Specific Downregulation of Spermatogenesis Genes Targeted by 22G RNAs in Hybrid Sterile Males Associated with an X-Chromosome Introgression

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ABSTRACT

Hybrid incompatibility (HI) prevents gene flow between species, thus lying at the heart of speciation genetics. One of the most common HIs is male sterility. Two superficially contradictory observations exist for hybrid male sterility. First, an introgression on the X chromosome is more likely to produce male sterility than on autosomes (so-called large-X theory); second, spermatogenesis genes are enriched on the autosomes but depleted on the X chromosome (demasculization of X chromosome). Analysis of gene expression in Drosophila hybrids suggests a genetic interaction between X chromosome and autosomes that is essential for male fertility. However, the prevalence of such an interaction and its underlying mechanism remain largely unknown. Here we examine the interaction in nematode species by contrasting the expression of both coding genes and transposable elements (TEs) between hybrid sterile males and its parental nematode males. We use two lines of hybrid sterile males each carrying an independent introgression fragment from C. briggsae X chromosome in an otherwise C. nigoni background, which demonstrate similar defects in spermatogenesis. We observe a similar pattern of downregulated genes that are specific for spermatogenesis between the two hybrids. Importantly, the downregulated genes caused by the X chromosome introgressions show a significant enrichment on the autosomes, supporting an epistatic interaction between the X chromosome and autosomes. We investigate the underlying mechanism of the interaction by measuring small RNAs and find a subset of 22G RNAs specifically targeting the downregulated spermatogenesis genes are significantly upregulated in hybrids, suggesting that perturbation of small RNA-mediated regulation may contribute to the X-autosome interaction.
INTRODUCTION

Hybrid incompatibility (HI) refers to any measurable reduction in fitness commonly seen in interspecific hybrids. One of the most extreme types of HI is hybrid male sterility that may block gene flow between species. However, genetic and molecular mechanisms underlying an observed HI vary dramatically between species, making it necessary to dissect HI across taxa to achieve a global view of genetic or genomic conflict in species hybrid that leads to HI. It is widely observed that heterogametic hybrid progeny are more likely to suffer HI than its homogametic siblings, which is dubbed Haldane’s rule (Turelli and Orr 1995; Schilthuizen et al. 2011). One explanation for this is the dominance theory, which proposes that if alleles causing HI are recessive and sex-linked, the heterogametic hybrid progeny will manifest full effects because of hemizygosity whereas homogametic hybrid progeny will not owing to compensation by a second copy of a wild-type allele (Turelli and Orr 2000). Dominance theory has gained wide support in genetic studies of hybrid sterility in both animal and plant species (Masly et al. 2006; Yang et al. 2012).

However, expression and genetic analyses have revealed seemingly opposite roles of the X chromosome in hybrid male sterility. For example, genes expressed in the germline are enriched on the autosomes but depleted on the X chromosome during spermatogenesis in both Drosophila and Caenorhabditis species (Reinke et al. 2004; Sturgill et al. 2007; Ortiz et al. 2014; Vicoso and Bachtrog 2015). Moreover, histone markers indicative of active gene transcription are enriched on autosomes but depleted on the X chromosome in C. elegans male germline, whereas an opposite pattern is observed for repressive histone markers (Kelly et al. 2002; Schaner and Kelly 2006). These observations support the hypothesis of X demasculization or sexual antagonism and X inactivation (SAXI) (Wu and Xu 2003). Nevertheless, the X chromosome is found to play a disproportionately larger role than autosomes in development of hybrid male sterility (Masly and Presgraves 2007; Bi et al. 2015). One possible explanation for this discrepancy is the presence of a genetic interaction between the X chromosome and autosomes to maintain the correct expression ratios between the X
chromosome and the autosomes (hereafter termed as X:A imbalance), which is essential for spermatogenesis (Wu and Xu 2003). The clearest way to test for such an interaction would be to use hybrid sterile males carrying a defined introgression from another species in an otherwise isogenic genetic background. Indeed, expression analyses of testis genes between hybrid D. simulans males that carry a fertile or sterile introgression revealed that autosomal genes were more likely to be misexpressed than those on the X chromosome (Lu et al. 2010). These hybrid males differ only in a small region of the X chromosome containing the Ods-site homeobox (OdsH) locus of hybrid sterility (Ting et al. 1998; Sun et al. 2004), minimizing the complications associated with heterogeneous genetic background as is the case in F1 hybrid. The study supports a genetic interaction between X and autosomes, but how the interaction is maintained and whether a similar interaction is present in other taxa are largely unknown.

The control of transposable elements (TEs) is another important factor contributing to HIs (Maheshwari and Barbash 2011). TEs may become aberrantly activated when introduced into a new host that lacks a specific control mechanism, leading to a widespread invasion of the host genome and hybrid sterility or lethality. For example, TE mobilizations have been observed in hybrids between different marsupial species (Metcalfe et al. 2007) and between Drosophila species (Shpiz et al. 2014; Erwin et al. 2015) though whether the TE activation is the only cause of HI remains to be determined. Two HI genes Hmr and Lhr genetically interact to cause hybrid lethality between D. melanogaster and D. simulans (Brideau et al. 2006). RNA-seq analyses revealed that Hmr and Lhr are required to repress transcription from satellite DNAs and many families of TEs in its native host (Satyaki et al. 2014). One possible cause of aberrant TE expression in hybrids is altered expression of Piwi-interacting small RNAs (piRNAs), a class of small RNAs that interacts with the Piwi family of Argonaute proteins to control the expression of TEs in the germline (Di Giacomo et al. 2013). This is because the piRNA population in a host rapidly adapts to the TE content through generation of new piRNA clusters, allowing de novo production of piRNA and other type of small RNAs for silencing of the invading TE (Shpiz et al. 2014; Senti et al. 2015). This rapid divergence can be seen clearly
through the phenomenon of hybrid dysgenesis, in which intraspecific crosses between different *Drosophila* lines with and without a particular TE produce sterile progeny due to a requirement for maternally deposited piRNAs for silencing a paternally derived TE (Brennecke et al. 2008). Consistent with this, *Drosophila* interspecific hybrids phenocopy piRNA pathway mutants (Kelleher et al. 2012). Whether de-silencing of TEs is directly involved in interspecific hybrid sterility remains largely unexplored in other species. It is worth noting that in most of the studies on TE-mediated hybrid sterility or lethality, total RNAs were extracted from hybrid F1 animals with a largely heterozygous genetic background. Consequently, both sets of piRNA loci on autosomes are present, making it difficult to demonstrate directly that misexpression of piRNAs is actually causative for TE dysfunction, and indeed whether TE dysfunction itself is responsible for sterility. This also means that it is unclear whether imbalance between TEs and piRNAs would occur to the same extent in species without maternal deposition of piRNAs.

Isolation of a *C. briggsae* sister species called *C. nigoni*, previously known as *C. sp.9*, (Felix et al. 2014) opens the possibility of using nematode species to study interspecific HI. Results from preliminary crossing between the two support Haldane’s rule (Woodruff et al. 2010; Kozlowska et al. 2012). To facilitate the species pair as a model for isolation of HI loci, we have recently developed random labeling of *C. briggsae* chromosomes with GFP markers and mapped them into defined genomic regions (Bi et al. 2015). Multiple independent introgressions from *C. briggsae* X chromosome produce sterile males in *C. nigoni* background (Bi et al. 2015), but the molecular mechanism underlying the observed sterilities remains unclear. Here we examined the molecular mechanisms of hybrid male sterility between *C. briggsae* and *C. nigoni* mainly through genomic approaches.

**RESULTS**

*A C. nigoni draft genome was generated with approximately 20X Illumina long reads*

A current draft assembly of *C. nigoni* genome was produced using paired end Next Generation
Sequencing (NGS) data with an average read length of approximately 100 bps, making it difficult to generate sequence contigs with sufficient length and accuracy for the downstream analysis (Kumar et al. 2012). It is worth noting that the draft genome is further complicated by sequence contamination from *C. afra* (previously known as *C. sp.7*) to an unknown extent (Felix et al. 2014). In particular, the short reads are problematic for assembling contigs that are rich in repetitive sequences, which will prevent accurate annotation of these sequences, including transposable element (TE) and some small RNAs that are to be addressed in this study. We previously demonstrated that Illumina synthetic long reads are not only able to recover non-repetitive sequences, but also are capable of recovering most types of repetitive sequence except for those arranged in a long stretch of tandem repeats (Li et al. 2015). To facilitate comparative analysis of expression of small RNAs, TEs and protein-coding genes between wild type and hybrid strains, we produced approximately 20X coverage of Illumina long reads for *C. nigoni* as described previously for *C. elegans* genome assembly (Li et al. 2015). Most of the reads are around 10 Kbp in length with a minimum size of 1.5 Kbp.

A total of 6882 contigs were assembled using the long reads with an N50 size of 57 Kbp. We next aligned the contigs against *C. briggsae* (AF16) reference genome “cb4” (Ross et al. 2011) using LAST (Kielbasa et al. 2011) to locate the syntenic regions between the two, the output of which was used to construct *C. nigoni* pseudo chromosomes. A total of 119-Mbp sequences were anchored into the six pseudo-chromosomes while the remaining 22 Mbp were retained as so-called unassigned sequences. Taking account of residual heterozygosity, we estimated the total size of the *C. nigoni* genome was around 130-140 Mbp. We used the draft genome that is called “cn1” hereafter for the subsequent analysis. We also performed preliminary annotation for the “cn1” genome, including prediction of structural genes, which were refined with our RNA-seq data along with their homolog in *C. briggsae* (Fig. 1, Supplemental Fig. 1 & Supplemental Table 1). To facilitate the use of the draft genome by the research community, we established an online genome browser that allows sequence retrieval, visualization of *C. nigoni* gene models and gene expression as well as synteny between *C. briggsae* and *C. nigoni* genome (see Data Access). At present, the pseudo-chromosomal assembly
strategy used means that this genome will have a limited use in determining complex structural rearrangements between *C. nigoni* and *C. briggsae*. Future work incorporating long-read single molecule, real-time (SMRT) sequencing technology from Pacific Biosciences (PacBio) or sequencing of mate pair genomic libraries may help to resolve structural variations through generating an unbiased contiguity (Huddleston et al. 2014). Nevertheless the “cn1” genome assembly is expected to cover majority of DNA elements, including small RNAs, TEs and protein-coding genes that will be analyzed in this study.

**Replacement of different fragments of the *C. nigoni* X chromosome with homologous regions from *C. briggsae* results in defective spermatogenesis and sterility**

Our previous efforts in systematic introgression of GFP-linked *C. briggsae* genomic fragment into *C. nigoni* background produced multiple independent introgression lines that demonstrate complete male sterility (Bi et al. 2015). Here we focused on two male sterile lines each of which carries an independent, non-overlapping fragment from the *C. briggsae* X chromosome (Fig. 2). The females carrying either of the introgressions as a heterozygote are fertile. One of the lines, ZZY10330, carries a fragment from the right arm of the *C. briggsae* X chromosome that is approximately 5.1 Mb in size while the other line, ZZY10307, contains a fragment of approximately 7.6 Mb in size from the middle of the *C. briggsae* X chromosome (Fig. 2A to 2C). The introgression was achieved by repeated backcrossing of the GFP linked *C. briggsae* genomic fragment into *C. nigoni* for at least 15 generations (Fig. 2D) (Bi et al. 2015), meaning the two male sterile lines essentially carry pure *C. nigoni* background except for the introgression region. To verify the introgression boundaries defined by single-worm PCR (Yan et al. 2012), we performed NGS to determine the introgression boundaries as described (Bi et al. 2015). As expected, the introgression boundaries defined by NGS agree well with those by the PCR-based genotyping results in both strains (Fig. 2B). No extra *C. briggsae* fragments were found in the introgression lines (Supplemental Fig. 2), indicating the genetic background of the two lines is essentially *C. nigoni* JU1421 except for a *C. briggsae* genomic
fragment that is tightly linked to a chromosomal insertion of cbr-myo-2::GFP.

To investigate whether the observed sterilities are caused by defective spermatogenesis, we first examined germline morphology in the hybrid males. We found that, instead of the typical “U” shape as seen for *C. nigoni* male germline, the ZZY10307 and ZZY10330 male germlines are disorganized and appear to lack an obvious turn (Supplemental Fig. 3A & B). In addition, sperm cells were displaced anteriorly comparable to those observed in the F1 hybrids between the two species (Ting et al. 2014), suggesting a defect in spermatogenesis. We next performed mating tests to examine whether the sterility is produced by incapability of sperm transfer using MitoTracker stained sterile males that were mated with *C. nigoni* virgin female. We found sperm transfer was successful for both ZZY10307 and ZZY10330 (Supplemental Fig. 4A & B), indicating the sterility is not caused by the failure in sperm transfer during copulation. We finally evaluated the morphology of the sperm cells and their competences for activation in both hybrid and *C. nigoni* males. We found that sperm cells in the two lines showed similar phenotypes. For example, overall sperm cell shapes tend to be irregular and their sizes are smaller in both hybrids than in *C. nigoni* (Fig. 2 E to J). Residual bodies were frequently found in the sperm cells from ZZY10330, suggesting a defect in spermatogenesis (Liu et al. 2013). We next evaluated the activation potential of the sperm in both JU1421 and the sterile males. Unlike *C. elegans* sperm, which can be activated by either of Pronase, zinc, monensin or TEA (Liu et al. 2013), *C. nigoni* sperm cells can only be activated by Pronase (Fig. 2H and data not shown). Despite the defective size and shape, the sperm cells from both sterile males also showed a sign of activation (Fig. 2I & J), but they may not be competent enough for fertilizing *C. nigoni* oocytes. Taken together, the data demonstrate that the sterilities in both ZZY10307 and ZZY10300 are mainly caused by defective spermatogenesis.

Independent introgressions on X chromosome produce a similar pattern of downregulation in genes that are significantly enriched on autosomes

Previous expression studies using microarray, RNA-seq or antibody staining demonstrate a biased
expression of spermatogenesis genes between X chromosome and autosomes, namely most genes expressed in male germline are located on autosomes but depleted from X chromosome (Albritton et al. 2014; Ortiz et al. 2014). We asked whether replacement of part of the C. nigoni X chromosome with its homologous sequence from C. briggsae would disrupt the “imbalanced expression” between X chromosome and autosomes, which could be associated with the observed sterilities. To test this, we performed RNA-seq analysis to quantify the mRNA transcripts in the two sterile male lines and its parental males from C. nigoni and C. briggsae (Supplemental Table 2). We produced around 8 million reads for each sample with three replicates each. All RNA reads were mapped against annotated coding sequences in C. briggsae (AF16) (Harris et al. 2014). The reads were also de novo assembled into transcripts using Trinity (Haas et al. 2013) for the purpose of small RNA mapping as detailed below.

We identified a total of 574 and 922 genes that are significantly upregulated in the males of ZZY10307 and ZZY10330 respectively and 1242 and 1317 genes that are significantly downregulated respectively in the two compared with C. nigoni males (Supplemental Table 3). A total of 358 and 860 genes that are significantly up- and downregulated respectively in both hybrid lines compared with C. nigoni males (Fig. 3A). Interestingly, the two independent introgressions lead to downregulation of a highly overlapping set of genes (p<1e-5 by random sampling), whereas the upregulated genes do not show the similar overlapping pattern (Fig. 3B). Overall similarity of gene expression between the two sterile lines is much higher than that between either of the two and JU1421 (Fig. 3C and 3F to H). Intriguingly, we observed that the ratio of downregulated genes is disproportionately higher on autosomes than on the X chromosome where the introgression fragments are located (p<1e-17, Fisher’s exact test). For example, 96% of the shared downregulated genes are located on the autosomes and only 4% on the X chromosome (Fig. 3D to E) though X chromosome carries around 17% of C. nigoni protein coding genes. Notably, such enrichment could not be observed for upregulated genes in the hybrids (Fig. 3E).

To evaluate whether genes located within and outside of the introgressions demonstrated differential
patterns in misregulation, we divided the *C. briggsae* X chromosome into three distinct regions, i.e., introgression from ZZY10330 and ZZY10307 and the remaining region, which are about 5.1, 7.7 and 8.7 Mbp in size respectively. We counted the number of *C. briggsae* genes within each introgression as well as the number of *C. nigoni* genes orthologous to the *C. briggsae* genes within the remaining 8.7 Mbp (X_other in Fig. 3D & E) that showed misregulation in the hybrid background. We found no significant enrichment of upregulated genes located within and outside of both introgressions on the X chromosome (Figs. 3D & E). However, genes within introgression region of ZZY10330 does show a significant enrichment compared with those outside of both introgressions on the X chromosome (p=1e-4, Fisher’s exact test, Fig. 3E). A pairwise comparison of genes located within the introgression regions between a hybrid and its native parental strain reveals that the genes within the ZZY10307 introgression seem to be expressed in a more similar way as those in *C. briggsae* than in *C. nigoni*, while those located within the ZZY10330 introgression show an expression pattern that is comparable to both parents (Supplemental Fig. 5). In addition, correlations of expression between genes within introgression and those in both parental species are higher in ZZY10307 (with a correlation coefficient of 0.817 and 0.903 respectively) than in ZZY10330 (with a correlation coefficient of 0.744 and 0.717 respectively), indicating there is a higher level of gene misregulation in the introgression of ZZY10330 than in the introgression of ZZY10307. In summary, independent replacements of X chromosome fragment lead to preferential downregulation of autosome-linked genes associated with male sterilities, supporting an interaction between X chromosome and autosomes, which is consistent with the hypothesis of demacuslization of X chromosome.

**Only spermatogenesis genes are significantly enriched in the downregulated gene set in both hybrid sterile males**

Given the primary phenotype associated with the two introgressions is the male sterility that seems to be a product of defective spermatogenesis (Fig. 2 E-J), we investigated the relationship between the sterility and the misregulated genes shared in both sterile strains. We took advantage of the
classification of *C. elegans* male- and female-specific germline genes as well as *C. briggsae* sex-biased genes defined in recent studies using RNA-seq (Thomas et al. 2012; Albritton et al. 2014; Ortiz et al. 2014). We examined whether the categories of spermatogenesis or male specific genes were enriched in our misregulated gene list (Supplemental Table 4, see Methods). We performed enrichment analysis separately for the down- and upregulated genes shared between the two hybrids against spermatogenesis or male-specific genes as stated above. Only spermatogenesis genes were significantly enriched in our downregulated gene list (FDR<0.01) whereas oogenesis and gender neutral ones show no enrichment in the list (Fig. 4A, Supplemental Table 4). However, no significant enrichment was found in upregulated genes. Similar enrichment analysis was performed using sex-biased genes defined by previously (Thomas et al. 2012; Albritton et al. 2014). Again, male specific genes were significantly enriched in the downregulated gene list (Fig. 4B, Supplemental Table 4). In addition, a category of “Low Male” also showed a significant enrichment in the upregulated list albeit at a much lower scale compared with that in the downregulated list. Gene ontology (GO) analysis demonstrates that genes involved in regulation of cell shape or protein phosphorylation are significantly enriched (Fig. 4C), suggesting that these genes may control spermatogenesis through these pathways. Notably, genes upregulated in both hybrids do not show the enrichment on either spermatogenesis (Fig. 4A) or autosomes (Fig. 3E), suggesting that these genes are involved in the pathways other than spermatogenesis, but in those of general developmental pathway or physiology. Consistent with this, gene ontology analysis reveals that these genes are primarily involved in response to nutrient levels or extracellular stimuli (Fig. 4C). Similar enrichment patterns were observed when the misregulated genes were analyzed separately for the two hybrid strains (Supplemental Fig. 6). Intriguingly, a significant enrichment on the autosomes versus the X chromosome was observed for the downregulated but not the upregulated genes (Supplemental Fig. 7). Taken together, non-overlapping replacements of X chromosome fragment produce male sterilities through preferential downregulation of spermatogenesis genes that are mainly located on autosomes.
Few transposable elements show aberrant expression in the hybrid sterile males compared with *C. nigoni* males

Overexpression of TEs is frequently found in F1 hybrids, and is associated with male sterility (Rozhkov et al. 2013; Erwin et al. 2015). We wondered whether misregulation of TEs might be associated with the observed male sterility in the introgression lines despite the fact that they were subjected to backcrossing for at least 15 generations (Bi et al. 2015). To analyze TE expression, we compiled a TE list *de novo* from both *C. briggsae* and *C. nigoni* genomes (Supplemental Fig. 8, see Supplemental Methods). We defined a total of 247 and 319 families of TE in *C. briggsae* and *C. nigoni* respectively (Supplemental Table 5). Interestingly, *C. nigoni* genome seems to carry all the TE families found in *C. briggsae* genome, but also contain unique TE families. Read mapping against TEs was performed in the similar way as that for protein-coding ones but with modified parameters (see Methods and Supplemental Methods). Overall expression of TEs in both hybrids was similar to those in *C. nigoni* (Fig. 5). Out of all the measured TE families, we detected a significant elevation of expression for only one family in ZZY10307 males (cb_rnd-3_family-789(DNA/PiggyBac)) and two families in ZZY10330 males (cb_rnd-3_family-381 (DNA/Merlin) and c_sp9_rnd-4_family-428 (LINE/CR1)) compared with *C. nigoni* males; a single TE family (c_sp9_rnd-5_family-1528(LINE)) showed significant downregulation in ZZY10307 (Fold change >2 and p<0.01) (Fig. 5C & D, Supplemental Table 5). Since a single TE family can contain multiple members, we further investigated whether the member count contributed to the significant increase in TE expression. The average numbers of normalized reads between three replicates in the hybrids that mapped to the three families were 38, 34 and 25 respectively with average fewer than two copies per member, arguing against a significant role of overexpression of the three families in the observed male sterilities. This is based on empirical data on TE misregulation in other species. For example, despite the significantly elevated expression of TEs relative to *C. nigoni*, the expression of these TEs in our hybrids is much lower than the misregulated
TEs caused by two HI proteins, Hmr and Lhr in sterile hybrids between D. melanogaster and D. simulans, where over 50 families demonstrated overexpression in F1 hybrids with a much higher read count (up to 1 million reads) relative to both parents (Satyaki et al. 2014). It should be noted that due to difficulty in isolating germlines from hybrid sterile males, the whole animals were used for RNA extraction. Therefore, all expression assays do not distinguish between germline and somatic tissues.

Misregulation of 22G RNAs but not piRNAs in the hybrid sterile males

Piwi-interacting small RNAs (piRNAs) are key regulators of genome stability via regulation of TEs. piRNA mutants in both D. melanogaster and M. musculus show defects in spermatogenesis, leading to sterility, which is associated with altered TE expression (Cox et al. 1998; Brennecke et al. 2007; Carmell et al. 2007). Moreover, due to their critical role in TE regulation differences in piRNA content cause hybrid dysgenesis between different Drosophila species (Kelleher et al. 2012). In C. elegans piRNAs, also known as 21U-RNAs, associate with the Piwi protein PRG-1 and target transposable elements for silencing via the induction of 22G RNAs (Batista et al. 2008; Das et al. 2008). piRNAs also target both transgenes and endogenous loci for heritable silencing. Loss of the piRNA pathway leads to progressive sterility associated with increased repetitive element expression (Batista et al. 2008; Das et al. 2008; Simon et al. 2014). We therefore tested whether differences in piRNA expression between sterile hybrids and the parental lines might underlie defective spermatogenesis and male sterility in these lines.

Given the highly similar phenotypes in sperm as well as in the mRNA and TE expression patterns between the two sterile lines, we focused on small RNAs in one of the two sterile lines, ZZY10330. In C. briggsae, as in C. elegans, piRNAs are produced from individual loci located in clusters, with each individual piRNA associated with a specific upstream promoter identified by a highly conserved motif (the Ruby motif) (Ruby et al. 2006; de Wit et al. 2009; Shi et al. 2013). The C. briggsae piRNA clusters have previously been shown to reside on chromosome I and IV (de Wit et al. 2009;
Shi et al. 2013). We therefore aligned small RNAs from both C. nigoni parent males (hitherto wild-type) and sterile hybrid males to the C. nigoni scaffolds “cn1” as described above. In both cases we found strong enrichment of piRNA alignments within the regions of the C. nigoni genome corresponding to the C. briggsae piRNA loci, with little difference in the alignment positions, as would be expected given that the region of the C. briggsae genome inserted in the hybrid strain is outside of these loci. Although there was a reduction in the total number of overall reads from piRNA loci in the hybrid (p<2e-16, Mann-Whitney U-test), this difference was rather small (median 8 reads/million for C. nigoni versus 7 reads/million in ZZY10330) (Fig. 6A & B).

Although the reduction in piRNA expression was very small, it is conceivable that this might lead to differences in silencing of TEs downstream. To test this, we examined whether 22G RNAs mapping to the TEs compiled above were different between the hybrid and wild type males. Overall there was excellent correlation between wild type and sterile hybrids in terms of the reads mapped to consensus TE sequences (Supplemental Fig. 9), indicating that 22G RNAs mapping to TEs were mostly unchanged in hybrids. There were 5 TEs showing >4-fold difference in read count, but none of these corresponded to the TEs showing differential expression by RNA-seq (Supplemental Table 5). Moreover, there were no significant shifts in the levels of 22G RNAs mapping to different classes of TEs between sterile hybrid and wild type (p>0.1 for each, Wilcoxon paired test, Fig. 6C). Taken together with the absence of differences in the expression of TEs, this suggests that altered silencing of TEs by small RNAs is unlikely to explain the male sterility phenotype in the hybrid.

In addition to TEs, 22G RNAs also map to genic loci and can be generated from many different primary small RNA triggers, both piRNA dependent and piRNA independent (Gu et al. 2009; de Albuquerque et al. 2015; Phillips et al. 2015). We therefore considered whether 22G RNAs mapping to coding genes might show differences between the hybrid and the wild type. Given the sterility phenotype of the hybrids we focused on germline genes. In C. elegans, there are two major classes of 22G RNAs that are found in the germline, CSR-1, which bind to the Argonaute CSR-1 (Claycomb et al. 2009) or WAGO, which bind to WAGO-family Argonautes (Yigit et al. 2006; Gu et al. 2009).
CSR-1 dependent small RNAs activate target gene expression whereas WAGO dependent small RNAs generally reduce target gene expression (Seth et al. 2013; Wedeles et al. 2013). In order to classify genes in *C. nigoni* we used previously published *C. briggsae* annotations of CSR-1 RNAs based on a direct immunoprecipitation approach for CSR-1 (Tu et al. 2015) and homology mapping from *C. elegans* for WAGO (Gu et al. 2009). We then examined levels of 22G RNAs mapping antisense to these genes in hybrid and wild types (Supplemental Table 6).

Interestingly, putative spermatogenesis genes showed a clear increase in median levels of 22G RNAs in hybrid relative to wild type *C. nigoni* (p=1.5e-10, Wilcoxon Signed Rank test) (Fig. 6D). This increase did not correspond to increased 22G RNAs for either CSR-1 target genes, which showed no significant change (p>1e-3, Wilcoxon Signed Rank test) or WAGO-dependent genes, which showed a decrease in 22G RNA levels (p=2e-4, Wilcoxon Signed Rank test), suggesting that neither CSR-1 nor WAGO-1 target genes could explain the increased levels of 22G RNAs mapping to spermatogenesis genes. Consistent with this interpretation, neither CSR-1 nor WAGO-1 targeted spermatogenesis genes showed altered levels of 22G RNAs, with the changes predominantly affecting genes that were not targeted by either Argonaute (Fig. 6E). However, importantly, these annotations refer specifically to hermaphrodites. In *C. elegans* males, CSR-1 was shown to target spermatogenesis genes (Conine et al. 2013). We therefore examined 22G RNAs mapping to male-specific CSR-1 targets. Intriguingly, orthologs of spermatogenesis genes that were targets of CSR-1 in *C. elegans* males were significantly upregulated in hybrids relative to *C. nigoni* (p=1e-10, Wilcoxon Signed Rank test). This did not affect the *C. nigoni* orthologs of CSR-1 targets in *C. elegans* males that were not sperm genes, which were not significantly different between hybrids and *C. nigoni* (Fig. 6F).

We wondered whether the downregulation of spermatogenesis gene expression we observed (Fig. 4A) could be associated with the upregulation of 22G RNAs. We therefore subdivided spermatogenesis genes into upregulated and downregulated genes. This analysis showed that only downregulated spermatogenesis genes demonstrated increased 22G RNAs mapping to them whilst upregulated
spermatogenesis genes showed no significant change in 22G RNAs (Fig. 6G). Importantly, the misregulation of 22G RNAs that we observe is not directly due to cis-acting differences in the X chromosome. None of the spermatogenesis genes with 22G RNAs increased by at least 2-fold and showed significantly altered gene expression by RNAseq are found within the introgressed region (Supplemental Table 6), and these genes indeed were more likely to be found on autosomes than expected even given the biased distribution of spermatogenesis genes with altered expression by mRNA-seq (Odds ratio >15 p=0.02 Fisher’s exact Test; Supplemental Table 6). Thus both increased 22G RNAs and the decreased expression in spermatogenesis genes are likely to be an indirect response to the introgression region. We speculate that misregulation of 22G-RNAs caused by disruption of X chromosome integrity might be a general response to the disrupted X:A imbalance that occurs in hybrids.

**Differential cis-acting regulation of miR-237 between C. briggsae and C. nigoni is associated with sterility in hybrid males**

In order to further examine possible molecular differences between sterile hybrids and the parent C. nigoni strain, we compared expression of miRNAs between males of hybrid and those of C. nigoni. Although some miRNAs are remarkably highly conserved across species, the short sequence lengths and relatively straightforward requirements for processing give them the potential to evolve rapidly (Marco et al. 2013). Moreover, their important roles in development make differences in miRNAs attractive potential causes of species-specific differences Slack (Niwa and Slack 2007; Tang et al. 2010), although examples of clear roles for miRNAs in speciation have yet to be described. In order to identify possible miRNAs that were different between C. briggsae and C. nigoni we first identified homologues to previously identified C. briggsae miRNAs (de Wit et al. 2009; Shi et al. 2013; Kozomara and Griffiths-Jones 2014) in the wild type C. nigoni. The majority of miRNAs had identical sequences between C. briggsae and C. nigoni, and of the 6 for which we found single nucleotide polymorphisms in the mature miRNA sequence, none showed any change in the critical
“seed region” between nucleotides 2-7 (Bartel 2004) responsible for targeting (Supplemental Table 7). Thus miRNA sequences are highly conserved between the two species. We next analyzed the expression of these miRNAs in wild type males and ZZY10330 sterile males. The majority of miRNAs showed highly consistent expression between the two strains; however, one miRNA, miR-237, showed >5-fold reduced reads in the hybrid ZZY10330 relative to wild type C. nigoni. (Fig. 7A; Supplemental Table 8). Intriguingly, when we examined the expression of this miRNA in C. briggsae, we found that C. briggsae males also showed strongly reduced reads relative to its counterpart in C. nigoni (Fig. 7B). Notably, this miRNA is found within the introgression region (Supplemental Table 8). We therefore speculated that the difference in expression of miR-237 might reflect sequence changes in the locus. The mature miRNA shows only one nucleotide difference, outside of the seed region, which is unlikely to directly affect expression. Moreover, using RNAfold, we predicted that miRNA-precursors would form with highly comparable free energy (-39kJ/mol for C. nigoni versus -41kJ/mol for C. briggsae; Fig. 7 C & D). Thus this difference in expression is unlikely to come from differences in the processing of the precursor miRNA by Drosha or Dicer. However, the region immediately 5’ to the miR-237 sequence showed appreciable sequence divergence (Fig. 7E). Thus it is possible that a sequence difference in the promoter of mir-237 might drive the differential expression of this miRNA in both C. briggsae and C. nigoni background. The differential expression we observed may contribute to the sterility of hybrid lines. In this regard, it is interesting that miR-237 is expressed in the somatic gonads of C. elegans (Harris et al. 2014), so might contribute to reproductive capacity in C. nigoni; testing the consequences of reduced miR-237 expression in C. nigoni males will be an interesting area for further research. However, because the two introgression regions on the X chromosome that give rise to sterility are non-overlapping, the cis-acting differences in the mir-237 promoter will only be found in one of the sterile lines arguing that this cannot be the sole cause of sterility.
**DISCUSSION**

Despite intensive studies of hybrid incompatibility by expression analyses, its regulatory mechanism across species remains poorly understood. Most of these studies concentrate on F1 hybrids incompatibilities, including hybrid male sterility. However, the mechanisms underlying F1 HI can be different from that in animals carrying an introgression genomic segment in an otherwise pure genetic background, which provides an opportunity for identifying region-specific interaction that could be responsible for a given HI. Consistent with this, a recent study demonstrated that lethality of hybrid F1 males between *C. nigoni* and *C. briggsae* could be suppressed by *cbr-him-8* (Ragavapuram et al. 2015). However, the gene did not show misregulation in either of the two hybrids used in this study (Supplemental Table 2). In addition, most of the F1 hybrid males between the two species are atypically small and 37% of them have no gonad (Woodruff et al. 2010; Kozlowska et al. 2012); whereas nearly all of the hybrid males used in this study have gonads with obvious sperm cells (Bi et al. 2015). Here we investigated the molecular mechanism of hybrid male sterilities between the nematodes *C. briggsae* and *C. nigoni* through detailed analysis of expression changes of coding-genes, transposable elements and small RNAs. The sterilities are caused by independent introgressions of an X-chromosome linked fragment from *C. briggsae* in an otherwise *C. nigoni* background. Our results support a genetic interaction between X chromosome and autosomes and suggest a role of the endogenous RNAi pathway in mediating the interaction.

It is widely held that misregulation of piRNAs and their associated TEs is one of the major contributors to hybrid incompatibility especially in F1 hybrids (Erwin et al. 2015). However, this is apparently not the case for the hybrid that carries a homogenous genetic background except for an X-linked introgression fragment because we detected few changes in expression for both piRNAs and TEs in the two hybrid lines (Figs. 5 & 6). It is possible that misregulation of piRNAs and their associated TEs could nevertheless occur in the F1 hybrid males between *C. briggsae* and *C. nigoni*, but the hybrid might develop immunity against these TEs during the subsequent introgression steps.
through generation of 22G RNAs downstream of piRNA targeting (Ashe et al. 2012; Shirayama et al. 2012). It would be informative to investigate these possibilities by assessment of transcriptomes of both TEs and piRNAs in the F1 hybrid males between the two species.

An alternative explanation for the observed hybrid male sterilities is the disruption of a genetic interaction between the X chromosome and the autosomes (X:A imbalance) as demonstrated previously in *Drosophila* species (Lu et al. 2010). It has long been observed that genes expressed in the male germline tend to be enriched on the autosomes but depleted on the X chromosome (Albritton et al. 2014; Ortiz et al. 2014), suggesting an interaction between X chromosome and autosomes which is presumed to be essential for proper spermatogenesis. So far direct evidence in favor of this hypothesis is limited to *Drosophila* species (Lu et al. 2010). Our transcriptome data of the hybrid sterile males and its parental *C. nigoni* males demonstrate the interaction in *Caenorhabditis* species. The two sterile lines each carry an independent introgression fragment from the X chromosome of another nematode species *C. briggsae* but in an otherwise *C. nigoni* background achieved through multiple generations of backcrossing (Fig. 2D), which minimizes the complications of the mixed genomes as is the case of F1 hybrids. However, in contrast to the *Drosophila* study, in which the sterility was attributed to one specific locus on the X chromosomes (Lu et al. 2010), the two *C. briggsae* introgressions in the *C. nigoni* background are quite large non-overlapping fragments (Fig. 2A to C), yet the two sterile lines demonstrate very similar defects in germline and sperm, suggesting they suffer from similar defects in spermatogenesis. In addition, spermatogenesis genes are downregulated in the similar way between two sterile lines and these expression changes are significantly enriched in autosomal genes (Figs. 3 & 4). Altogether this suggests that a correct expression balance between the X-linked and autosomal genes is important to ensure proper spermatogenesis. The fact that distinct non-overlapping regions can produce the similar effect suggests that maintaining the correct balance requires broad regions of the X chromosome and is not limited to one or two “master regulators”. Instead, a perturbed X:A imbalance caused by disrupted X chromosome integrity involving multiple loci is a plausible
explanation for the sterilities as a result of introgression of the *C. briggsae* X into the *C. nigoni* background. Evidence of an interaction between X chromosome and autosome also comes from studies of dosage compensation (Meyer 2005). Whether the disrupted X chromosome perturbs dosage compensation in *cis* remains an open question.

How the interaction between autosomes and X chromosome is maintained forms a longstanding enigma. Our analysis of small RNAs provides insights into the interaction. In sterile hybrids, 22G RNAs targeted to spermatogenesis genes were specifically upregulated. Importantly, upregulated small RNAs corresponded well to downregulated spermatogenesis genes in the hybrids (Fig. 6D). Thus upregulation of 22G RNAs mapping to spermatogenesis genes may cause failure of spermatogenesis by reducing expression of spermatogenesis genes possibly through inducing epigenetic change in chromatin. In *C. elegans*, CSR-1 and WAGO type 22G RNAs are the two major classes of small RNAs enriched in germline. CSR-1 dependent small RNAs are enriched for spermatogenesis genes (Conine et al. 2013), thus we might expect that the spermatogenesis genes showing increased 22G RNAs are CSR-1 dependent. Indeed, the spermatogenesis genes we identified with increased 22G RNAs are enriched for genes homologous to CSR-1 targets previously annotated as enriched in *C. elegans* males (Fig. 6F) (Conine et al. 2013). However, CSR-1 functions to protect gene expression, in contrast to WAGO dependent 22G RNAs, which act to silence their targets (Seth et al. 2013; Wedeles et al. 2013). Thus we speculate that 22G RNAs targeted to spermatogenesis genes that bind to CSR-1 in wild type *C. nigoni* males and thus normally support gene expression become rerouted in sterile hybrids into the WAGO pathway and result in silencing.

As “non-self” DNA triggers stable 22G RNA mediated silencing of the “non-self” region in *C. elegans* (Ashe et al. 2012; Lee et al. 2012; Shirayama et al. 2012), it is interesting to speculate that the 22G RNA upregulation that occurs as a result of disrupted X:A imbalance might be a more widespread response to foreign sequences that leads to global misregulation of spermatogenesis genes. Definitive tests of such a possibility will require further experiments, for example pull-down assays using various types of Argonantes followed by RNA-seq for small RNAs, which are beyond...
the scope of this study.

Our improved *C. nigoni* draft genome “cn1” will become an invaluable resource for the community, which will not only facilitate the studies of HIs between the two nematode species, but also provide a foundation for comparative analysis of other fundamental biological processes. One of the fascinating biological processes is sex determination, which is known to have been subjected to fast evolution between *Caenorhabditis* species (Ellis and Lin 2014). Despite *C. briggsae* is the closest relatives of *C. nigoni* and their hybrid progeny are viable, the two nematode species have distinct modes of sex determination, with the former adopting a hermaphroditic mode of reproduction while the latter using a dioecious mode. Availability of an improved *C. nigoni* genome as well as transcriptomes produced in this study will facilitate the study of sex determination pathway and its modifiers.

**METHODS**

**Worm strains and maintenance**

All worms were maintained at 25 °C on NGM plates seeded with OP50 as food source. Strains AF16 and JU1421 were used as wild isolate of *C. briggsae* and *C. nigoni* respectively throughout the manuscript. Hybrid male sterile strains ZZY10307 and ZZY10330 were produced previously by backcrossing with GFP-labeled *C. briggsae* strains with JU1421 for at least 15 generations (Bi et al. 2015). The GFP-containing introgression fragments were maintained as a female heterozygote.

**Genotyping of introgression boundaries by NGS**

Introgression boundaries of ZZY10307 and ZZY10330 were genotyped as described (Bi et al. 2015). A total of 1.5M and 1.7M reads were generated for ZZY10307 and ZZY10330, respectively, which were mapped back to combined *C. briggsae* (“cb4”) and *C. nigoni* reference genomes (“cn1”). Read coverage was visualized with *C. briggsae* genome as shown in Fig. 2 and Supplemental Fig. 2.
C. nigoni genome assembly

Around 20X coverage of Illumina long reads were produced for C. nigoni (JU1421) and its genome assembly was produced as described (Li et al. 2015). C. nigoni pseudo chromosomes were produced using syntenic information with C. briggsae and preliminary genome annotation was performed using BAKER1 pipeline (Hoff et al. 2015) (see Supplemental Methods).

Compilation of TEs in C. nigoni and C. briggsae genome

To de novo annotate putative TEs, RepeatModeler (version 1.0.7, http://www.repeatmasker.org/RepeatModeler.html) was run with the genome sequences of C. nigoni (“cn1”, this study) and C. briggsae (“cb4”) as an input, respectively using default parameters. The consensus sequences of TEs produced were used as an input to RepeatMasker (version 4.0.4, http://www.repeatmasker.org) to annotate all the possible TE loci over the two genomes. RMblast (version 2.28) was used in the two annotation pipelines to align genome sequence against possible TEs.

Sequencing of mRNAs by NGS

300 young adult males were picked for mRNA extraction for each sample for C. briggsae (AF16), C. nigoni (JU1421), ZZY10307 and ZZY10330 with three replicates each. For collection of ZZY10307 and ZZY10330 young adult males, fertile females that carry the introgression were mated with JU1421 males. The male progeny that carry the introgression as judged by the expression of cbr-myo-2::GFP were picked under a fluorescence microscope for mRNA extraction. Initial attempts of using dissected germlines for RNA extraction were made, but dissecting of germline from hybrid sterile males turned out to be impractical as hydrostatic pressure in the sterile males is not comparable to parental males (data not shown). Total RNAs were extracted using TRIzol (Invitrogen) following the manufacturer's instructions. mRNA purification and fragmentation, cDNA synthesis, end repairing, adapters ligation, and DNA fragment enrichment were performed using Illumina’s TruSeq Stranded mRNA Library Preparation Kit based on the Kit's manual. Each library was bar-coded and sequenced to obtain pair-end (2×150 bps) reads using Illumina MiSeq. We obtained
approximately 8 million reads of high-quality score (>30 mean quality score) on average per sample.

**Sequencing of small RNAs by NGS**

RNA was extracted from *C. briggsae* (AF16), *C. nigoni* (JU1421) and hybrid (ZZY10330) males expressing GFP for small RNA sequencing. RNA was treated with 2ul RppH (NEB) to remove both 5’ caps and 5’ triphosphate and enable cloning of 22G RNAs. Libraries were constructed using the Illumina TruSeq small RNA library prep kit according to the manufacturer’s instructions.

**Data analysis of mRNA-seq reads**

All the mRNA-seq reads were mapped against *C. briggsae* genome (“cb4”) using CLC genomic workbench 8.0. To ensure the same mappability between *C. nigoni* and *C. briggsae* reads against *C. briggsae* ORFs, a relaxed mismatch cutoff was used for reads derived from *C. nigoni* and hybrids than for those derived from *C. briggsae* (see Supplemental Methods). The gene with fold change >2 and FDR<0.01 between *C. nigoni* (JU1421) and a hybrid was considered as a DEG (differentially expressed gene). For quantification of TE expression, we mapped all reads from mRNA-seq against the TE consensus sequences as described above using CLC genomic workbench 8.0, with a cutoff of at least 75% similarity and 75 % length. Total number of the mapped reads was counted for calculation of the differential expression between hybrids and JU1421 in the similar way as that for mRNAs.

**Enrichment analysis of misregulated genes**

Gene ontology (GO) terms for *C. elegans* and the *C. elegans/C. briggsae* ortholog table were fetched from WormBase (WS250). The gonad-specific gene categories were previously defined by the comparison of RNA-seq results between *C. elegans* male and female specific gonads (Ortiz et al. 2014). The *C. briggsae* GO term and gonad-specific gene categories were built from *C. elegans* data, based on the ortholog table. The sex-biased gene categories for *C. briggsae* were fetched from the comparison of RNA-seq results of *C. briggsae* male and female worms (Albritton et al. 2014). The enrichment analysis for GO terms, gonad-specific and sex-biased gene categories were carried out using ClusterProfiler (Yu et al. 2012) by hypergeometric test in R statistical computing environment.
(R Core Team 2016). The p value and FDR were calculated for each category, and the categories with FDR < 0.05 were reported.

**Data analysis on 22G RNAs**

To identify piRNAs, 21U-RNAs that did not map either exactly or with up to 1 mismatch to a *C. briggsae* miRNA were aligned to the *C. nigoni* genome assembly (“cn1”) produced in this study using bowtie, allowing 0 mismatches and reporting only one alignment per sequence. We visually examined plots of the density of piRNA loci across the *C. nigoni* genome using histograms made in R, verifying that piRNAs were strongly enriched at the syntenic regions to the *C. briggsae* piRNA clusters described previously (Shi et al. 2013). We then prepared plots of the piRNA clusters to compare the number of unique sequences per million total unique sequences. To assess the difference in the overall piRNA read counts we used the Wilcoxon test (unpaired), which makes no assumption about the distribution involved.

To analyze 22G RNAs we aligned 22G RNAs to either TEs or transcripts from *C. nigoni* (See supplemental files at http://158.182.16.70:8080/share). We then assessed the total counts for 22G RNAs mapping to individual genes or TEs and this total read count was normalized to the total number of mapped 22G RNAs. To compare the 22G RNAs mapping to different gene classes we first used annotations of 22G RNAs mapping to *C. briggsae* (Claycomb et al. 2009). This study reports CSR-1 targets in *C. briggsae* hermaphrodites by a direct immunoprecipitation approach, and also adds information about WAGO targets by homology. We supplemented these annotations with identification of CSR-1 targets in *C. elegans* males (Conine et al. 2013). We then used blast to identify the best matching *C. nigoni* transcript to the *C. briggsae* transcriptome, discarding genes that failed to map with an e-value of less than $10^{-4}$, for which we could not assign a homologue with a better than $10^{-4}$ e-value. The significance of up or downregulation of 22G RNAs was assessed using the Wilcoxon test (paired between individual loci), which does not make any assumption of the underlying distribution. Data analysis on miRNAs and piRNAs was outlined in Supplemental Methods.
DATA ACCESS

The Illumina Synthetic long-reads and the “cn1” genome assembly for C. nigoni have been submitted to the NCBI BioProject database (http://www.ncbi.nlm.nih.gov/bioproject/) under BioProject ID PRJNA306403 with accession number LWKT00000000 for the assembly and SRR3031106 for the Illumina long reads. The mRNA and small RNA sequencing data have been submitted to the NCBI Gene Expression Omnibus (GEO; http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE76306 and GSE75763, respectively. The DNA sequencing data for introgression boundary mapping of ZZY10307 and ZZY10330 have been submitted to Sequence Read Achieve (SRA; http://www.ncbi.nlm.nih.gov/sra) with accession number SRR3081358 and SRR3081363, respectively. A genome and synteny browser for this project is available at http://158.182.16.70:8080/.

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AUTHOR CONTRIBUTIONS

RL, XR, YB, VWSH performed introgressions and worm preparation. XR conducted mRNA sequencing, RL and ZZ performed data analysis on mRNA-seq and TEs. CH, AY and ZZhang provided long reads. TL, ZY and LM characterized sperm phenotypes. PS performed small RNA sequencing, data analysis and contributed to manuscript writing. ZZ conceived the project, provided
the reagents and wrote the manuscript.

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**FIGURE LEGENDS**

**Figure 1. Overview of *C. nigoni* genome.** Shown are densities of coding genes (blue), transposable element (TE) (red) and \(d_\text{KD}/d_\text{DS}\) ratio (cyan) of orthologous pairs between *C. nigoni* and *C. briggsae* over chromosome I (A) and X (B) in a window size of 100 Kbp.

**Figure 2. Characteristics of two hybrid male sterile lines.** (A-C) Confirmation of introgression boundaries by NGS for ZZY10307 (A) and ZZY10330 (C), each carrying an introgression derived from *C. briggsae* X chromosome in an otherwise *C. nigoni* background. Read coverage (y axis) is shown across *C. briggsae* X chromosome coordinate (x axis). Genotyping results by single-worm PCR are shown in the middle (B), with PCR positive and negative corresponding to the presence and absence of *C. briggsae*-specific amplification respectively. (D) Schematic diagram showing steps of generating two hybrid sterile lines. (E-J) Sperm morphology and activation in *C. nigoni* (JU1421) or hybrid males. (E-G) DIC micrographs showing sperm morphologies for JU1421, ZZY10307 and ZZY10330 respectively. (H-J) DIC micrographs showing sperm activations for JU1421, ZZY10307 and ZZY10330 respectively. Residual bodies and sign of activation in hybrids are indicated with an arrow head and arrow respectively. SM: sperm media.

**Figure 3. Expression profiling of coding genes in male hybrids and their parental males.** (A) Venn diagrams showing shared numbers of up- or downregulated genes between both hybrid males and *C. nigoni*. (B) The bootstrap test for the intersection of misregulated genes between two hybrids. The observed number of intersection from downregulated genes (860, blue line) is significantly higher than random, while the observed number of intersection from upregulated genes (358, red line)
is not higher than random. The bootstrap sampling was performed 100000 times with one-tailed p-value shown on top. Green: p<0.05; red: p>0.05. (C) A heat map showing hierarchical clustering of normalized expression for the shared up-(red) and downregulated (green) genes in (A) for each RNA-seq sample of *C. briggsae* (AF16), *C. nigoni* (JU1421), ZZY10330 and ZZY10307. Expression of each gene is normalized against the average of 12 samples. (D) Chromosomal distribution of up- (brown) and downregulated genes (green) as defined in (C). (E) Percentages of down- (green) or upregulated genes (red) on autosomes or X chromosome. Note the percentage of downregulated genes is significantly higher on the autosomes than on the X chromosome regardless the genes are located within or outside the introgressions; whereas no significant enrichment is found for upregulated genes. (F, G) A pairwise comparison of overall expression in males between ZZY10307 (F) or ZZY10330 (G) and *C. nigoni* respectively. Correlation coefficient (R) is indicated. (H) A pairwise comparison of overall expression in males between ZZY10307 and ZZY10330.

**Figure 4. Enrichment analysis of down- or upregulated genes against sex-biased genes.** Analysis was performed separately for downregulated and upregulated genes found in both hybrids against the gene categories defined by two previous studies (see text). Note a significant enrichment of spermatogenesis genes (A) or male specific genes (B) in the downregulated gene list. “Low male” category also shows a significant enrichment for upregulated genes, but its ratio is not comparable to those in downregulated list. (C) Gene ontology analysis of down- or upregulated genes shared by the two sterile lines.

**Figure 5. Comparison of transposable-element (TE) expression between hybrid and *C. nigoni* (JU1421) males.** (A and B) Boxplots showing the comparison of overall expression of TEs in the males at TE class level between ZZY10307 or ZZY10330 and JU1421 respectively. (C and D) Volcano plots showing the comparison of overall expression of TEs in the males at TE family level between ZZY10307 or ZZY10330 and JU1421 respectively. Note there are only one and two TE...
families showing a significantly higher expression (highlighted in red) in ZZY10337 (C) or ZZY10330 (D) than in JU1421 for respectively. LINE: long interspersed nucleotide elements; LTR: long terminal repeats; RC: rolling circle.

Figure 6. Comparison of expression of piRNAs and 22G RNAs between the males of ZZY10330 and C. nigoni (JU1421). (A) Comparison of read counts for piRNAs between C. nigoni males and the hybrid males (ZZY10330). (B) Distribution of piRNAs along the piRNA clusters on Chromosome I and Chromosome IV for C. nigoni (top) and Hybrid (bottom). The y axis shows the number of unique piRNA sequences per million in each genomic window of 100 Kbp. The x axis shows position along the chromosome in base pairs. (C) Boxplot showing differences in 22G RNAs mapping antisense to different classes of TEs between hybrid males and C. nigoni males. Box shows interquartile range with a line at the median and the whiskers show the furthest point <=1.5 times the IQ range. (D) Boxplot showing differences between hybrid and C. nigoni in 22G RNAs mapping antisense to C. nigoni genes categorized by annotations from C. briggsae (CSR-1) or C. elegans (other categories). (E) Breakdown of spermatogenesis genes from (D) into CSR-1 and WAGO targets. (F) Differences between hybrid and C. nigoni either in all male-specific CSR-1 targets as defined in C. elegans or the male-specific CSR-1 targets that overlap with spermatogenesis genes from (D). herm: hermaphrodite. Boxplot parameters as in (C). (G) Boxplot showing differences between hybrid and C. nigoni 22G RNAs mapping antisense to spermatogenesis genes found either upregulated or downregulated in hybrid males by mRNA-seq analysis (Fig. 3). Boxplot parameters as in (C).

Figure 7. miR-237 is differentially expressed between males of ZY10330 and C. nigoni. (A and B) Scatterplots showing miRNA read counts in C. nigoni (X axis) against either hybrid males (A) or C. briggsae (B). miR-237 mature (5p) and star (3p) are highlighted. (C and D) Secondary structure predictions of the miRNA precursors in C. briggsae and C. nigoni. Predictions were made by RNAfold. (E) Alignment of mir-237 genomic loci in C. briggsae and C. nigoni showing some
sequence divergence in the putative promoter region.
C. *briggsae* miR-237 precursor
mfe -41.5 kJ/mol

C. *nigoni* miR-237 precursor
mfe -39.7 kJ/mol
Specific downregulation of spermatogenesis genes targeted by 22G RNAs in hybrid sterile males associated with an X-Chromosome introgression

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