High-throughput sequencing reveals extraordinary fluidity of miRNA, piRNA, and siRNA pathways in nematodes

Zhen Shi, Taiowa A. Montgomery, Yan Qi, et al.

Genome Res. published online January 30, 2013
Access the most recent version at doi:10.1101/gr.149112.112
High-throughput sequencing reveals extraordinary fluidity of miRNA, piRNA, and siRNA pathways in nematodes

Zhen Shi,1,2 Taiowa A. Montgomery,1,3 Yan Qi,1,3 and Gary Ruvkun1,3,4

1Department of Molecular Biology, Massachusetts General Hospital, Boston, Massachusetts 02114, USA; 2Biological Sciences in Dental Medicine Program, Harvard Medical School, Boston, Massachusetts 02115, USA; 3Department of Genetics, Harvard Medical School, Boston, Massachusetts 02115, USA

The nematode Caenorhabditis elegans contains each of the broad classes of eukaryotic small RNAs, including microRNAs (miRNAs), endogenous small-interfering RNAs (endo-siRNAs), and piwi-interacting RNAs (piRNAs). To better understand the evolution of these regulatory RNAs, we deep-sequenced small RNAs from C. elegans and three closely related nematodes: C. briggsae, C. remanei, and C. brenneri. The results reveal a fluid landscape of small RNA pathways with essentially no conservation of individual sequences aside from a subset of miRNAs. We identified 54 miRNA families that are conserved in each of the four species, as well as numerous miRNAs that are species-specific or shared between only two or three species. Despite a lack of conservation of individual piRNAs and siRNAs, many of the features of each pathway are conserved between the different species. We show that the genomic distribution of 26G siRNAs and the tendency for piRNAs to cluster is conserved between C. briggsae and C. elegans. We also show that, in each species, 26G siRNAs trigger stage-specific secondary siRNA formation. piRNAs in each species also trigger secondary siRNA formation from targets containing up to three mismatches. Finally, we show that the production of male- and female-specific piRNAs is conserved in all four species, suggesting distinct roles for piRNAs in male and female germ lines.

[Supplemental material is available for this article.]
sequenced small RNAs from *C. elegans*, *C. briggsae*, *C. remanei*, and *C. brenneri*, and using a comparative genomics approach, we examined the conservation and evolution of each of the small RNA pathways. We identified hundreds of new miRNAs in *C. briggsae*, *C. remanei*, and *C. brenneri*, as well as thousands of piRNAs and each of the classes of endogenous siRNAs present in *C. elegans*. Although there is essentially no conservation of individual small RNAs among any of the four species aside from miRNAs and a very limited number of CSR-1 class siRNAs, many features such as their genomic distribution, expression patterns, and tendency for 26G siRNAs and piRNAs to trigger secondary 22G siRNA production are conserved. We also identified a new subfamily of RdRPs absent in *C. elegans* but conserved in each of the other three species. Finally, we discovered that production of distinct sex-specific populations of piRNAs is conserved across nematode species, suggesting different roles for piRNAs in male and female germlines.

**Results**

Expansion and diversification of Argonautes and RNA-dependent RNA polymerases

To examine small RNA evolution in nematodes, we subjected small RNAs from *C. elegans* and three other nematode species, *C. briggsae*, *C. remanei*, and *C. brenneri*, to high-throughput sequencing (Fig. 1A; Supplemental Table S1). Each of the four species is morphologically similar; however, their genomic sequences are highly divergent, with common ancestry ~110 million generations ago (Fig. 1B; Cutter et al. 2009). We constructed 18- to 28-nt small RNA

![Figure 1](Shi et al. Cold Spring Harbor Laboratory Press on February 1, 2013 - Published by genome.cshlp.org)
libraries from synchronized populations of young gravid adult hermaphrodites for the androdioecious (male/hermaphrodite) species \textit{C. elegans} and \textit{C. briggsae} and from mixed populations of adult males and females for the gonochoristic (male/female) species \textit{C. remanei} and \textit{C. brenneri}. For each species, we also sequenced small RNAs from embryos and young adult males. In each library $\sim$80\%–90\% of small RNAs were 21–23 nt and contained either a 5' U or 5' G, features indicative of piRNAs, miRNAs, and 22G siRNAs (Fig. 1A). Each of the libraries, but particularly male and embryo libraries, also had a small fraction of 26-nt 5' G-containing small RNAs (26G siRNAs) (Fig. 1A).

In \textit{C. elegans}, small RNAs can be classified by their genetic requirements, particularly their Argonaute binding partners, but for siRNAs, the RNA-directed RNA polymerase that produces them is also indicative of their function. To determine the potential for each class of \textit{C. elegans} small RNAs to be present in other nematodes and, thus, in our small RNA libraries, we identified all Argonautes and RdRPs in \textit{C. elegans}, \textit{C. briggsae}, \textit{C. remanei}, and \textit{C. brenneri} and examined their phylogenetic relatedness. For each of the 25 Argonaute proteins in \textit{C. elegans}, we identified at least one, but often multiple, likely orthologs in each of the three nematode species, suggesting that a similar repertoire of small RNA pathways exists in each species (Table 1; Supplemental Fig. S1).

There are four RdRPs in \textit{C. elegans}: RRF-3 functions in the production of 26G siRNAs (Gent et al. 2010), EGO-1 and RRF-1 have partially overlapping roles in 22G siRNA formation (Gu et al. 2009), and RRF-2, which is closely related to RRF-1, has an unknown role (Sijen et al. 2001). From a phylogenetic analysis of nematode RdRPs, we identified likely orthologs of the three RdRP subfamilies, RRF-1, RRF-3, and EGO-1 in each of the other three nematode species (Fig. 1C). We also identified a fourth subfamily of RdRPs, RRF-4, in each of the three nematode species but not in \textit{C. elegans} (Fig. 1C). By RT-PCR assays from mixed-stage animals, we could readily detect expression of \textit{Cbr-rrf-4} (CBG04006, \textit{Cre-rrf-4.1}/\textit{CRE23681}, and \textit{Cbn-rrf-4.1}/\textit{CBN18731}, but not \textit{Cre-rrf-4.2}/\textit{CRE24272} or \textit{Cbn-rrf-4.2}/\textit{CBN30970} (Fig. 1D), suggesting that at least one gene in the \textit{rrf-4} family is expressed in each species except \textit{C. elegans}.

The majority of \textit{Caenorhabditis} miRNA families within each species are conserved

miRNAs in \textit{C. elegans} are well-characterized (Lau et al. 2001; Lee and Ambros 2001; Grad et al. 2003; Lim et al. 2003; Ruby et al. 2006; Kato et al. 2009; Gerstein et al. 2010); however, miRNAs in \textit{C. briggsae} and \textit{C. remanei} have been only partially characterized (de Wit et al. 2009), and miRNAs in \textit{C. brenneri} have not been examined. To identify miRNAs in each of the four species, we used the miRDeep2 program (Mackowiak 2011; Friedlander et al. 2012). To conclusively call a miRNA, we required a candidate predicted by miRDeep2 to also be predicted by MiReNA (Mathelier and Carbone 2010) and/or contain a seed sequence (positions 2–7) conserved between \textit{Caenorhabditis} species. As validation of the approach, we identified the majority of annotated \textit{Caenorhabditis} miRNAs from our deep sequencing libraries (Supplemental Table S2). We identified 37 new miRNAs in \textit{C. briggsae}, 48 new miRNAs in \textit{C. remanei}, and 215 new miRNAs in \textit{C. brenneri} (Supplemental Table S3) but did not identify any new miRNAs in \textit{C. elegans}. We grouped all known and newly identified miRNAs into families, based on seed sequence. To date, there are 106 miRNA families annotated in \textit{C. elegans}, 84 in \textit{C. briggsae}, 85 in \textit{C. remanei}, and 87 in \textit{C. brenneri}, which constitute 176 distinct miRNA families (Fig. 2A,B; Supplemental Table S4). Fifty-four miRNA families are conserved in all four species, 61 families are present in three or more species, and most miRNA families within each species have homologs in at least two other species (Fig. 2C). Few (<5\%) miRNA families are conserved between only two or three nematode species (Fig. 2C). However, in each species, at least 20\% of miRNA families are unique, suggesting that miRNAs are born at relatively high rates (Fig. 2C).

Among conserved miRNA families, ~40\%–60\% have multiple members within a species, whereas <13\% of nonconserved families contain multiple members, suggesting that ancient miRNA families expand to confer robustness in gene regulatory networks (Fig. 2D). A striking example of this is the miR-35 family which has expanded to contain at least eight members in each species and as many as 27 members in \textit{C. brenneri} (Fig. 2A). The miR-35 family is one of the few miRNA families essential for development (Alvarez-Saavedra and Horvitz 2010). Each of the other families essential for development, including miR-51, miR-58, and let-7, also contain multiple (>4) members in each species (Fig. 2A).

Most miRNAs are processed from primary transcripts in sequential steps involving the ribonucleases Drosha and Dicer. However, some miRNAs are, instead, derived from short intronic hairpins called mirtrons during splicing, thereby bypassing Drosha cleavage (Okamura et al. 2007; Ruby et al. 2007). We found that out of the 15 \textit{C. elegans} annotated mirtrons (Chung et al. 2011), only

\begin{table}[h]
\centering
\caption{Argonautes and their associated small RNAs in \textit{Caenorhabditis}}
\begin{tabular}{|l|l|l|l|l|}
\hline
Class & Clade & Associated small RNAs & \textit{C. elegans} & \textit{C. briggsae} & \textit{C. remanei} & \textit{C. brenneri} \\
\hline
ALG-1 & AGO & miRNAs & 1 & 1 & 1 & 1 \\
ALG-2 & AGO & miRNAs & 1 & 1 & 3 & 1 \\
HPO-24 & AGO & Unknown & 1 & 1 & 1 & 2 \\
ALG-3/4 & AGO & Sperm 26G siRNAs & 2 & 1 & 1 & 1 \\
ERGO-1 & PIWI & Oocyte/embryo 26G siRNAs & 1 & 4 & 4 & 9 \\
PRG-1/2 & PIWI & piRNAs/21U-RNAs & 2 & 1 & 4 & 1 \\
CSR-1 & WAGO & 22G siRNAs & 1 & 1 & 1 & 3 \\
RDE-1 & WAGO & Exogenous siRNAs, miR-243 & 1 & 1 & 2 & 2 \\
CO4F12.1 & WAGO & 22G siRNAs & 1 & 1 & 3 & 2 \\
WAGO-1-5 & WAGO & 22G siRNAs & 5 & 3 & 4 & 7 \\
WAGO-6-8 & WAGO & 22G siRNAs & 4 & 5 & 6 & 11 \\
NRDE/HRDE & WAGO & Nuclear 22G siRNAs & 5 & 4 & 7 & 6 \\
\hline
Total & & & 25 & 24 & 37 & 46 \\
\hline
\end{tabular}
\end{table}
Figure 2. Conservation of miRNAs in nematodes. (A) Table of conserved miRNAs classified by family. Seed sequences are positions 2–7, relative to the 5’ end of the miRNA. The number in each row represents the number of miRNAs in each family in each species. Shading indicates presence of at least one member of a family. (B) As in A, but nonconserved miRNAs. (C) Venn diagram shows the number of miRNA families and their overlap in each of the four nematode species. (D) The percentage of families having the indicated number of members is shown for conserved and nonconserved miRNAs in each species. (E) Table of mirtrons classified by the gene hosting the mirtron.

*Numbers in columns are total family members.

**N**umbers in columns are total family members.
miR-62, embedded in the third intron of ugt-50, is conserved in the other three nematodes (Fig. 2E). miR-62 has 100% sequence conservation in all four nematodes, and the conservation of the intron sequence itself is much higher than that of other ugt-50 introns (Supplemental Fig. S2). Although the role of miR-62 is unknown, the strong selective pressure to maintain it hints at an important function. In addition to miR-62, we identified one nonconserved mirtron in \textit{C. briggsae}, one in \textit{C. remanei}, and two present in \textit{C. briggsae}, \textit{C. remanei}, and \textit{C. brenneri} but not in \textit{C. elegans} (Fig. 2E).

\textbf{piRNAs} share common features in their biogenesis and mechanism of action

\textit{C. elegans} piRNAs (21U-RNAs) are 21 nt long and contain a 5’ U. Although the mechanism by which piRNAs are formed is unclear, they map downstream from conserved sequence motifs that may function as transcriptional promoters (Ruby et al. 2006; de Wit et al. 2009; Cecere et al. 2012). To identify piRNAs from \textit{C. briggsae}, \textit{C. remanei}, \textit{C. brenneri}, and \textit{C. elegans}, we filtered our data sets for small RNAs that were not identified in our miRNA analysis and that were 21 nt long and contained a 5’ U. Consistent with previous studies (Ruby et al. 2006; de Wit et al. 2009), we found that a GTTTC core motif was strongly overrepresented between −37 to −47 nt upstream of the starting T in candidate piRNA coding sequences from each species. The nucleotide composition and the length of spacer sequence between the large and small motifs were nearly identical in all species, consistent with previous findings in \textit{C. briggsae} and \textit{C. remanei} (Supplemental Fig. S3; de Wit et al. 2009).

Using a scoring matrix based on the consensus motif, spacer sequence length, and 5’ U features of piRNAs, we scanned each of the four genomes for putative piRNA loci. The numbers of both predicted and observed piRNA loci were similar between \textit{C. elegans}, with 17.6K predicted and 9.9K observed, and \textit{C. briggsae}, with 14.8K predicted and 7.5K observed (Fig. 3A). \textit{C. remanei} and \textit{C. brenneri} possess far more piRNA loci than \textit{C. elegans} and \textit{C. briggsae}. In \textit{C. remanei}, 33.8K piRNAs were predicted and 23.7K were observed, and in \textit{C. brenneri}, 54.8K piRNAs were predicted and 29.8K were observed (Fig. 3A). A saturation analysis indicates that we captured the majority of piRNAs produced at these particular developmental stages (Supplemental Fig. S4). The genomes of \textit{C. remanei} and \textit{C. brenneri} are ∼35% larger than those of \textit{C. elegans} and \textit{C. briggsae}; however, this alone does not account for the 2–3 times more piRNAs identified in \textit{C. remanei} and \textit{C. brenneri}. Interestingly, \textit{C. elegans} and \textit{C. briggsae} are both androdioecious (male/hermaphrodite) species, whereas \textit{C. remanei} and \textit{C. brenneri} are both gonochoristic (male/female). This suggests that species that mate at every generation require a more complex repertoire of piRNAs for genome surveillance.

Figure 3. Functional and genomic features of piRNAs are conserved. (A) The numbers of piRNA loci predicted based on regulatory motifs and the numbers identified from high-throughput sequencing. (B) The percentage of protein-coding genes in each genome that could be targeted by piRNAs if zero, one, two, or three mismatches are allowed. (C) Density of small RNAs within a 100-nt window centered on the predicted target sites of the top 20% most abundant piRNAs. (Blue) Small RNAs that are antisense to the predicted targets; (red) those that are sense to the targets. (D) Distribution of observed (red) and predicted (blue) piRNA loci per 100-kb window in \textit{C. elegans} (top) and \textit{C. briggsae} (bottom). There are two piRNA clusters on \textit{C. briggsae} chromosome IV: the 0 to 6.9 Mb region largely in synteny with the two \textit{C. elegans} piRNA clusters (highlighted in lines with arrows); and the 13.1 to 15.1 Mb cluster. In addition, \textit{C. briggsae} has a third piRNA cluster on chromosome I at position 9.9 to 11.3 Mb.
Although we identified nearly 70,000 total piRNAs in our deep sequencing data sets, not a single piRNA sequence was present in more than one species. We also assessed piRNA sequence conservation when one, two, or three mismatches were allowed. Even with this less stringent criterion, only 0%, 0.01%, and 0.1% of C. elegans piRNAs have potential homologs in C. briggsae allowing for one, two, and three mismatches, respectively.

To identify potential piRNA targets, we aligned piRNA sequences with all annotated protein-coding genes from each species. piRNA-target recognition is thought to be permissive to around three mismatches (Bagijn et al. 2012; Lee et al. 2012); thus, we did the analysis allowing for zero, one, two, or three mismatches (Fig. 3B). When up to three mismatches are allowed, ~30% of C. elegans and ~20% of C. briggsae genes are potential targets (Fig. 3B). Although C. remanei and C. brenneri contain a substantially larger repertoire of piRNAs, the proportions of genes with potential piRNA targets are similar to that of C. elegans (Fig. 3B).

C. elegans piRNAs can trigger the production of RdRP-dependent secondary siRNAs centered on and antisense to piRNA target sites (Bagijn et al. 2012; Lee et al. 2012). To determine if piRNAs trigger siRNA formation in the other three nematodes, we assessed both sense and antisense siRNA abundance at candidate piRNA target sites. When all observed piRNAs were included in the analysis, there was only a slight enrichment of siRNAs at predicted piRNA target sites (data not shown). However, when only the top 20% most abundant piRNAs were considered, we observed a substantial enrichment of siRNAs antisense, but not sense, to the predicted piRNA target sites in each species (Fig. 3C). Of these siRNAs, 70%–80% are 22G siRNAs.

These results suggest that, although individual piRNAs are not conserved, the mechanism in which they are formed and their propensity to trigger secondary siRNA formation are conserved (Bagijn et al. 2012; Lee et al. 2012).

C. elegans piRNAs are primarily derived from two broad clusters on chromosome IV (Fig. 3D; Ruby et al. 2006). We asked whether piRNA loci are also clustered in other species. We restricted our analysis to C. briggsae because C. remanei and C. brenneri DNA sequences have not yet been assembled into chromosomes. There is extensive conservation of chromosome organization and synteny between C. elegans and C. briggsae (Stein et al. 2003; Hillier et al. 2007). In C. briggsae, the syntenic regions of the two major C. elegans piRNA clusters on chromosome IV also produced high levels of piRNAs (Fig. 3D; Ruby et al. 2006; de Wit et al. 2009). The two regions that give rise to the C. elegans piRNA clusters on chromosome IV are rearranged in C. briggsae such that they are separated from one another by only ~1 Mb. Interestingly, the ~1-Mb region separating the two clusters also contains a high abundance of piRNA loci. Together, the region forms a continuous 6.9-Mb piRNA cluster. We also identified a second piRNA cluster on chromosome IV (13.1–15.1 Mb) and another on chromosome I (9.9–11.3 Mb) specific to C. briggsae (Fig. 3D). The regions that give rise to these two C. briggsae-specific piRNA clusters lack continuous synteny with C. elegans, as determined by pairwise alignments. Two C. briggsae piRNA clusters previously identified on chromosomes I (7.8–9.5 Mb) and III (0–0.3 Mb) were represented by only average numbers of reads in our libraries and may have been artifacts of low sequencing depth (de Wit et al. 2009). That piRNAs in both C. elegans and C. briggsae tend to cluster and that conservation of these clusters appears to depend on long regions of continuous synteny suggests that the clusters are important for the birth of new piRNAs.

Distinct populations of piRNAs in males and females

In C. elegans, differences in the upstream regulatory motif drives male- and female-specific piRNA expression (M Freeberg and J Kim, pers. comm.). In each of the four Caenorhabditis species we analyzed, we observed a highly significant positive correlation between piRNA populations in early embryos and adult hermaphrodites or adult hermaphrodites + males (Fig. 4A). In contrast, piRNA levels between adult males and adult hermaphrodites or adult females + males are only modestly correlated and show a biphasic pattern of distribution indicative of two distinct classes (Fig. 4B). In C. elegans, we identified 9307 distinct piRNAs in hermaphrodites and 6065 distinct piRNAs in males. Of these, 5044 were enriched greater than threefold in hermaphrodites and 3336 were enriched greater than threefold in males. Only 1493 piRNAs had similar expression levels in hermaphrodites and males (Fig. 4C). Each of the other species also had distinct sets of piRNAs that were enriched in either males or hermaphrodites/females + males (Fig. 4D), suggesting that the production of distinct classes of male and female piRNAs is conserved in nematodes. The upstream motifs and length of spacer between the large and small motifs are nearly identical between the two classes of piRNAs, although several positions in the large motif and surrounding sequence show a stronger bias for a particular nucleotide in one class relative to the other (Supplemental Fig. S5).

The genomic distributions of C. elegans hermaphrodite- and male-enriched piRNAs are similar, although hermaphrodite-enriched piRNAs are somewhat more biased toward the cluster on the right arm of chromosome IV than are male-enriched piRNAs (Fig. 4D; Kato et al. 2009). In contrast to C. elegans, all C. briggsae piRNAs enriched in hermaphrodites are clustered on chromosome IV 0–6.9 Mb, the region in synteny with C. elegans piRNA clusters. However, almost all C. briggsae male-enriched piRNAs are located in the chromosome IV 13.1–15.1 Mb cluster or the chromosome I 9.9–11.3 Mb cluster (Fig. 4D). The nonoverlapping genomic distributions of C. briggsae hermaphrodite- and male-enriched piRNAs not only suggest that they have distinct evolutionary trajectories but also points to the possibility that differences in chromosomal-scale epigenetic marks may contribute to the sex-biased expression of piRNAs.

Given that the majority of piRNAs are differentially expressed, we asked whether the hermaphrodite/female- and male-enriched piRNAs have different sets of target genes. Indeed, when we allowed for up to three mismatches, genes containing target sites for hermaphrodite- or male-enriched piRNAs are largely nonoverlapping in C. elegans (Fig. 4E). In microarray-based studies, ~5000 genes have been identified that are highly expressed in the germline, during either oogenesis or spermatogenesis (Reinke et al. 2000, 2004). We found that genes containing target sites for hermaphrodite- or male-enriched piRNAs were neither enriched nor depleted for germline or oogenesis genes. However, genes containing target sites for male-enriched piRNAs were significantly depleted of spermatogenesis genes (Fig. 4E). This suggests a selective pressure on male-enriched piRNAs to avoid targeting genes that are important for spermatogenesis. It is unclear why we do not observe a similar depletion of hermaphrodite-enriched piRNA target sites in genes important for oogenesis.

Conservation of 22G siRNA pathways

There are two distinct classes of 22G siRNAs in C. elegans which can be classified by their Argonaute binding partners. WAGO class 22G siRNAs silence certain protein-coding genes, transposons, pseu-
dogenes, and cryptic loci to protect genome integrity (Gu et al. 2009). CSR-1 class 22G siRNAs have a role in regulating chromosome segregation (Claycomb et al. 2009; van Wolfswinkel et al. 2009) and might provide a memory of self to prevent endogenous genes from being silenced by the piRNA surveillance pathway, although data supporting this is lacking (Lee et al. 2012; Shirayama et al. 2012).

Approximately 85% of CSR-1 targets in *C. elegans* have a 1:1 ortholog in at least one of the other three *Caenorhabditis* species we analyzed, whereas ~60% of the total coding genes in *C. elegans* have 1:1 ortholog in at least one other species. Similar to the average conservation of protein-coding genes, ~60% of WAGO targets have a 1:1 ortholog in at least one of the other species analyzed (Fig. 5A). Approximately 50%–70% of *C. elegans* CSR-1 target genes also produce siRNAs in orthologs in each of the other species (Fig. 5A). We also observed a modest but significant ($P < 10^{-11}$) positive correlation in 22G siRNA levels among orthologous CSR-1 target genes between *C. elegans* and each of the other species (Fig. 5B). In contrast, targeting by WAGO class siRNAs is more poorly conserved. Only ~20%–30% of WAGO targets in *C. elegans* produce siRNAs in one of the other species analyzed, and there is lower correlation in siRNA levels among orthologous WAGO target genes between species ($P = 0.002–0.1$) (Fig. 5A,C). Because 22G siRNAs are biochemically indistinguishable from one another, we cannot rule out the possibility that in *C. briggsae, C. remanei, and C. brenneri*, a fraction of *C. elegans* CSR-1 targets are, instead, WAGO targets and vice versa. Nonetheless, these results point to relatively high conservation of targeting by CSR-1 class siRNAs and poor conservation of targeting by WAGO class siRNAs. WAGO class siRNAs, which are likely produced through successive rounds of

Figure 4. Two distinct classes of piRNAs in each species. (A) Scatter plots display the levels of piRNAs in adult hermaphrodites or adult females+males (x-axis) and early embryos (y-axis). (B) Scatter plots display the levels of piRNAs in adult hermaphrodites or adult females+males (x-axis) and adult males (y-axis). (C) Venn diagrams show the number of piRNAs that are greater than threefold enriched in hermaphrodites/females+males (left) or males (right). piRNAs shown in the overlapping section have similar expression levels in hermaphrodites/females and males. (D) Distribution of hermaphrodite- or male-enriched piRNA loci per 100 kb in *C. elegans* and *C. briggsae*. (E) The bar diagram displays the percentage of *C. elegans* genes containing hermaphrodite- or male-enriched piRNA target sites that are expressed in the germline, during oogenesis or during spermatogenesis. Asterisks indicate that the difference is statistically significant ($P < 0.001$, $\chi^2$ test). The Venn diagram shows the numbers of *C. elegans* genes containing target sites for hermaphrodite- or male-enriched piRNAs in purple and green, respectively.
siRNA amplification, are more abundant than CSR-1 class siRNAs in *C. elegans* (Claycomb et al. 2009; Gu et al. 2009; Phillips et al. 2012). We observed that WAGO target genes, on average, produce higher levels of 22G siRNAs than CSR-1 target genes, regardless of whether or not the genes are conserved. Moreover, nonconserved WAGO targets tend to produce more siRNAs than conserved WAGO targets \((P < 0.0001)\), whereas CSR-1 targets tend to produce similar levels of siRNAs regardless of their conservation \((P = 0.13)\) (Fig. 5D).

Embryo and sperm class 26G siRNA pathways are conserved in *Caenorhabditis* species

There are two distinct classes of 26G siRNAs in *C. elegans*. One class is enriched in oocytes and embryos and associates with ERGO-1 (Han et al. 2009; Vasale et al. 2010; Fischer et al. 2011). The other class associates with ALG-3 and ALG-4 during spermatogenesis (Han et al. 2009; Conine et al. 2010). Both classes are thought to function by triggering 22G siRNA formation and subsequent silencing of target mRNAs. Because each class of 26G siRNAs is sex- and stage-specific, our embryo and adult male small RNA libraries are each enriched for one of the two classes.

From our *C. elegans* embryo small RNA library, we defined a set of 29 26G siRNA-yielding genes, which produced at least 10 26G siRNA reads per million total reads (RPM), 26 of which were previously annotated as ERGO-1 targets (Han et al. 2009; Vasale et al. 2010; Fischer et al. 2011). In addition, we identified 121 embryo 26G siRNA-yielding genes in *C. briggsae*, 71 in *C. remanei*, and 272 in *C. brenneri*. None of the 26G siRNA sequences are conserved among species. Although 2%–12% of embryo 26G siRNA-yielding genes are conserved (Fig. 6A), in each instance only one ortholog yields 26G siRNAs.

**Figure 5.** Conservation of CSR-1 and WAGO class 22G siRNA pathways. (A) The percentages of *C. elegans* CSR-1 or WAGO target genes that contain an ortholog in *C. briggsae*, *C. remanei*, or *C. brenneri*, as well as the percentage of orthologous genes that produce 22G siRNAs in each species, is shown. (B) Scatter plots display conserved CSR-1 target genes having at least one siRNA read as a function of the log2 22G siRNA reads in *C. elegans* versus each of the other species. Pearson’s \(r\) values and corresponding \(P\)-values for significant correlation are shown. Degree of freedom (df) = number of orthologs analyzed – 2 was used for \(P\)-value calculation. (C) Same as B, but conserved WAGO target genes are plotted. (D) Box plot displays the levels of 22G siRNAs generated from conserved and nonconserved CSR-1 and WAGO target genes in *C. elegans*.

Embryo and sperm class 26G siRNA pathways are conserved in *Caenorhabditis* species

There are two distinct classes of 26G siRNAs in *C. elegans*. One class is enriched in oocytes and embryos and associates with ERGO-1 (Han et al. 2009; Vasale et al. 2010; Fischer et al. 2011). The other class associates with ALG-3 and ALG-4 during spermatogenesis (Han et al. 2009; Conine et al. 2010). Both classes are thought to function by triggering 22G siRNA formation and subsequent silencing of target mRNAs. Because each class of 26G siRNAs is sex- and stage-specific, our embryo and adult male small RNA libraries are each enriched for one of the two classes.

From our *C. elegans* embryo small RNA library, we defined a set of 29 26G siRNA-yielding genes, which produced at least 10 26G siRNA reads per million total reads (RPM), 26 of which were previously annotated as ERGO-1 targets (Han et al. 2009; Vasale et al. 2010; Fischer et al. 2011). In addition, we identified 121 embryo 26G siRNA-yielding genes in *C. briggsae*, 71 in *C. remanei*, and 272 in *C. brenneri*. None of the 26G siRNA sequences are conserved among species. Although 2%–12% of embryo 26G siRNA-yielding genes are conserved (Fig. 6A), in each instance only one ortholog yields 26G siRNAs.
In contrast to the poor conservation of embryo class 26G siRNA targets, ~80% (~50% 1:1 orthologs) of *C. elegans* sperm class 26G target genes have a potential ortholog in *C. briggsae*, compared to ~70% (~60% 1:1 orthologs) of all coding genes. This suggests that sperm class 26G target genes are not particularly conserved or fast evolving. Despite average conservation (~70%) of sperm class 26G siRNA targets, we observed a disproportionately low tendency for more than one ortholog to produce 26G siRNAs. Only ~38% of *C. elegans* genes that produce sperm class 26G siRNAs are conserved and also produce >1 RPM 26G siRNAs in *C. briggsae* (Fig. 6B). In *C. remanei* and *C. brenneri*, <15% of *C. elegans* genes are conserved and also produce 26G siRNAs (Fig. 6B). These results indicate that distinct cohorts of genes are targeted by 26G siRNAs during spermatogenesis and embryogenesis in each of the different nematode species.

In all four nematode species, embryo class 26G siRNA target genes also produce 22G siRNAs (Fig. 6C). In each species, these secondary 22G siRNAs are enriched in embryos relative to adults (Fig. 6C). Similarly, sperm class 26G siRNA target genes produce relatively high levels of 22G siRNAs that are enriched in adult males relative to embryos or hermaphrodites/males-females in each species (Fig. 6C). Thus, 26G siRNAs trigger stage-specific production of 22G siRNAs across nematode species.

Although 26G siRNA targets do not appear to be conserved, we nonetheless asked whether embryo class 26G siRNA chromosomal distribution is conserved. In *C. elegans*, oocyte/embryo class 26G siRNAs tend to be generated from gene-poor arms of chromosomes (Fig. 6D; Vasale et al. 2010; Fischer et al. 2011). In *C. briggsae*, for which the DNA sequence has been assembled on chromosomes, embryo class 26G siRNAs also tend to derive from chromosome arms (Fig. 6D). In contrast, sperm class 26G siRNAs are more or less evenly distributed along each chromosome in both *C. elegans* and *C. briggsae* (Fig. 6D). Thus, despite very little conservation of 26G siRNAs among species, the genomic distribution is very similar.

Embryo and sperm class 26G siRNAs are produced from nearly completely nonoverlapping targets in *C. elegans* (Fig. 6E; Han et al. 2009). However, in each of the other three nematodes, we observed substantial overlap between the embryo and sperm class 26G siRNA-yielding genes (Fig. 6E). Furthermore, the distributions of 26G siRNAs on the common targets were very similar in embryos and males within each species (data not shown). The common target genes are nonconserved, and in *C. briggsae* they are located on the chromosomal arms. Thus, the features of 26G siRNA targets that produce siRNAs in both males and embryos are more similar to ERGO-1 class 26G siRNA targets than to ALG-3/4 26G siRNA targets. It is interesting to note that ergo-1 appears to have undergone extensive gene duplications in *C. briggsae*, *C. remanei*, and *C. brenneri* after diverging from their common ancestor (Supplemental Fig. S1; Table 1). It is plausible that ergo-1 paralogs ac-
required new expression patterns or functions that account for the relatively high overlap between sperm and embryo 26G siRNA targets in species other than *C. elegans*.

**Discussion**

Using a deep-sequencing comparative genomics approach, we identified ~25 million distinct small RNAs from gravid adults and developing embryos of the four nematode species *C. elegans*, *C. briggsae*, *C. remanei*, and *C. brenneri*. Our results indicate that each of the major classes of small RNAs is conserved across *Caenorhabditis* species, although the degree of conservation differs tremendously. miRNAs are by far the most highly conserved class of small RNAs based on sequence. CSR-1 class 22G siRNAs are typically not conserved at the sequence level, yet they derive from a largely overlapping panel of genes in each species, indicating that the features that route CSR-1 targets into the siRNA pathway are conserved. piRNAs, WAGO class 22G siRNAs, and each of the classes of 26G siRNAs are not individually conserved but share many similarities in their biogenesis and function among the different species.

Although piRNA sequences are not at all conserved, other features, including the upstream motifs (de Wit et al. 2009), genomic clustering, and the tendency to trigger secondary 22G siRNA production from target genes, are conserved. *C. elegans* piRNAs are thought to play a role in genome surveillance and protection against nonself nucleic acids (Bagijn et al. 2012; Lee et al. 2012; Shirayama et al. 2012). Our results suggest that the roles of piRNAs are conserved across nematodes. That there is essentially no overlap in piRNA sequences among different nematode species indicates that piRNAs themselves are under very little stabilizing selection. The sheer number of piRNAs (likely >15,000 in each species) would allow them to target most, if not all, genes if multiple mismatches are permissible and would, therefore, not require sequence conservation of individual piRNAs. Indeed, rapid evolution of piRNAs would be advantageous in an arms race against invading nucleic acids.

We found that gonochoristic species that mate at every generation possess much larger numbers of piRNAs than androdioecious species that primarily self-fertilize. In *Drosophila*, maternally deposited piRNAs can silence paternal transposons (Brennecke et al. 2008). Moreover, crosses between *Drosophila* strains that differ in the presence of a particular transposon result in sterile progeny only if the transposon is inherited from the paternal genome (Brennecke et al. 2008). Similarly, desilencing of paternal transposons was observed in interspecies *Arabidopsis* crosses (Josefsson et al. 2006). It is possible that gonochoristic nematodes require a larger repertoire of piRNAs than androdioecious nematodes in order to defend against the greater diversity of paternal transposons encountered during mating. Moreover, the gonochoristic nematodes also have nearly twice as many Argonautes (37 in *C. remanei* and 47 in *C. brenneri*) as the androdioecious nematodes (24 in *C. briggsae* and 25 in *C. elegans*) and have undergone extensive proliferation of the PIWI clade Argonautes ERGO and PRG (Table 1; Supplemental Fig. S1).

Most piRNAs are differentially expressed in hermaphrodites/females and males in each of the four species tested. Differences in the piRNA repertoire of *C. elegans* males and hermaphrodites may result from distinct upstream sequence motifs that drive sex-specific expression (M Freeberg and J Kim, pers. comm.). The male and female/hermaphrodite piRNAs target a largely nonoverlapping set of genes. Spermatogenesis genes are depleted of male-enriched piRNA targets, suggesting selective pressure on male-enriched piRNAs to avoid targeting genes that are highly expressed during spermatogenesis. It will be important to distinguish the specific roles of male and female piRNAs.

If the role of CSR-1 class siRNAs is to provide a memory of self as has been proposed (Bagijn et al. 2012; Lee et al. 2012; Shirayama et al. 2012), then one would expect targeting by CSR-1 to be highly conserved. This is, indeed, what we observed. Targeting by CSR-1 class 22G siRNAs does appear to be highly conserved despite the individual siRNAs lacking conservation. In contrast, WAGO class 22G siRNAs are thought to target aberrant and foreign RNAs, and targeting of annotated protein-coding genes by this class of small RNAs is more poorly conserved.

Embryo class 26G siRNAs primarily target recently evolved genes with very little sequence conservation, and thus, it is not surprising that their targets are completely distinct in each species. Sperm class 26G siRNAs target many conserved genes, yet orthologous genes often do not produce 26G siRNAs. In spite of this, 26G siRNAs share many features in common among different nematode species, including genomic distribution, low abundance, sex specificity, and the propensity to trigger stage-specific 22G siRNA production.

Through phylogenetic analysis, we identified a distinct subfamily of RdRPs, RRF-4, which is present in each of the *Caenorhabditis* species we analyzed except for *C. elegans*. We were unable to link a distinct class of siRNAs to RRF-4 from analysis of our small RNA libraries, likely because RRF-4-dependent siRNAs are biochemically indistinguishable from other classes of siRNAs present in these libraries. It will be important to dissect the role of RRF-4 once the necessary genetic resources are available to do so in *rrf-4*-containing species.

The evolutionary fluidity of the different classes of small RNAs and their effectors in *Caenorhabditis* underscores their importance in the regulation, surveillance, and evolution of the genome.

**Methods**

**Nematode strains**

Nematode strains used in this study: *C. elegans* N2, *C. briggsae* AF16, *C. remanei* PB4641, and *C. brenneri* PB2801. Worms were cultured with bacterial strain OP50 on modified nematode growth medium (Andersen et al. 2012) containing 1% agar and 0.7% agarose to prevent burrowing of *C. brenneri*. All strains were grown at 20°C.

**Phylogenetic analyses of Argonautes and RdRPs**

To obtain a comprehensive and nonredundant list of Argonaute proteins and RdRPs in *C. briggsae*, *C. remanei*, and *C. brenneri*, we did BLASTP searches in the National Center for Biotechnology Information (NCBI) database using the protein sequences of *C. elegans* alg-1 and *rrf-1* as queries for Argonautes and RdRPs, respectively. We obtained Argonaute orthologs with *E*-values <10−10 and query coverage >35%, and RdRP orthologs with *E*-values <10−11 and query coverage >70%. Using different Argonautes or RdRPs as queries did not affect the outcome. Multiple sequence alignments were then performed using CLUSTALW slow/accurate alignments with default settings (Larkin et al. 2007). We constructed unrooted trees with the outputs of multiple sequence alignments, using the PHYLIP DRAWTREE program.

**High-throughput sequencing and data analysis**

For the construction of adult hermaphrodite/male+female, male or embryo small RNA libraries, animals were grown at 20°C for 72–74 h post-L1 synchronization and harvested as day one gravid adult.
hermaphrodites for *C. elegans* and *C. briggsae* and mixed populations of adult males and females for *C. remanei* and *C. brenneri*. For male isolation, 150–200 day one adult males were hand-picked from the NGM plate. Embryos were harvested by bleach treatment of −15,000 gravid adults. Total RNA was isolated by dounce homogenization of worms in TRI Reagent, followed by chloroform extraction and isopropanol precipitation. Small RNA high-throughput sequencing libraries were prepared as described (Montgomery et al. 2012). Briefly, 18- to 28-nt small RNAs were size-selected and treated with 20 U Tobacco Acid Phosphatase (Epicenter) at 37°C for 2 h to digest 5’ tri- and diphosphates to monophosphates. Small RNAs were then ligated to the 3’ adapter using T4 RNA ligase 2 truncated (NEB) for 16 h at 16°C. 5’ ligations were done with T4 RNA ligase 1 (NEB) for 16 h at 16°C. Adapter-ligated RNAs were then reverse-transcribed and PCR-amplified using Illumina’s TruSeq RNA Indexing PCR primers. Small RNA amplicons were size-selected by gel purification and subjected to Illumina HiSeq sequencing. Small RNA sequences were parsed using a custom Python program to remove adapter sequences and then mapped to the corresponding nematode reference genome using Bowtie software (Langmead et al. 2009). For sequences mapping to *C. elegans* WAG0, CSR-1, ERGO-1, and ALG-3/4 class siRNAs, the total number of millions of mapped reads (i.e., reads per million) were calculated using the Welch’s *t*-test. For the comparison of 22G siRNA levels among orthologous CSR-1 or WAGO target genes between *C. elegans* and each of the other three species, Pearson’s *r* correlation and corresponding *P*-values were calculated using R. *P*-values for comparing 22G siRNA levels produced from *C. elegans* CSR-1 and WAGO target genes were calculated using the Welch’s *t*-test.

Data access

All high-throughput sequencing data have been submitted to the NCBI Gene Expression Omnibus (GEO) (http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE41461.

Acknowledgments

We thank the Ruvkun lab members for constructive suggestions and thoughtful critiques of the work. We thank Allison Billi, Mallory Freeberg, and John Kim for sharing unpublished information. We thank the Caenorhabditis Genetics Center (CGC) for worm strains. Z.S. is supported by an American Heart Association predoctoral fellowship. T.A. M. was supported by the Damon Runyon Cancer Research Foundation (DRG 2029-09) and is a Massachusetts General Hospital Executive Committee of Research Tosteson postdoctoral fellow. Y.Q. is an Ellison Medical Foundation/AFAR postdoctoral fellow supported by the Life Sciences Research Foundation. This work was supported by a grant from the NIH to G.R. (R01-GM44619).

Small RNA classification

*C. elegans* small RNAs were classified as described (Zhang et al. 2011). Briefly, published data sets were parsed for classifying targets of *C. elegans* WAG0, CSR-1, ERGO-1, and ALG-3/4 class siRNAs. WAGO class was defined as the siRNAs produced from 1112 genes depleted of siRNAs in WAGO-1,2, 3-1, and 4-1 (Gu et al. 2009). CSR-1 class was defined as siRNAs produced from the 4191 genes that yield siRNAs enriched in CSR-1 immunoprecipitates (Claycomb et al. 2009). ALG-3/4 (401 genes) class and ERGO-1 (75 genes) class were defined as 26G siRNAs enriched during spermatogenesis and oogenesis/embryogenesis, respectively (Han et al. 2009; Conine et al. 2010; Vasale et al. 2010). miRNAs were predicted using mirDeep2 with default settings (Mackowiak 2011; Friedlander et al. 2012). To identify new miRNAs with high confidence, we further required a candidate miRNA predicted by miRDeep2 to also be predicted by MiRENA (Mathelier and Carbone 2010) and/or possess a seed sequence conserved between *Caenorhabditis* species. piRNAs/21U-RNAs were predicted as described (Ruby et al. 2006), using a scoring matrix based on the consensus motif, spacer sequence length, and 5’ U features of piRNAs. piRNA saturation analysis was performed by taking a random subset of sequencing reads with increasing size and calculating the percentage of piRNAs identified from this sublibrary.

Gene orthology analyses

Orthologous genes were identified using InParanoid4, which allows both one-to-one and many-to-many orthology cases (O’Brien et al. 2005). For clarity, we only kept one-to-one orthologs to analyze the conservation of different classes of small RNA target genes.

Statistical analyses

For analysis of *C. elegans* genes containing hermaphrodite- or male-enriched piRNA target sites in Figure 4E, the χ^2 test (df = 1) was performed. For the comparison of 22G siRNA levels among orthologous CSR-1 or WAGO target genes between *C. elegans* and each of the other three species, Pearson’s *r* correlation and corresponding *P*-values were calculated using R. *P*-values for comparing 22G siRNA levels produced from *C. elegans* CSR-1 and WAGO target genes were calculated using the Welch’s *t*-test.

**References**


Friedlander JR, Lehrbach NJ, Sapetschnig A, Cutter AD, Dey A, Murray RL. 2009. Evolution of the *Caenorhabditis* family of genes containing hermaphrodite- or male-enriched piRNA target sites in *C. elegans*. The χ^2 test (df = 1) was performed. For the comparison of 22G siRNA levels among orthologous CSR-1 or WAGO target genes between *C. elegans* and each of the other three species, Pearson’s *r* correlation and corresponding *P*-values were calculated using R. *P*-values for comparing 22G siRNA levels produced from *C. elegans* CSR-1 and WAGO target genes were calculated using the Welch’s *t*-test.
at the first stage of an siRNA amplification pathway that targets recent gene duplications. PLoS Genet 7:e1002569.


Shi et al.

Received September 13, 2012; accepted in revised form December 6, 2012.