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Genomic basis of recombination suppression in the hybrid between *Caenorhabditis briggsae* and *C. nigoni*

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**ABSTRACT**

DNA recombination is required for effective segregation and diversification of genomes and for the successful completion of meiosis. Recent studies in various species hybrids have demonstrated a genetic link between DNA recombination and speciation. Consistent with this, we observed a striking suppression of recombination in the hybrids between two nematodes, the hermaphroditic *Caenorhabditis briggsae* and the gonochoristic *C. nigoni*. To unravel the molecular basis underlying the recombination suppression in their hybrids, we generated a *C. nigoni* genome with chromosome-level contiguity and produced an improved *C. briggsae* genome with resolved gaps up to 2.8 Mb. The genome alignment reveals not only high sequence divergences but also pervasive intra- and inter-chromosomal sequence re-arrangements between the two species, which are plausible culprits for the observed suppression. Comparison of recombination boundary sequences suggests that recombination in the hybrid requires extensive sequence homology, which is rarely seen between the two genomes. The new genomes and genomic libraries form invaluable resources for studying genome evolution, hybrid incompatibilities and sex evolution for this pair of model species.

**INTRODUCTION**

Genome stability is essential for maintaining species identity. Different populations of the same species may possess a certain level of sequence divergence, but in most cases they can still effectively exchange their genetic information through crossing and DNA recombination (1). However, this exchanges of genetic information between populations may be severely compromised or lost between species, a phenomenon called reproductive isolation, which ultimately leads to speciation. Reproductive isolation can take place before or after mating, which are referred to as pre- and post-zygotic hybrid incompatibility, respectively. The prevailing explanation for the post-zygotic hybrid incompatibility is that the differential divergence of genes between separate populations lead to disrupted interactions, resulting in hybrid sterility or death (2). A handful of such gene pairs have been identified whose interactions cause hybrid sterility or lethality (3,4).

Recent studies in hybrid mice support a genetic link between DNA recombination and speciation (5). For example, the hybrid sterility locus on chromosome X was found to regulate the meiotic recombination rate in mice. Recombination is required for effective segregation and diversification of the genomes during meiosis (6). Compromised recombination severely affects fertility. It is well established that chromosome rearrangements such as inversion or translocation inhibit DNA recombination (7,8). It is also possible that chromosome sequences demand substantial homology to allow chromosome pairing or crossover during meiosis. During chromosome pairing, the homolo-
ous copies of each chromosome find each other through an active search process that enables chromosomes to distinguish ‘self versus non-self’ and assume a side-by-side alignment (6). This pairing is essential for crossover formation and thus the successful completion of meiosis. However, the precise requirements for effective recombination remain elusive, especially in the case of hybrids.

We have recently characterized the post-zygotic hybrid incompatibilities by genome-wide introgression between two closely related nematode species, *C. briggsae* and *C. nigoni* (9,10), whose hybrids are partially viable (11). Backcrossing the *C. briggsae* X chromosome into an otherwise *C. nigoni* background frequently leads to hybrid male sterility. The gene *chr-him-8* was identified as a possible recessive maternal-effect suppressor of F1 hybrid male-specific lethality in crosses with *C. briggsae* as the mother (12), although these results remain controversial (13). To develop a molecular understanding of this male sterility, we previously generated a *C. nigoni* draft genome using Illumina synthetic long reads (SLRs) with a contig N50 size of 57 kb (14). We demonstrated that spermatogenesis genes were significantly downregulated in the sterile males compared to the wild type. Intriguingly, the down-regulated genes were targeted by a subset of 22G RNAs that were significantly upregulated in the sterile males compared with the wild-type males (15). To facilitate genome wide comparisons, we generated so called *C. nigoni* pseudo chromosomes by chaining the contigs together using the homology to *C. briggsae* genome. However, this approach could not identify genome rearrangements between the two species because this step forces the *C. nigoni* to have a similar syntenic block to that of *C. briggsae*, but does not improve the actual size of the contigs themselves. Therefore, *C. nigoni* genome is rather fragmented given its small size of contig N50.

Although the recombination frequency shows variations along chromosome arms and between autosome and sex chromosome in *C. briggsae* (16), the overall frequency does not exhibit a significant drop compared with other phylogenetic groups. We referred to recombination frequency drop as large chromosomal regions being refractory to recombination rather than the reduction of recombination frequency to less than one recombination event per meiosis. Surprisingly, the recombination frequency is severely compromised in the hybrids between the two species, compared with the intraspecific crosses in *C. briggsae*. In addition, the two species use different mating systems, with *C. briggsae* being androdioecious (coexistence of males and hermaphrodites) and *C. nigoni* being gonochoristic (11). The mechanism through which recombination becomes suppressed in the hybrids remains largely unknown. However, the widespread recombination suppression in these hybrids between *C. briggsae* and *C. nigoni* provides an excellent opportunity to unravel the sequence requirements for effective recombination. The key to addressing these questions relies on the existence of high quality of genome sequences for both parental species. Here we first produced an improved *C. briggsae* genome using SLRs and high-throughput chromosome conformation capture (Hi-C) data. We next generated a *C. nigoni* genome assembly with chromosome-level contiguity using a combination of SLRs, Hi-C, Nanopore sequencing and mate-pair sequencing of the ends from both fosmid and BAC genomic libraries. Alignment between these two high-quality genomes allowed us to pinpoint the molecular bases of recombination suppression in the hybrids between *C. briggsae* and *C. nigoni*.

**MATERIALS AND METHODS**

**Strains, maintenance and backcrossing**

All strains were maintained at 25°C on NGM plates seeded with OP50 as food source. Strain AF16 and JU1421 were used as a wild isolate for *C. briggsae* and *C. nigoni*, respectively, for genomic DNA preparation. The strain JU1421 was subject to inbreeding for at least 70 generations by crossing a single young adult male with a single L4 female in three replicates before extraction of genomic DNA for preparation of sequencing or genomic libraries. Backcrossing was performed as described previously (9).

**Generation of SLRs for *C. briggsae* genome**

Genomic DNA was extracted from mixed-staged *C. briggsae* animals using PureLink® Genomic DNA Mini Kit (Invitrogen). The sequencing libraries were prepared according to Illumina’s protocol as described previously (14). 500 ng high quality genomic DNA was sheared in g-Tube by Covaris sonicator and went through end repair, dA-tailing and adaptor ligation. Products after ligation were size-selected for 8–10 kb by agarose gel followed by qPCR quantification. In the first stage, the sequencing and PCR errors of the reads were corrected and read depths were normalized across fragments. In the second stage, the String Graph Assembler (SGA) was used to construct a string graph based on the pre-processed reads, which was used to produce an initial set of contigs with the paired-end information from the short reads. In the third stage, the contigs were further scaffolded to resolve repeats and fill gaps. After examination of possible sequence errors and mis-assemblies, the assembled long-reads were generated as FASTQ files and used in the subsequent analysis.

**Construction of *C. nigoni* fosmid and BAC library**

For construction of fosmid genomic library, DNA extraction was performed by using PureLink® Genomic DNA Mini Kit (Invitrogen). Fosmid library was prepared according to the protocol of NxSeq® 40 kb Mate Pair Cloning Kit (Lucigen) with modifications. The genomic DNAs were end-repaired and run at 0.7% agarose gel at 40 V overnight. The end-repaired genomic DNAs were size-selected for fragments pf around 40 kb. Size-selected genomic DNAs were recovered by using GELase Enzyme and GELase 50X Buffer provided in CopyControl™ Fosmid Library Production Kit with pCC1FOS™ Vector (Epicentre). The recovered genomic DNAs were ligated into pNGS FOS Cloning Vector provided in NxSeq® 40 kb Mate Pair Cloning Kit (Lucigen), which contains NGS primer binding sites. The recombinant pNGS FOS fosmid clones were packaged into MaxPlax Lambda Packing Extract provided in CopyControl™ Fosmid Library Production Kit with pCC1FOS™
Vector (Epicentre). The packed fosmid clones were transfected into Replicator FOS strain provided in NsSeq® 40 kb Mate Pair Cloning Kit (Lucigen) and spread on LB agar plates with 12.5 μg/ml chloramphenicol. The clones were inoculated into 96-well plates with LB broth and 12.5 μg/ml chloramphenicol, cultured and then stored in −80°C. Construction of BAC genomic library was outsourced to Genome Resource Laboratory of Huazhong Agricultural University using the methods described previously (17).

Paired-end sequencing of *C. nigoni* fosmid or BAC library by NGS or nanopore sequencing

The fosmid clones were inoculated into 96-well plates with LB broth and 12.5 μg/ml chloramphenicol. They were cultured in 37°C for 16 h and pooled with a four-dimension strategy by columns, rows, plates with the same last digit and plates with the same second last digit. The fosmids were extracted from those pools and subject to CviQI enzyme digestion. The digested DNAs were run at 0.7% agarose gel at 85 V for 50 min and were size-selected for 8–9 kb. The recovered fosmid backbones carrying two ends of genomic DNA insertions were self-circularized and subject to PCR amplification with barcoded Illumina compatible primers (Supplementary Table S1). The PCR products were sequenced on MiSeq at 2 × 250 bp mode or Nanopore sequencing and the end sequences generated were used in the subsequent analysis. Mate-pair sequencing of BAC ends and its data analysis were performed as described previously (17).

*C. briggsae* genome repair and *de novo* genome assembly

15× SLRs were aligned against *C. briggsae* genome (‘cb4’) using BWA-MEM (18). If at least four SLR showed 100% sequence identity and spanned both ends of a gap with any number of ‘N’, the ‘Ns’ were replaced and the missing sequences within the gap were added with the sequences within the SLR. If the end of a gap could be aligned to two separate SLRs, the gap was filled as described previously (14). The process was reiterated for 12 times till no more ‘Ns’ could be replaced by any SLRs, which generated an assembly called ‘cb4 improve’. De novo genome assembly using SLRs was also performed basically as described (14). Scaffolding of the SLR-derived contigs using Hi-C data was described below to generate an assembly called ‘cb_SLR_Hi-C’.

Hi-C sequencing and data analysis

Hi-C sequencing was performed as described previously (19) with modifications. Briefly, approximately 10 000 synchronized L1 animals were fixed with 2% formaldehyde, snap frozen in liquid nitrogen for one hour, then quenched by glycine at room temperature followed by brief centrifugation. The remaining DNA ligation and Illumina sequencing library preparation steps were performed basically according to those described earlier (19), without size selection.

For Hi-C data analysis, a duplicate-free list of paired alignments of Hi-C reads against a given contig assembled with SLRs was generated using Juicer pipeline (20). The paired alignments were used as input for 3D *de novo* assembly (3D-DNA) pipeline (20) with the following parameters: haploid model, three rounds of mis-join correction and six expected chromosomes. To validate the genome assembly using the Hi-C reads, a *de novo* assembled *C. briggsae* genome was generated using ∼100× Hi-C data and 15× contigs generated using SLRs with an N50 of 61 kb. Comparison of chromosomal collinearity between the *de novo* generated *C. briggsae* assembly (cb_SLR_Hi-C) and the existing *C. briggsae* genome assembly (‘cb4’) was performed to provide confidence level of the newly generated genome with Hi-C data (Figure 2D).

Nanopore sequencing and data analysis

Genomic DNAs were prepared from mix-staged and starved *C. nigoni* animals using Qiagen DNeasy Blood & Tissue Kit followed by shearing to 8–10 kb using Covaris G-TUBE. Nanopore sequencing was performed with MinION using Ligation Sequencing Kit 1D from Oxford Nanopore Technologies according to the manufacturer’s description. Around 30× reads (assuming *C. nigoni* genome size of ∼130 Mb) were generated and were subject to self-correction using Canu assembler (22). The corrected reads were used for *de novo* genome assembly using Canu or scaffolding using SSpace-3.0 (21) with default parameters.

Analysis of synteny between *C. briggsae* and *C. nigoni*

One-to-one orthologous gene pairs were defined by mutual best BLASTP hits. The orthologous pairs were used as an input to MCScan package (23), to scan for syntenic block, which demanded at least ten contiguous orthologous genes as an seed. At most five gene rearrangements between the two seeds were allowed for extending the syntenic block.

*C. nigoni* gene prediction

The gene prediction were carried out using Marker 3.0 with default EViidenceModeler (EVM) parameters (24). Three *ab initio* gene prediction tools, the Augustus (25), SNAP (26) and GeneMark (27) were incorporated into the Maker pipeline. *De novo* mRNA assembly with Trinity and the pre-trained Augustus parameters were inherited from our previous study. The smallest size of a single exonie gene was set as ten amino acids for SNAP. Protein and CDS sequences from *C. briggsae*, *C. elegans*, *C. sinica* and *C. remanei* were used for sequence alignment using BLASTX or BLASTN, which were input for Maker pipeline.

Identification of orthologs and phylogenetic analysis

Protein coding genes among 10 *Caenorhabditis* species from Wormbase (WS252) were used to identify ortholog groups...
using OrthoMCL (v.2.0.9) (28) based on all-vs-all BLASTP with an E-value cutoff of $1 \times e^{-5}$. The aligned protein sequences of a total of 248 single-copy orthologs among the 10 species were used for phylogenetic analysis. The maximum-likelihood (ML) analysis was performed by using the bootstrap RAxML pipeline (29) implemented in ETI package (30). The divergent time estimation was calculated using RelTime method (31), with calibration by including two established pair-wise divergence time between species (32).

Gene family (Pfam domain) analysis

All protein coding genes from all 14 species were scanned against Pfam A domains using ‘hmmscan’ (hmmer.org). A domain with multiple copies in a protein was counted only once. The Pfam domain counts in C. nigoni were compared against C. briggsae. Significance of the difference between the two species were compared with Fisher’s exact test in R (V3.1.1). Only gene families with a $P$ value <0.05 were considered as significantly expanded/contracted. The gain or loss of a domains in the evolution history of all the 10 species were calculated using Count (33). The numbers of gene gain or loss on each branches of the phylogenetic tree were calculated using Wagner parsimony (34).

Analysis of conservation of the sequences flanking recombination sites

A total of 19 introgression boundaries were mapped with NGS previously (9). 12 out of the 19 boundaries were able to be mapped with up to 250-bp resolution (approximately the read size of MiSeq sequencer), which were used in this analysis. Mapping resolution for the remaining boundaries was too poor to be used for the same analysis. The poor resolution was mainly due to the low sequencing coverage ($<3x$) in the relevant regions.

The 6-kb flanking genomic regions centering on the 12 mapped boundaries were retrieved from C. briggsae genome. Corresponding 6-kb syntenic regions from C. nigoni were retrieved using the C. briggsae 6-kb sequences as a query. Pairwise Smith-Waterman (SW) alignment was implemented using SIMD Smith–Waterman Library (35) with default parameters. SW alignment scores were calculated using the following parameters, i.e. match = 2, mismatch = −1, gap open = −3, gap extend = −1.

For permutation analysis, one thousand 6-kb genomic regions without any gap were randomly sampled from C. briggsae chromosomes, which were used to retrieve their syntenic regions in C. nigoni genome. Pairwise alignment and its score calculation were performed in the similar way as that for the boundary sequences. Statistical difference in alignment scores between the boundary and permuted sequences was inferred with Wilcoxon ttest implemented in R package.

**RESULTS**

Recombination suppression in the hybrids between C. briggsae and C. nigoni

We previously performed backcrossing of 96 chromosomally integrated GFP markers in the C. briggsae genome into C. nigoni using the GFP as a selection marker (9). We observed an unusually large introgression size after 15 generations of backcrossing compared with that expected from intraspecific crossing in C.briggsae (16). The number of backcrossing generations was arbitrarily chosen based on Drosophila backcrossing data (36). To examine whether more backcrossing steps could reduce the introgression size, we chose a subset of these strains carrying a GFP-labeled introgression fragment for further backcrossing followed by genotyping with single-worm PCR. Unfortunately, most of these strains demonstrated little or no reduction in introgression size even after 50 generations of backcrossing (Figure 1). Many of the introgressions retained over one-fourth of the original chromosome, and approximately half of them contain nearly half of the entire tagged chromosome after 15 or more generations of backcrossing, indicating that recombination became more difficult for the marked DNA fragment in the hybrids than in its parental species, and the size of the introgression fragment could not be reduced simply by more backcrossing steps. However, the
mechanism behind the suppressed recombination was not clear.

Highly accurate SLRs allow gap filling in the existing *C. briggsae* genome

To unravel the molecular basis of the recombination suppression in the hybrids, we set out to improve the existing *C. briggsae* genome (‘cb4’), which contains 5997 gaps represented by an arbitrary number of ‘Ns’ on six chromosomes and 361 ‘unassigned contigs’ that were not anchored to any chromosome (16). We previously used the SLRs to generate a *de novo* assembly of the *C. elegans* genome (14). The SLRs were generated from next-generation sequencing (NGS) reads but with a much higher read length, i.e. ∼9.2 kb. We demonstrated that the reads were highly accurate and were able to recover most types of repetitive sequences except tandem repeats. Here, we generated approximately 15X *C. briggsae* (AF16) SLRs with an N50 size of ∼8.8 kb by a similar method to that used for *C. elegans* (14). As expected, the overall alignment of the *C. elegans*-derived SLRs (14) against the *C. elegans* genome (37) showed a very low ratio of discrepancies, whereas a similar alignment of the *C. briggsae*-derived SLRs against the *C. briggsae* genome (‘cb4’) revealed a much higher level of discrepancy (Figure 2A and B), indicating the presence of many more errors in the ‘cb4’ genome assembly than in the *C. elegans* genome. The N50 size of the contigs assembled with the SLRs was expected to be significantly improved compared with those obtained by *de novo* genome assembly using NGS reads only (38,39).

We previously demonstrated that the SLRs were not reliable in detecting extra inserted sequences that were incorrectly included in the *C. elegans* reference genome, but they were highly accurate at the nucleotide level and were fairly reliable in identifying missing sequences in this genome (14). We therefore decided to repair the *C. briggsae* genome in a similar manner to fill the gaps (i.e., those represented by ‘Ns’ in the ‘cb4’ assembly) or recover missing sequences using the SLRs. In total, we closed a total of 2304 out of the 5997 gaps by replacing a total of roughly 240 kbs on six chromosomes (Supplementary Table S2). We also partially closed gaps in ‘unassigned contigs’ by identifying and filling ‘self-mapped’ or ‘cross-mapped’ gaps as we did previously (14), which allowed extension of roughly 800-kb sequences. In total, we filled gaps by replacing or adding a total of 1040 kb of sequences across the ‘cb4’ genome using the SLRs (Figure 2C). We validated a subset of five cases in which the missing sequences were more 100 bps in length by PCR as we did previously (14). All the missing sequences were found to be real as judged by the expected sizes of the PCR products (data not shown). We next repaired the *C. briggsae* genome by anchoring a total of 124 out of 361 ‘unassigned’ contigs listed in the ‘cb4’ assembly with a cumulative size of ∼1.8 Mb back to individual chromosomes. This was achieved by alignment between ‘cb4’ assembly and the *de novo* assembly produced by a combination of SLRs and Hi-C data as described below. Seven out of the 124 were merged with our updated genome with DTools (https://jgi.doe.gov/data-and-tools/bbtools/) using the cutoff of 98% sequence identity (Supplementary Table S3). In
summary, we were able to update a total of 2.8 Mb of the
C. briggsae genome, including the replaced ‘Ns’ or the
recovered missing sequences and the ‘unassigned’ contigs
that were anchored back to chromosome (Table 1).

To independently validate the repaired sequences by
SLRs, we examined two cases that were located within an
existing gene model and caused gene model change after
sequence repairing (Supplementary Figure S1). To verify the
corrected gene models, we mapped the existing RNA-seq
sequence repairing (Supplementary Figure S1). To verify the
corrected gene models, we mapped the existing RNA-seq
data (15,40,41) against the revised sequence. We observed
perfect alignments between some RNA-seq reads and the
newly added sequences, indicating a missing exon and its
flanking sequences in the existing gene model.

Evaluation of Hi-C data for contig scaffolding

We previously produced a C. nigoni draft genome using
approximately 22X SLRs (15). As the SLRs could not fur-
ther extend a contig when they encounter tandem repeats,
the overall contig size of the assembly was relatively small
with an N50 size of 57 kb. To facilitate the molecular char-
acterization of the recombination suppression in the hy-
brids and empower the species pair as a model for speci-
cation genetics and sex evolution, we decided to generate a
C. nigoni genome assembly with chromosome-level contigu-
ity using a combination of data consisting of the previous
SLRs, Hi-C, Nanopore sequencing reads and the mate-pair
end-sequences of both fosmid and BAC genomic libraries.

To gain confidence of the Hi-C data for de novo genome
assembly, we first produced C. briggsae Hi-C data with
around 100X coverage. We then used these data to perform
de novo C. briggsae genome assembly by using the SLRs.
The resulting assembly was 108 Mb in size (Table 1), which
was comparable with the size of ‘cb4’. We called this assembly
‘cb_SLR_Hi-C’ (Figure 2C). The total size of contigs
that could be assembled into chromosome was only 95 Mb,
in contrast to the 105 Mb of sequences assigned to chromo-
somes in the ‘cb4’ assembly, leading to 13 Mb of contigs that
could not be assigned onto any chromosome (referred to as
‘unassigned’) (data not shown). This relatively large portion
of ‘unassigned contigs’ was likely due to the tandem repeats
present at their ends.

The alignment of the newly generated C. briggsae assem-
bly against ‘cb4’ revealed an excellent collinearity, albeit
with some obvious inversions on the chromosomes II, IV
and X (Figure 2D), suggesting potential errors in the ‘cb4’
assembl above. Importantly, the N50 size of the scaffolded con-
tigs increased from 57 kb to over 10 Mb after incorporation of
the Hi-C data (Figure 3B). These results indicate that Hi-
C data are highly reliable for scaffolding of SLR-derived
contigs. The comparison of ‘cb_SLR_Hi-C’ with ‘cb4’ al-
lowed a total of 124 ‘unassigned’ contigs with a combined
size of 1.8 Mb present in the ‘cb4’ assembly to be anchored
back to individual chromosomes (Supplementary Table S3).
As a result, we generated an improved C. briggsae genome
assembly called ‘cb4_improve’ after the incorporation of
both SLRs and the Hi-C data. It is worthy of note that the
‘cb4_improve’ retains the ‘cb4’ scaffolds, but with updated
sequences by SLRs and a subset of ‘unassigned’ contigs an-
chored back to individual chromosomes (see Materials and
Methods).

A high-quality of C. nigoni genome with chromosome-level of
contiguity

We next produced Hi-C data using C. nigoni synchronized
L1 animals (see Materials and Methods) with a genome
coverage of up to 2000× of its genome. Modeling analysis
showed that the contribution of the Hi-C data to the scaf-
foiling of the SLR-derived contigs reached a plateau with
approximately 50X Hi-C reads (Figure 3A), demonstrating
that an 100X coverage of Hi-C data is sufficient for scaffold-
ing purposes when used with SLR-derived contigs. Given
the relatively small proportion of the contigs that could be
assigned into individual chromosomes compared with that
using genetic mapping data in C. briggsae, i.e. 95 versus
105 Mb (16), to further improve scaffolding results, we pro-
duced roughly 15× fosmid and 10× BAC (Bacterial Artifi-

Table 1. Statistics of newly assembled C. nigoni genome (‘cn2’) and improved C. briggsae genome ‘cb4_improve’

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<th>Size (bp)*</th>
<th>Gene count</th>
<th>% repeat</th>
<th>C. nigoni (‘cn2’)</th>
<th>Size (bp)</th>
<th>Updated sequence (bp)#</th>
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<td>108 384 165</td>
<td>21 863 23</td>
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</table>

*including ‘Ns’ estimated with BAC or fosmid interval. # including replaced ‘Ns’, newly added sequences and ‘unassigned’ contigs anchored back to
chromosome. Number of the ‘Ns’ in some gaps seems overestimated as judged by the sequences present in the SLRs covering these gaps. LG, linkage
group. NA, not applicable.
to obtain six large contigs, each representing an individual chromosome. Despite the modest contribution of the fosmid end-sequences to the contig N50 size, the incorporation of the BAC end-sequences more than doubled the N50 size, from 7 to 17 Mb, which is essentially the chromosome size of the ‘cb4’ assembly (16). As an alternative genome assembly strategy, we performed scaffolding of the SLR-derived contigs first using the Nanopore sequencing data. This resulted in an increase in the N50 size from 57 kb to approximately 1 Mb. Further scaffolding with the fosmid and BAC end-sequences led to a linear increase in the N50 size, to approximately 1.5 and 3.2 Mb, respectively (Figure 3B). We also explored another strategy by performing de novo genome assembly first with the Nanopore sequencing reads followed by scaffolding using the fosmid and BAC end-sequences. Using this method, we were able to achieve an N50 size of approximately 1.8 Mb. Given the different scaffold sizes achieved with the various assembly strategies, we decided to use the C. nigoni genome assembly with the first strategy with a genome size of approximately 135 Mb (Table 1) for the subsequent analysis, which was termed as ‘cn2’. The assembly consisted of 124.0 Mb of chromosomal contigs and 11.8 Mb of small contigs that could not be assigned to any chromosome (Table 1). The total size was ∼27 Mb bigger than that of the C. briggsae genome (‘cb4’) (Table 1). To examine whether the size difference is contributed by repetitive sequence, we defined repetitive sequences in both genomes using RepeatMasker (repeatmasker.org). We found extra repetitive sequences in C. nigoni genome accounted for roughly 17 out of the 27 Mb genomic sequences, while the remaining 10 Mb genome sequences could not be explained by repetitive sequences. However, it is plausible that these regions may encode quite a few genes, but they
are missed in our gene list due to under-prediction as described below.

A total of 22,462 protein-coding genes were predicted in the newly assembled genome (Table 1), compared with the 21,863 C. briggsae genes annotated in Wormbase (WS252) (42). It should be noted that the genes were likely be under-predicted in C. nigoni. This was because the EVM parameters we used to finalize the gene model demanded a satisfaction score from all inputs, including ab initio gene prediction results from all independent gene-prediction algorithms, and the results from de novo assembled mRNAs as well as gene-model conservation across species (see Materials and Methods). Therefore, the predicted genes may only represent the highly reliable ones. However, there might be a substantial number of genes that were missed out in our prediction, for example, those highly divergent ones or those with simple gene structure. Comparison of one-to-one orthologs (mutual best hits) allowed us to infer a phylogenetic distance of between Caenorhabditis. Using an estimated divergence time of ~30 million years between C. briggsae and C. elegans (32) as a reference, we estimated the divergence time between C. briggsae and C. nigoni is ~3.7 million years (Supplementary Figure S2). This is roughly comparable with what was estimated previously between the two species using synonymous site divergence (43).

Chromosomes of C. briggsae and C. nigoni show overall collinearity but demonstrate substantial sequence divergences and pervasive rearrangements

To pinpoint the molecular basis of the recombination suppression, we first examined the overall DNA sequence conservation between the two species. We then evaluated the synteny between the two genomes using one-to-one orthologous genes. Specifically, we performed genome-wide sequence alignment between the two genomes (see Materials and Methods) and extracted the best hits as their corresponding genomic regions for visualization of genomic collinearity and arrangement. In principle, most of the repetitive sequences should be excluded from the orthologous regions because few of them could find a single best hit. The results show that overall genome sequences were alignable and are located within the corresponding chromosomal parts (Figure 4A and C). However, considering that their female hybrid progeny were fertile, the level of sequence divergence between the two genomes was surprisingly high. For example, the coding sequences of their one-to-one orthologs exhibited only around a 90% identity, and most of their introns and intergenic regions were not alignable (Figure 4A and data not shown).

To examine the genomic collinearity and rearrangement, we extracted all of the best hits between the two genomes that were >1 kb in size. We observed over 400 interchromosomal translocations across the genomes (Figure 4B). Many more translocations were found within a single chromosome than between chromosomes (data not shown). We postulated that the presence of these pervasive genomic rearrangements and widespread unalignable sequences prevents effective chromosome crossover, thus inhibiting recombination in the hybrids. Given that hybrid females between C. briggsae and C. nigoni are fertile, it is unexpected that the two genomes have undergone such a high level of sequence divergence and rearrangement.

Despite the apparent recombination suppression in the hybrids between C. briggsae and C. nigoni, we were able to produce homozygous viable introgressions representing ∼28% of the C. briggsae genome (9), indicating that over one quarter of the genome was interchangeable between the two species. To examine whether the inviability of the remaining C. briggsae genome in the C. nigoni background as homozygote was caused by gene rearrangements leading to abnormal gene loss or gain, we produced one-to-one orthologous gene pairs between C. briggsae and C. nigoni, as defined by the mutual best hits using BLASTP. This resulted in a total of 15,157 orthologous gene pairs (Supplementary Table S6). To contrast the genomic synteny between C. briggsae and C. nigoni and that between C. briggsae and C. elegans, we retrieved C. briggsae’s one-to-one orthologous gene pairs with C. elegans from WormBase (WS252). The pairwise comparisons between C. briggsae and C. nigoni revealed an excellent genomic synteny between the two species. In contrast, frequent intrachromosomal rearrangements were observed between C. briggsae and C. elegans (Figure 5). These results demonstrated that the large C. briggsae chromosome fragments that were not viable in the C. nigoni background were unlikely to be caused by major inversions or translocations that prevent chromosome alignment, but possibly by local rearrangements or sequence divergences that complicate the inheritance of orthologous gene pairs.

In addition to the one-to-one orthologs, many genes were found to have undergone substantial species-specific amplification or loss (Supplementary Table S7). Therefore, these could not be unambiguously assigned as orthologous pairs. Many of the C. nigoni-specific genes seemed to originate from transposable elements (Supplementary Figures S3A and S4). Others encoded putative proteins but with unknown functions. In contrast, many of the C. briggsae-specific genes encoded transcription factors or proteins containing domains of unknown function (Figures S3B and S4). It is not clear why these transcription factors underwent specific amplification in C. briggsae. Despite the species-specific amplification or loss of genes, the overall number of protein domains gained or lost was roughly comparable for the two species (Supplementary Figure S5). The C. nigoni genome contained a substantially higher proportion of repeats than the C. briggsae genome (Table 1). The difference in repeat content accounted for 57% of the differences in genome size difference between the two species.

Recombination in hybrids appears to demand an extended region of high sequence identity

Availability of precise collinearity of the genomic sequences between C. briggsae and C. nigoni permitted pinpointing of the detailed sequence requirements for recombination in the hybrids. We previously mapped the recombination boundaries for all of the homozygous introgressions using NGS with up to 100-bp resolution (9). We first focused on one particular introgression, ZZY10331, for which the NGS data had the highest coverage (~13×) (Figure 6A) to characterize the boundary sequences of recombination site. This introgression was mapped onto the right arm of
Figure 4. Genomic collinearity and rearrangement between *C. briggsae* and *C. nigoni*. (A) Diagram showing corresponding sequences defined by mutual best hits using LASTAL with a size cutoff of one Kb. Note the white regions are not alignable. Names of *C. briggsae* and *C. nigoni* chromosomes are labeled in black and red, respectively. (B) Diagram showing inter-chromosomal translocations that contain sequence over 1 kb in size between the two species. Chromosome names are similarly labeled as in (A). (C) Dotplot of corresponding chromosomes between *C. briggsae* (‘cb4 improve’, black) and *C. nigoni* (‘ca2’).

*C. briggsae* chromosome III, with its left and right boundaries extending to the middle and the very end of the chromosome, respectively. We retrieved the sequence of a 6-kb *C. briggsae* genomic fragment centered on the mapped recombination site of the left boundary. All of the sequence and syntenic information between *C. briggsae* and *C. nigoni* can be found at [http://158.182.16.70:8080/](http://158.182.16.70:8080/). Most parts of the sequences within the region were alignable against their *C. nigoni* syntenic regions, albeit with a 100-bp gap (Figure 6B). Surprisingly, in addition to the coding sequences of the gene present within the region, the intergenic sequences and introns were also alignable, which seems contradicting our prediction that most of these regions are not alignable. We next asked whether the intronic sequences were conserved between orthologous pair. To this end, we performed alignment of syntenic 6-kb regions between *C. briggsae* and *C. nigoni* that each contained an orthologue of *abce-1*, an extremely conserved and essential gene in *C. elegans*. This gene was found to be essential in protein translation and its depletion produced 100% embryonic lethality. Importantly, it was present as a single copy across metazoan genomes (10,44). As expected from our genome-wide sequence alignment, we observed excellent alignment only between exonic regions, but not between intronic regions (Figure 6C), demonstrating that most of the intronic regions were barely alignable. We finally assess whether the recombination boundary sequences are more likely to be conserved than genomic background systematically. To this end, we performed pairwise alignments of the 6-kb syntenic regions centering on the 12 recombination sites which were mapped to resolution up to 100 bp. We calculated their alignment scores using SIMD Smith–Waterman Library (35). To calculate alignment score for background sequences, we performed permutation analysis by randomly choosing one thousand 6-kb syntenic regions between the two species followed by alignment score calculation. We observed a significantly higher conservation level in recombination sites than in genomic background (*P* = 0.002129, Wilcoxon test) (Figure 6D). The results suggested that the conservation of sequences extending up to 6 kb in length in the recombination site was likely to be an exception in the non-coding region, and that such a conserved block of multiple kbs in length was possibly the minimal requirement for recombination to take place in a hybrid.

**DISCUSSION**

DNA recombination is required for the correct segregation of genetic material and successful completion of meiosis. It is regulated by both *cis*-acting elements and *trans*-acting factors, which rapidly evolve across species (45). The highly divergent *cis* elements and *trans* factors presumably serve to
reduce the recombination compatibility between species. Indeed, recent studies in mice and other species indicate a genetic link between recombination and speciation (46). Consistent with this, we found DNA recombination was severely suppressed in the hybrids between *C. briggsae* and *C. nigoni* compared with that in the parental species (16). The availability of high-quality genomes for both species allowed us to pinpoint the molecular basis behind the suppressed recombination in the hybrids.

First, the *cis* elements and *trans* factors might be too divergent to be compatible with each other for recombination to take place in the hybrids between *C. briggsae* and *C. nigoni*. Consistent with this, there are only 15,157 one-to-one orthologs between the two, although a total of 21,863 (WS252) and 22,462 genes (Table 1, this study) were predicted in the genomes of *C. briggsae* and *C. nigoni*, respectively. Second, most parts of the two genomes might be too divergent to allow the pairing or crossover of homologous chromosomes that is critical for recombination to take place (48). For example, the overall sequence identity of the coding sequences from the one-to-one orthologous genes was <90%, and the vast majority of intronic and intergenic regions were not alignable (Figures 4A and 6C). Third, despite the overall synteny of orthologous protein-coding genes between the two species, there are pervasive intra- and inter-chromosomal sequence rearrangements between the two genomes, which are expected to impair recombination. Fourth, the size of *C. nigoni* genome was substantially larger than that of *C. briggsae*, i.e. 136 versus 108 Mb, although some residual heterozygosity could have contributed to overestimation of the *C. nigoni* genome size. Some of the divergent alleles could be erroneously treated as paralogs, especially for those sequences present as ‘unassigned’ contigs, which are highly enriched for repetitive fragments. Genome shrinkage in a selfing species (e.g. *C. briggsae*) versus its outcrossing relative, for example, *C. nigoni*, seems not uncommon (49). Nearly half of the reduced genome in *C. briggsae* could be explained by its lost repeats relative to the *C. nigoni* genome. These differences in genome size and repeat content are expected to create gaps in the sequence alignment or make the syntenic sequences not alignable, thus decreasing recombination ef-

Figure 5. Comparison of synteny consisting of gene blocks (see Materials and Methods) between *C. briggsae* and *C. nigoni* or *C. elegans*. A total of 15,157 (Supplementary Table S6) and 7,679 (WS252) orthologous gene pairs were used for comparison between *C. briggsae* and *C. nigoni* or *C. elegans*, respectively, with a bin size of 30 genes. (A) Syntenic view. (B and C) Dotplot view. Gene count is indicated on each axis along with its associated chromosome (differentially color coded).
Figure 6. Conservation of sequences flanking recombination site between *C. briggsae* and *C. nigoni*. (A) The left and right boundaries of an introgression (ZZY10331) were mapped onto the right arm of *C. briggsae* chromosome III by NGS. Shown is the coverage of sequencing reads (Y axis) derived from hybrid mapped against *C. briggsae* chromosome III (X axis). The left boundary is indicated with an arrow and the right boundary is mapped to the very end of the chromosome. (B) Dotplot of the 6-kb syntenic sequences flanking the recombination site (indicated with arrow) between *C. briggsae* and *C. nigoni*. Gene models within the region are shown in scale. (C) Dotplot of the 6-kb homologous sequences spanning the genomic regions of a highly conserved gene, *abce-1*, between *C. briggsae* and *C. nigoni*. Gene models are shown in scale for both species. Note that only exons are conserved. (D) Boxplot of alignment scores for the 6-kb syntenic genomic DNAs of recombination regions (left) or randomly sampled genomic regions (right). SW score, Smith–Waterman alignment score.

Nanopore sequencing is expected to perform better in *de novo* genome assembly than it did in our study. One potential issue is that the read coverage was not deep enough. We produced only 30× reads of the *C. nigoni* genome. Given the high error-rate of its read sequence, some remaining errors in its self-corrected reads may prevent it from effective scaffolding and contig assembly. In addition, we performed size selection for the DNA library, which was optimized to recover genomic fragments of ~8–10 kb in length. This would constrain the power of their reads for scaffolding or *de novo* sequencing. The generation of a higher coverage of reads and size selection of fragments with greater size may boost their performance in *de novo* genome assembly.

We did not expect that Hi-C data to be more valuable than Nanopore reads for scaffolding the SLR-derived contigs (Figures 2D and 3). The superior performance relative to the Nanopore reads was likely the result of the higher coverage (~100×) and the highly accurate read sequences generated by NGS. In summary, the improved *C. briggsae* genome and the new *C. nigoni* genome provide insights into the unusual low rate of recombination for the large *C. briggsae* DNA fragments in *C. nigoni* background. The avail-

fiency. Nevertheless, it remains possible that bombard-
ment may have introduced some balanced chromosomal re-
arrangements into our GFP lines that are expected to pre-
vent effective recombination (7,8) but are not detected by
our genotyping steps. Alternatively, the presence of the GFP
transgene may disrupt synteny completely in an unnatural
way, which could exert some sort of ‘shadow’ of recombi-
nation suppression around itself that would not be observed
in unmarked DNA.

Genome instability was proposed to be intimately linked
to speciation and evolution (50). The newly generated *C.
nigoni* genome and the improved *C. briggsae* genome will
facilitate the studies of genome evolution and speciation
genetics. They should also facilitate the studies of sex evo-
lution given the distinct reproductive modes of the two
species. Our end-sequenced fosmid and BAC libraries pro-
vide invaluable resources for the functional characterization
of hybrid incompatible loci and other DNA elements. The
availability of boundary-resolved introgressions in the con-
text of syntenic chromosomal regions should form a foun-
dation for further studying the molecular and genetic con-
trol of recombination frequency.
ability of both genomes forms invaluable resources for elucidation of the molecular mechanisms underlying the hybrid incompatibilities between the two species.

**AVAILABILITY**

*C. briggsae* SLRs and Hi-C data and *C. briggsae* genome assemblies were deposited into SRA database under BioProject number PRJNA396670. *C. nigoni* Hi-C, Nanopore sequencing data, BAC and fosmid end-sequencing data as well as *C. nigoni* genome assembly 'cn2' and its annotation files were deposited into SRA database under BioProject number PRJNA306403. Synteny and genomic positions of BAC and fosmid clones can be accessed under designated website, http://158.182.16.70:8080/.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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