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Genome sequencing of *C. elegans* balancer strains reveals previously unappreciated complex genomic rearrangements

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Genetic balancers in *Caenorhabditis elegans* are complex variants that allow lethal or sterile mutations to be stably maintained in a heterozygous state by suppressing crossover events. Balancers constitute an invaluable tool in the *C. elegans* scientific community and have been widely used for decades. The first/traditional balancers were created by applying X-rays, UV, or gamma radiation on *C. elegans* strains, generating random genomic rearrangements. Their structures have been mostly explored with low-resolution genetic techniques (e.g., fluorescence in situ hybridization or PCR), before genomic mapping and molecular characterization through sequencing became feasible. As a result, the precise nature of most chromosomal rearrangements remains unknown, whereas, more recently, balancers have been engineered using the CRISPR-Cas9 technique for which the structure of the chromosomal rearrangement has been predesigned. Using short-read whole-genome sequencing (srWGS) and tailored bioinformatic analyses, we previously interpreted the structure of four chromosomal balancers randomly created by mutagenesis processes. Here, we have extended our analyses to five CRISPR-Cas9 balancers and 17 additional traditional balancing rearrangements. We detected and experimentally validated their breakpoints and have interpreted the balancer structures. Many of the balancers were found to be more intricate than previously described, being composed of complex genomic rearrangements (CGRs) such as chromoanagenesis-like events. Furthermore, srWGS revealed additional structural variants and CGRs not known to be part of the balancer genomes. Altogether, our study provides a comprehensive resource of complex genomic variations in *C. elegans* and highlights the power of srWGS to study the complexity of genomes by applying tailored analyses.

[Supplemental material is available for this article.]

Balancer chromosomes typically contain chromosomal rearrangements that cover a large portion of genomes in model organisms (Edgley et al. 2006). They permit the maintenance of lethal or sterile mutations affecting essential genes in the heterozygous state by either suppressing recombination during meiosis, preventing the recovery of recombinant chromosomes, or leading to inviable recombinants.

Balancers in *Caenorhabditis elegans* implicating large structural variants (SVs) or chromosomal rearrangements stem from diverse origins. Some were created decades ago by applying physical mutagens such as X-ray, UV, and gamma radiation on *C. elegans* strains. They are referred to in this manuscript as “traditional balancers.” The mutagenesis processes generate random DNA breakages and chromosomal rearrangements. Rarely, rearrangements can also occur naturally (Edgley et al. 2006). Altogether, these traditional balancers cover ~70% of the *C. elegans* genome. More recently, the advent of CRISPR-Cas9 technology has allowed the creation of a new set of targeted genetics balancers, balancing pre-

viously uncovered regions (Chen et al. 2018) and covering ~85% of the genome (Iwata et al. 2016; Dejima et al. 2018), referred to as “CRISPR-Cas9 balancers” in this study.

C. elegans balancers with randomly generated chromosomal rearrangements were created before DNA sequencing was available. Their structure has mostly been explored with low-resolution labor- and resource-intensive techniques such as genetic linkage mapping, fluorescence in situ hybridization, PCR analysis (Zhao et al. 2006), SNP mapping (Kadandale et al. 2005), and oligonucleotide array comparative genomic hybridization (Maydan et al. 2007; Jones et al. 2009, 2007). As such, the exact base-pair-resolution DNA location of most balancer breakpoints is unknown. In addition, the structure of the balancers remains poorly characterized as previous approaches were unable to fully resolve complex rearrangements (Jones et al. 2007, 2009). Furthermore, we and others (Jones et al. 2007, 2009; Edgley et al. 2021; Flibotte et al. 2021;

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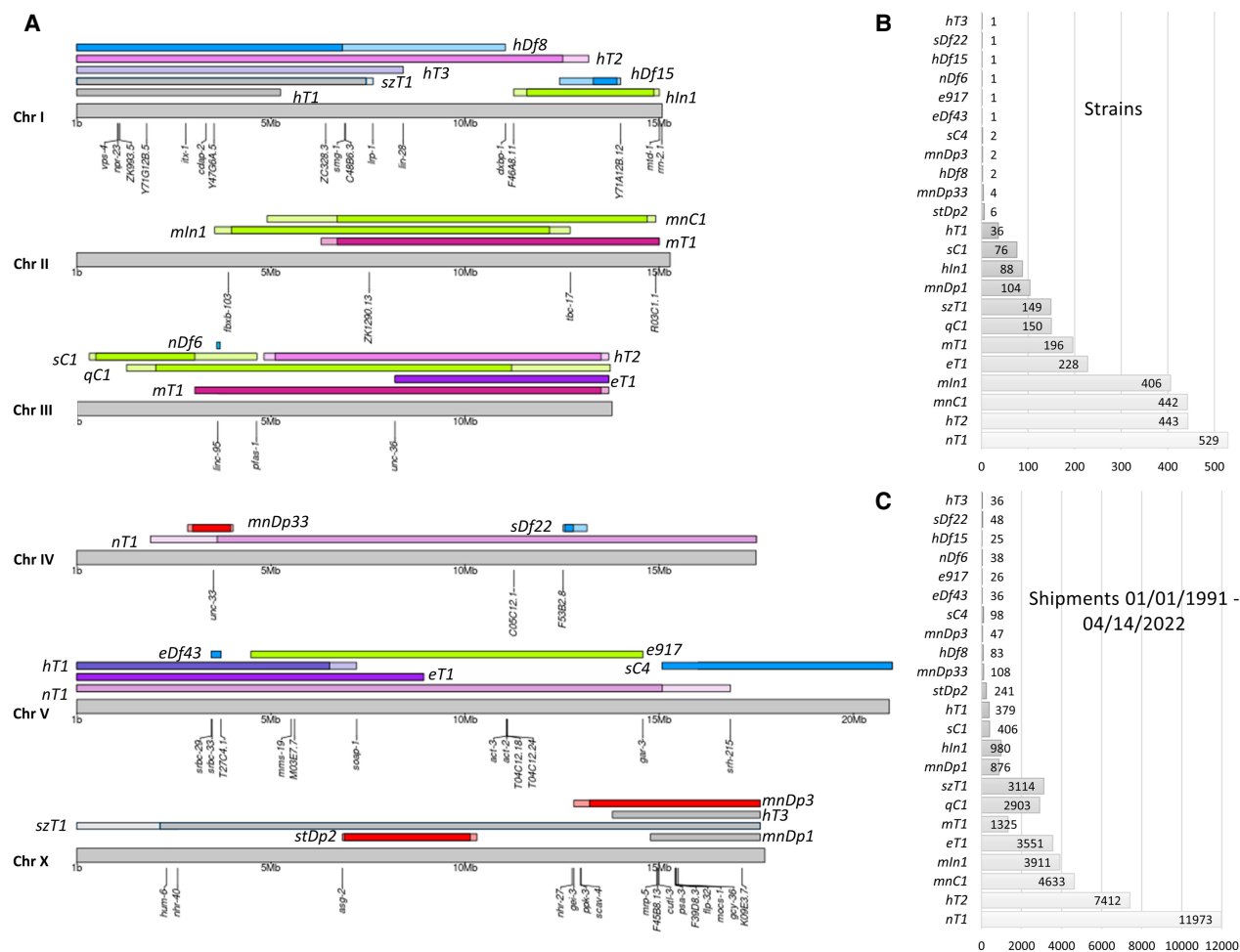


Figure 1. Overview of balancers. (A) Map of the *C. elegans* genomic regions balanced by rearrangements previously characterized at molecular level by our group (Edgley et al. 2021; Flibotte et al. 2021; Maroille et al. 2021) or included in this study. Blue bars are for deletions. Green bars are for inversions or complex inversions. Purple and pink bars are for rearrangements involving translocations without large variation in copy number. Red bars are for rearrangements with copy number gains. Gray bars are for balancers not retrieved in this study, probably owing to breakage of balancers, or for which data are supporting the presence of a derivative. Darker colors represent the region covered by each balancer as described in previous studies. Lighter colors are the additional regions found balanced in this study. (B) Chart representing the number of strains available at the CGC carrying each balancer. (C) Chart representing the number of shipments between 1991 and 2022 of strains containing each balancer.

Maroille et al. 2021) have shown that the molecular structure of genetic balancers is often more complex and larger than the descriptions made decades ago when high-resolution techniques were not available. They frequently combine several overlapping SVs and, in some instances, compounding to create chromoanagenesis-like complex rearrangements.

In *C. elegans* genetic studies, an incomplete characterization of balancer rearrangements prevents their reliable use in mapping experiments and can confound data analysis (Kadandale et al. 2005). Further, given that most *C. elegans* laboratories rely on balancers for studying essential genes, it is important to decipher the precise molecular structure of the rearrangements occurring in each *C. elegans* balancer.

Short-read whole-genome sequencing (srWGS) is currently the most cost-effective sequencing technology. Recently, Miller et al. (2016a,b,c) used genome sequencing to characterize genetic balancers in *Drosophila melanogaster*, illustrating the utility of this technology. Similarly, we recently showed that srWGS contains enough information to accurately detect breakpoints and charac-

terize complex genome rearrangements of various types in *C. elegans* balancer strains (Edgley et al. 2021; Flibotte et al. 2021; Maroille et al. 2021). In human diseases, long reads are often preferred (Gorzynski et al. 2022; Pei et al. 2022). However, recent studies have also benefitted from the power of srWGS to diagnose rare disease patients with complex genomic rearrangements (CGRs) (Ostrander et al. 2018; Minoche et al. 2021; Singer et al. 2021; Nicholas et al. 2022; Zenagui et al. 2022). However, in practice, it remains challenging to detect and characterize the whole spectrum of genomic variations with the available sequencing technologies as there are no standardized analysis pipelines. In addition, a complete characterization of the complexity of events like chromoanagenesis is especially difficult when the length of the reads precludes the overlap of the entire rearrangement. Indeed, chromoanagenesis events are characterized by sudden chromosome shattering and religation. They can be divided into three types based on their molecular properties: chromothripsis, chromoanagenesis, and chromoplexy (Holland and Cleveland 2012; Collins et al. 2017; Pellestor et al. 2022). Chromothripsis and

Table 1. Breakpoints and genes identified in CRISPR-Cas9 balancer strains

Event	Strain(s)	Type	Left Breakpoint	Right Breakpoint	Gene at left breakpoint	Gene at right breakpoint
<i>tmC20</i>	FX30235	INV-INV ^a	I:114579 I:4709755	I:8210593 I:8256469	<i>F53G12.8</i> <i>gsp-3</i>	<i>T02E1.7</i> <i>sre-23</i>
<i>tmC24</i>	FX30237	INV-INV	X:8467140 X:12460309	X:15829281 X:15842089	<i>mec-10</i> <i>odr-7</i>	<i>Y7A5A.20</i> <i>serp-1.1</i>
<i>tmC25</i>	FX30257	INV-INV	IV:749886 IV:888732	IV:3418445 IV:7203027	<i>kvs-5</i> <i>mak-2</i>	<i>lgc-9</i> <i>unc-8</i>
<i>tmC3</i>	FX30133	INV-INV	V:6484110 V:8938061	V:12142846 V:12197761	<i>unc-83</i> <i>unc-23</i>	<i>C27A7.3</i> <i>ogdh-2, lon-3</i>
<i>tmC6</i>	FX30144	INV-INV	II:2331869 II:2671237	II:2758154 II:6912298	<i>ZK1240.1</i> <i>sri-57</i>	(<i>F29A7.8..fbxb-107</i>) <i>asm-1</i>
Additional events	{ FX30133 FX30144 FX30235 FX30237 FX30257 FX30133 FX30144 FX30133 FX30237 FX30144 }	Homozygous deletion	I:12586101	I:12586223	(<i>F33E2.7..F33E2.6</i>)	(<i>F33E2.7..F33E2.6</i>)
		Homozygous deletion (1)	V:18429890	V:18465644	<i>srw-83</i>	(<i>ZK228.4..ZK228.12</i>)
		Heterozygous tandem duplication (2)	V:6236672	V:6239149	<i>K08B12.3, K08B12.4</i>	(<i>K08B12.3..K08B12.11</i>)
		Heterozygous deletion	I:11849276	I:11851612	(<i>dhhc-2..ced-1</i>)	<i>ced-1</i>
		Homozygous deletion (3)	V:15826754	V:15886806	<i>srbc-51</i>	<i>cgt-1</i>
		DUP-DEL-LOSS (4)	II:5131837 II:5136381 II:5805620	II:5805620 II:5806749 II:5806749	<i>lgc-56</i> (<i>ggr-3..F58A6.2</i>) (<i>K05F1.10..K05F1.12</i>)	(<i>K05F1.10..K05F1.12</i>) (<i>K05F1.12..msp-64</i>) (<i>K05F1.12..msp-64</i>)
			X:2932305	X:2951620	(<i>F52B10.13..fard-1</i>)	(<i>Y41G9A.15..Y41G9A.13</i>)
		DUP-DUP (5)	X:2939368	X:2940583	<i>Y71H10A.3,</i> <i>Y71H10A.4</i>	(<i>Y71H10A.3..pfk-1.1</i>)
		CGR (6)	IV:16464403 IV:16464489 IV:16469112 IV:16462855	IV:16465989 IV:16465367 IV:16510345 IV:16478758		

The first five lines report the breakpoints of the CRISPR-Cas9 double-inversions. The second part of the table reports additional independent structural variants (SVs) found in the genomes at other loci. In some instances, an SV is shared by different strains such as the “homozygous deletion I:12586101–12586223” present in all five strains. For each event (rearrangement or SV), we also report if the breakpoints are localized in a gene (“gene at left breakpoint” for a left breakpoint mapping in a gene and “gene at right breakpoint” for a right breakpoint mapping in a gene). For intergenic breakpoints, we report the two closest genes between parentheses [e.g., (gene-1..gene-2)]. DUP-DEL-LOSS is a double-event complex genomic rearrangement (CGR) involving a duplication and a deletion, with an addition loss of copy as a secondary event of the rearrangement. DUP-DUP is a double duplication. CGR stands for CGR with more than two DNA-breaking events. Events 1–6 can be visualized on Figure 2A.

^aStructure as published by Dejima et al. (2018).

chromoplexy events are often balanced with little to no change in copy number and arise from nonhomologous end joining rejoining after chromosome pulverization. Chromoplexy involves several chained translocations, whereas chromothripsis is mainly characterized by inversions and rearrangements in often one or occasionally a few chromosomes. However, chromoanasythesis displays multiple copy number variations (CNVs) that emerge during replication from defective DNA repair mechanisms owing to microhomologies at breakpoints (Pellestor et al. 2022).

Here, we set out to use a srWGS approach (Maroille et al. 2021) to systematically detect and characterize the molecular structure and genomic mapping of *C. elegans* genetic balancers. In addition, we report specific SVs and CGRs for each strain included in this study, constituting the first catalog of molecular characterization of complex genomic variation in *C. elegans*. We anticipate that this data set will provide a valuable resource for the extensive userbase of traditional genetic balancers in *C. elegans* genetic studies. In addition, it will give researchers a better appreciation of the complexity of genomic variations, often overlooked in genomic studies (Maroille and Traill-Graovac 2019).

Results

Balancers in *C. elegans* genetic studies

The first balancers in *C. elegans* were reported as early as 1976 (Herman et al. 1976). As previously performed for four traditional balancers (Edgley et al. 2021; Flibotte et al. 2021; Maroille et al. 2021), in this study, we report breakpoints for 17 tested and reliable traditional balancers described by Edgley et al. (2006), which are used to balance over half of the genome (Fig. 1A). The list of selected strain(s) for each balancer is detailed in the Methods and Supplemental Table S1. They were ordered from the *Caenorhabditis* Genetics Center (CGC). Over time, morphologically marked and lethal variants have been generated for many of these balancers. Currently, extremely stable balancers such as *nT1* are carried by more than 500 different strains available from the CGC (Fig. 1B; Supplemental Table S2), suggesting that balancers are a prominent tool in *C. elegans* genetics studies. Since 1991, strains with each balancer have been ordered from the CGC by the *C. elegans* research community many times (Fig. 1C; Supplemental Table S2), illustrating the great use that has been made of the balancers over the past decades. For instance, strains

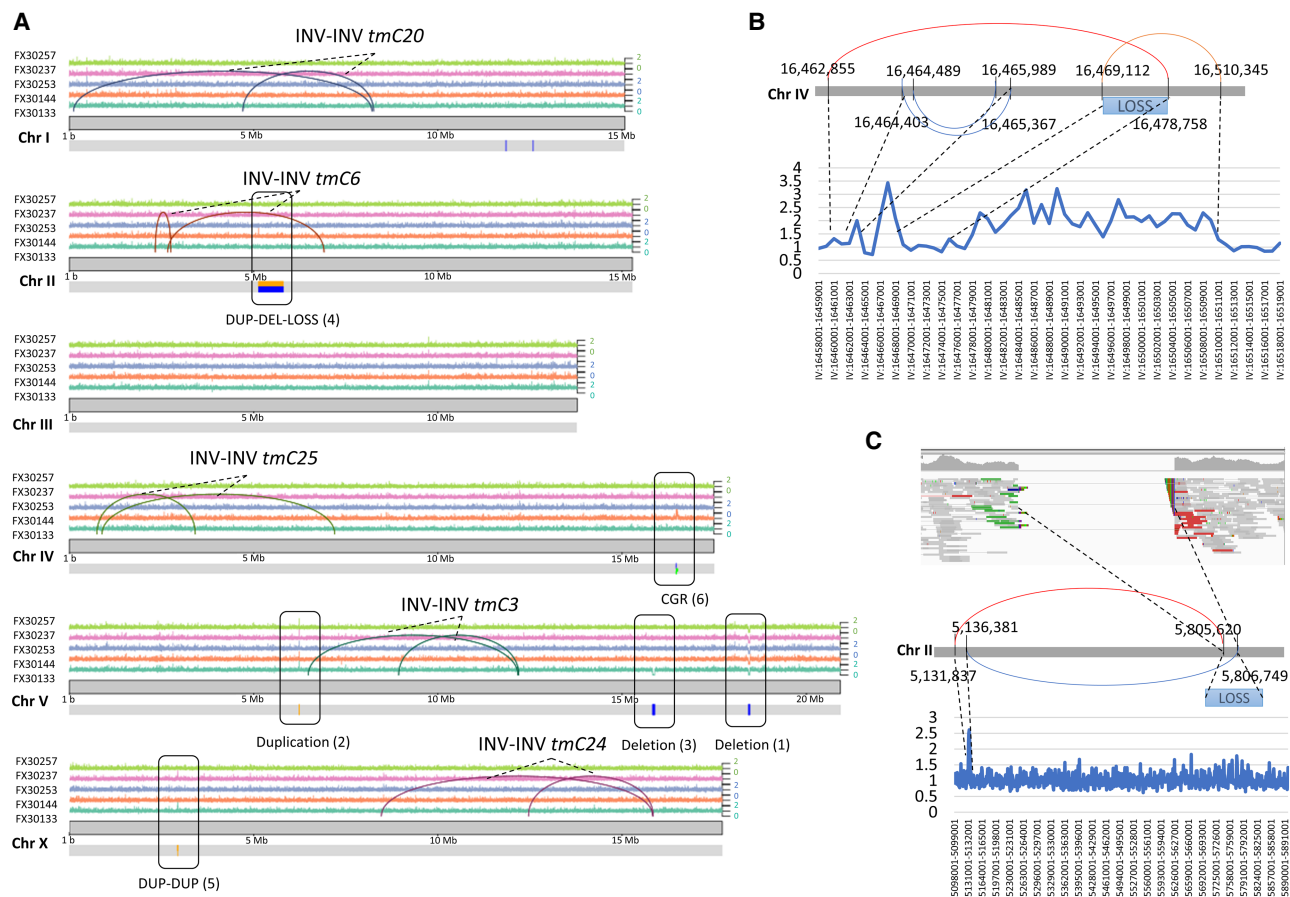


Figure 2. Overview of variants in CRISPR-Cas9 balancers. (A) Map of the CRISPR-ed double inversions and additional rearrangements detected. The upper tracks represent the normalized coverage for each strain. The lower track shows the position of structural rearrangements detected. The color code is green for FX30257, pink for FX30237, blue for FX30235, orange for FX30144, and blue-green for FX30133. The additional events 1–6 are detailed in Table 1. (B, C) Complex genomic rearrangements (CGRs) on Chr IV and Chr II detected in FX30144. Red link represents a copy gain. Blue link represents a loss of copy. Orange link represents a gain of copy with inversion (possible inverted tandem duplication). In addition, we observed loss of copies (LOSS) by visualizing alignments using the Integrative Genomics Viewer (IGV) tool. The LOSS manifested by a complete absence of coverage (no read aligned on that segment of chromosome) but was not supported by split reads. It is possible that the loss of those regions is a consequence of the rearrangement, not a primary event. The graph on the top is a linear schematic representation of the breakpoints and junctions. Blue links represent a loss of copy. Orange link represents a gain of copy with inversion (possible inverted tandem duplication). In addition, we observed loss of copies (LOSS) by visualizing alignments using the Integrative Genomics Viewer (IGV) tool. The LOSS manifested by a complete absence of coverage (no read aligned on that segment of chromosome) but was not supported by split reads. It is possible that the loss of those regions is a consequence of the rearrangement, not a primary event. The scatter plots are a representation of the normalized coverage along the rearrangement calculated by windows of 1 kb. Both scatter plots were adapted to the size of the CGR. In C, the screenshot on the top shows how the reads aligned along the genome (IGV screenshot).

with *nT1* have been ordered at the CGC almost 12,000 times. With over 1500 *C. elegans* laboratories worldwide (as listed in CGC and WormBase as of May 8, 2022), one could estimate that every research group working with lethal or extremely deleterious mutations uses balancers.

Using srWGS to detect known and unexpected high complexity in genomic variation

We first sequenced five balancer strains with well-defined and structurally complicated intrachromosomal rearrangements to reassess our approach with srWGS as published by Maroille et al. (2021). The workflow is summarized in Supplemental Figure S1. These balancers were CRISPR-Cas9 engineered by applying two sequential and overlapping inversions (Dejima et al. 2018).

We successfully retrieved the breakpoints of these CRISPR-crossover suppressors previously described by Dejima et al. (2018): *tmC3* V (FX30133), *tmC6* II (FX30144), *tmC20* I (FX30235), *tmC24* X (FX30237), and *tmC25* IV (FX30257). The precise genomic position of each breakpoint is described in Table

1 and Figure 2 and corroborates the data of their original study (Dejima et al. 2018). Experimental validation data (primers, PCR gels, and sequences from Sanger sequencing) are available in Supplemental Information.

We found eight additional SVs and CGRs not reported before (Fig. 2A–C; Table 1), and possibly specific to these strains. Some were shared by all five strains and are therefore likely originated in the parental strain. But others were unique to only one strain, such as the two CGRs in FX30144 (Fig. 2B,C). These specific SVs were likely secondary targets of CRISPR or the result of spontaneous mutations.

For *tmC20*, the signature of reads for the larger inversion (I:114,579–8,210,593) is suggestive of a CGR involving a duplication and a deletion, both happening at the same breakpoint. Indeed, a usual inversion signature shows pair mate reads with the same orientation, displayed in blue by Integrative Genomics Viewer (IGV) (see Fig. 3A,B; Supplemental Fig. S2) and split reads aligned at the breakpoint junction with opposite directions. Here, we observed two distinct groups of reads. One presented characteristics of a duplication: pairs of reads (marked as green

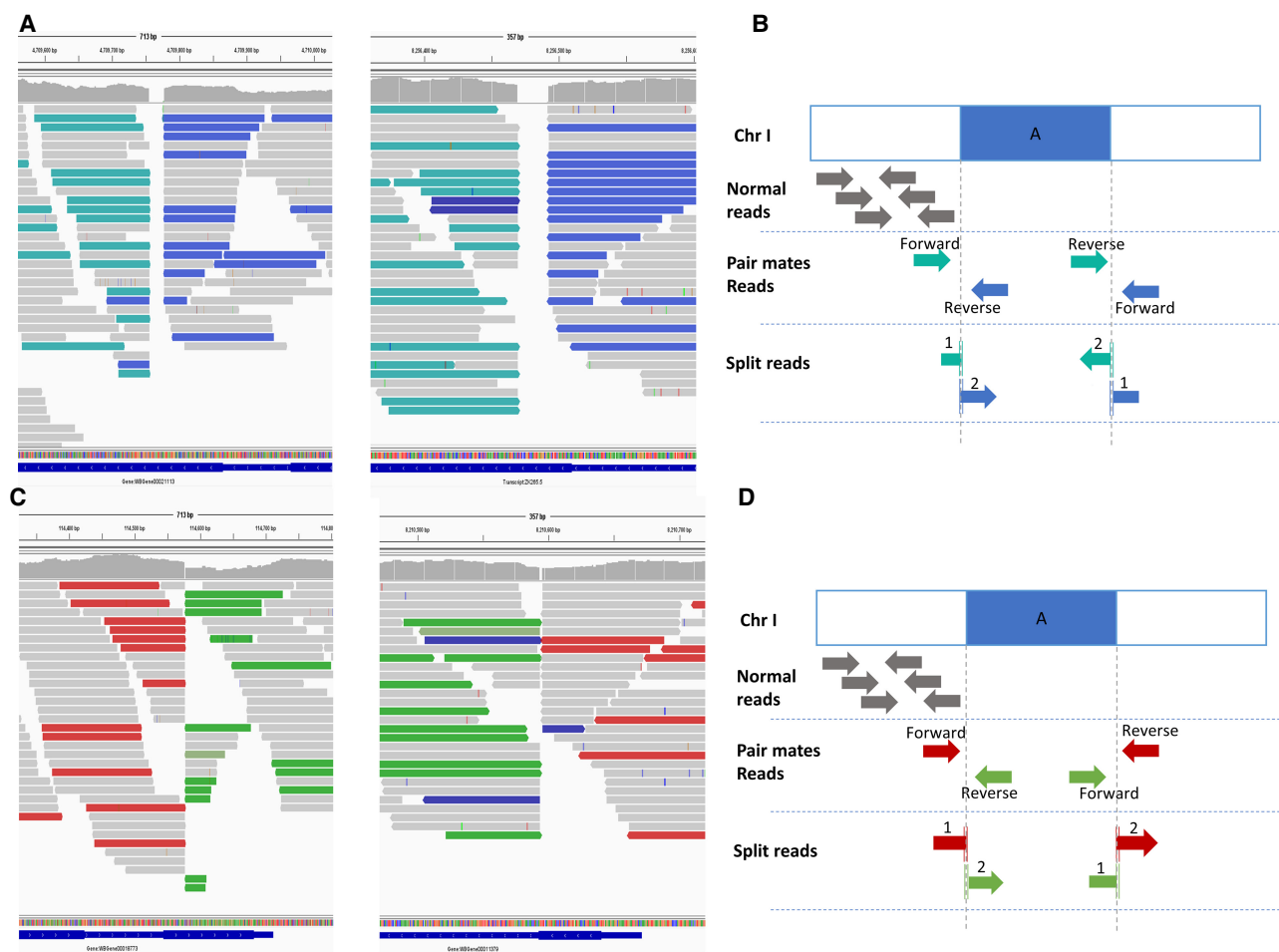


Figure 3. Read signature when the alignments are visualized with IGV and representation of the split reads for inversions in *tmC20* (FX20235). (A,C) IGV screenshots of the regions surrounding breakpoints. (B–D) Schematic representation of split read signatures. (A,B) The small inversion (4.7–8.2 Mb) shows a “normal” inversion signature: The reads in blue and blue-green are marked by IGV as the pair mates (forward and reverse read) have the same orientation instead of opposite orientation. (C,D) The larger inversion (0.1–8.2 Mb) shows a DEL-DUP read signature: Reads in green are marked as such by IGV because the reverse read aligned before the forward. Reads in red are marked as such by IGV as the inner distance between pair mates (forward and reverse reads) is longer than normal inner distance (usually 300–350 bp).

on IGV) (see Fig. 3C,D), for which the reverse read of a pair aligned before the forward read, and split reads, for which the second part of the read aligned before the first part of the read, in the same orientation (Supplemental Fig. S2). The second group of reads had characteristics of a deletion (marked as red on IGV) (see Fig. 3C, D), because the inner distance between pair mate reads is shorter than usual, and split reads for which the second part of the read was aligned further away on the genome, with the same orientation (Supplemental Fig. S2). The origin of such a signature is unclear.

Mapping breakpoints of traditional balancers helps elucidate their effect on phenotypes

We then applied the same methods to decipher the molecular basis of 21 balancers. Four have been already published (Edgley et al. 2021; Flibotte et al. 2021; Maroilley et al. 2021), and we reported breakpoints for 17 additional traditional balancers in this study. Information regarding strains, sequencing, and experimental validation is available in the Methods and Supplemental Tables S1, S3, and S4.

For nine balancers (including *eT1*, previously published by Maroilley et al. 2021), our analyses only detected one large event responsible for balancing one region of the *C. elegans* genome (Fig. 4A; Table 2), as previously described in the literature. For most of these, we are reporting for the first time the exact genomic location, at base-pair resolution, of the breakpoints of each rearrangement and the structure at the junction of the breakpoints: strict, microduplications, or inserted sequence (Fig. 4B).

Often, we determined that balancers are larger than expected (Fig. 1A). For instance, *mT1* was described as a translocation (II;III) balancing the right portion of Chr II from the right end through *dpy-10* (mapped at II:6,710,149–6,712,227) and the right portion of Chr III from the right end to between *daf-2* (III:2,994,514–3,040,846) and *unc-93* (III:3,644,375–3,648,822). We analyzed the genome of SS746 [*k1p-19(bn126)/mT1[dpy-10(e128)] III*]. We detected and confirmed the translocation breakpoints at II:6,296,872 and III:3,635,354, expanding the balanced region of Chr II by ~400 kb.

The balancer *sC1* is described as effectively balancing a 15-map unit portion of Chr III from *unc-45* (III:491547–502625) to near *daf-2* (III:2994514–3040846) (Edgley et al. 2006).

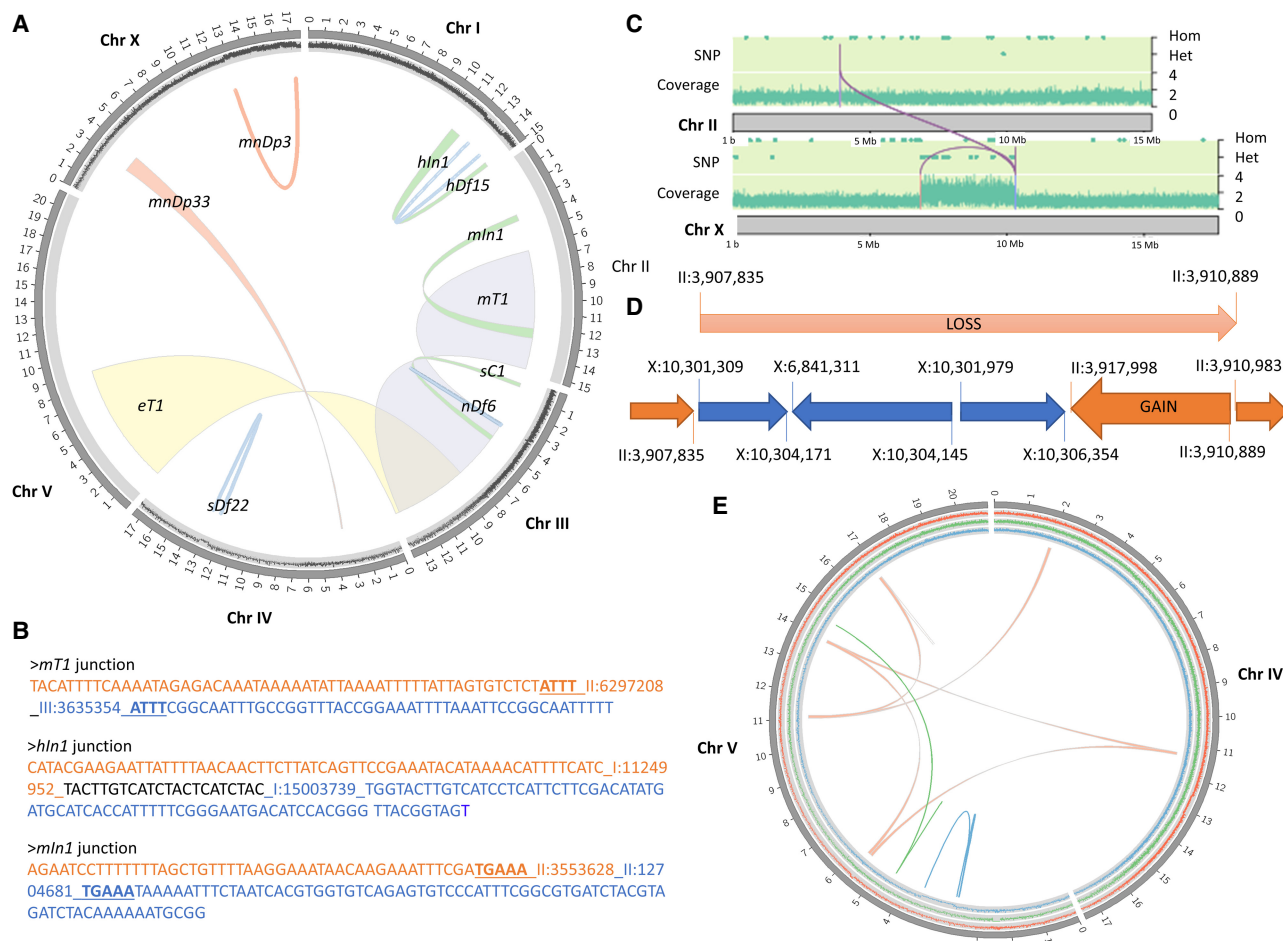


Figure 4. Overview of balancers. (A,B) Single event rearrangements. (A) Circos plot representing the regions affected by single-rearrangement balancers. Green links represent inversions (*hln1*, *mln1*). Blue links represent deletions (*nDf6*, *sDf22*, *hDf15*). Red links represent duplications (translocated duplication *mnDp33* and free duplication *mnDp3*). Ribbons represent translocations (*eT1*, *mT1*). Scatter plots in the outer layer represent the read coverage calculated by window of 1 kb and normalized for KR2839's Chr I, MT690's Chr III, BC1217's Chr IV, and SP123's Chr X. It shows an increase in coverage in case of duplication and a decrease in case of deletion. (B) Breakpoints' junctions for *mT1*, *hln1*, and *mln1*, as confirmed by Sanger sequencing. Orange and blue sequences are from different sides of the junction (different chromosomes for a translocation or different regions of one chromosome in case of inversion). Sequence in black is an inserted sequence at the junction. Bold underlined sequences are potential microduplications at the junction as those bases aligned on both sides of the junction. (C–E) Complex rearrangements. (C,D) *stDp2*. (C) The graph of the upper layer represents by a dot a single-nucleotide variant (SNV) detected in the RW6002 genome. Dots on the higher layer are homozygous SNVs with two copies of the alternate allele. Dots on the middle layer are heterozygous SNVs with only one copy of the alternate allele. Stretches of heterozygous SNVs suggest that the region is balanced by a complex rearrangement. The coverage graph is a scatter plot representing the read coverage of the genome along the chromosome, calculated by window of 1 kb. The graph shows an increase of read coverage of a short region of Chr X. The purple lines represent the breakpoints detected. (D) Interpretation of the structure of *stDp2* based on srWGS analyses. Gain and loss events were detected by coverage analysis only and not supported by split reads, suggesting that they are consequences of the rearrangement, and not the main event. (E) Circos plot representing breakpoints for *nT1* (red), *eDf43* (blue), and *e917* (green).

However, the structure was yet to be described. We identified a large inversion on Chr III from III:0.3 Mb to III:4.6 Mb. Our analyses of srWGS data showed the right breakpoint at the position III:4,641,137. However, the left breakpoint falls into a large repetitive region, and two breakpoints are equally plausible—III:323,321 or III:330,985 (Supplemental Fig. S2), perhaps explaining why the balancer was never molecularly characterized.

We found lesions at some rearrangement breakpoints that could be responsible for observed phenotypes. For example, *mT1* homozygotes are sterile, and we found that the translocation disrupts *linc-95* (Table 2), a long intervening noncoding RNA that may be important for fertility. In addition, the *mln1* inversion disrupts the essential gene *tbc-17* (Table 2), which may explain why homozygous *mln1* animals do not survive freezing well (Edgley and Riddle 2001), although *mln1* also carries *dpy-10(e128)*, and

dpy-10 homozygotes were already known to poorly survive freezing (Levy et al. 1993).

Uncovering CGRs in balancers

The structure of 10 of the balancing rearrangements (including *hT2* [Flibotte et al. 2021], *sC4* [Maroille et al. 2021], and *qC1* [Edgley et al. 2021]) was revealed to be more complex than expected (Table 3).

The balancer *stDp2* (X;II) has been described as a translocated duplication, effectively balancing a small region of the center of Chr X, from around *unc-58* to around *unc-6* (Meneely and Wood 1984; Edgley et al. 2006). Our genomic analysis of the strain RW6002 [*stDp2* (X;II)/+II; *unc-18(e81)* X] revealed a more complex crossover suppressor structure, with 13 breakpoints localized on

Table 2. Breakpoints and structure of one-event traditional balancers

Balancer	Strain sequenced	Left breakpoint	Right breakpoint	Gene at left breakpoint	Gene at right breakpoint	Genes overlapped
Reciprocal translocations						
<i>eT1</i> ^a	BC986, CB4281, VC109	III:8200764	V:8930675	<i>unc-36</i>	(<i>H14N18.7..H14N18.2</i>)	
<i>mT1</i>	SS746	II:6296872	III:3635354	(<i>21ur-14881..C29F5.5</i>)	<i>linc-95</i>	
Inversions						
<i>hIn1</i>	KR2839, VC528	I:11249952	I:15003739	<i>F46A8.11</i>	<i>mtd-1</i>	
<i>mIn1</i>	VC1771	II:3553628	II:12704681	(<i>W10G11.19..dnc-3</i>)	<i>tbc-17</i>	
<i>sC1</i>	DP246	III:323321 or III:330985	III:4641137		<i>pfas-1</i>	
Deletions						
<i>nDf6</i>	MT690	III:3607207	III:3695411			21 genes (<i>ptp-1</i> to <i>npr-15</i>)
<i>sDf22</i>	BC1217	IV:12526376	~IV:13140000	<i>F53B2.8</i>		189 genes
<i>hDf15</i>	KR2839	I:12431742	I:14006044		<i>Y71A12B.12</i>	459 genes
Translocated duplication						
<i>mnDp33</i>	SP309	IV:3518557 ~X:2853000	X:2833653 ~X:4028000	<i>unc-33</i>	(<i>ssr-2..Y71H10B.1</i>)	

For SVs, we report if the breakpoints are localized in a gene (“gene at left breakpoint” for a left breakpoint mapping in a gene and “gene at right breakpoint” for a right breakpoint mapping in a gene). Such genes will be disrupted in case of inversions and translocations. In case of copy number variations (deletions and duplications), overlapped genes will be disrupted (deleted or duplicated). For intergenic breakpoints, we report the two closest genes between parentheses [e.g., (gene-1..gene-2)].

^a(Maroille et al. 2021).

Chr II and Chr X (Table 3; Fig. 4C,D). To summarize, the region on Chr X from around X:6.8 Mb to 10.3 Mb is represented by four copies, instead of two, with an inversion at one of the extremities of the duplicate. Those two additional copies were inserted in Chr II ~3.9 Mb, in tandem, with one copy inverted. This caused rearrangements on Chr II, including the complete loss of the region from II:3,907,835 (first breakpoint linking Chr II and Chr X) to II:3,910,889, with an inversion from II:3,910,889 to II:3,910,983 and a gain of copy of the region until II:3,917,999 (second breakpoint between Chr II and Chr X).

The *nT1* balancer was previously described as a stable reciprocal translocation, effectively balancing the right end of Chr IV through *unc-17* and the left portion of Chr V through *unc-76* (Ferguson and Horvitz 1985; Clark et al. 1988; Rogalski and Riddle 1988; Edgley et al. 2006). We detected a more complex type of rearrangement resembling chromoplexy, with four pairs of breakpoints linking Chr IV and Chr V (Table 3; Fig. 4E), suggesting a shuffling of several segments of those two chromosomes, as well as additional rearrangements at the breakpoints’ junctions.

The balancer *mnC1* still harbored an unknown structure, probably because its complexity had made the characterization of its structure challenging with classical molecular biology techniques (FISH, G-band, and aocGH). *mnC1* has been reported as a very effective balancer for the right portion of Chr II from around *dpy-10* to around *unc-52* (Herman 1978). Overall, we identified five pairs of breakpoints (Table 3), suggesting a complex inversion (three inversions and a deletion) to be responsible for suppressing crossover from II:4,904,692 to II:14,909,258, a region larger than previously described with the gene *dpy-10* being localized at II:6,710,149–6,711,866 and *unc-52* at II:14,647,325–14,684,456 (Supplemental Fig. S3).

The *hDf8* balancer has been described as a deficiency of Chr I and balancing probably from around *unc-13* to at least *dpy-5*, ex-

tending to the left end of the chromosome (McKim et al. 1992; Edgley et al. 2006). Clucas et al. (2002) located the left breakpoint of *hDf8* between the predicted genes *R10A10.1* and *rbx-2*. By the coverage analysis, it seems that the left end of the Chr I is indeed deleted (Supplemental Fig. S4) up to ~800 kb; however, our deeper analysis has also revealed 12 pairs of breakpoints. The structure of this rearrangement involves complex inversions and large copy number gains (Supplemental Fig. S5).

The balancer *mnDp3* is reported as a free duplication of the right end of the Chr X (Herman et al. 1976, 1979; Herman 1984; Edgley et al. 2006). Analyzing the coverage of the SP123 strain (*unc-3(e151) X; mnDp3 (X;f)*) showed an increase at the right end of the Chr X, with a coverage ratio of 1.2, no abrupt increase in coverage, and no split reads around the edges, suggesting indeed a free duplication (Supplemental Figs. S6, S7). By further analyzing the duplication, we retrieved 14 pairs of breakpoints (Table 3; Supplemental Fig. S8). Among them, we identified two junctions between Chr I and Chr X, including one confirmed by PCR and Sanger sequencing. This could suggest that the duplicate region is in fact inserted around the right end of Chr I (Supplemental Fig. S8). However, *mnDp3* has been previously examined and observed by immunofluorescence as free duplication (Lamelza and Bhalla 2012). In addition, both breakpoints fall into a repetitive and telomeric Chr I region and at the extremity of the free duplicate of Chr X. So, such breakpoints could be an artifact owing to the repetitive character of telomeres and the telomeric-like region that allows structural stability to the duplicate without being physically linked to any chromosome.

srWGS uncovers balancer derivatives

For the strains RM2431 and CB3475, the interpretation of their sequenced genome differs from the expected genotype, suggesting

Table 3. Breakpoints (BKPs) of high-complexity traditional balancers

Balancer	Strain (s)	Left BKP	Right BKP	Interpretation	Gene at left BKP	Gene at right BKP
<i>stDp2</i>	RW6002	X:6841311 II:3907835 II:3917998 X:10301979 II:3910889 II:3907835 II:3910984	X:10304171 X:10301309 X:10306354 X:10304145 II:3910983 II:3910889 II:3917999	Translocated duplication and additional rearrangements	<i>asg-2</i> <i>fbxb-103</i> (<i>F53C3.1..plpp-1.1</i>) (<i>F41E7.19..fip-5</i>) (<i>bcmo-2..F53C3.1</i>) <i>fbxb-103</i> (<i>bcmo-2..F53C3.1</i>)	(<i>F41E7.20..fipr-21</i>) (8 bp from <i>F41E7.19</i>) (<i>fipr-21..cnc-8</i>) (<i>F41E7.20..fipr-21</i>) (<i>bcmo-2..F53C3.1</i>) (<i>bcmo-2..F53C3.1</i>) (<i>F53C3.1..plpp-1.1</i>)
<i>nT1</i>	BC1217	IV:1901208 IV:11255068 IV:11254917 IV:1901518 V:11078600 V:16831759 V:5622382 V:11084443	V:11072802 V:13866656 V:5515476 V:11077404 V:16831547 V:16832779 V:13861523 V:16821528	Chromoplexy	(<i>F55A8.6..egl-18</i>) <i>C05C12.1</i> <i>C05C12.1</i> (<i>F55A8.6..egl-18</i>) <i>srh-215</i> <i>srh-215</i> <i>M03E7.1</i> (<i>T04C12.22..T04C12.19</i>)	<i>act-3</i> (<i>srx-137..nhr-175</i>) <i>mms-19</i> <i>act-2</i> <i>T04C12.18, T04C12.24</i> (<i>srh-215..phy-3</i>) (<i>srx-135..srx-137</i>) (<i>T20B3.14..srh-216</i>)
<i>eDf43</i>	CB4281	V:3466642 V:3466718 V:3466648 V:3467157	V:3483655 V:3714673 V:3469548 V:3467175	Duplication Deletion Duplication Insertion	(<i>slc-28.1..gst-33</i>) (<i>slc-28.1..gst-33</i>) (<i>slc-28.1..gst-33</i>) (<i>slc-28.1..gst-33</i>)	<i>srbc-33</i> <i>T27C4.1</i> <i>srbc-29</i> (<i>slc-28.1..gst-33</i>)
<i>e917</i>	CZ1072	V:4489199 V:4489202 V:4488516 V:4489199	V:14577009 V:4488422 V:14577008 V:14577013	Large inversion and rearrangements at breakpoints	(<i>exc-15..unc-62</i>) (<i>exc-15..unc-62</i>) (<i>exc-15..unc-62</i>) (<i>exc-15..unc-62</i>)	<i>gar-3</i> (<i>exc-15..unc-62</i>) <i>gar-3</i> <i>gar-3</i>
<i>hdf8</i>	KR1233	I:836490 I:1113085 I:1802224 I:1801798 I:1966186 I:2814292 I:3294801 I:3294795 I:3315217 I:3535046 I:6419925 I:6420479	I:1063262 I:1119781 I:1802589 I:1802159 I:6412411 I:6915195 I:3326157 I:3294917 I:11034814 I:3536437 I:6910541 I:6915192	Complex genomic rearrangement	(<i>zfh-2..ZC123.1</i>) <i>ZK993.5</i> <i>Y71G12B.5</i> <i>Y71G12B.5</i> <i>Y71G12B.5</i> (<i>YS1F10.11..YS1F10.3</i>) <i>itx-1</i> (<i>W01B11.6..W01B11.1</i>) (<i>W01B11.6..W01B11.1</i>) (<i>Y44E3A.7..trx-4</i>) <i>Y47G6A.5</i> (<i>R10A10.1..rbx-2</i>) (<i>R10A10.1..rbx-2</i>)	<i>npr-23, vps-4</i> (<i>ZK993.5..ZK993.4</i>) <i>Y71G12B.5</i> <i>Y71G12B.5</i> <i>ZC328.3</i> <i>C48B6.3</i> <i>cdap-2</i> (<i>W01B11.6..W01B11.1</i>) <i>dxbp-1</i> <i>Y47G6A.5</i> <i>smg-1</i> <i>C48B6.3</i>
<i>mnC1</i>	SP127	II:4904692 II:4904677 II:7535181 II:9692491 II:7535124	II:14909258 II:7535185 II:14909251 II:9695040 II:14909253	Complex inversion	(<i>R05F9.14..btbd-10</i>) (<i>R05F9.14..btbd-10</i>) <i>ZK1290.13</i> (<i>nlp-11..Y53C12A.8</i>) <i>ZK1290.13</i>	<i>R03C1.1</i> <i>ZK1290.13</i> <i>R03C1.1</i> (<i>nlp-11..Y53C12A.8</i>) <i>R03C1.1</i>
<i>mnDp3</i>	SP123	~X:12805000 X:12983109 X:15422113 X:15419295 I:15064452 X:15437541 X:14998958 X:17132213 X:17132318 X:15434592 X:13175974 X:12974200 X:14943711 X:14954977 X:12806419	Right end LG X X:12983352 X:15796219 X:15421670 X:15484795 X:15438062 X:15009370 X:17133014 X:17133117 X:15436210 X:13382739 X:12984629 X:14954885 X:14943581 I:15069556	Complex rearrangement of a free duplication	<i>nhr-27, gei-3</i> <i>scav-4</i> <i>F39D8.3</i> <i>psa-3</i> <i>rrn-2.1</i> <i>mocs-1</i> <i>cutl-3</i> <i>K09E3.7</i> <i>K09E3.7</i> <i>flp-32</i> (<i>mob-2..nlp-3</i>) <i>ppk-3</i> <i>mrp-5</i> <i>F45B8.13</i> <i>nhr-27, gei-3</i>	<i>scav-4</i> (<i>Y7A5A.10..Y7A5A.t3</i>) <i>F39D8.3</i> <i>gcy-36</i> <i>mocs-1</i> (<i>ergj-2..C18B12.12</i>) <i>K09E3.7</i> <i>K09E3.7</i> <i>mocs-1</i> (<i>C49F8.6..C49F8.3</i>) <i>scav-4</i> <i>F45B8.13</i> <i>mrp-5</i> <i>rrn-1.2</i>
<i>hT2^a</i>	VC386	5 breakpoints	5 breakpoints		Complex inversion	
<i>sC4^b</i>	BC4586	4 breakpoints	4 breakpoints		Large deletion and chromosome fusion	
<i>qC1^c</i>		8 breakpoints	8 breakpoints		Complex inversion	

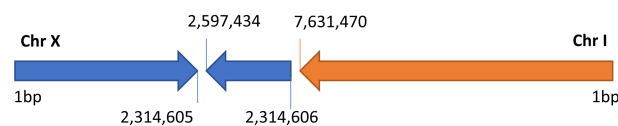
If a breakpoint is localized in a gene and disrupting it, we report such gene (“gene at left breakpoint” for a left breakpoint mapping in a gene and “gene at right breakpoint” for a right breakpoint mapping in a gene). For intergenic breakpoints, we report the two closest genes between parentheses [e.g., (gene-1..gene-2)].

^a(Flibotte et al. 2021); ^b(Maroilley et al. 2021); ^c(Edgley et al. 2021).

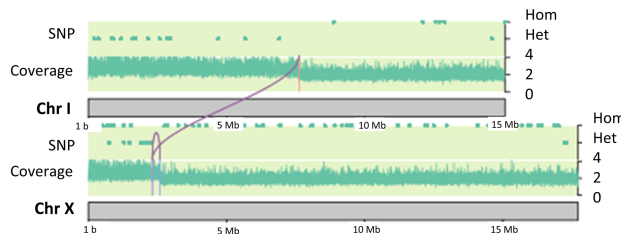
A

Balancer	Strain(s)	Left breakpoint	Right breakpoint	Interpretation	Gene at left breakpoint	Gene at right breakpoint
<i>hT1</i> (derivative)	RM2431	I:8409987	V:7207631	Translocation	<i>lin-28</i>	<i>soap-1</i>
		Chr I Left end	~I:8.4 Mb	Free duplication		
		Chr V Left end	~V:7.2 Mb	Loss		
<i>szT1</i> (derivative)	CB3475	I:7631470	X:2314605	Translocation	<i>lrp-1</i>	<i>hum-6</i>
		X:2314606	X:2597434	Inversion	<i>hum-6</i>	<i>nhr-40</i>
		Chr I Left end	I:7.6 Mb	Free duplication		
		Chr X Left end	X:2.5 Mb			

B



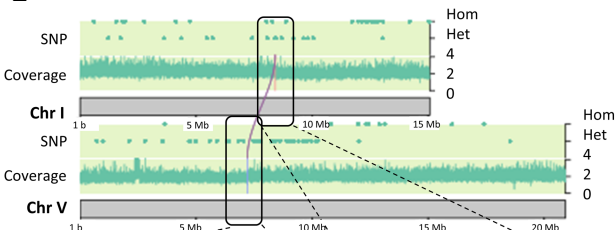
C



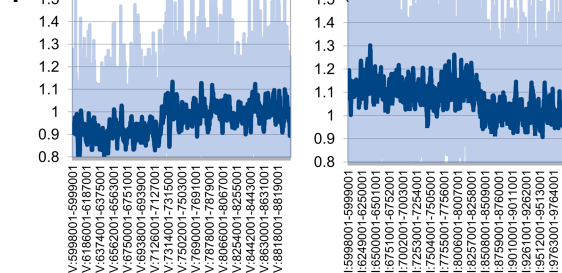
D

>*szT1* (derivative) junction
 TGTGATAGAAAGGATGATTGTGGAGATGCTTCTGATGAAATGGATGCTGTAAGTTTCAAT
 GATTCAGTACTGTGAGAAATTT_I:7631470_TCAA_X:2314605_ATCAAATAATCACCC
 ATCAAANAATAGGGAAGTGGCATTATAGATTGTTAAAGTTATTTTTTTCTTTCTT

E



F



G

>*hT1* (derivative) junction
 ACAGAGAAACCGACAGGAGGGCATTGGCGGAGGAGCCGTGTGCGCGCACT
 _I:8409987_V:7207631_ACTGCTGCAGTCTCGGAGGATAGTTCGATTCCCGAC
 GGTAGTCAAAGT

Figure 5. Overview of potential derivative balancers. (A) Breakpoints of *hT1* and *szT1* derivatives. If the breakpoints are localized in a gene and disrupting it, we report such gene (“gene at left breakpoint” for a left breakpoint mapping in a gene and “gene at right breakpoint” for a right breakpoint mapping in a gene). (B–D) *szT1* derivative. (B) Interpretation of the structure of *szT1* derivative in CB3475 based on srWGS analyses. (C) In the graph of the *upper* layer, SNVs detected in the CB3475 genome are denoted by dots. Dots on the higher layer are homozygous SNVs with two copies of the alternate allele. Dots on the *middle* layer are heterozygous SNVs with only one copy of the alternate allele. Stretches of heterozygous SNVs suggest that the region is balanced by a complex rearrangement. The coverage graph is a scatter plot representing the read coverage of the genome along the chromosome, calculated by window of 1 kb. The graph shows an increase in read coverage of left parts of Chr I and X. The purple lines represent the breakpoints detected. (D) Sequence at the junction of the breakpoint (I;X) obtained by Sanger sequencing. (E,F) *hT1* derivative. (E) In the graph of the *upper* layer, SNVs detected in the RM2431 genome are denoted by dots. Dots on the *higher* layer are homozygous SNVs with two copies of the alternate allele. Dots on the *middle* layer are heterozygous SNVs with only one copy of the alternate allele. Stretches of heterozygous SNVs suggest that the region is balanced by a complex rearrangement. The coverage graph is a scatter plot representing the read coverage of the genome along the chromosome, calculated by window of 1 kb. The graph shows an increase in read coverage of left parts of Chr I and X. The purple lines represent the breakpoints detected. (F) Scatter plots focused on the regions where the ratio of coverage changes on Chr I and Chr V. The normalized coverage along the rearrangement calculated by windows of 1 kb. The darker line represents the moving average (trend line, calculated over 20 points). (G) Sequence at the junction of the breakpoint (I;V) obtained by Sanger sequencing.

that such strains might present a derivative of traditional balancers.

The rearrangement *szT1* balances the left portion of Chr I through *unc-13* and nearly all of Chr X from the right end to around *dpv-3*. It was previously described as a reciprocal translocation (I;X), with rare exceptional progeny carrying one half-translocation as a complex free duplication (Fodor and Deák 1985; McKim et al. 1988; Howell and Rose 1990; McKim and Rose 1990; Edgley et al. 2006). We analyzed the genome of the strain CB3475 (*mcm-4* (*e1466*)/*szT1* [*lon-2*(*e678*)]I; +/*szT1* X). Our initial analysis showed a signature of a translocation (I;X) with breakpoints at I:7,631,470 and X:2,314,605 and an inversion X:2,314,606–2,597,434 (Fig. 5A–C; Table 3).

By coverage analysis (Fig. 5C), we determined that both segments of Chr I (from left end to ~I:7.6 Mb) and Chr X (from left end to ~X:2.5 Mb) were duplicated, with a coverage ratio of around 1.2, which suggests that both can occur as free duplications. It is supported also by the stretch of heterozygous variants on the left end of Chr I with unbalanced representation of the alternative allele. Indeed, heterozygosity is usually determined as ~50% reads supporting the reference allele; ~50%, the alternative allele. Visual inspection of the alignment for variants on the right ends of Chr I and X (listed in Supplemental Table S7) showed a proportion of reads supporting the reference allele different from the expected 50/50. That reflects the presence of an additional mosaic copy of the region.

Table 4. Subset of SVs and CGRs detected in *C. elegans* genomes and validated by PCR and Sanger sequencing

Type	Strain	Left breakpoint	Right breakpoint	Length	Genes	
Homozygous deletion	DP246	I:12824847	I:12825125	278 bp	<i>gsk-3</i>	
	GE2722	I:11148805	I:11149851	1046 bp	<i>clec-95</i>	
	DP246	IV:3052310	IV:3052413	103 bp	<i>Y67D8C.23</i>	
	RW6002 ^a	IV:5370371	IV:5371720	1349 bp	<i>F41H10.3</i>	
Heterozygous deletion	SP127	V:4694692	V:4694952	260 bp	<i>T22B11.1</i>	
	SP127	IV:16029231	IV:16029350	119 bp	<i>21ur-14347, 21ur-2569</i>	
	BC1217	IV:12417282	IV:12418048	766 bp	<i>F08G5.t2, F08G5.21, F08G5.3</i>	
	RM2431 ^a	I:14338214	I:14338233	20 bp	<i>F49B2.6</i>	
	DP246	III:5402384	III:5402454	69 bp	<i>Y32H12A.8</i>	
	RM2431	X:7095702	X:7095968	266 bp	<i>C36B7.5</i>	
Homozygous direct tandem duplication	CZ1072 ^a	I:10565986	I:10577022	11,042 bp	<i>rfa-0, F25H2.16, tct-1, F25H2.14, F25H2.12, rtel-1</i>	
	BC1217	X:14974538	X:14993793	19,255 bp	<i>tbx-1, F20D1.3, mir-1829.2, emre-1, arl-7, plp-2, rbg-1, iglr-1, F20D1.16</i>	
	BC1217	III:5059444	III:5063035	3591 bp	<i>C45G9.10, C45G9.4</i>	
	KR2839	I:10565986	I:10577022	11,036 bp	<i>pas-5, rfa-0, F25H2.16, tct-1, F25H2.14, F25H2.12, rtel-1</i>	
Heterozygous tandem duplication	CZ1072	III:3891219	III:3898488	7279 bp	<i>tir-1</i>	
	BC1217	X:1509128	X:1515794	6666 bp	<i>toca-1</i>	
	MT690	X:11845164	X:11852389	7225 bp	<i>F54F7.2, F54F7.11, F54F7.9, F54F7.3, F54F7.14</i>	
	SP309	II:3495711	II:3513352	17,641 bp	<i>Y49F6B.22, Y49F6B.9, Y49F6B.21, col-71, Y49F6B.15, Y49F6B.2, cyh-1, linc-71, Y49F6B.19, Y49F6B.17, Y49F6B.16, srz-5</i>	
	BC1217	IV:14576313	IV:14582569	6256 bp	<i>Y57G11B.223, Y57G11B.8, irdl-18</i>	
	RW6002	III:11830810	III:11834895	4085 bp	<i>rsp-8, ntl-3</i>	
	DP246	V:16914510	V:16928379	13,869 bp	<i>nhr-116, F09C6.10, fbxa-206, Y102A5C.2, fbxa-207, Y102A5C.4, Y102A5C.35, Y102A5C.36, Y102A5C.5</i>	
	DP246	II:12235220	II:12305725	70,505 bp	<i>19 genes</i>	
	RM2431	IV:12055597	IV:12069818	14,221 bp	<i>spt-5, K08E4.12, K08E4.2, K08E4.3, K08E4.10, K08E4.11, K08E4.8, bath-37</i>	
	RM2431 ^a	I:1307000	I:1308796	1796 bp	<i>dapk-1, K12C11.8</i>	
	GE2722	V:4405080	V:4417828	12,747 bp	<i>nhr-118, Y47D7A.3, Y47D7A.17, grl-5, Y47D7A.18, Y47D7A.6</i>	
	KR2839, VC528	I:14573853	I:14580325	6472 bp	<i>Y105E8A.25, Y105E8A.53, Y105E8A.55</i>	
	Inverted Tandem Duplication	CZ1072	II:15271369	II:15273155	1786 bp	<i>rga-9</i>
		RW6002	III:165133	III:171148	6019 bp	<i>F40G9.7, F40G9.18, F40G9.21, F40G9.22, F40G9.9</i>
RM2431		V:18866326	V:18866821	495 bp	<i>emb-4</i>	
SP423 ^a		V:16305	V:17193	888 bp		
Inversion	BC1217 ^a	IV:4399382	IV:4401161	1783 bp	<i>ell-1, Y24D9A.6</i>	
	BC1217	IV:8065618	IV:8066587	969 bp	<i>sax-7</i>	
	RM2431	II:8513281	II:10124446	1,611,165 bp	<i>232 genes</i>	
TD-DEL-DEL	VC1771	II:11113451	II:11113761	310 bp	<i>apc-1</i>	
	GE2722	II:6758980	II:6763714	4738 bp	<i>F41G3.3, agr-1</i>	
	GE2722	II:6760512	II:6761141	629 bp	<i>F41G3.3, agr-1</i>	
DUP-DEL	GE2722	II:6761122	II:6761844	722 bp	<i>agr-1</i>	
	DP246	III:11120259	III:11121729	1471 bp	<i>Y48A6C.4, Y48A6C.12</i>	
	DP246	III:11112143	III:11129064	16,921 bp	<i>15 genes</i>	

TD-DEL-DEL is a CGR involving a tandem duplication and two deletions. DUP-DEL is a CGR with one duplication and one deletion overlapping.

^aSee Integrative Genomic Viewer screenshot in Supplemental Figure S2.

After validation of the breakpoints by PCR and Sanger sequencing (Fig. 5D), we then interpreted this structure as both duplicates being attached together, with a 280-kb inversion at the extremity of Chr X (Fig. 5A–C). However, this structure does not corroborate that *szT1* balances Chr X from the right end. Indeed, our data showed no stretches of heterozygous SNVs from the right end of Chr X to the translocation breakpoint (Supplemental Table S7). In addition, a derivative of *szT1*, called *szDp1*, has been previously reported (Edgley et al. 2006), with one of the half translocations *szT1(X)* carrying a portion of Chr I that was present as a duplication in addition to two normal copies of Chr I and two normal copies of Chr X. It is then possible that the genome of CB3475 sequenced in this study carries a derivative of *szT1*.

The balancer *hT1* was described as balancing the left portion of Chr V from the left end through *dpy-11* (V:6511583–6515240) (McKim et al. 1988; Howell and Rose 1990). By analyzing the strain RM2431 [*unc-13(md2415)/hT1 I; +/hT1 V*], we showed that the breakpoint is localized ~0.8 Mb further away (V:7,207,631) in the *soap-1* gene. In addition, *hT1* was described as balancing the left portion of Chr I from the left end through *let-80*, a gene with no known genomic position. Such lack of information could restrict the usage of this balancer. Now, we have completed the description by mapping the second breakpoint of the reciprocal translocation in the *lin-28* gene at the position I:8,409,987.

In our hands, the strain selected for sequencing (RM2431) also shows on the left end of Chr I an increase in read coverage until around I:8,735,001 (coverage ratio of 1.2). Ratio of coverage was

calculated by dividing the read coverage by window of 1 kb, divided by the average of coverage of the entire genome, and normalized against the ratio in control (N2 strain; see Methods). Usually, for regions with no variations of copy number, the ratio equals one. For deletion, the ratio equals zero (homozygous) or 0.5 (heterozygous; loss of one copy). And for tandem duplications, the ratio is 1.5 (heterozygous) or two (homozygous). A ratio of 1.2 is only observed in case of free duplications and translocated duplications, which is, in this case, probably suggestive of a free duplication (Fig. 5E,F). It also showed a decrease in coverage for the left end of Chr V (Fig. 5E,F). With the experimental validation of the inter-chromosomal breakpoint junction (Fig. 5G), our data show that the strain RM2431 is not a clean translocation but presents a free duplication. The rearrangement *hDp133*, a derivative of *hT1*, has been previously reported to produce rare exceptional progeny that could carry one half-translocation as a complex free duplication (Edgley et al. 2006). Altogether, this suggests that the genome of RM2431 sequenced in our study carries a derivative of *hT1*.

Unsuspected complex variation of *C. elegans* balancer genomes beyond balancing variants

Besides balanced regions previously partially characterized, we explored the entire genomes of all the strains (for complete list and phenotypes, see Supplemental Information). Overall, we found an average of six additional pairs of breakpoints per strain (range 1–17), suggesting the existence of large variation in *C. elegans* genomes owing to SVs and CGRs being understudied. We ignored breakpoints related to insertion events (e.g., mobile element insertions).

Earlier in this paper, we described eight SVs and CGRs in the five CRISPR-Cas9 strains (Fig. 2; Table 1). In addition, beyond the regions balanced by the traditional balancers they carry, we explored the entire genomes of 20 strains (18 with one balancer, two with two balancers) (Supplemental Table S1). We report tandem duplications (direct and inverted), deletions, inversions, and CGRs of all sizes and on all chromosomes. Table 4 displays a subset of SVs and CGRs detected in traditional balancer strains, further validated by PCR and Sanger sequencing (PCR gels are available in Supplemental Information, and Sanger sequences in Supplemental Table S5). The complete list of the 96 additional rearrangements detected by RUFUS (Ostrander et al. 2018) is available in Supplemental Table S5 (32 homozygous deletions, 10 heterozygous deletions, four homozygous tandem duplications, 22 heterozygous tandem duplications, six inverted tandem duplications, three inversions, and 19 CGRs).

Of note, we have only reported SVs that were not present in our N2 reference strain and CB4856 strain. Still, some SVs were found in several strains such as the tandem duplication I:14573853–14580325 detected in KR2839 and VC528, or the homozygous deletion II:10440527–10440558 present in the genomes of the strains SP123, KR1233 and VC528.

Discussion

In this study, we have generated the first molecular map of *C. elegans* balancers with genomic locations of their breakpoints and characterized for the first time the structure of a subset of these balancers. We also elucidated higher complexity for other balancers, which is an unsuspected finding based on previous analysis with low-resolution genetic techniques. Accumulation of spontaneous

short mutations has been previously observed in balanced regions (Rosenbluth et al. 1983), and here, we generated the first atlas of molecularly dissected and experimentally confirmed genomic variations based on SVs and CGRs, still understudied in model organisms.

By applying our sequence analysis protocol published by Maroille et al. (2021), we successfully used the short-read WGS approach to map breakpoints for all types of SVs (inversions, deletions, tandem duplications, inverted tandem duplications, translocated duplications, reciprocal translocation, insertions) and even complex rearrangements (chromoanagenesis-like). In addition, we were able to detect such rearrangements as a free duplication, which would translate in human studies as a mosaic complex rearrangement. We then showed that short reads with tailored bioinformatic downstream analyses can contribute more to sequence analysis than what is appreciated in most previous studies, which tend to focus srWGS applications to SNVs, short insertions, and deletions (Maroille and Tarailo-Graovac 2019). With such an approach, whole-genome sequencing based on short reads constitutes an interesting way to explore, in a high-throughput manner, the entire spectrum of genomic variability in *C. elegans* and other model organisms.

In some instances, the analyses of the genomes of strains selected for specific balancers, such as *szT1* but also *hT3* and *mnDp1* (for discussion, see Supplemental Information), showed a different structure than previously described and, more importantly, a discrepancy regarding the regions balanced. It has been well described that some balancers are unstable (Edgley et al. 2006). Our work further supports this and advocates that balancer strains be either sequenced by the laboratories or at least PCR-tested (facilitated by primers designed in our study) to validate the expected strains and the balanced region when using the balancer strains.

Despite the successful application of the srWGS in our work to detect a diverse spectrum of SNVs as well as SVs and CGRs, we acknowledge that identification of certain types of complex variants is quite challenging using short reads, considering the known limitations of the technology. First, one read cannot overlap the entire rearrangement. Detection and interpretation of breakpoints rely then on the presence of split reads at breakpoint junctions and the ability of bioinformatics processes to recognize split reads, reliant on tailored algorithms but also the coverage and the quality of sequencing data (Supplemental Fig. S9). Second, the alignment of the reads is influenced by the aligner used and the genome reference that might not be ideal for every sample. For instance, the *C. elegans* reference genome was based on N2 strain, which may only partially represent the variability of other strains and create challenges during read alignment. Third, the resolution of the structure of complex rearrangements with short reads is subjected to one's interpretation of the variation of coverage and split reads at breakpoint junctions, which may be challenging. Finally, there is a possibility that some breakpoints located in the highly repetitive regions may be missed (Supplemental Fig. S10; Supplemental Table S8). Therefore, in more complex genomes, such as those of wild isolates (Lee et al. 2021) or even those of some challenging rearrangements in our study (e.g., *szT1*), it may be useful to apply long reads to resolve the complex variants that are within highly variable and repetitive regions.

We detected several rearrangement breakpoints mapping in genes, including some essential genes (e.g., *soap-1*), and even in potentially essential, previously unexplored long intervening non-coding RNAs (*linc-95*). It is challenging to interpret the effect of such disruption on the expression of genes and resulting

phenotypes. Indeed, although various tools that interpret the effects of single-nucleotide variants and indels in coding regions have been developed, only a few tools exist that predict in silico the effect of SVs. It is also important to consider the impact of breakpoints in noncoding regions, as they could disrupt genes by falling in introns or disrupt regulatory mechanisms by falling into a topological associated domain, thereby disrupting the link between a gene and an enhancer.

We have also reported additional SVs and CGRs identified with srWGS in the strains included in this study. Our findings revealed that most of these balancer strains are carrying various types of large events, some of them apparently owing to mutagenesis but others likely as part of the spontaneous genetic variation in *C. elegans*. Like Newman et al. (2015) have reported, we found translocated duplications and free duplications to be quite rare compared with tandem duplications. In addition, we observed that CGRs, resulting from several overlapping SVs, are more frequent in these balancer genomes than previously appreciated. It is tempting for investigators to assume that balancer strains behave essentially as wild type in all aspects except their balancing behavior. However, by showing that balancer strains also carry various SVs, beyond the expected rearrangement that results in balancing, our study indicates that some SVs might cause additional phenotypes or interact with the balanced mutations. Then, variations of genetic background in balancer strains should be considered, and balanced mutations should be analyzed in more than one background. Beyond the mutagen-induced balancers, our analysis of CRISPR-Cas9-induced balancers revealed some interesting findings. First, some inversions induced by CRISPR-Cas9 showed a signature different from the typical inversion signature. The nature and origin of this signature will need further investigation. Furthermore, we identified additional SVs that were not previously known. Some were shared by all five strains and thus were probably inherited from the background genome used to create those balancers. But other SVs were only present in one or two of the CRISPR-Cas9 balancer genomes. It would be interesting to investigate whether either of these variants occurred naturally or whether they could be off-site targets of the RNA guide.

Overall, this atlas of molecular structure of genetic balancers is very important for the *C. elegans* community because balancers are widely used genetic tools. Indeed, by describing their structures, we gain a better understanding of how the balancers are working. It also facilitates balancer usage by mapping exactly their breakpoints for an easy confirmation of their presence in strains and better awareness of the capabilities of each balancer (specifically, which exact region can be balanced by each rearrangement) (Fig. 1A; Supplemental Table S6). Finally, the description of the exact position of breakpoints disrupting genes in rearrangements reported to be recessively lethal could be very important for *C. elegans* genetic studies of currently understudied genes. On a larger scale, this study clearly shows the power of srWGS in resolving genomic complexity and paves the way to tailored bioinformatic workflows for both model organism and human studies. Such workflows would better take advantage of srWGS data sets, reduce cost, and reduce the multiplication of techniques necessary to explore the whole complexity of the genomes.

Methods

CGC data

For each rearrangement as balancer, we used the CGC database to retrieve the list of strains carrying the balancer of interest. We used

the CGC query system “search strain” and filtered by phenotype. We manually reviewed the final lists to avoid picking up genotypes that would contain a marker/gene with an identification similar enough to the balancer to be picked up by the query. We then sent those lists of strains to the CGC to obtain the number of orders of each strain over time. We calculate the number of orders by balancer by adding up order for each strain carrying each balancer.

Worm maintenance

All worm strains in this study were obtained from the CGC. All strains were maintained at 16°C and kept on standard NGM plates streaked with OP50.

Sequencing

We sequenced the genomes of 25 strains, for a total of 30 genomes: genomes from homozygous individuals of the strains BC1217, CB3475, CB4281, CZ1072, DP246, FX30133, FX30144, FX30235, FX30237, FX30257, GE2722, KR1233, KR1876, KR2839, MT690, RM2431, RW6002, SP123, SP127, SP309, SP423, SP998, VC1771, VC471, VC528; genomes from heterozygous worms of the strains DP246, SS746, VC1771, VC471, and VC528. We used the sequencing data of N2 and CB4856 strains published by Maroilley et al. (2021). The strains FX30133, FX30144, FX30235, FX30237, and FX30257 were engineered by CRISPR-Cas9 (Dejima et al. 2018). The strain BC1217 was used to explore *sDf22* and *nT1*. The strain GE2722 was used to retrieve *qC1* breakpoints published by Edgley et al. (2021). The strain VC471 carried *hT2*, already described by Flibotte et al. (2021). The strain KR2839 was used to describe *hDf15* and *hIn1*. The strains KR1876 and SP423 were originally selected to study *hT3* and *mnDp1*, respectively, but the sequencing data did not allow the resolution of those two balancers (see Supplemental Information). All genomes sequenced in this study were used to build the catalog of structural variations.

DNA extraction and library preparation

Genomic DNA was collected from ~100 mg of worm tissue using the Qiagen blood and tissue kit (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. Paired-end short-read WGS were obtained for all strains with PCR-free library preparation protocol and NovaSeq 6000 Illumina sequencing technology.

Primary data analyses

We checked the quality of the FASTQ files using FastQC (<http://www.bioinformatics.babraham.ac.uk/projects/fastqc>). The reads were on average 151 bp long. We trimmed the reads and removed the adapters using Trimmomatic v0.36 (Bolger et al. 2014). For each sample, we aligned between 16 million and 34 million reads using the BWA-MEM v0.7.17 (Li 2013) algorithm to the *C. elegans* reference genome WS265. It resulted in a 31× read coverage per strain on average (Supplemental Table S3). We then sorted the reads according to their coordinates with “samtools sort” (SAMtools v1.5, Li et al. 2009).

For all genomes, we performed an analysis of coverage by calculating the read depth by window of 1 kb. We normalized the average of read coverage in each 1-kb window by dividing it by the average of read depth across the entire genome and by the read depth of the same window in the reference strain (N2). If N2 ratio

was equal to zero in a specific window (no reads in the N2 strain sequenced in our lab), the ratio reported for further analyses was then only normalized by the average coverage of the genome. The ratio obtained was then plotted according to its genomic position on a graph using the R package “karyoploteR” (Gel and Serra 2017; https://bernatgel.github.io/karyoploteR_tutorial/; <https://github.com/bernatgel/karyoploteR>). The Circos plots on Figure 4 were realized using Circos (Krzywinski et al. 2009; <http://circos.ca/>).

Breakpoints were identified as in the work of Maroille et al. (2021). Briefly, we used the *k*-mer approach implemented in RUFUS (<https://github.com/jandrewfarrell/RUFUS>) to detect de novo breakpoints, using N2 and Hawaiian strains as “parents.” Additional breakpoints were detected using GRIDSS (<https://github.com/PapenfussLab/gridss>) when a CGR was suspected. All breakpoints were assessed visually using IGV, which allows the visualization of the reads aligned along the reference genome (see Supplemental Information; Robinson et al. 2011). We performed a read inspection for each breakpoint. We extracted reads around each breakpoint using “samtools view” (SAMtools v1.5) and realigned them using BLAT (UCSC, portable version). The presence of several split reads supporting the breakpoints was decisive whether or not to send the breakpoint for experimental validation. We extracted the sequence of one split read for each breakpoint to design primers for PCR and Sanger sequencing. In the absence of split read at the breakpoint, we looked for discordant pair of reads.

Independent analyses

For the balancers *nT1* and *mln1*, an independent analysis was performed as in the work of Flibotte et al. (2021) and Edgley et al. (2021). In both cases, the same breakpoints were retrieved.

Experimental validation

We confirmed breakpoints of SVs and complex rearrangements by PCR and Sanger Sequencing. Primers were design with PrimerQuest IDT tool (<https://www.idtdna.com/PrimerQuest>). All primers and sequences are available in Supplemental Tables S4 and S5.

Data access

The data generated in this study have been submitted to the NCBI BioProject database (<https://www.ncbi.nlm.nih.gov/bioproject/>) under accession number PRJNA838950. The codes for Figures 1A, 2A, 4C, 5C, and 5E are available in the Supplemental Material and at GitHub (<https://github.com/MTG-Lab/C-elegans-balancers>).

Competing interest statement

The authors declare no competing interests.

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