Deciphering complex genome rearrangements in *C. elegans* using short-read whole genome sequencing

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Genomic rearrangements cause congenital disorders, cancer, and complex diseases in human. Yet, they are still understudied in rare diseases because their detection is challenging, despite the advent of whole genome sequencing (WGS) technologies. Short-read (srWGS) and long-read WGS approaches are regularly compared, and the latter is commonly recommended in studies focusing on genomic rearrangements. However, srWGS is currently the most economical, accurate, and widely supported technology. In *Caenorhabditis elegans* (*C. elegans*), such variants, induced by various mutagenesis processes, have been used for decades to balance large genomic regions by preventing chromosomal crossover events and allowing the maintenance of lethal mutations. Interestingly, those chromosomal rearrangements have rarely been characterized on a molecular level. To evaluate the ability of srWGS to detect various types of complex genomic rearrangements, we sequenced three balancer strains using short-read Illumina technology. As we experimentally validated the breakpoints uncovered by srWGS, we showed that, by combining several types of analyses, srWGS enables the detection of a reciprocal translocation (eT1), a free duplication (sDp3), a large deletion (sC4), and chromoanagenesis events. Thus, applying srWGS to decipher real complex genomic rearrangements in model organisms may help designing efficient bioinformatics pipelines with systematic detection of complex rearrangements in human genomes.

Structural variations (SVs) are genomic rearrangements such as copy number alterations, inversions, and translocations. More complex events, known as chromoanagenesis, combine a cascade of chromosomal rearrangements1. Over the past few years, structural variants and complex genomic rearrangements have been implicated in various phenotypes: cancer2, rare disorders3–9 and common diseases10 in humans, reproduction traits in pigs11, virulence traits in plant pathogenic fungi12, local adaptation in maize13, and behavior in *Caenorhabditis elegans* (*C. elegans*)14. However, the technologies and methods used to identify SVs and complex rearrangements are still multifaceted and no approach has yet been recognized as standard. Short-read and long-read whole genome sequencing (WGS) technologies, as well as their respective tools and pipelines, are often assessed and compared in their ability to detect structural variants and complex rearrangements15–21. The read length of short-read technologies is often reported as a limitation for detecting larger and more complex events22. Meanwhile, long-read sequencing and linked-reads approaches are gaining popularity23–25, especially when the analysis of short-read sequencing data fails to uncover SVs and complex rearrangements of interest26,27. Here, we focused on short-read WGS of *C. elegans* strains known to harbor SVs and show that short-read WGS provides enough data to decipher SVs of various types and complex genomic rearrangements in these genomes when tailored workflows are used.

In *C. elegans*, SVs and complex rearrangements have been used for decades to balance large parts of the genome by suppressing crossover events and maintaining heterozygosity. It facilitates the investigation of lethal mutations, the construction of new strains, and the screening of mutations28. While some balancers are spontaneous, like the reciprocal translocation *nT1(IV;V)*28, most were created via random mutagenesis processes, such as X-ray mutagenesis, chemical mutagens (acetaldehyde, ENU, EMS), gamma irradiation, and more recently

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by using CRISPR-Cas9 methods\(^{29,30}\). For most of the mutagen-induced balancers, the implicated chromosomal rearrangements are uncharacterized at the molecular level (i.e., precise genomic position and nature of the rearrangement are unknown. Thus, C. elegans balancers constitute an interesting source of various genomes and complex genomic rearrangements to assess the ability of short-read PCR-free WGS Illumina technologies and tailored bioinformatics workflows to detect and characterize complex structural variants. Here, we sequenced the genomes of three C. elegans balancers, ranging from a well-characterized SV \(eT1(III;V)\), a reciprocal translocation to an uncharacterized and molecularly unknown balancer \([sC4 (BC4586)]\). Beyond the successful proof-of-concept detection of \(eT1(III;V)\), we deciphered the structure and genomic positions of \(sDp3\) and \(sC4\), as well as additional rearrangements not previously known to exist in the balancer strains selected for this study (BC4586, BC986, and VC109).

In our study, we found that short-read WGS datasets can be used to detect, identify, and characterize SVs and complex genomic rearrangements in C. elegans genomes. The knowledge gained from the analytical methods used on C. elegans balancers may help optimize detection and characterization of complex variants in humans using short-read WGS.

**Results**

**Short-read WGS can be used to detect homozygous and heterozygous reciprocal translocations.** The strains BC986 and VC109 carry the reciprocal translocation \(eT1(III;V)\). In C. elegans, the reciprocal translocation \(eT1(III;V)\) balancer has been well studied and it is described as balancing LGV, from the left chromosome end through \(unc-23\), and LGIII, from the right end to \(unc-36\)\(^{31}\). Its genomic breakpoints were more recently localized in the second intron of \(unc-36\) on LGIII and between \(rol-3\) and \(unc-42\) on LGV\(^{32}\). Therefore, we first focused our efforts on retrieving \(eT1(III;V)\) breakpoints, to assess the ability of short-read WGS to decipher reciprocal translocations as a proof of concept for our approach.

Reads were aligned to the C. elegans reference genome (WS265) and candidate breakpoints were predicted using an ensemble of tools (see “Methods”). Two sets of breakpoints related to a translocation between LGIII and LGV were correctly identified by several tools in these \(eT1\) strains, but not in controls. The breakpoints we identified agreed with the locations previously described by Zhao and colleagues\(^{32}\): III:8,200,762–V:8,930,675 and III:8,200,764–V:8,930,675 (Fig. 1A and Supplemental Fig. S1). As a first validation step, we used the Integrative Genomics Viewer (IGV) to review the visual signature of reads aligned around those locations (Fig. 1B). In homozygous genomes, we observed that no read was overlapping the position of the breakpoint (i.e., the reads mostly aligned either on the left or the right of the breakpoint, with little or no read sequence aligning across the breakpoint position). In heterozygous genomes, half of the reads were displaying this signature (Fig. 1B).

Then, we amplified the genomic loci around those breakpoints by PCR and submitted the PCR products for Sanger sequencing (Fig. 1C–E). By analyzing the Sanger sequences, we confirmed that the breakpoint on LGIII was in the second intron of \(unc-36\) at 8,200,764 Mb and that the breakpoint on LGV was intergenic, localized at 8,930,675 Mb. Additionally, we characterized microhomologies at the breakpoint on LGIII, composing a 43-bp sequence inserted at the junction containing several sequences flanking the breakpoints. The main part of the inserted sequence (27 bp) has been duplicated from the LGV flanking region. Two additional sequences, respectively 5 bp and 1 bp long, are duplicated from the LGIII flanking region (Fig. 1D).

One of the strains (VC109) was viable in both heterozygous and homozygous states\(^{33}\). We prepared genomic DNA from both \(eT1\) heterozygous (wild-type looking worms) and \(eT1\) homozygous (phenotypically \(unc-36\) worms) and sequenced them. We were able to identify (Fig. 1B) and confirm the \(eT1\) breakpoints in both cases (Fig. 1C), demonstrating that the short-read WGS approach is effective at deciphering position and structure of the breakpoints for reciprocal translocations regardless of the zygosity status.

**Short-read WGS contains enough information to identify short and large copy number variations.** By combining calls from various tools, coverage analysis, and read inspection, we detected an assorted set of copy number variations. We confirmed their nature and positions by PCR and Sanger sequencing. Overall, we observed five deletions specific to BC986, spanning from 69 bp to 8,779 bp (Supplemental Figs. S2, S3). In VC109 genomes, we also detected four additional deletions ranging from 86 to 255 bp in size. Some were heterozygous, others were homozygous (Supplemental Figs. S4, S5, S6). We also identified two direct tandem copy number gain events in VC109. The first one, localized on LGI, was a homozygous direct tandem duplication in both heterozygous (phenotypically wild-type) and homozygous (phenotypically \(unc-36\)) worms. The second direct tandem duplication mapped on LGV and was both heterozygous and homozygous in heterozygous and homozygous worms, respectively (Supplemental Fig. S6). More information regarding these reported CNVs is available in Supplemental Table S1.

**Short-read WGS can uncover a free duplication.** The \(sDp3\) balancer, also present in BC986 along with \(eT1(III;V)\), has been described as a free duplication on LGIII effectively balancing the left portion of LGIII from around \(unc-86\) through to at least \(dpy-1\), but does not extend to \(unc-45\)\(^{34}\). So far, 22 genes have been described to be overlapped by \(sDp3\) and, by analysis of the coverage, we confirmed that their sequences were duplicated (Fig. 2A). None of the tools we applied (see “Methods”) reported breakpoints or structural variants that could fit the \(sDp3\) description. However, we observed heterozygous SNVs from the left end of LGIII until at least the \(eT1\) breakpoint (III:8,200,675), corroborating the presence of an event balancing this part of LGIII, and maintaining heterozygosity (Supplemental Fig. S2 and Supplemental Table S3). An unbiased analysis of the sequencing read depth on LGIII helped us map the duplication to two different loci: between III:1.4 Mb–2.4 Mb and III:3.6 Mb–8.6 Mb (Fig. 2B). To confirm this structure, we inspected the reads aligned around III:2.4 Mb and III:3.6 Mb. We identified read pairs for which the forward read was aligned to the first segment of the duplication.
and the mate aligned along the second segment, thus corroborating our hypothesis. To experimentally validate it, we identified the breakpoint linking the two parts of the duplication (III:2,452,252 and III:3,693,056) and confirmed by PCR and Sanger sequencing (Fig. 2C and D).

Short-read WGS can reveal and characterize unexpected complex rearrangements. By comparing variants and breakpoints in the three eT1 strains and controls—strains without eT1(III;V) including N2 and BC4586, we built an "eT1 haplotype" composed of variants specific to the eT1 strains. Interestingly, along with eight SNVs (list available in Supplemental Table S2), we also characterized two unexpected and undescribed complex rearrangements.

The first one could have been interpreted at first sight as a classic large copy number gain in tandem (direct) spanning from V:2,144,217 to V:2,156,311 (Supplemental Fig. S7A). It overlapped seven intact genes: srbc-20, C45H4.t1, C45H4.21, C45H4.13, C45H4.19, srbc-24 and, srbc-23, as well as partially spanning srbc-52 (exon 1 only) and srbc-21 (up to intron 4). PCR and Sanger sequencing confirmed the duplication breakpoints and structure in direct tandem (Supplemental Fig. S7B). Both BC986 and unc-36 VC109 worms [eT1(III;V) homozygous] were homozygous for the direct tandem duplication (ratio of coverage = 2) while wild-type looking VC109 [eT1(III;V) heterozygous] was heterozygous (ratio of coverage = 1.5) (Supplemental Fig. S7C). The analysis of the coverage however showed a discontinuity in the coverage (ratio of coverage dropping back to 1) between V:2,148,200 and V:2,148,630 which corresponds to the three last exons of srbc-20 and a part of the last exon of C45H4.21 (Supplemental Fig. S7C). An inspection of the reads revealed that this variant is complex, with an inversion overlapping the copy number gain. We confirmed the inversion V:2,148,056–2,148,630 by PCR and Sanger sequencing (Supplemental Fig. S7B and D).

The second eT1 specific complex rearrangement was localized on LGV around 1.1 Mb. The complex rearrangement described here overlapped with the gene Y50D4B.1, a non-essential gene in C. elegans. Between

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**Figure 1.** Overview of eT1 genomes and validation of eT1(III;V) reciprocal translocation breakpoints. (A) Circos plot created with Circos62 with information regarding BC986 (yellow), wild-type looking VC109 (red; VC109 Het), and unc-36 VC109 (blue; VC109 Hom) genomes. The outer section is composed of three-line charts (one per strain) representing the ratio of coverage calculated by windows of 1 Kb and divided by the strain-specific genome coverage. The middle section is composed of three scatter plots. Each dot represents the genomic position of heterozygous SNPs identified in each strain. The inner section highlights, with links and ribbons, SVs and complex rearrangements identified in each strain. The eT1(III;V) reciprocal translocation is displayed by a blue ribbon. For better resolution, see Supplemental Fig. S1. (B) Signature of reads aligned around the eT1 breakpoints, visualized with IGV, BC986, wild-type looking VC109, and unc-36 VC109 genomes. (C) PCR agarose gels validating eT1 breakpoints and heterozygous and homozygous genotypes in VC109. (D, E) Sequences at eT1 junctions on LGIII (D) and LGV (E) resolved by Sanger Sequencing. The junction on LGIII shows microhomologies with flanking regions. The sequence in red is from LGIII, the sequence in blue is from LGV. The sequence in black is the de novo sequence inserted at the junction, composed of microhomologies of the surrounding sequences. We represented the microhomologies between the junction sequence and the surrounding sequences with bold, italic and underlined characters.


Figure 2. Free duplication sDp3 in BC986. (A) Ratio of coverage for the 22 genes overlapped by the free duplication sDp3 in BC986 (orange) and N2 (blue). (B) Coverage of LGIII in BC986 and sDp3 boundaries. The map on the top displays the breakpoints identified for sDp3 along LGIII. The line chart represents the ratio of coverage on the entire LGIII. The coverage was evaluated by a sliding window of 10 kb and divided by the average coverage for the entire genome. This ratio was then divided by the ratio of coverage for the same window in N2. (C) PCR gel confirming the breakpoints III:2,452,252 and III:3,693,056 linking the two duplicated segments of LGIII. (D) Sequence of the PCR product obtained by Sanger sequencing. The red sequence belongs to the first part of the free duplication (III:1.4–2.4 Mb) and the blue sequence belongs to the second portion of the duplication (III:3.6–8.6 Mb).

| Variant                  | Breakpoint 1       | Breakpoint 2       | Gene        |
|--------------------------|--------------------|--------------------|-------------|
| Inverted tandem duplication | V:1,118,539       | V:1,118,853       | Y50D4B.1    |
| Deletion                 | V:1,118,855       | V:1,128,457       | Y50D4B.1    |
| Deletion                 | V:1,126,382       | V:1,126,983       | Y50D4B.1    |
| Deletion                 | V:1,127,007       | V:1,127,020       | Y50D4B.1    |
| Inverted tandem duplication | V:1,127,471       | V:1,129,752       | Y50D4B.1    |
| Inverted tandem duplication | V:1,128,827       | V:1,129,264       | Y50D4B.1    |
| Deletion                 | V:1,129,264       | V:1,129,753       | Y50D4B.1    |
| Inversion                | V:1,126,322       | V:1,128,612       | Y50D4B.1    |

Table 1. Breakpoints of the complex rearrangement present in eT1 containing strains on LGV.

V:1.118 Mb and V:1.130 Mb, we identified 15 different breakpoints (Table 1). By inspecting the reads, we identified three short deletions (homozygous in BC986 and VC109 unc-36 worms, heterozygous in VC109 wild-type worms), one inversion, one large deletion and three inverted tandem duplications (Fig. 3). We confirmed experimentally all breakpoints by PCR and Sanger sequencing.

In the strain VC109 only, we detected eight breakpoints on LGIII around 10 Mb (Table 2 and Fig. 4). Based on coverage analysis and PCR, we further characterized this complex rearrangement as being composed of one direct tandem duplication, one inverted tandem duplication, and two deletions. Because of the presence of copy gains in the rearrangement and microhomologies at breakpoint junctions, this complex rearrangement could be characterized as chromoanasynthesis. It overlapped the non-essential gene tbc-8, so it is not expected to have an important effect on the fitness of the worms.

**Short-read WGS to characterize BC4586, an uncharacterized genetic balancer strain.** The strain BC4586 contains the sC4 rearrangement that has been used to balance the right end of LGV, from rol-9

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to \textit{unc-76}. It was also reported that it reduces the genetic distance between the genes \textit{unc-76} and \textit{rol-9} to 1.8\%, suggesting the presence of a deletion\textsuperscript{28}. To the best of our knowledge, the rearrangement \(sC4\) remains molecularly uncharacterized. We used short-read WGS to determine the nature of the \(sC4\) rearrangement and to report additional genomic variants in BC4586.

We first performed “\(sC4\) haplotype” analysis (Supplemental Table S3) and observed stretches of heterozygous SNVs only on LGV from ~12 to ~16 Mb and from ~19 Mb to its right end. This suggested that \(sC4\) might be able to balance further than \textit{unc-76}. We detected a deletion on the right portion of LGV between 16 and 19 Mb (Fig. 5A) that explains the reduced genetic distance previously reported between \textit{unc-76} and \textit{rol-9}. We confirmed the deletion by PCR (Fig. 5B). We have also detected a non-reciprocal translocation of the right arm of LGV to the right arm of LGIV (Table 3). We hypothesized that this has led to a fusion of the two chromosomes, by their right ends. The breakpoint was supported by several reads. However, the region surrounding the breakpoint on LGV is highly repetitive, and despite our best efforts, we could not design a unique set of primers to validate this hypothesis by Sanger. Therefore, we assessed the karyotypes of the diakinetic oocytes using DAPI staining. The wild-type oocytes typically have six pairs of DAPI-stained bivalent diakinetic chromosomes (Fig. 5C), whereas in the BC4586, we frequently observe five pairs (Fig. 5C), confirming \(sC4\) chromosome fusion.

On LGIV, we also characterized a complex homozygous rearrangement combining two deletions, one inversion and one direct tandem duplication localized around 9.8 Mb (Fig. 5D). We confirmed the breakpoints for both complex rearrangements by PCR (Fig. 5D). We also reported and validated a deletion on LGV and a direct tandem duplication on LGIII (Fig. 5D, Supplemental Table S1, Supplemental Fig. S8). The Circos plot in Supplemental Fig. S9 summarizes our findings.

**Figure 3.** Complex rearrangement in \(\varepsilon T1\) strains on LGV. (A) Schematic representation of the complex rearrangement combining three inverted tandem duplications (red), four deletions (blue), and one inversion (green) along LGV around 1.1 Mb. (B) LGV screenshot of the VC109 genome (\textit{unc-36}) for the region overlapped by the complex rearrangements. (C) PCR gels. 1 = Deletions V:1,126,582–1,126,983 and V:1,127,007–1,127,020; 2 = Deletion V:1,129,264–1,129,753, inverted tandem duplication V:1,127,471–1,129,752 and inverted tandem duplication V:1,128,857–1,128,895. 3 = Deletion V:1,118,853–1,128,457, inverted tandem duplication V:1,118,539–1,118,853, and inversion V:1,126,322–1,128,612. (D) Sanger sequencing of the PCR product #3 confirming several breakpoints.

**Table 2.** Complex rearrangement breakpoints in VC109 on LGIII.

| Variant                  | Length | Breakpoint 1       | Breakpoint 2       | Gene |
|--------------------------|--------|--------------------|--------------------|------|
| Inverted tandem duplication | 8315 bp | III:10,362,573     | III:10,370,888     | tbc-8 |
| Deletion                 | 4387 bp | III:10,366,492     | III:10,370,879     | tbc-8 |
| Direct tandem duplication | 9409 bp | III:10,366,037     | III:10,375,446     | tbc-8 |
| Deletion                 | 2500 bp | III:10,368,666     | III:10,371,166     | tbc-8 |
Discussion

Short-read whole genome sequencing (WGS) has often been used to retrieve structural variants and more complex rearrangements among other variations in humans\(^\text{17,34,35}\), \textit{D. melanogaster}\(^\text{36}\), as well as \textit{C. elegans}\(^\text{37–40}\). Here, by reporting the precise breakpoints of complex rearrangements in \textit{C. elegans} \([eT1(III;V), sC4, sDp3]\), we describe the molecular structure of widely used balancers, most of them for the first time to the best of our knowledge. We also show that short-read WGS enables identification and characterization of large SVs and complex rearrangements, by deep analysis of short-read WGS datasets.

Every breakpoint that we uncovered with deep analysis of the short reads was validated experimentally. However, the interpretation of the structure of the rearrangements could necessitate further exploration. Still, despite the limited ability of short-read WGS to span large genomic rearrangements fully or to explore repetitive regions such as telomeres, we characterized the balancer \(sC4\) as a large deletion and a chromosome fusion \((IV;V)\). This rearrangement could reflect a telomere crisis\(^\text{41}\) occurring as an end-to-end chromosome fusion associated with telomere shortening. This type of event has been previously studied in \textit{C. elegans}\(^\text{38,42,43}\). We also uncovered a free duplication, composed of two genomic segments \((sDp3)\), along with chromosomal rearrangements combining several various events that present features of chromoanagenesis. Our analyses of \(eT1\) strains confirmed the \(eT1\) breakpoints as identified by Zhao and colleagues\(^\text{32}\). Interestingly, we placed the LGIII breakpoint 3-bp anterior to the one previously reported. Although we retrieved the junction sequence described as a 35-bp duplication, our approach with short-read WGS showed a more complex scenario with microhomologies of several flanking sequences, suggesting the involvement of a replication-based DNA mechanism repair such as fork-stalling and template switching, or microhomology-mediated break-induced replication.

In \textit{C. elegans}, short-read WGS has been employed in only a few studies to describe SVs. For instance, Meier and colleagues\(^\text{38}\) and Volkova and colleagues\(^\text{39}\) reported mutational signatures (SNVs and SVs) created by carcinogen exposure on strains with DNA repair deficiency. Itani and colleagues\(^\text{37}\) characterized a complex rearrangement created by ENU-mediated mutagenesis. In 2017, Cook and colleagues\(^\text{44}\) published the database CeNDR (\textit{C. elegans} Natural Diversity Resource) that regroups genomic variations uncovered by genome sequencing in wild \textit{C. elegans} strains. Other than insertions of transposable elements\(^\text{45}\), SVs and complex rearrangements are not reported for the natural isolates in the CeNDR. Our study shows that short-read sequencing is a viable option for future studies to explore the natural variation of \textit{C. elegans} species beyond SNVs, especially by re-analyzing datasets already available in CeNDR, for which those complex variants might have been overlooked.

It is quite common in human studies to assess analysis pipelines of short-read WGS using either generic genomes (Genome in a Bottle) or simulated data\(^\text{46,47}\), especially because real-life cases emerge anecdotally. However, in our study, we assessed several tools and approaches on real biological data from model organism genomes. This approach presents two main advantages. First, in model organisms like \textit{C. elegans}, balancers are
widely used and well-known as being genomes containing SVs and complex rearrangements, largely comparable to humans. Thus, they constitute good surrogates of real-life cases, without the limitation related to a low frequency of those events. Second, as shown here, real biological data allows us to uncover unexpected events, of various natures and complexity. Thus, there is a probability that simulations might not be able to cover the wide diversity of chromosomal rearrangements or might not simulate the complexity of read signatures. Thus, we reasoned that tool assessment would be more accurate if they were performed on real data, human or not, combined with experimental validation.

**Conclusions**

In our study, we showed that short-read data provides enough information to detect a spectrum of complex variants with tailored bioinformatics approaches. Thus, to improve the detection and characterization of SVs and complex rearrangements, it is important to also optimize pipelines and analyses to get the best out of the short-read datasets. Indeed, short-read sequencing is the most widely used approach and the most cost-effective
technology available. Also, as there are more tools and pipelines available to analyze short-read data than for long-read or linked-read data, it facilitates pipeline tailoring by using different tools and approaches. Additionally, short-reads permit the detection of both single nucleotide variants and larger ones, whereas long-read approaches are error prone and thus, limited, in their ability to accurately detect SNVs. This constitutes quite an advantage for short-read approaches as it avoids the necessity to resort to another assay for small variants. Moreover, public databases on human variation such as TopMed and gnomAD have been built upon calls from short-read datasets. Therefore, in the context of human rare disease unsolved cases, where population databases are a major asset to decipher rare and pathogenic variants from common and benign ones, short-read sequencing remains the main approach. Thus, improving short-read sequencing pipelines to maximize the detection of variants is of utmost importance.

Methods
Worm maintenance and strains. Strains Bristol N2 wild type, Hawaiian strain wild type CB4856, BC986 (sDp3(III); +/eT1 (III; V)), VC109 (apc-11(gk37)/eT1 III; +/eT1 V) and BC4586 (unc-76(e911) rol-9(sc148)/ sCa3(s2172) [dp-21(e428)] V) were used in this study. Strains were obtained from the CGC (Caenorhabditis Genetics Center). N2 was used as the wildtype strain. All strains were maintained at 16 °C and kept on standard NGM plates streaked with OP50.

DNA extraction. Genomic DNA was collected from approximately 100 mg of worm tissue using the Qiagen Blood and Tissue kit (Cat #: 13323) following the manufacturer’s recommendations. DNA was eluted with 10 mM Tris–HCl (pH 8.0). Samples were quality-checked to ensure a minimum quantity of 1500 ng and a 260/280 ratio of 1.8 before submitting for sequencing.

Library preparation, sequencing and data pre-processing. Paired-end short-read WGS were obtained for all strains with PCR-free library preparation protocol and NovaSeq6000 Illumina sequencing technology. We checked the quality of the fastq files using FastQC. The reads were 151 bp long. We trimmed the reads and removed the adapters using TRIMMOMATIC v0.36. For each sample, we aligned between 16 and 34 million reads using BWA-MEM v0.7.17 algorithm to the C. elegans reference genome WS265. It resulted in a 30X read coverage per strain on average (Supplemental Table S4). We then sorted the reads according to their coordinates with ‘samtools sort’ (samtools v1.5)’.

SV and complex rearrangement detection. We called and characterized SNVs, indels, SVs, and complex rearrangements for each strain in this study using a collection of published tools and downstream in-house designed analysis methods. Strain N2 was used as a control. The SNVs and indels genotype of each strain was established using RUFUS. The analysis of SNV heterozygosity along the genome of each strain was used to highlight balanced genomic regions. For SVs and complex variants, we initially ran nine different tools with default parameters: BreakDancer v.BreakDancerMax-1.1r11254, CNVnator v0.4.1, CNVnator v0.4.1, DELLY v0.7.8, DELLY v0.7.8, GRIDSS v2.8.0, GRIDSS v2.8.0, SeekSV v1.2.3, SeekSV v1.2.3, Tardis v1.0.7, Tardis v1.0.7, TIDDIT v2.12.0, TIDDIT v2.12.0, Manta v1.6.0, Manta v1.6.0, Biltong CompGen/tardis, Biltong CompGen/tardis, ScilifeLab/TIDDIT and RUFUS. For complex variants, breakpoints were defined combining RUFUS and GRIDSS calls and custom methods (visual assessment, coverage analysis, reads inspection):

1. Visual assessment consists in reviewing the visual signature of reads aligned around each breakpoint with IGV. A breakpoint is represented by accumulation of split reads, with little or no read sequence aligning across the breakpoint position. The visual signature gives information to characterize the type of rearrangement.
2. Read inspection consists in a “manual” re-alignment of reads aligned at each breakpoint junction. Reads are extracted from bam files with “samtools view” and re-aligned using Blast (UCSC – Feb. 2013; WBcel235/ce11). This analysis aims to identify split reads supporting the breakpoint junctions (as described by Iwata et al.). Such reads are fundamental to the design of PCR primers for further validation.
3. To characterize copy number variations (stand-alone CNVs or as part of complex variants), we estimated the average genome coverage and read depth by intervals of 1–10 kb (depending on the length of the CNV) using the ‘samtools depth’ function.

The circular visualizations were produced using Circos. The line charts were prepared in Excel. The data relating to the genomic variations are available in Supplementary information for BC4586 (Supplemental Table S5) and VC109 (Supplemental Tables S6 and S7) genomes. The complete list of additional SVs identified and confirmed by PCR, but not discussed in this paper, is available in Supplementary Table S1. Circos plots, PCR gels, and IGV screenshots are available in Supplemental Figures S1, S2, S3, S4, S5, S6, S7, S8, S9.

Experimental validation. We confirmed breakpoints of SVs and complex rearrangements by PCR and Sanger Sequencing. All primers and sequences are available in Supplementary Information (Supplemental Tables S8, S9, S10, S11). For the cytological assessment of bivalent diakinetic oocyte karyotypes, 1-day-old adult hermaphrodite worms were washed once in M9 medium, fixed in cold methanol, rehydrated in PBS (0.01% Tween) and mounted using SlowFade Gold antifade reagent with DAPI (Invitrogen S36938). Images were...
acquired using a Zeiss Imager M2. Raw counts can be found in Supplemental Table S12. The p-value was calculated using two-tailed Z-test.

Data availability
The sequencing data generated in this study have been submitted to the NCBI BioProject database (https://www.ncbi.nlm.nih.gov/bioproject/) under accession number PRJNA728090.

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Author contributions
T.M. and M.T.G. conceptualized and designed the study. T.M., M.O. and M.T.G. analyzed and interpreted the WGS data. X.L. performed the experimental validation. S.S. and F.I. prepared the strains and DNA. T.M. drafted the manuscript and M.T.G. helped finalize it. Every author read, commented, and validated the manuscript.

Competing interests
The authors declare no competing interests.

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