supplementary Figures S2 and S3).
Next, we tested four gonochoristic species. For three species (C. nigoni, C. portoensis, and C. sp. 33), only one extrachromosomal line was generated, and in all cases we observed skew (all $P < 0.001$, $\chi^2$-test, df = 1; Fig. 2).

We also observed skew in C. brenneri for three extrachromosomal lines (antEx3, antEx4, antEx117), but a fourth one did not (antEx5). The TBRs for these extrachromosomal arrays in these four species ranged from 0.99 to 3.82 (Fig. 2).

Skew appears specific to the male germline. Thus far, all tests for skew assayed males which are XO and only produce sperm. In the following experiments, we tested if skew might be restricted to sperm or to males. Wild-type hermaphrodites also produce sperm, but since they are XX, testing for preferentially segregation of insertions away from the X is not possible. Thus, we used fem-3 and her-1 mutants where XO individuals are transformed into females (producing only oocytes) or self-fertile hermaphrodites, respectively.

We first tested whether oocytes exhibited skew by crossing fem-3 XO pseudo-females heterozygous for a GFP transgene (gfp/+; fem/fem) to males with an X-linked RFP (rfp/O) and scoring their progeny. The X-linked RFP transgene was necessary to identify sons that inherited the paternal X (RFP) as opposed to the maternal X (non-RFP). This cross yielded six viable offspring genotype classes (of eight possible) which we grouped into the "preferred" or "anti" gamete combinations (see Methods and Supplementary Figure S4). Of 177 individuals scored, 92 and 85 inherited the "preferred" and "anti" gamete combinations, respectively, which was not different from random assortment (Fig. 3A, $P = 0.33$, binomial test).

To confirm that this lack of skew in oocytes was not specific to the fem-3 mutant background, we tested her-1 mutants similarly. We crossed XO hermaphrodites (her-1 mls10/her-1 +; XO) to rfp/O males and assayed the progeny phenotypes. Again, we found that oocytes did not exhibit skew (Fig. 3A; 132 preferred, 150 anti, $P = 0.31$, binomial test).

Finally, we tested if the sperm of XO her-1 individuals (her-1 mls10/her-1 +; XO) exhibited skew by looking for deviations from random segregation in self-progeny. Of the self-progeny the most discriminatory cases are the GFP/GFP homozygotes. If there is no skew in the hermaphrodite sperm (i.e., random segregation of chromosomes), then the predicted ratio of XO:XX individuals would be 2:1. In contrast, a transmission bias ratio of ~4:1 for mls10 (above) yields a predicted ratio of ~5:1 for XO:XX individuals (see Supplementary Figure S5). Examination of the self progeny revealed that the segregation pattern was not random (Fig. 3B; 155 XO: 27 XX; $P = 2.3e-8$, binomial test with likelihood of success = 2/3). This corresponds to a TBR of 4.74 for hermaphrodite sperm, which is not different from the ~4 TBR for mls10 in normal males ($P = 0.55$, binomial test with likelihood of success = 5/6). Together these data indicate that skew is not specific to the soma sex, per se, but is limited to male meiosis or some subsequent step of spermatogenesis.

Discussion
We found that the transmission of transgenes preferentially toward sons is present in all Caenorhabditis species tested. Skew occurs in C. elegans whereby fathers transmit autosomes that are heterozygous in length to their offspring in a non-Mendelian fashion15. In this study, we used integrated transgenes to demonstrate that all three
known hermaphroditic species and three gonochoristic species of *Caenorhabditis* exhibit skew (Fig. 1). We also showed that extrachromosomal arrays are preferentially transmitted from fathers to sons rather than daughters in six species (Fig. 2). Because of the similar transmission patterns and DNA composition for both integrated and extrachromosomal transgenes, we refer to this transmission bias in both as skew. In total, we observed skew in all eight of the *Caenorhabditis* species tested (Fig. 4).

To our knowledge, this study is the first to systematically examine skew in multiple species in the *Caenorhabditis* genus. However, skew or likely cases of it have been noticed previously. Several early *C. elegans* studies observed that free duplications of partial chromosomes were transmitted preferentially from fathers to sons in *C. elegans*23–26. More recently, a study examining hybrid incompatibility found that *C. nigoni* males heterozygous for a *C. briggsae* introgression (strain ZZY10028) preferentially transmitted the *C. nigoni* orthologous genomic fragment to sons27. As the *C. nigoni* genome is larger than that of *C. briggsae*28, the focal homologous genomic fragment is presumably longer than the introgressed *C. briggsae* counterpart, and thus the chromosome transmission patterns are consistent with skew27.

With the current evidence, it is very likely that all the species in the Elegans subclade display skew. Our study also included one species, *C. portoensis*, that is outside of the Elegans subclade. This would suggest that skew is conserved among all *Caenorhabditis* species and was likely already present in the ancestor of the *Caenorhabditis* genus.

The finding that all *Caenorhabditis* species have skew is important because this observation supports the possibility that smaller genomes in hermaphrodite (all circa 100 Mbp) compared to gonochoristic species (all circa 130 Mbp)9,10,13 could be due, in part, to skew.

How might skew have led to genome reduction in hermaphrodite species? With skew, every time a male carrying a heterozygous deletion (or insertion) mates to a hermaphrodite, the daughter hermaphrodites will tend to inherit the shorter allele. Because a few or even a single hermaphrodite individual is sufficient to colonize a patch, an opportunity for a ratchet-like reduction in genome size could occur after every such mating involving a deletion. In contrast, for gonochoristic species although daughters would preferentially inherit the shorter chromosomes, sons would preferentially inherit the longer one. On average, matings should mix the slightly smaller female genomes with the slightly longer male ones resulting in no directional change in genome size. This model is supported by simulations17. Thus, the current *Caenorhabditis* genome size patterns could partially reflect skew operating over evolutionary time scales.

Reduction of hermaphrodite genome sizes due to skew requires mating. Direct observations of males in hermaphrodite species are rare5,10,21 and genetic estimates of the natural mating rate in extant hermaphrodite species is low18,19,29. However, matings were likely more frequent at the onset of hermaphrodite evolution, potentially implying that the rate of genome size reduction may have been faster at that time. The evolution of
hermaphroditism can occur with mutations in only two genes\textsuperscript{30,31}, which do not cause reproductive isolation. Thus, in any potential cases of abrupt evolution of hermaphroditism, matings may still occur frequently with sympatric males of the originating gonochoristic sister species before speciation has occurred. A more gradual evolution of hermaphroditism, perhaps via an intermediate mating system with males, females, and hermaphrodites (e.g., the free-living Rhabditis sp. SB34732–34 and the entomopathogenic \textit{Heterorhabditis} species\textsuperscript{35,36}), would imply more mating occurrences, and again more opportunity for genome reduction.

While skew could cause genome reduction, whether it has is unclear. Genome size can be affected by many factors including directional bias in mutations, effective population size, and transposable element accumulation\textsuperscript{37–39}. A recent study comparing the genomes of the three hermaphroditic species with five gonochoristic species came to the conclusion that the main driver for hermaphrodite genome reduction was probably the loss of genes associated with traits no longer needed\textsuperscript{9}. Some gene losses may have been beneficial because the associated mating or sex-specific traits were costly to maintain; more investment could be made in other aspects of its biology. Other, and possibly most, genes and traits were probably lost because of relaxed selection. Genome reduction in this model would not require mating, and hence, would not require skew.

Nevertheless, skew is still compatible with both positive and relaxed selection because any deletion would need to spread throughout the population in order to reach fixation. In principle, a deletion strain experiencing positive selection could completely sweep through the population in a clonal fashion. With matings the spread of a beneficial deletion allele would be accelerated because the direction of selection and skew are coincident. During relaxed selection, drift would play a stronger role than selection, and for truly neutral deletion alleles fixation or loss is proportional to its allele frequency. In these cases, skew could be a driver, providing directionality for the fixation of neutral or nearly neutral deletion alleles.

Quantifying the contribution, if any, of skew on genome size reduction in hermaphroditic species may be challenging. One possibility would entail long-term experimental evolution that contrasted an obligately mating strain (e.g., \textit{fog-2} or \textit{spe-27} mutants\textsuperscript{40,41}) to a hermaphroditic strain experiencing episodic matings. Genome comparisons between true sister species pairs of hermaphroditic and gonochoristic species may also be enlightening.

In addition to a potential impact on genome size differences, skew may have an impact on sexually antagonistic alleles\textsuperscript{42}, which are alleles that benefit one sex but harm the other. When an allele is found equally often in males and females (as for typical autosomal genes), the allele’s beneficial or harmful effects often need to be balanced between the sexes. However, if an allele is more frequently found in one sex, then sexual antagonism may be easier to evolve. The sex chromosomes in heterogametic organisms (e.g., X and Y chromosomes) are an extreme example. Given skew in \textit{Caenorhabditis} species, selection may be possible for male beneficial (and female detrimental) alleles on the longer chromosomes and female beneficial (and male detrimental) alleles on the shorter chromosomes\textsuperscript{43}. As there is natural variation in gene copy number in \textit{C. elegans} and other nematodes\textsuperscript{45–46}, it is likely that wild variation in copy numbers of genes is common. Thus, an interesting future avenue will be to test if there are any such sexually antagonistic alleles, especially in the gonochoristic species where the two sexes are always present.

Figure 4. Skew across the \textit{Caenorhabditis} genus. Summary of skew results placed on the \textit{Caenorhabditis} phylogeny. All eight species tested exhibited skew for integrated (Is) arrays, extrachromosomal (Ex) arrays, or both (blue names and circles). Blue lines indicate inferred ancestral, evolutionary lineages with skew. Other species and clades within the \textit{Caenorhabditis} are likely to have skew but are untested (gray text and triangles; numbers indicate number of known species within). Phylogeny adapted from\textsuperscript{12,70} with the position of \textit{C. sp 33} placed next to \textit{C. remanei} based on ITS2 sequence (unpublished data).
Meiotic cells can segregate chromosomes that are not joined by chiasmata (sites of crossovers during homologous recombination) through a process called achiasmate segregation. Achiasmate segregation is often considered a "backup" mechanism to segregate homologous chromosomes that fail to form crossovers. This is also the primary mechanism to segregate some autosomes that never form crossovers such as the 4th chromosome in females and all chromosomes in males in Drosophila melanogaster. Achiasmate segregation of heterologous chromosomes also occurs, for example for the heterogametic sex chromosomes that do not have shared regions of homology (i.e., lack pseudoautosomal regions)\(^4\), compound chromosomes in D. melanogaster\(^49,50\), and artificial chromosome pairs in yeast\(^51\). Skew in C. elegans is also a form of heterologous achiasmate segregation (i.e., the solitary X and a longer autosome, which itself is synapsed with its shorter homolog). Achiasmate segregation has also been observed in C. elegans for both free duplications\(^23–28\) and autosomes\(^52\).

In the C.elegans autosomal case, homologous achiasmate segregation appears to occur only in sperm and not in oocytes\(^52\). Our experiments revealed a similar difference for heterologous achiasmate segregation between the gametes for skew. During spermatocyte meiosis, an integrated transgene preferentially segregated away from the X, while during oocyte meiosis (in fem-3 or her-1 XO animals) it segregated randomly with respect to the solitary X. The difference in achiasmate segregation between spermatocyte and oocyte meiosis could be due to the presence of centrioles in spermatocytes and their absence in oocytes\(^53\), differences in the organization of the meiotic spindles\(^54,55\), accelerated meiotic progression kinetics in spermatocytes\(^54\), distinct morphologies of the meiotic chromosomes\(^54\), the presence of a solitary X in males, and/or other unknown differences. From an adaptive perspective, sperm-specific achiasmate segregation may imply that Caenorhabditis meiosis is more prone to failure to form chiasmata in spermatocytes than oocytes.

Finally, during the course of this study, we noticed that all seven non- C. elegans species produced at least one cross with female-biased sex ratios (11 crosses with \(P < 0.0029\); Bonferroni adjusted threshold; binomial test; Fig. 1 and Supplementary Figure S2). Sperm competition or temporally-delayed usage of O-bearing sperm has been implicated previously for C. briggsae\(^56\) and a similar mechanism may explain the observed pattern.

In summary, we report that skew is likely the ancestral state in Caenorhabditis species, which may contribute to the patterns of genome size in this genus. We also show that skew is specific to spermatocyte meiosis. Finally, depending on the extent of conservation of the meiotic machinery, it is possible that skew extends to other nematode taxa.

Materials and Methods

Nematode culture and Caenorhabditis strains. All strains were maintained on Nematode Growth Medium (NGM) plates seeded with E. coli OP50 at 20°C in a TKS Low-Temp incubator LTI 603 or at room temperature (\(-22°C\)). The following strains were obtained from the Caenorhabditis Genetics Center (CGC) or generated in this study. Strains are ordered as follows: wild-type, mutants, transgenics.

C. elegans: N2 var. Bristol (wild-type (wt)), BRC174 (unc-24(e138) fem-3(e1996) unc-24(e138) dpy-20(e1282) IV), BRC183 (unc-24(e138) dpy-20(e1282) IV), BRC189 (unc-119(ed3) III; trit5605 II [EG4322 out-crossed 10 times with N2]), CB3843 (fem-3(c1996) unc-24(e138) dpy-20(e1282) IV), DR108 (dpy-11(c224) unc-42(c270) V), XA8150 (dpy-11(c224) her-1(e1518) V), BRC148 (unc-119(ed3) III; him-8(e1489) IV; antEx50 Puro-R, Neo-R, myo-2::GFP, sur-5::GFP, myo-3::mCherry)), BRC153 (unc-119(ed3) III; him-8(e1489) IV; antEx51 [Puro-R, Neo-R, myo-2::GFP, sur-5::GFP, myo-3::mCherry]), BRC237 (unc-42(c138) fem-3(c1996) unc-24(e138) dpy-20(e1282) IV), mls10 V, OH4460 (atEx2572 [unc-97::NLS::GFP]), PD4793 (mls10[myo-2::GFP, pes-10::GFP, F22879::GFP]) V, WHIR10 (mls10 V; [mls10 in PD4793 was out-crossed 10 times with N2]), and XA8160 (dpy-11(e224) her-1(e1518) mls10 V).

C. briggsae: AF16 (wt), BRC1983 (Cbr-dpy(1281)), JU929 (Cbr-dpy-18 (mf104) III), JU1018 (mf112(Ce-sid-2, Ce-myo-2::DsRed))\(^22\), PS9392 (syb803[Ce-def-4(+) myo-2::GFP])\(^80\) and PS9396 (syb807[Ce-def-4(+) myo-2::GFP]) IV\(^82\).

C. brenneri: CB5161 (wt), JU1397 (wt), BRC467 (antIs2[myo-2::GFP, sur-5::GFP]), BRC532 (antIs3[myo-2::GFP, sur-5::GFP]), BRC536 (antIs17[myo-2::GFP, sur-5::GFP]), BRC567 (antIs5[myo-2::GFP, sur-5::GFP]), and BRC579 (antIs4[myo-2::GFP, sur-5::GFP]).

C. remani: BRC20108 (wt, isolate from Okinawa), PB4641 (wt), and BRC534 (mf183[Cel-sid-2 + Cel-myo-2::DsRed]; allele previously known as mflEx34, M. A. Félix personal communication) (mf183 in JU1184 was out-crossed 8 times with BRC20108)\(^22\).

C. tropicalis: JU1373 (wt), BRC461 (Cir-dpy(ant27)), BRC493 (antEx12[myo-2::mCherry]), and BRC555 (antIs1[mio-2::mCherry]).

C. portoensis: EG4788 (wt), BRC313 (antEx2[myo-2::GFP, sur-5::GFP]) and BRC585 (antIs7[myo-2::GFP]).

C. nigoni: BRC10094 (wt, isolate from Taiwan) and BRC340 (antEx29[myo-2::GFP, sur-5::GFP]).

C. sp. 33: BRC10016 (wt, isolate from Taiwan) and BRC311 (antEx1[myo-2::GFP]).

Generating transgenic strains. All plasmids (p destDD04neo and pDD04neo\(^38\), pBCN21-R4R3\(^39\), pDD158.87\(^60\), pCFJ151 and pCFJ104\(^61\)) were prepared for injection using standard protocols. In brief, each plasmid was transformed into DH5α E. coli cells. Plasmid containing bacteria were cultured in LB broth containing 50 ng/ml ampicillin overnight at 37°C and then purified with the QIAprep Spin Miniprep kit (Qiagen) with final elution using 50 μl of distilled water.

To make transgenic strains for the hermaphroditic species, C. elegans (N2) and C. tropicalis (JU1373), we established extrachromosomal lines using standard C. elegans protocols (Supplementary Table S1)\(^62–64\).

For the gonochoristic Caenorhabditis species (C. brenneri (CB5161), C. portoensis (EG4788), C. nigoni (BRC10094), and C. sp. 33 (BRC10016)), we adapted the microinjection methods for C. elegans\(^62–64\) and C. rema- neri\(^22\) as follows. We used a Narishige IM300 microinjection system attached to a Zeiss Vert A1 microscope to inject the plasmid mixture into the distal arms of the anterior and posterior gonads of virgin wild-type (WT).
young adult females (females were isolated at the L4 stage). Every injected female was placed individually with three to four WT L4 males on an OP50-seeded NGM plate. The F1 progeny were examined daily under an epifluorescent microscope for individuals that expressed one or more of the injected transgenic markers. Specifically, we looked for green fluorescent protein (GFP) or mCherry in the pharynx driven by the Cel-myO-2 promoter or in the hypodermis and intestine driven by the Cel-sur-5 promoter (pPD158.87). Subsequently, single L4 transgenic F1 individuals were placed with three WT L4 individuals (except for C. sp. 33, see below) of the opposite sex.

We enriched for transgenic individuals by selection on antibiotic NGM plates. We prepared NGM plates containing 0.35 mg/ml G418 (Sigma-Aldrich, A1720-25G) seeded with 100 µl of OP50 that had been cultured in LB broth overnight and concentrated by centrifugation58. Single transgenic F2 individuals were crossed to three to five WT individuals of the opposite sex. This was repeated until we were confident that the transgenic line was established, which in all cases was at least 7 generations.

The myo-2::GFP signal was hard to detect at the L3, L4 and adult stages for C. sp. 33. Thus, each L1 or L2 transgenic F2 larva was transferred to a separate NGM plate. Upon reaching the L4 stage, two WT L4 individuals of the opposite sex were mated to each transgenic worm. Subsequently, matings between brothers and sisters were conducted for multiple generations until the transgenic line was established.

Generating integration lines. Integrated lines were already available for C. elegans and C. briggae. In addition, myoEx34 in C. remanei appears to be a spontaneously integrated transgene22 (and our observations); thus this allele has been renamed mfs83 (see above). We generated integrated lines for three additional species: C. brenneri, C. portoensis, and C. tropicalis.

For C. brenneri, several lines appeared to have very high transmission rates of the extrachromosomal transgenes after culture on G418-containing NGM plates, possibly indicative of spontaneous integration. To identify integrants, we selected four to five transgenic gravid females (P0) from each putatively integrated line to separate NGM plates. From the P0 females that produced all or nearly all GFP F1 progeny, we crossed single L4 transgenic F1 males to single L4 transgenic F1 females on NGM plates without G418 and found two strains with 100% transmission of GFP in the F2 generation, suggesting that they likely carried spontaneously transposed transgenes. We conducted a second test to verify integration in these two apparently 100% transmission lines. If transmission is due to homozygosity rather than exceptionally high transmission of an extrachromosomal array, then crossing to WT should yield a heterozygous gfp/+ individual that should bear ~50% GFP cross progeny when crossed again to WT individuals. One strain fulfilled this criteria, and we renamed the transgene antIs2.

For C. portoensis, approximately 30 L4 extrachromosomal array carrying females (BRC313 (antEx-2[myo-2::GFP])) were irradiated using a Spectrolite UV crosslinker at a power setting of 250 (×100 µJ/cm²). Single irradiated P0 females were transferred to E. coli-seeded NGM plates and crossed to five WT L4 males. Each F1 GFP individual was crossed to two WT L4 individuals of the opposite sex. If the array was integrated onto an autosome (presumably as a heterozygote), then the predicted GFP:non-GFP ratio in the F2 generation would be approximately 50:50. For such broods, single GFP F2 males were crossed to single F2 GFP females (both putatively gfp/+). We then selected the F2 families producing approximately 75% GFP (both gfp/gfp and gfp/+), and 25% non-GFP progeny (+/+) in the F3 generation. To eliminate potential mutations caused by the integration procedure, we out-crossed single transgenic individuals (gfp/gfp or gfp/+), two to five WT worms of the opposite sex. This was repeated for six generations. Next, we crossed (inbred) single L4 GFP brothers to single L4 GFP sisters repeatedly for three to four generations. In the end, we obtained one homozygous line (100% GFP progeny; BRC585 (antIs7)).

For C. tropicalis, approximately 60 L4 extrachromosomal array carrying hermaphrodites (BRC493 (antEx-116[myo-2::mCherry])) were UV-irradiated as above, and then two to five irradiated P0 individuals were transferred to E. coli-seeded NGM plates. Subsequently, F1 mCherry individuals (n = 209) were singly to new plates. The F2 progeny of potential F1’s with heterozygous integrants should segregate approximately 75% mCherry, and from these candidate plates, two to four F2 potentially homozygous mCherry individuals were singly to new plates. Picking of candidate homozygous mCherry individuals was repeated in the F3 and subsequent generations, resulting in one homozygous line. This strain was then outcrossed to WT (JU1373) three times to obtain BRC555 (antIs1). While outcrossing, the integration status of antIs1 was confirmed by candidate heterozygous males producing approximately 50% mCherry cross progeny, in contrast to a higher rate of transmission indicative of an extrachromosomal array.

Generating mutants for C. tropicalis. To aid assessing cross progeny during crosses with the hermaphroditic species C. tropicalis, we conducted a standard F2 EMS screen in C. tropicalis (JU1373) for strains with clear recessive morphological mutant phenotypes. After screening ~2,000 genomes, we isolated five Ctr-dpy (ant23, 25, 26, 27, 32), four Ctr-unc (ant20, 22, 24, 28), one Ctr-egl (ant29), and one Ctr-rol (ant21) mutants with 100% penetrance and strong phenotypes. We outcrossed some of these strains 3-3 times back to JU1373 to remove mutations before choosing the 3x outcrossed Ctr-dpy (ant27) strain BRC461 for testing cross progeny. Backcrossing revealed that Ctr-unc’s (ant20, 28) and Ctr-dpy (ant32) are on the X-chromosome.

Testing for skew of integrated transgenes. For the hermaphroditic species, single homozygous transgene (e.g., gfp/gfp) L4 males were crossed to WT L4 hermaphrodites. Their heterozygous F1 L4 males were crossed singly to two or three L4 Dpy or Unc hermaphrodite testers. Each set of parents was transferred together to new OP50-seeded NGM plates every day until they stopped producing F2 cross progeny. All F2 cross progeny were counted for transgenic (Green/GFP or Red/mCherry) and non-transgenic (non-Green or non-Red/mCherry) females and males at the L4 or adult stage; the Dpy or Unc self progeny were not counted. We used the
Alternately, if \( mI_{s10} \) binomial tests for departure from these ratios. Some segregating with \( nullo-X \); “anti” was 1) transgene chromosome segregating with \( X \) and 2) \( \text{wt} \) chromosome most discriminatory cases are the study, there was a bimodal pattern of brood sizes with all broods having either brood size to distinguish \( XX \) and \( XO \) individuals. \( XX \) have many progeny whereas \( XO \) have none or few. In this testing sex-biased transmission of extrachromosomal transgenes. For the \( \text{C. elegans} \) strain \( \text{OH4460 (otEx2572)} \), we crossed single transgene carrying males to two or three \( dpy-11 \ unc-42 \) mutant hermaphrodites and then scored all nonDpyUnc individuals for transgenic (\( \text{Green} \)) and non-transgenic (\( \text{non-Green} \)) females and males at the \( L4 \) or young adult stage. For two other \( \text{C. elegans} \) strains, \( \text{BRC148 (unc-119(ed3); antEx50)} \) and \( \text{BRC153 (unc-119(ed3); antEx51)} \), we crossed transgene carrying males to three \( unc-119(ed3) \) hermaphrodites. The \( F1 \) cross progeny inheriting \( \text{antEx50} \) or \( \text{antEx51} \) carry \( Cb-unc-119(+) \), and thus, were non-Unc. The numbers of non-Unc males to non-Unc hermaphrodites in the \( F1 \) generation were counted. We did not consider Unc progeny because we could not distinguish between self and cross progeny.

For the gonochoristic species, we tested \( \text{C. breneri} \), \( \text{C. nigoni} \), \( \text{C. portoensis} \), and \( \text{C. sp. 33} \). Because many of the transgenic lines had poor fecundity, possibly due to inbreeding of the original \( \text{WT} \) strain, we selected the healthier transgenic lineages. We crossed single \( P0 \) males carrying an extrachromosomal array to \( \text{WT} \) females. Next, we crossed single \( F1 \) transgenic males to one or two \( \text{WT} \) virgin female testers. All \( F2 \) progeny at the \( L4 \) or adult stage were counted for transgenic (\( \text{Green/GFP} \)) and non-transgenic (\( \text{non-Green/non-GFP} \)) females and males.

All of the transgenic \( \text{C. breneri} \) strains were extremely unhealthy (brood sizes often <20), and thus, we decided to backcross to a second wild isolate \( \text{JU1397} \). We backcrossed each transgenic line seven to 12 times resulting in strains that produced larger broods (Supplementary Table S1). We used these strains to test for skew.

Testing for skew in \( \text{fem-3} \) and \( \text{her-1} \) oocytes. For the \( \text{fem-3} \) oocyte skew experiments, the test individuals were pseudo-females with the following genotype: \( unc-24 \ fem-3 IV; mls10(+) \text{+}/V; \text{XO} \). To simplify obtaining individuals of this genotype, we chose to construct two intermediate strains (\( \text{BRC174 and BRC237} \)) as follows. First, from strain \( \text{CB3843 (+fem-3+/+ unc-24 + dpy-20 IV)} \), we picked \( \text{F1 Unc-nonDpy} \) individuals. We then identified and kept an \( unc-24 \ fem-3 IV; mls10(+) \text{+}/2 \) males (strain \( \text{BRC174} \)) based on it being \( \text{Unc} \) and segregating three classes of individuals: fertile \( \text{Unc} \), sterile \( \text{Unc} \) pseudo-females (seen as an accumulation of oocytes in gonads), and fertile \( \text{UncDpy} \). Next, we crossed an \( F2 unc-24 \ fem-3 IV; mls10(+) \text{+}/2 \) males (strain \( \text{BRC174} \)) to \( \text{WT} \) females to test for skew. The \( F3 \) cross progeny males (\( unc-24 \ fem-3 IV; mls10(+) \text{+}/2 \text{F1} \)) were then crossed to \( \text{unc-24 dpy-20} \) hermaphrodites to yield \( F4 unc-24 \ fem-3 IV; mls10(+) \text{+}/2 \text{F2} \) males based on the \( \text{Unc} \) non-Dpy phenotype. Subsequently, bright GFP Unc \( F5 \) individuals were picked and the \( unc-24 \ fem-3 IV; mls10(+) \text{+}/2 \text{F3} \) hermaphrodites that produced \( \text{F4} \text{+} \text{F5} \) females were counted for transgenic (\( \text{Green/GFP} \)) and non-transgenic (\( \text{non-Green/non-GFP} \)) females and males.

For the \( \text{her-1} \) experiments, the test individuals were \( \text{XX} \) mothers (both type 2 and 3) because they had large brood sizes and also because they never produced RFP males; such crosses were discarded.

Testing for skew in \( \text{her-1} \) sperm. We selfed the \( dpy-11 \ her-1 \ mls10(+) \) hermaphrodites (above) and scored the brood sizes of single, cloned, strong GFP individuals, which were likely \( gfp/gfp \). Of the self-progeny, the most discriminatory cases are the \( gfp/gfp \) homozygotes, which can be determined by intensity for \( mls10 \) (see Supplementary Figure S5). We further verified that all progeny (when present) expressed bright GFP. We used brood size to distinguish \( \text{XX} \) and \( \text{XO} \) individuals. \( \text{XX} \) have many progeny whereas \( \text{XO} \) have none or few. In this study, there was a bimodal pattern of brood sizes with all broods having either >40 or <30 progeny, which we scored as \( \text{XX} \) and \( \text{XO} \), respectively.

If the \( mls10 \) transgene segregated randomly, then the expected ratio of \( \text{XO to XX} \) progeny would be 2:1. Alternatively, if \( mls10 \) segregated away from the \( \text{X} \) at approximately the same rate as for normal males (\( \sim 4:1 \) for \( \text{nullo-X}:X \)), then the expected ratio of progeny would be \( \sim 5:1 \) (see Supplementary Figure S5). Thus, we conducted binomial tests for departure from these ratios.

Transmission bias ratio (TBR). We defined the TBR as the ratio of the sex-normalized “preferred” to “anti” gamete combinations. “Preferred” was 1) \( \text{wt} \) chromosome segregating with \( X \) and 2) transgene chromosome segregating with \( \text{nullo-X} \); “anti” was 1) transgene chromosome segregating with \( X \) and 2) \( \text{wt} \) chromosome.
seggregating with nullo-X. For crosses with integrated arrays, “preferred” corresponded to non-transgenic daughters and transgenic sons while “anti” corresponded to transgenic daughters and non-transgenic sons. For the fem-3 and her-1 oocyte crosses, there were six viable offspring genotype classes (of eight possible) of which three corresponded to the “preferred” gamete combinations and three to the “anti” (see Supplementary Figure S4). For crosses with extrachromosomal arrays, “preferred” was extrachromosomal transgene segregating with nullo-X (i.e., transgenic sons) and “anti” was extrachromosomal transgene segregating with X (i.e., transgenic daughters). Non-transgenic progeny were excluded in the calculation because not inheriting an extrachromosomal array could be due to either standard chromosome segregation or spontaneous loss. Because some species have female-biased sex ratios, we normalized gamete combinations to its percent within each respective sex.

**Insertion size and transmission bias.** For the comparison between WHR10 and PD4793, we estimated insertion sizes as previously described39 (Supplementary Table S2). For the interspecies comparisons, we could not use the same set of reference primers for DNA qPCR due to sequence divergence among the species. Therefore, we identified a putative highly conserved element, LG2_6028, using the iHCE software package40 (Supplementary Text S2 and Supplementary Table S2). We then conducted genomic DNA qPCR assays on bla41 (plasmid marker) and LG2_6028 for six strains from four species (C. elegans, C. briggsae, C. tropicalis, and C. brenneri). Additional details are in Supplementary Text S2.

**Statistical analysis.** Statistical analyses were conducted in R (version 3.4.0)42. Some figures were plotted using ggplot243. For the skew tests in which both sex and transgene status were scored, we used the Chi-squared test (chisq.test function in R) with the null hypothesis being equal segregation of transgenes between hermaphrodites (or females) and males (i.e., 50% in both sexes for integrated arrays and a constant but equal number for extrachromosomal arrays). For the fem-3 oocyte, her-1 oocyte, and two extrachromosomal line crosses, we used the binomial test (binom.test function in R) to test for departure from equality between the preferred and anti categories. For her-1 sperm, we tested against departure from expected progeny ratios with (5:1) and without the binomial test (binom.test function in R) to test for departure from equality between the preferred and anti oocyte, and two extrachromosomal line crosses, we used the binomial test (binom.test function in R) to test for departure from equality between the preferred and anti categories. For her-1 sperm, we tested against departure from expected progeny ratios with (5:1) and without skew (2:1, see above). Confidence intervals were calculated using the multinomialCI function44 in R. For the correlation tests between the TBRs and estimated insertion sizes (after adjusting for genome size) we used the Spearman’s rank correlation test.

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Author Contributions
T.S.L. and J.W. conceived and designed the experiments and wrote the manuscript. T.S.L., F.-J.Y., and C.-Y.K. generated transgenic strains. T.C.C. conducted qPCR assays. All authors contributed to data collection. All authors reviewed the manuscript.

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