# Engineering the Caenorhabditis elegans genome using Cas9-triggered homologous recombination 

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#### Abstract

Study of the nematode Caenorhabditis elegans has provided important insights in a wide range of fields in biology. The ability to precisely modify genomes is critical to fully realize the utility of model organisms. Here we report a method to edit the C. elegans genome using the clustered, regularly interspersed, short palindromic repeats (CRISPR) RNA-guided Cas9 nuclease and homologous recombination. We demonstrate that Cas9 is able to induce DNA double-strand breaks with specificity for targeted sites and that these breaks can be repaired efficiently by homologous recombination. By supplying engineered homologous repair templates, we generated gfp knock-ins and targeted mutations. Together our results outline a flexible methodology to produce essentially any desired modification in the C. elegans genome quickly and at low cost. This technology is an important addition to the array of genetic techniques already available in this experimentally tractable model organism.


The ability to precisely modify the genome of an organism by adding, deleting or mutating genes is a critical tool for experimental biology. The type II CRISPR-associated (Cas) system is a powerful tool for genome editing in a variety of experimental systems. The Cas9 nuclease and two small noncoding RNAs comprise an adaptive immune system in prokaryotes ${ }^{1}$. A chimeric fusion of the two RNAs, termed a single guide RNA (sgRNA), supports site-specific cleavage of target DNA by Cas9, with target specificity determined by base-pairing between the $5^{\prime}$ end of the sgRNA and the target DNA ${ }^{1}$ (Fig. 1a). The only specific sequence requirement for cleavage is an NGG nucleotide sequence (the protospacer adjacent motif) at the $3^{\prime}$ end of the target DNA sequence (Fig. 1a). Changing the targeting sequence at the $5^{\prime}$ end of the sgRNA can allow an exceptionally broad variety of DNA substrates to be targeted without the need to reengineer the Cas9 nuclease ${ }^{1-13}$. Compared to zinc-finger nucleases and transcription activator-like effector nucleases (TALENs) that can be used to produce double-strand breaks ${ }^{14}$, the CRISPR-Cas9 system is substantially less expensive and is easier to program for new target sites. Cas 9 has been used to produce targeted insertion or deletion (indel) mutations, which are generated via error-prone
repair mechanisms, in a wide range of species ${ }^{2-13}$. In addition, homology-directed repair of Cas9-induced double-strand breaks has been demonstrated in bacteria ${ }^{7}$, yeast ${ }^{3}$, cultured human and mouse cells ${ }^{5,11}$, fruit flies ${ }^{4}$, zebrafish ${ }^{10}$ and mice ${ }^{13}$.

The nematode C. elegans is a valuable and widely used experimental system owing to its rapid growth, ease of handling and transparency (which facilitates microscopy). Recent reports demonstrated that Cas9 can induce double-strand breaks in the C. elegans germ line, which leads to mutations via errorprone repair mechanisms ${ }^{8,15}$. Here we demonstrate that Cas9induced double-strand breaks can be repaired efficiently by homologous recombination. By supplying engineered homologous repair templates, we generated in-frame $g f p$ insertions and targeted mutations. We refer to this method as Cas9-triggered homologous recombination.

## RESULTS

Design of a CRISPR-Cas9 system for C. elegans
To establish Cas9 as a tool for genome editing in C. elegans, we expressed Cas9 and sgRNA in the C. elegans germ line. In mammalian systems, sgRNAs were expressed from a U6 small nuclear RNA promoter, which drives transcription by RNA polymerase III ${ }^{5,11}$. To our knowledge, no germline RNA polymerase III promoters had been described in C. elegans when we initiated these experiments. Aligning ten C. elegans U6 genes revealed a region of 200-300 base pairs (bp) upstream of the transcriptional start site that is partially conserved and may function as the promoter (Fig. 1b and Supplementary Fig. 1). We therefore expressed sgRNAs using 350 bp upstream of the highly expressed R07E5.16 U6 gene ${ }^{16}$. A different U6 promoter was independently identified by Friedland et al. ${ }^{8}$. To express Cas9 in the germ line, we used the eft-3 promoter and tbb-2 $3^{\prime}$ UTR, which have been used successfully for genome modifications employing the Mos1 transposon ${ }^{17}$. We built a Cas9-sgRNA plasmid containing both the Peft-3::Cas $9:: t b b-2-3^{\prime}$ UTR and PU6 $:: s g R N A$ constructs (Fig. 1c). This plasmid, available through Addgene, can be engineered to target any desired sequence by using site-directed mutagenesis to insert the appropriate targeting sequence.

[^0]
b



Figure 1 | Adaptation of the CRISPR-Cas9 system for C. elegans. (a) Schematic of the Cas9 nuclease and sgRNA. Formation of a double-strand break requires (i) base-pairing between the sgRNA and the target DNA sequence and (ii) the presence of the NGG motif (protospacer adjacent motif, PAM) immediately adjacent to the target sequence ${ }^{1}$. Cleavage occurs $3 \mathrm{bp} 5^{\prime}$ of the PAM. The guanine ( G ) residue at the $5^{\prime}$ end of the sgRNA is required for transcription initiation by the U6 promoter. (b) Sequence conservation of the ten U6 RNA genes that we identified in C. elegans. The blue trace is a rolling average produced using locally weighted scatter-plot smoothing. The green bar indicates the region of R07E5.16 that we used as the promoter in the Cas9-sgRNA construct. (c) Schematic of the Cas9-sgRNA plasmid.

## Homologous repair of Cas9-induced double-strand breaks

Several existing methods for modifying the C. elegans genome rely on homologous repair of double-strand breaks generated by excision of a Mos1 transposon ${ }^{18-21}$, but these techniques are limited by the relative scarcity of Mos1 insertion sites in the genome ${ }^{22}$. In principle, a much wider range of genome modifications could be made by using Cas 9 to generate double-strand breaks. We therefore tested whether Cas9-induced double-strand breaks could be repaired by homologous recombination in C. elegans. We designed an sgRNA targeting a sequence adjacent to the $t t T i 5605$ Mos1 insertion site on chromosome II and compared the efficiency of singlecopy transgene insertion into this site using either Mos1-mediated single-copy insertion (MosSCI ${ }^{20}$ ) or Cas9-triggered homologous recombination (Fig. 2a). The efficiency of both approaches varied among individual experiments, but the overall efficiency of the two methods was similar (Fig. 2b and Supplementary Table 1). We examined expression of the resulting single-copy $g f p$ transgenes under the control of the mex-5 promoter and tbb-2 $3^{\prime}$ UTR. The transgenes were expressed in the germ line, and the pattern

of expression was indistinguishable regardless of the method used for transgene insertion (Fig. 2c). These data demonstrate that Cas9-induced double-strand breaks can stimulate homologous recombination in the C. elegans germ line.

## Integration of $g f p$ into endogenous loci

In C. elegans, fluorescent fusion proteins are often expressed via microinjection of DNA into the gonad, which generates semistable extrachromosomal arrays that contain many copies of the injected DNA ${ }^{23}$. Transgenes generated in this way are typically overexpressed in somatic tissues and silenced in the germ line and early embryo ${ }^{24}$. Microparticle bombardment can be used to generate low-copy transgenes, which are expressed at closer to endogenous levels ${ }^{25,26}$, but this approach is expensive and time consuming. MosSCI can be used to generate single-copy transgenes ${ }^{20}$, but for many genes, the regulatory sequences needed to recapitulate the native expression pattern are unknown. All of these approaches also leave the endogenous copy of the gene of interest intact, which makes it difficult to assess the function of the fusion protein genetically and can introduce complications in quantitative experiments because less than $100 \%$ of molecules of the protein of interest are labeled. Inserting genetically encoded tags

Figure 2 | Efficiency of Cas9-triggered homologous recombination in C. elegans.
(a) Schematic for homologous recombination (HR) mediated by either Mos1 transposon excision (left) or Cas9 (right). (b) Efficiency of single-copy transgene insertion for three different transgenes using either Mos1-mediated single-copy insertion (MosSCI) or Cas9. $n$ values at the bottom of each bar indicate the number of successfully injected animals (those that yielded non-Unc progeny). Percent efficiency is the fraction of successfully injected animals that yielded integrated transgenes. See
Supplementary Table 1 for raw data.
(c) Images of germline GFP expression from Pmex-5::gfp ::tbb-2-3' UTR transgenes generated using MosSCI or Cas9. Images were acquired, processed and displayed with identical settings. Results are representative of five animals of each strain. Scale bars, $20 \mu \mathrm{~m}$.


Figure 3 | Tagging of endogenous nmy-2 with gfp. (a) Strategy for producing $n m y-2:: g f p$ knock-ins. Cas9 cleavage of the $3^{\prime}$ end of $n m y-2$ stimulates homologous recombination, resulting in insertion of gfp and unc-119(+) into the genome. After isolating recombinants, we excised the unc-119(+) selectable marker by expressing Cre recombinase. (b) PCR genotyping of the nmy-2 locus in the indicated strains, using primer pairs as indicated and as schematized in a. Results are representative of three independently isolated knock-in strains. (c) PCR genotyping of the nmy-2 locus before and after excision of the unc-119(+) marker with Cre. Results are representative of five independent Cre-mediated unc-119(+) excision experiments. (d) Western blot showing NMY-2 levels in embryonic lysates in N2 (wild type), a strain carrying zuIs45 and strains carrying three independent knock-in alleles. Coomassie staining of total protein is shown as a loading control. Results are representative of three independent experiments. (e) Stage-matched images of NMY-2-GFP localization in an nmy-2::gfp knock-in strain compared to in zuIs45. The embryos shown were placed side-by-side on the same coverslip and imaged simultaneously. The images in the four left columns are maximum-intensity projections of two $0.5-\mu \mathrm{m}$ sections at a cortical focal plane and are taken from Supplementary Video 1. The far-right panels are single confocal sections from a different pair of embryos at gastrulation stage. Arrowheads indicate apical accumulation of NMY-2-GFP in gastrulating endodermal precursors. Results are representative of 14 independent experiments. Scale bars, $10 \mu \mathrm{~m}$.
such as $g f p$ into endogenous genes ensures $100 \%$ labeling and expression under the control of native regulatory elements and in the normal chromatin context.
To test whether Cas9-triggered homologous recombination could be used to insert protein tags into endogenous genes, we targeted the nmy-2 gene, which encodes nonmuscle myosin II (Fig. 3). We built a homologous repair template comprising the C-terminal 1.5 kilobases (kb) of $n m y$ - 2 fused in-frame to $g f p$, which was followed by the nmy-2 $3^{\prime}$ UTR, an unc-119(+) selectable marker and 1.5 kb of downstream genomic sequence (Fig. 3a). The unc-119(+) gene was flanked by loxP sites, which allowed it to be removed by subsequent expression of Cre recombinase. We also generated a Cas 9 -sgRNA plasmid targeted to cleave at the $3^{\prime}$ end of $n m y-2$. Co-injection of the Cas9-sgRNA plasmid and homologous repair
template into unc-119 worms resulted in integration of $g f p$ and unc-119(+) into the $3^{\prime}$ end of the $n m y$ - 2 locus (three independent knock-in alleles from 60 total injected animals). We confirmed the correct integration of $g f p$ at the $3^{\prime}$ end of the $n m y-2$ gene in all three lines by PCR (Fig. 3b) and sequencing.

We examined the expression and localization of the NMY-2GFP fusion protein in these three homozygous knock-in lines. For comparison, we analyzed a strain carrying zuIs45, a wellestablished transgene generated by microparticle bombardment ${ }^{27}$. The knock-in strains expressed NMY-2-GFP at levels similar to those of endogenous NMY-2, whereas the zuIs 45 strain expressed NMY-2-GFP at lower levels (Fig. 3d). NMY-2-GFP localized to the cell cortex in $n m y-2:: g f p$ knock-in embryos, with a pattern that was indistinguishable from that in embryos carrying zuIs 45


Figure 4 | Tagging of endogenous his-72 with gfp. (a) PCR genotyping of the his-72 locus in the indicated strains using a PCR strategy similar to that outlined in Figure 3a,b. (b) his-72 mRNA expression levels in the indicated strains, as measured by quantitative reverse transcription PCR (qRT-PCR). Results are the average of three independent experiments, and error bar shows $95 \%$ confidence interval. N.S., not significant ( $P=0.07$, two-tailed $t$-test). (c) HIS-72-GFP fluorescence in whole worms at the indicated stages. Results are representative of seven animals imaged. Scale bars, $50 \mu \mathrm{~m}$.
(Fig. 3e and Supplementary Video 1). However, the knock-in embryos showed consistently brighter fluorescence, which is consistent with the higher expression of NMY-2-GFP in the knock-ins.

Because nmy-2 is an essential gene ${ }^{28}$, we tested whether $g f p$ insertion disrupted protein function by assaying for lethality. Two of three $n m y-2:: g f p$ knock-in strains were $100 \%$ homozygous viable, and a third strain was $99 \%$ viable ( $n>100$ embryos for each strain). Animals of all three knock-in strains displayed wild-type movement and had no discernable phenotypes. We conclude that insertion of $g f p$ at the endogenous locus does not affect nmy-2 gene function.
For some applications, the insertion of the unc-119(+) selectable marker into the genome may be problematic. We therefore developed a simple procedure to remove the unc-119(+) marker by injecting a plasmid encoding Cre recombinase under the control of the eft-3 promoter and tbb-2 3' UTR (Online Methods) and picking uncoordinated (Unc) worms from the $\mathrm{F}_{2}$ progeny (Fig. 3c). We isolated Unc animals in 5 of 5 independent experiments (12-20 animals injected in each experiment). The high success rate achieved with relatively few injected animals indicates that Cre-mediated excision of unc-119(+) is highly efficient. We then outcrossed to wild-type worms to remove the unlinked unc-119 mutation required for unc-119(+) selection. This resulted in a strain containing no known genomic modifications except for insertion of $g f p$ into the $n m y-2$ gene and a 34 -bp loxP site in the intergenic region downstream of $n m y-2$ (Fig. 3a). NMY-2-GFP expression in these animals was not altered by removal of the unc-119(+) selection marker (data not shown), suggesting that this marker does not affect fluorescent fusion protein expression when it is inserted into an intergenic region.
To test whether this strategy is likely to be broadly applicable, we used a similar approach to endogenously tag the his-72 gene,
which encodes histone H3, with GFP. We obtained one knockin strain from ten successfully injected animals. We were able to amplify the left and right insertion junctions, confirming insertion of $g f p$ into the his- 72 locus (Fig. 4a). However, we were unable to amplify across the insertion, and a more detailed PCR characterization showed that a rearrangement involving a duplication of the unc-119(+) cassette had occurred (data not shown). Such rearrangements have been reported to occur in other studies involving homologous recombination ${ }^{20,29}$ and occurred at a low frequency in our study ( 1 of 16 strains generated by Cas9triggered homologous recombination, 1 of 7 MosSCI strains). Of note, his-72 mRNA levels in the his-72::gfp knock in-strain were indistinguishable from those of wild type, indicating that the rearrangement did not affect the his-72 gene itself (Fig. 4b). These animals were healthy and showed bright nuclear GFP fluorescence in a wide range of tissues including the germ line, consistent with labeling of endogenous histone (Fig. 4c).

## Generation of multiple point mutations in a single step

Another application of Cas9-triggered homologous recombination is the generation of targeted mutations at endogenous loci. To demonstrate this, we made mutations in lin-31, which encodes a FOXB transcription factor required for vulval development ${ }^{30}$. LIN-31 forms a complex with a LIN-1 (a homolog of mammalian Ets transcription factor), and this complex is thought to repress the primary vulval fate ${ }^{31}$. MPK-1, a mitogen-activated protein kinase (MAPK), can phosphorylate LIN-31 on four C-terminal threonines, and addition of active MAPK disrupts the interaction between LIN-31 and LIN-1 in vitro ${ }^{31}$. Overexpression of nonphosphorylatable LIN-31 causes a vulval phenotype ${ }^{31}$, but whether MAPK phosphorylation affects the function of endogenous LIN-31 has not been tested directly.

We used Cas9-triggered homologous recombination to mutate the four MAPK phosphorylation sites at the C terminus of LIN-31 to either alanine (nonphosphorylatable) or glutamic acid (phosphomimetic) (Fig. 5a,b). We used a Cas9-sgRNA plasmid targeting a site $5^{\prime}$ of these four residues to ensure that the Unc-119(+) animals we isolated would contain mutations at all four sites (Fig. 5b). We obtained two independent $\operatorname{lin}-31(4 T \rightarrow A)$ alleles from 60 total injected animals and three independent $\operatorname{lin}-31(4 T \rightarrow E)$ alleles from 62 total injected animals. We confirmed the desired mutations by PCR (not shown) and sequencing (Fig. 5c).

Mutation of the four MAPK phosphorylation sites to alanine caused a distorted vulval morphology at the L4 stage and a partially penetrant protruding vulva phenotype in the adult. A similar phenotype was observed when these residues were mutated to glutamic acid, mimicking constitutive phosphorylation (Fig. 5d,e and Supplementary Table 2). Identical phenotypes were observed for two independently isolated $\operatorname{lin}-31(4 T \rightarrow A)$ lines and for three independently isolated lin-31 $(4 T \rightarrow E)$ lines. These results suggest that dynamic regulation of LIN-31 phosphorylation is important for normal vulval development.

## Assessment of Cas9 specificity in C. elegans

Cas9 has been reported to produce off-target mutations in mammalian cells, which raises concerns about the specificity of this enzyme in genome-editing applications ${ }^{32}$. To assess the specificity of Cas9 in C. elegans, we identified the genomic sequences most similar to the targeting sequences that we used to modify $n m y-2$

Figure 5 | Targeted mutations in an endogenous gene. (a) Model for how MAPK phosphorylation affects LIN-31 function ${ }^{31}$ and the predicted effects of mutating these residues to either alanine or glutamic acid. (b) Strategy for simultaneous mutagenesis of four threonine residues (T145, T200, T218 and T220) in lin-31 to either alanine or glutamic acid (asterisks). Cas9 is targeted to the $5^{\prime}$ end of the last exon of lin-31, ensuring mutation of all four threonine residues. (c) Sequence confirmation of the induced mutations in lin-31 mutant strains. Results are representative of two independently isolated $\operatorname{lin}-31(4 T \rightarrow A)$ strains and three independently isolated lin-31 ( $4 T \rightarrow E$ ) strains. (d) Vulval morphology in L4 and adult animals of the indicated genotypes. Images are representative of 15 N 2 animals, $35 \mathrm{lin}-31(4 T \rightarrow A)$ animals and $36 \mathrm{lin}-31(4 T \rightarrow E)$ animals. Scale bars, $20 \mu \mathrm{~m}$. (e) Quantification of the Pvl phenotype of lin-31 adults. Scoring of phenotypes was done in a blinded fashion. Results from multiple isolates of each lin-31 mutation are combined; $n=406$ for $\mathrm{N} 2, n=267$ for lin-31 (4T $\rightarrow A$ ) and $n=217$ for lin-31 ( $4 T \rightarrow E$ ). See Supplementary Table 2 for the raw data.

and lin-31. We focused on candidate offtarget sequences that closely matched the $3^{\prime}$ end of our targeting sequences, as Cas9 activity has been shown to be most sensitive to mismatches in the $3^{\prime}$ half of sgRNA targeting sequences ${ }^{1,5,7,32}$. We PCR-amplified and sequenced ten candidate off-target sites for the $n m y-2$ sgRNA and four for the lin-31 sgRNA (Supplementary Table 3). We found no mutations at any of these sites in any of the strains we isolated. We note that although Fu et al. detected off-target activity of Cas9 toward sequences with up to five mismatches to the sgRNA sequence ${ }^{32}$, the closest matches that existed to our nmy-2 and lin-31 sgRNAs contained six or more mismatches each (Supplementary Table 3). The small size of the C. elegans genome compared to that of mammalian genomes may reduce the odds of closer off-target matches to particular target sequences.

## DISCUSSION

We have demonstrated a method to efficiently induce essentially any desired modification in the C. elegans genome. This approach was robust and cost-effective: for example, we obtained three independent nmy-2 $2: g f p$ knock-in lines from a single set of injections, requiring less than 4 weeks total time (of which less than 2 d was handson time) and less than $\$ 200$ worth of materials (Supplementary Fig. 2 and Supplementary Protocol). In 6 of 8 experiments, we obtained multiple independent lines on the first attempt. In the other two cases, the first set of injections failed for trivial technical reasons (Online Methods), and we readily obtained homologous recombinants upon reinjection. Thus, we achieved an overall success rate of $100 \%$ without extensive reengineering or optimization, which implies that our strategy is likely to be broadly applicable. In addition, the ability to remove the selectable marker in a single step using Cre recombinase should facilitate the use of unc-119 or other selectable markers for a wide variety of genome-editing strategies.
Our method relies on double-strand break repair using an engineered homologous template, similar to earlier methods that used

Mos1 transposon excision to generate double-strand breaks ${ }^{18-21}$. The use of Cas9 in our system overcomes several important limitations of Mos1-based methods. First, Cas9 target sites occur once every 32 bp in random DNA sequence; we estimate that there are over 1 million potential Cas9 target sites in the C. elegans genome, compared to approximately 14,000 Mos1 insertion sites ${ }^{22,33}$. This greatly expands the range of modifications that can be made because homologous recombination is most efficient within 0.5 kb of a double-strand break ${ }^{18}$. Indeed, none of the alleles presented in this study could have been made using Mos1based methods because appropriate transposon insertion alleles do not exist. Second, by carefully choosing the positions of the Cas9 cleavage site and selectable marker relative to the desired modification, the investigator can ensure that the desired genome modification is present in every isolated recombinant (Fig. 5b and Supplementary Protocol), in contrast to Mos1-based methods, in which the length of a gene conversion track is stochastic and decays rapidly with increasing distance from the transposon insertion site ${ }^{18}$. Third, Cas9 could in principle be used in any genetic background, whereas Mos1-based genome modifications must be generated in a strain carrying a Mos1 insertion. Finally, mobilization of Mos1 generates an average of 2-3 new Mos1 insertions at random sites elsewhere in the genome ${ }^{33,34}$, which could cause undesired phenotypes.

Cas9-triggered homologous recombination also has advantages over conventional transgenic approaches for generating fluorescent-protein fusions. Generating and maintaining a knockin line with our approach is cheaper and less labor intensive than either microparticle bombardment or extrachromosomal arrays. Also, it is expected to maintain the endogenous pattern of expression in most cases, and fusion-protein function can be easily assessed genetically owing to the absence of untagged protein. Knock-in strategies are the standard approach for
generating fluorescent-protein fusions in yeast for these reasons, and our methodology will allow widespread application of this approach in C. elegans.

Despite these advantages, there are some potential limitations associated with the use of Cas9-triggered homologous recombination to endogenously tag genes. First, fusion of $g f p$ to the $3^{\prime}$ end of a gene may compromise protein function, though this can be detected if it causes a loss-of-function phenotype. Second, GFP-coding sequences are sometimes recognized as 'nonself' and silenced by the Piwi Argonaute RNA pathway ${ }^{35}$. Such a silencing event should again be detectable if it produces a loss-of-function phenotype and/or loss of GFP fluorescence. Third, the DNA repair mechanism involved in homologous recombination can sometimes generate complex rearrangements, as we observed for $h i s-72:: g f p^{20,29}$. Because these events are uncommon, a straightforward solution is to isolate multiple knock-in alleles for each gene of interest. We obtained multiple independent alleles from a single set of injections for 7 of the 8 Cas9-triggered homologous recombination experiments performed for this study (his- $72:: g f p$ was the exception). Finally, for some genes with low endogenous expression levels, fluorescence of $g f p$ knock-ins may be too dim to visualize. Development of brighter fluorescent proteins ${ }^{36-38}$ may facilitate knock-in approaches for genes with lower endogenous expression levels. If fluorescence in a knock-in strain is too dim to be useful, then the investigator may choose to use an overexpression strategy, accepting the caveat that overexpression artifacts are possible.
In using Cas9-triggered homologous recombination to generate targeted mutations in endogenous genes, we chose to make point mutants; but we expect that insertions, deletions or other modifications could be made with similar ease, as has been done previously using Mos1 excision ${ }^{20,21}$. Of note, single point mutants have also been made in C. elegans using TALEN cleavage and ssDNA oligo-mediated repair ${ }^{15}$, but this approach is limited by the length of a synthetic DNA oligonucleotide and would have required at least two sequential injection steps to produce the four lin-31 point mutations that we were able to generate in a single step.

Interestingly, the phenotype we observed in lin-31 mutants was different from that reported by Tan et al. ${ }^{31}$, who found that expression of lin-31 $(4 T \rightarrow A)$ from an extrachromosomal array inhibited vulval fate specification. We suggest that repression of vulval fate in earlier experiments may have been due to overexpression of LIN-31 protein rather than solely its inability to be phosphorylated by MPK-1. The ability to quickly and efficiently induce mutations in endogenous genomic loci renders the use of multicopy extrachromosomal arrays unnecessary and should greatly simplify the interpretation of reverse genetic experiments.
In summary, we have developed a flexible, inexpensive and robust strategy for genome editing in C. elegans using Cas9 targeted cleavage and homologous recombination. Given the ease with which our approach can be adapted to new targets, we suggest that the ability to modify the C. elegans genome is now limited only by the imagination of the investigator.

## METHODS

Methods and any associated references are available in the online version of the paper.

Accession codes. Addgene: Cas9-sgRNA plasmid targeting a site near $t t T i 5605,47550$; Cas9-sgRNA plasmid with no targeting sequence, 47549; Peft-3::Cre $:: t b b-2-3^{\prime}$ UTR construct, 47551.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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## AUTHOR CONTRIBUTIONS

D.J.D. and J.D.W. jointly conceived of the project, and all authors discussed and contributed to the experimental design. D.J.D. performed the experiments. D.J.D. and D.J.R. analyzed the data. D.J.D. prepared the manuscript, and all authors discussed and contributed to the final version.

## COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.
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## ONLINE METHODS

Strains and nomenclature. New genetic nomenclature for genome editing applications has been developed by the WormBase Gene Name Curators (J. Hodgkin, University of Oxford, and T. Schedl, Washington University School of Medicine, personal communication). Briefly, edited loci are assigned conventional allele designations, with the nature of the modification described in brackets after the allele name. For example, one of our nmy-2::gfp knockin alleles is $n m y-2(c p 7[n m y-2:: g f p+L o x P$ unc-119(+) LoxP]) I, and a lin-31 mutant lin-31(cp2[T145A T200A T218A T220A + LoxP unc-119(+) LoxP]) II. Note that each independently isolated mutant line is given its own allele designation, even if the molecular lesion is identical. When the unc-119(+) selectable marker has been removed using Cre recombinase, a new allele designation is assigned. For example, an $n m y-2:: g f p$ knock-in allele with the unc-119(+) cassette removed is designated $n m y-2(c p 13[n m y-2:: g f p+L o x P])$ I. The $c p 13$ allele was derived from $c p 7$ by Cre-mediated recombination.

Supplementary Table 4 lists all strains generated and used in this study. All strains were kept at $25^{\circ} \mathrm{C}$ and fed Escherichia coli strain OP50 except where noted below and were handled using standard techniques ${ }^{39}$.

Plasmid construction. Plasmids have been deposited in Addgene with the following accession numbers: Cas9-sgRNA plasmid targeting a site near $t t$ Ti5605, 47550; Cas9-sgRNA plasmid with no targeting sequence, 47549; and the Peft-3::Cre $:: t b b-2-3^{\prime}$ UTR construct, 47551 . All other plasmids used in this study are available from the authors upon request.

To construct the Cas9-sgRNA expression plasmid shown in Figure 1c, we first designed a synthetic gene encoding Cas9, with C. elegans coding bias and synthetic C. elegans introns, using the C. elegans Codon Adaptor ${ }^{40}$. Our Cas9 sequence includes a nuclear localization signal and an HA tag at the C terminus. The synthetic gene was produced as a series of overlapping 500-bp gBlocks (Integrated DNA Technologies), assembled using Gibson Assembly (New England BioLabs) and inserted into the vector pCFJ601 (Peft-3 ::Mos1 Transposase ::tbb-2-3' UTR) ${ }^{17}$ in place of the Mos1 transposase. Next, a gBlock containing the U6 promoter and sgRNA sequence was inserted $3^{\prime}$ of the $t b b-23^{\prime}$ UTR. Genomic targets of Cas 9 conform to the target sequence $\mathrm{GN}_{19} \mathrm{NGG}$, where N is any base. The initial G is a requirement for transcription initiation by the U6 promoter, and the NGG (PAM) motif is required for Cas9 activity (note that the NGG motif must be present in the genomic target but is not included in the sgRNA sequence). To target Cas9 to different genomic sequences, we inserted the desired targeting sequence into the Cas9-sgRNA construct using the Q5 site-directed mutagenesis kit (New England BioLabs) with forward primer $5^{\prime}-\mathrm{N}_{19}$ GTTTTAGAGCTAGAAATAGCAAGT- $3^{\prime}$, where $\mathrm{N}_{19}$ is replaced by the desired 19-bp targeting sequence, and reverse primer $5^{\prime}$-CAAGACATCTCGCAATAGG- $3^{\prime}$.
Supplementary Table 5 lists the targeting sequences used in this study.

Targeting vectors for single-copy transgene insertion on chromosome II were constructed in the pCFJ150 vector backbone ${ }^{20}$ using Gateway cloning. We used site-directed mutagenesis with the Q5 site-directed mutagenesis kit to delete a short region of the $3^{\prime}$ recombination arm comprising the Cas9 target sequence to prevent the homologous repair templates from being cleaved by Cas9.

Homologous repair templates for $g f p$ insertion and lin-31 mutagenesis were constructed in two steps. First, we PCR-amplified a 3- to $4-\mathrm{kb}$ region centered on the desired modification from N 2 genomic DNA and cloned the resulting fragment into the pCR-Blunt vector using the ZeroBlunt TOPO Cloning Kit (Life Technologies). Second, we modified this genomic clone by inserting $g f p$ (for $g f p$ knock-ins) or a $3^{\prime}$ exon containing point mutations (for lin-31 mutagenesis), along with the unc-119(+) rescue gene flanked by loxP sites. $g f p$ and $u n c-119(+)$ fragments were generated by PCR, and LoxP sites were included in the unc-119(+) primers. The mutated lin-313' exons were synthesized as gBlocks. These fragments were integrated into the genomic clones using Gibson assembly, which allows for seamless fusion of DNA fragments without the need to include any extra sequence (for example, restriction sites). To avoid cleavage of the repair templates by Cas9, we deleted or mutated the Cas9 target site in all repair templates. Complete plasmid sequences of all targeting vectors are available from the authors upon request.

To construct the Peft-3::Cre $:: t b b-2-3^{\prime}$ UTR plasmid used for removal of selectable markers with Cre recombinase, we first amplified the Cre ORF from the plasmid pEM3 (ref. 41) and cloned it into the Gateway donor vector pDONR221. We then performed a three-fragment gateway reaction using our Cre donor vector, pCFJ386 (Peft-3; a gift from C. Frøkjær-Jensen), pCM1.36 (tbb-2 $3^{\prime}$ UTR) $)^{42}$ and the destination vector pCFJ212 (ref. 17), which contains an unc-119(+) rescue gene.

Supplementary Table 6 lists all primers used in this study.
Single-copy transgene insertion with MosSCI. We inserted transgenes into the ttTi5605 Mos1 site by following a published MosSCI protocol ${ }^{17}$. We prepared an injection mix containing $10 \mathrm{ng} / \mu \mathrm{L}$ targeting vector, $50 \mathrm{ng} / \mu \mathrm{L}$ pCFJ601 (Peft-3::Mos1 Transposase), $10 \mathrm{ng} / \mu \mathrm{L}$ pMA122 (heat-shock driven PEEL-1 negative selection), $10 \mathrm{ng} / \mu \mathrm{L}$ pGH8 (Prab-3::mCherry neuronal co-injection marker), $5 \mathrm{ng} / \mu \mathrm{L}$ pCFJ104 (Pmyo-3 ::mCherry body-wall muscle co-injection marker) and $2.5 \mathrm{ng} / \mu \mathrm{L}$ pCFJ90 (Pmyo- $2::$ mCherry pharyngeal co-injection marker). The mixture was microinjected into the gonads of Unc young adults of strain EG6699 (ttTi5605 II; unc-119(ed3) III), which were raised on HB101 bacteria at $15^{\circ} \mathrm{C}$. After injection, single worms were picked to new plates and maintained at $25^{\circ} \mathrm{C}$ until starvation (10-12 d). Plates containing non-Unc worms were counted as successfully injected. Occasionally, a batch of 50-60 injected animals yielded fewer than five successful injections, and we concluded that the injections had failed (usually for technical reasons, such as a bad needle) and repeated the injections. Following successful injections, we applied heat shock to plates with non-Unc worms at $34{ }^{\circ} \mathrm{C}$ for 4 h in an air incubator to activate the PEEL-1 negative-selection marker, which kills animals carrying extrachromosomal arrays. After overnight recovery at $25^{\circ} \mathrm{C}$, plates were visually screened to identify non-Unc animals that survived heat shock and did not express the red fluorescent co-injection markers. Single worms from these plates were picked to establish lines, and the presence of single-copy inserts was confirmed by PCR using primers listed in Supplementary Table 6.

Cas9-triggered homologous recombination. To modify the genome using Cas9-triggered homologous recombination, we followed a protocol similar to that for MosSCI (above). A mix
containing $10 \mathrm{ng} / \mu \mathrm{L}$ homologous repair template, $50 \mathrm{ng} / \mu \mathrm{L}$ Cas9-sgRNA plasmid, and the negative selection and co-injection markers listed above was injected into young adults of strain DP38 (unc-119(ed3) III) ${ }^{43}$. Note that although we used DP38 for the experiments reported in this study, we are recommending the use of the outcrossed derivative HT1593 for future experiments (see Supplementary Protocol). The procedure for selecting insertions and eliminating extrachromosomal arrays was identical to that described above. We were able to isolate strains that were either homozygous or heterozygous for all of our modifications (D.J.D., unpublished observations); for the experiments presented in this study, only the homozygous lines were kept.

Removal of unc-119(+) using Cre recombinase. An injection mix containing $50 \mathrm{ng} / \mu \mathrm{L}$ of pDD104 (Peft-3::Cre $:: t b b-2-3^{\prime}$ UTR) and $2.5 \mathrm{ng} / \mu \mathrm{L}$ pCFJ90 (Pmyo-2::mCherry pharyngeal marker) was injected into the gonads of young adult animals carrying an unc-119(+) cassette flanked by loxP sites. In each experiment, $15-20$ animals were injected and placed at $25^{\circ} \mathrm{C}$. We picked single $F_{1}$ progeny ( $10-20$ per experiment) expressing the red pharyngeal marker, which represent progeny of successful injections. We then selected Unc animals from among the $F_{2}$ progeny of these mCherry-positive animals. Because our Cre expression construct also carries unc-119(+), only animals that have (i) excised both genomic copies of the unc-119(+) cassette and (ii) lost the extrachromosomal array generated by injecting the Cre expression construct will be Unc. We verified that animals picked during this step were mCherry-negative and segregated only Unc progeny. Excision of unc-119(+) was then confirmed by PCR (see Fig. 3c).

Antibodies and western blotting. Embryos were isolated from gravid adult worms by bleaching and were lysed by sonication for 20 min in a bath sonicator filled with boiling water. Lysates were separated on 3-8\% NuPAGE Tris-acetate gels (Life Technologies) and transferred to nitrocellulose. The following antibodies were used: rabbit anti-NMY-2 (ref. 28) at 1:1,000 dilution of crude serum and Alexa Fluor 790 goat anti-rabbit (Life Technologies, catalog number A11369) at 1:1,000 dilution. Blots were scanned on an Odyssey imaging system (LI-COR Biosciences).

Microscopy. DIC and fluorescence imaging of whole worms was performed using a Nikon Eclipse E800 microscope equipped with epifluorescence and Nomarski DIC optics. Worms were mounted on $2.5 \%$ agar pads containing 10 mM sodium azide as a paralytic.

For NMY-2-GFP imaging, early one-cell embryos were mounted on poly(L-lysine)-coated coverslips and gently flattened using $2.5 \%$ agar pads. Images were captured using a Nikon Eclipse Ti microscope equipped with a $60 \times$, 1.4-NA objective and a Yokogawa CSU-X1 spinning disk head.

Maximum-intensity projection and adjustment of brightness and contrast were done using Fiji. No other image manipulations were performed.
qRT-PCR. RNA was isolated from gravid adult worms as follows. Worms were picked into Trizol reagent (Life Technologies) and lysed by repeated freeze-thaw cycles. Chloroform was added to a final concentration of $20 \%$ to separate phases, and the upper aqueous phase was mixed with an equal volume of ethanol and loaded on a RNeasy spin column (Qiagen). On-column DNase digestion was performed with the Qiagen on-column DNase digestion kit, and then RNA was washed and eluted according to the manufacturer's instructions. 25-75 ng total RNA was used for cDNA synthesis with the Superscript III reverse transcriptase kit (Life Technologies). qPCR was performed using a Viia 7 realtime PCR instrument and SYBR Green PCR Master Mix (Life Technologies). his-72 transcripts were detected with forward primer $5^{\prime}$-TCGTTCGTGAGATTGCCCAG- $3^{\prime}$ and reverse primer 5'-GAGTCCGACGAGGTATGCTT-3'. Y45F10D. 4 was used for normalization ${ }^{44,45}$. Data were analyzed to determine his-72 expression levels in Viia 7 software; we used the default settings for a relative standard-curve experiment.

Screening for off-target mutations induced by Cas9. Candidate off-target cleavage sites for each sgRNA were identified by BLAST searches ${ }^{46}$ against the C. elegans genomic sequence. For each candidate off-target site, we PCR-amplified an $\sim 1-\mathrm{kb}$ fragment centered on the candidate site from genomic DNA isolated from N2 control animals and from each modified strain we generated. The PCR products were sequenced by Eton Bioscience using primers binding to each end of the fragment. The sequence reads were aligned to the genomic sequence to look for insertion or deletion mutations at the putative cleavage site.

Reproducibility. Our sample sizes were chosen to allow confidence in the results while maintaining feasibility, and they are consistent with established norms for C. elegans research and the developmental biology community more broadly. No randomization was necessary for these studies. Phenotypes presented in Figure 5d,e were scored blindly.
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## Supplementary Information For

# "Engineering the Caenorhabditis elegans Genome Using Cas9-Triggered Homologous Recombination" 

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## Supplementary Figure 1: Multiple alignment of C. elegans U6 genes

Residues that are $>50 \%$ conserved are highlighted, and the U6 RNA sequence is indicated above the alignment. The alignment was produced using CLC Sequence Viewer software.


Cas9 + sgRNA plasmid: Insert targeting sequence using site-directed mutagenesis


Homologous repair tempate:

1) Clone $3-4 \mathrm{~kb}$ genomic fragment
2) Modify as desired and insert selectable marker


Day 9: Heat shock to kill animals with arrays


Mixture of knock-in
 and array animals

Day 10: Pick candidate knock-in animals
 and confirm knock-in by PCR

Day 0: Inject

Injection mix: - Peft-3::Cre plasmid - Fluorescent co-injection marker

Day 2: Pick
fluorescent F1's

Days 4-7: Pick Unc animals


Supplementary Figure 2: Workflow and timetable for modifying the C. elegans genome using Cas9-triggered homologous recombination
The basic steps in the procedure are shown, using the $n m y-2:: g / g$ knock-in experiment as an example. Please refer to the Supplementary Text for a detailed protocol.

| Transgene | DSB Method | Total Injections | Successful Injections | Transgene Insertions | Insertion Efficiency |
| :--- | :---: | :---: | :---: | :---: | :---: |
| Pmex-5::4xYPET::tbb-2 3'UTR | Mos1 | 55 | 17 | 6 | $35 \%$ |
| Pmex-5::GFP::tbb-2 3'UTR | Mos1 | 54 | 27 | 1 | $4 \%$ |
| Phis-72::GFP::tbb-2 3'UTR | Mos1 | 64 | 26 | 4 | $15 \%$ |
| Total (Mos1) | Mos1 | 173 | 70 | 11 | $16 \%$ |
|  |  |  |  |  |  |
| Pmex-5::4xYPET::tbb-2 3'UTR | Cas9 | 66 | 13 | 2 | $15 \%$ |
| Pmex-5::GFP::tbb-2 3'UTR | Cas9 | 79 | 37 | 2 | $32 \%$ |
| Phis-72::GFP::tbb-2 3'UTR | Cas9 | 72 | 22 | 12 | $9 \%$ |
| Total (Cas9) | Cas9 | 217 | 72 | $22 \%$ |  |

## Supplementary Table 1: Efficiency of MosSCI and Cas9-triggered homologous recombination for single-copy transgene insertion

This table presents the raw data depicted in Fig. 2b. "Total injections" is the number of animals that survived injection. "Successful injections" is the number of animals that produced non-Unc progeny (which could be due to either single-copy transgene insertion or extrachromosomal array formation). "Transgene insertions" is the number of animals that produced progeny carrying single copy integrated transgenes and lacking extrachromosomal array markers.

| Genotype | Number Scored | Number Pvl | Percent Pvl |
| :--- | ---: | ---: | ---: |
| Wild Type (N2) | 406 | 1 | $0.2 \%$ |
|  |  |  |  |
| lin-31(cp1[4T->A]) unc-119(+) II; unc-119(ed3) III | 114 | 59 | $52 \%$ |
| lin-31(cp2[4T->A]) unc-119(+) II; unc-119(ed3) III | 153 | 86 | $56 \%$ |
| Total lin-31(4T->A) | 267 | 145 | $54 \%$ |
|  |  |  |  |
| lin-31(cp3[4T->E]) unc-119(+) II; unc-119(ed3) III | 48 | 22 | $46 \%$ |
| lin-31(cp4[4T->E]) unc-119(+) II; unc-119(ed3) III | 54 | 24 | $44 \%$ |
| lin-31(cp5[4T->E]) unc-119(+) II; unc-119(ed3) III | 115 | 59 | $51 \%$ |
| Total lin-31(4T->E) | 217 | 105 | $48 \%$ |

Supplementary Table 2: Scoring of Pvl phenotype in lin-31 mutants
This table presents the raw data depicted in Fig. 5e.

| Target | Sequence | \# of Mismatches | Mutations? |
| :--- | :--- | :---: | :---: |
| nmy-2 (3' end) | GTTAGTTGCGAACTGAGTCGCGG | 0 |  |
| Between inx-2 and F08G12.2 | AGTACTGTCATACTGAGTCGAGG | 6 | None Detected |
| sex-1 | GCTAAGAGCTTACTGAGTCGCGG | 6 | None Detected |
| Between F07A5.4 and sue-1 | TATCATTTCGAACCGAGTCGGGG | 6 | None Detected |
| C30F12.2 | ATCATACGAGAACTGAATCGTGG | 7 | None Detected |
| Between Y102A5C.2 and Y102A5C.4 | TGTCGACACGAAATGAGTCGCGG | 7 | None Detected |
| nhr-211 | CATTTTCCGGAACTGGGTCGGGG | 8 | None Detected |
| E01G6.1 | GAAATACACTTACTGAGTCGAGG | 8 | None Detected |
| Between F58H10.1 and nsf-1 | CTGAAAAAGATACTGAGTCGAGG | 9 | None Detected |
| ceh-89 | AAGGAATCGGAACTGTGTCGCGG | 9 | None Detected |
| lin-24 | TTAGTGACTGAACTGAGTGGCGG | 9 | None Detected |
|  |  |  |  |
| lin-31 (5' end of last exon) | GAGAGATTTTCGATGGAGCCGGG | 0 |  |
| Y51H1A.2 | CAGATCGGTGCGATGGAGCCCGG | 6 | None Detected |
| Between F15A4.10 and T21B4.15 | GGAATAAAAAGGATGGAGCCTGG | 8 | None Detected |
| nhl-3 | TTTTGTTGTATGATGGAGCCGGG | 8 | None Detected |
| Y57A10A.7 | TTGCTGAACTTGATGGAGCCGGG | 9 | None Detected |

## Supplementary Table 3: Candidate off-target cleavage sites sequenced

This table lists the putative off-target cleavage sites for Cas9 with our sgRNAs that we sequenced to look for off-target mutations. Mismatches to the sgRNA sequence are shown in red, and the PAM is underlined.

| Strain Name | Genotype | Reference |
| :---: | :---: | :---: |
| DP38 | unc-119(ed3) III | 43 |
| EG6699 | ttTi5605 II; unc-119(ed3) III | 17 |
| JJ1473 | unc-119(ed3) III; zuls45[Pnmy-2::NMY-2::GFP + unc-119(+)] V | 27 |
| LP116 | cpSi4[Pmex-5::4xYPET::3xFlag::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP117 | cpSi5[Pmex-5::4xYPET::3xFlag::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP118 | cpSi6[Pmex-5::4xYPET::3xFlag::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP119 | cpSi7[Pmex-5::4xYPET: 3xFlag::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP120 | cpSi8[Pmex-5::4xYPET::3xFlag::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP121 | cpSi9[Pmex-5::4xYPET::3xFlag::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP122 | cpIs5[Pmex-5::4xYPET::3xFlag::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP123 | cpls6[Pmex-5::4xYPET::3xFlag::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP126 | lin-31(cp1[T145A T200A T218A T220A + LoxP unc-119(+) LoxP]) II; unc-119(ed3) III | This Study |
| LP127 | lin-31(cp2[T145A T200A T218A T220A + LoxP unc-119(+) LoxP]) II; unc-119(ed3) III | This Study |
| LP128 | lin-31(cp3[T145E T200E T218E T220E + LoxP unc-119(+) LoxP]) II; unc-119(ed3) III | This Study |
| LP129 | lin-31(cp4[T145E T200E T218E T220E + LoxP unc-119(+) LoxP]) II; unc-119(ed3) III | This Study |
| LP130 | lin-31(cp5[T145E T200E T218E T220E + LoxP unc-119(+) LoxP]) II; unc-119(ed3) III | This Study |
| LP132 | nmy-2(cp7[nmy-2::gfp + LoxP unc-119(+) LoxP]) I; unc-119(ed3) III | This Study |
| LP133 | nmy-2(cp7[nmy-2::gfp + LoxP unc-119(+) LoxP]) I; unc-119(ed3) III | This Study |
| LP134 | nmy-2(cp7[nmy-2::gfp + LoxP unc-119(+) LoxP]) I; unc-119(ed3) III | This Study |
| LP135 | cpSi10[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP136 | cpls10[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP137 | cpls11[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP138 | cpls12[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP139 | cpls13[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP140 | cpls14[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP141 | cpls15[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II;; unc-119(ed3) III | This Study |
| LP142 | cpls16[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP143 | cpls17[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP144 | cpls18[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP145 | cpls19[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP146 | cpls20[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP147 | cpls21[Pmex-5::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP148 | unc-119(ed3) his-72(cp10[his-72::gfp+ LoxP unc-119(+) LoxP]) III | This Study |
| LP149 | cpSi11[Phis-72::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP150 | cpSi12[Phis-72::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP151 | cpSi13[Phis-72::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP152 | cpSi14[Phis-72::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP153 | cpls23[Phis-72::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP154 | cpls24[Phis-72::GFP::tbb-2 3'UTR + unc-119(+)] II; unc-119(ed3) III | This Study |
| LP157 | lin-31(cp11[T145A T200A T218A T220A + LoxP]) II; unc-119(ed3) III | This Study |
| LP158 | lin-31(cp12[T145E T200E T218E T220E + LoxP]) II; unc-119(ed3) III | This Study |
| LP159 | nmy-2(cp13[nmy-2::gfp + LoxP]) I; unc-119(ed3) III | This Study |

## Supplementary Table 4: C. elegans strains used in this study

| Target | Genomic Sequence | sgRNA targeting sequence |
| :--- | :--- | :--- |
| Chromosome Il near <br> ttTi5605 Mos1 insertion site | GATATCAGTCTGTTTCGTAACGG | GATATCAGTCTGTTTCGTAA |
| nmy-2 (3' end) | GTTAGTTGCGAACTGAGTCGCGG | GTTAGTTGCGAACTGAGTCG |
| his-72 (3' end) | GAGCTTAAGCACGTTCTCCGCGG | GAGCTTAAGCACGTTCTCCG |
| lin-31 (5' end of last exon) | GAGAGATTTTCGATGGAGCCGGG | GAGAGATTTTCGATGGAGCC |

Supplementary Table 5: sgRNA targeting sequences used in this study
The PAM is underlined.

| Primer Description | Primer Sequence |
| :---: | :---: |
| Cas9-sgRNA plasmid |  |
| Cas9 For | TCAGTTGGGAAACACTTTGCT |
| Cas9 Rev | GCTTGAAAGGATTTTGCATTTATC |
| Peft-3 Rev (Vector) | TTTTTCTAGAGCAAAGTGTTTCCCAAC |
| tbb-2 3'UTR For (Vector) | ctaaactagtGATAAATGCAAAATCCTTTCAAGC |
| PU6::sgRNA | (Synthetic gBlock fragment) |
| sgRNA vector Rev | GTTTTCCCAGTCACGACGTT |
| sgRNA vector For | CTGGCGTAATAGCGAAGAGG |
|  |  |
| Deletion of the Cas9 target from MosSCl constructs |  |
| pCFJ150 4 target For | CCGTATCACAACTGAGAAAAAAG |
| tbb-2 3'UTR Rev | gcatcgcgcgcaccgtacgTGAGACTTTTTTCTTGGCGG |
|  |  |
| PCR genotyping of the ttTi5605 locus |  |
| ttTi5605 Upstream | AGGCAGAATGTGAACAAGACTCG |
| ttTi5605 Downstream | ATCGGGAGGCGAACCTAACTG |
| unc-119(+) 5' out | CAATTCATCCCGGTTTCTGT |
| unc-119(+) 3' out | TTCGCTGTCCTGTCACACTC |
|  |  |
| nmy-2::gfp repair template |  |
| nmy-2 5' arm For | TCTCAAACTAGAGATTGACAATTTGG |
| nmy-2 5' arm Rev | tcctttactcatggctccgctagctcctgagttacggacagagtctctaTCTTCATCGGTGTTGCC |
| GFP For | tcaggagctagcggagccATGAGTAAAGGAGAAGAACTTTTCACTG |
| GFP Rev | tctcttgtcatcgtcatccttgtaatcggatccggatccacgcttatcgtcatcgtccttgtagtcgcatgcTTTGTATAGTTCATCCATGCCATG |
| nmy-2 3'UTR For | gattacaaggatgacgatgacaagagaggatctggatctgactacaaggacgatgacgataagcgttaaCCTCTAAAGTTTTTAATTCCGGA |
| nmy-2 3'UTR Rev | TGCGGACATGTCAATTTGT |
| nmy-2::LoxP::unc-119(+) For | tatttaatgtaacaaattgacatgtccgcaataacttcgtatagcatacattatacgaagttatCTTTGAGCCAATTTATCCAAG |
| nmy-2::LoxP::unc-119(+) Rev | ataatccggaattaaaaacttagaggttaataacttcgtataatgtatgctatacgaagttatCCTAGTTCTAGACATTCTCTAATG |
| nmy-2 3' arm For | TAACCTCTAAAGTTTTTAATTCCGGAT |
| nmy-2 3' arm Rev | TGACAAGGACATTCTCGGA |
|  |  |
| Cas9-sgRNA plasmid targeting nmy-2 |  |
| nmy-2 sgRNA For | ttagttgcgaactgagtcgGTTTTAGAGCTAGAAATAGCAAGT |
| sgRNA Rev | CAAGACATCTCGCAATAGG |
|  |  |
| PCR genotyping of the nmy-2 locus |  |
| nmy-2 Upstream | TGACGCACAAGAAAAGATCG |
| nmy-2 Downstream | ATCAACAAAACATGCGTCCA |
| GFP Rev | TTTGTATAGTTCATCCATGCCATG |
| unc-119(+) 3' out | TTCGCTGTCCTGTCACACTC |
|  |  |
| his-72::gfp repair template |  |
| his-72 5' arm For | GACCCCACAAAATCGATACG |
| his-72 5' arm Rev | tcctttactcatggctccgctagctcctgagGCACGTTCTCCtctGATGCGTCTGGCGAG |
| GFP For | tcaggagctagcggagccATGAGTAAAGGAGAAGAACTTTTCACTG |
| GFP Rev | tctcttgtcatcgtcatccttgtaatcggatccggatccacgcttatcgtcatcgtcettgtagtcgcatgcTTTGTATAGTTCATCCATGCCATG |
| his-72 3'UTR For | gattacaaggatgacgatgacaagagaggatctggatctgactacaaggacgatgacgataagcgttaaGCTCCATCACCAATTCTCG |
| his-72 3'UTR Rev | ttacaaggacttggataaattggctcaaagataacttcgtataatgtatgctatacgaagttatGCGGCGTGGAATATAGTT |
| unc-119(+) For | CTTTGAGCCAATTTATCCAAGTC |
| his-72::LoxP::unc-119(+) Rev | taaaagtgcttcgagaattggtgatggagcataacttcgtataatgtatgctatacgaagttatCCTAGTTCTAGACATTCTCTAATG |
| his-72 3' arm For | GCTCCATCACCAATTCTCGA |
| his-72 3' arm Rev | AGGCTGAATTTGCTCTGAG |
|  |  |
| Cas9-sgRNA plasmid targeting his-72 |  |
| his-72 sgRNA For | agcttaagcacgttctccgGTTTTAGAGCTAGAAATAGCAAGT |
| sgRNA Rev | CAAGACATCTCGCAATAGG |
|  |  |
| PCR genotyping of the his-72 locus |  |
| his-72 Upstream | GTGGATTCAGGATCCTTCG |
| his-72 Downstream | TTTTGAGCTGAAGCTTGTATGG |
| GFP Rev | TTTGTATAGTTCATCCATGCCATG |
| unc-119(+) 3' out | TTCGCTGTCCTGTCACACTC |
|  |  |


| lin-31(4T->A) and lin-31(4T->E) repair templates |  |
| :--- | :--- |
| lin-31 5' arm For | ATTTGGTTCAGCCGAAATTG |
| lin-31 5' arm Rev | gggagcagtggaagaggattttgcgagagatttgcttggggctggaTGGTGGAAGTCGTCGT |
| lin-31 4T->A 3' end | (Synthetic gBlock fragment) |
| lin-31 4T->E 3' end | (Synthetic gBlock fragment) |
| lin-31 3'UTR For | CAAATGCGCTCTGTTAACC |
| lin-31 3'UTR Rev | ttacaaggacttggataaattggctcaaagataacttcgtataatgtatgctatacgaagttatAAAAACGTTATTTGGTTGTTTATGACTAA |
| unc-119(+) For | CTTTGAGCCAATTTATCCAAGTC |
| lin-31::LoxP::unc-119(+) Rev | actggcaacttttcattattggagaaaaataacttcgtataatgtatgctatacgaagttatCCTAGTTCTAGACATTCTCTAATGAAAA |
| lin-31 3' arm For | TTTTCTCCAAATAATGAAAAAGTTGCC |
| lin-31 3' arm Rev | GTCGAACCCAGAAATCAAC |
|  |  |
| Cas9-sgRNA plasmid targeting lin-31 |  |
| lin-31 sgRNA For | agagatttcgatggagccGTTTTAGAGCTAGAAATAGCAAGT |
| sgRNA Rev | CAAGACATCTCGCAATAGG |
|  |  |
| PCR genotyping and sequencing of the lin-31 locus |  |
| lin-31 Upstream | GCTCAGGGTTAGGCTTAGGG |
| lin-31 Downstream | ATATGTGGCACTCGCTTCG |
| unc-119(+) 5' out | CAATTCATCCCGGTTTCTGT |
| unc-119(+) 3' out | TTCGCTGTCCTGTCACACTC |
| lin-31 Seq | CGGGATGTCTCTGTCTCTTCAACT |
|  |  |
| Cloning of Cre recombinase into pDONR221 |  |
| Cre For attB1 | ggggacaagttgtacaaaaaagcaggctCCATGGGCGCACCC |
| Cre Rev attB2 | ggggaccactttgtacaagaaagctgggtaCTAATCGCCATCTTCCAGCA |

## Supplementary Table 6: Primers used in this study

Primers are listed for each construct. For each primer, the region that anneals to the target is capitalized. LoxP sites are underlined.

## Supplementary Protocol: Cas9-triggered homologous recombination

We have provided the following protocol in order to facilitate rapid adoption of our methodology by members of the C. elegans research community. Supplementary Fig. 2 shows a graphical representation of the steps in this procedure and the time required. All necessary plasmids are available from Addgene.

## Construct Design and Cloning

## Homologous repair template

We describe our cloning strategy in general terms, but this can be substituted with another approach if desired.

- Design the desired genomic modification and make a sketch of the modified locus. Using a DNA sequence editor, generate a file that contains the sequence of the locus in its desired final state, following the guidelines below and using Figs. 3 a and 5 b as examples.
- PCR-amplify a 3-4 kb genomic region, centered on the desired modification, from N 2 genomic DNA. We aim to have $\sim 1.5 \mathrm{~kb}$ of unmodified sequence at each end of our homologous repair template to allow homologous recombination. Clone the genomic fragment into any suitable vector (we use the ZeroBlunt TOPO kit from Life Technologies).
- Build the modifications you want into your cloned genomic fragment. The exact steps will vary depending on the genome modification you want to make, but for GFP knock-ins we insert GFP between the last amino acid codon of the gene and the stop codon, and the unc-119(+) rescue gene downstream of the 3'UTR. We have only tested unc-119 selection, but other selectable markers could likely be substituted. If you wish to remove the selectable marker later, flank it with LoxP sites (5’-ATAACTTCGTATAGCATACATTATACGAAGTTAT-3'). We have built our homologous repair templates using Gibson Assembly Master Mix from NEB, but other cloning methods could be used instead.
- IMPORTANT: Ensure that the Cas9 target sequence (see below) is NOT present in the homologous repair template. Otherwise, Cas9 will cleave your repair template in addition to the genomic DNA, and recombination efficiency will suffer. If the targeting sequence is part of a coding region then you will need to introduce silent mutations to block cleavage. If possible, the simplest and most effective approach is to mutate the PAM (NGG motif), since this motif is
absolutely required for cleavage of a substrate by Cas9. If a PAM mutation is not feasible, introduce at least 5-6 mutations in the target sequence. Mutations closer to the 3 ' end of the targeting sequence are more likely to prevent cleavage.

Choose the Cas 9 target site

- Target sites conform to the pattern 5'G-N ${ }_{19}-N G G-3$ ', where N is any base. The initial G is required for transcription by the U6 promoter, and the NGG motif is required for Cas9 cleavage. Locate a target sequence near the site you wish to modify.
- IMPORTANT: In general, the Cas9 target sequence and the selectable marker should flank the desired genome modifications. The presence of these two landmarks fixes the location of the homologous recombination to the homology arms at the ends of the repair template, ensuring that all of the isolated recombinants will contain the desired modifications. For example, for Cterminal GFP insertions, we use a Cas 9 target sequence at or near the 3 ' end of the coding sequence, upstream of the $3^{\prime} U T R$, and place the unc-119(+) cassette downstream of the $3^{\prime}$ UTR in the repair template (see Fig. 3a for an example). The desired modification (GFP insertion) lies between the Cas 9 target site and the unc-119(+) marker. For point mutations, the target sequence should be located at or near the 5 ' end of the region containing the desired mutation(s), and the unc-119(+) cassette should be 3 ' of the desired mutations in the repair template (see Fig. 5b for an example).
- Aside from these general constraints, the target sequence may be oriented on either strand, and its exact location relative to the site to be modified is not critical. $\mathrm{GN}_{19}-\mathrm{NGG}$ motifs occur every 32 bases in random sequence, and we were able to find target sites matching the $\mathrm{GN}_{19^{-}}$ NGG sequence requirement within 50-100 bp of the desired insertion site (and much closer in most cases) for all of our homologous recombination experiments.
- At this stage, it is advisable to check the uniqueness of the target sequence by BLASTing against the C. elegans genome. We recommend avoiding targeting sequences with fewer than 5-6 mismatches to the next most similar site in the genome. Remember that if a putative off-target site has high homology to the $\mathrm{GN}_{19}$ targeting sequence you choose, the N in the NGG motif does not have to match the target in order to be cleaved: for example, if your target sequence is GAAAACCCCCGGGGGTTTTT-AGG, then the sequence GAAAACCCCCGGGGGTTTTT-CGG would also be cleaved. Although Cas9 can cleave off-target sites under some conditions ${ }^{32}$, the determinants of its specificity are still not
completely understood. We therefore keep a list of the closest matches for each targeting sequence we design, to facilitate later sequencing for identification of any off-target mutations. Although we found no evidence for a high non-specific activity of Cas9 in C. elegans, we advocate isolating multiple lines for each genomic modification, sequencing of candidate offtarget loci, and outcrossing as prudent measures to be adopted until the factors governing Cas9 specificity are more completely understood.


## Cas9-sgRNA construct

- We use NEB's Q5 Site-Directed Mutagenesis Kit to insert the targeting sequence into our Cas9sgRNA construct (Addgene \#47549). Use forward primer 5'-
$\mathrm{N}_{19}$ GTTTTAGAGCTAGAAATAGCAAGT-3', where $\mathrm{N}_{19}$ is replaced by the desired 19 bp targeting sequence, and reverse primer 5'-CAAGACATCTCGCAATAGG-3'. Note that the initial $G$ in the $\mathrm{GN}_{19}$ sequence is included in the reverse primer.
- IMPORTANT: Do not include the PAM (NGG motif) in your primers for the Cas9-sgRNA construct. The NGG motif must be present in the target DNA, but it is not part of the sgRNA (Fig. 1a).


## Injection and Selection

Our selection protocol is based on, and is highly similar to, the MosSCI protocol developed by Christian Frøkjær-Jensen and colleagues ${ }^{17}$. We are greatly indebted to Dr. Frøkjær-Jensen for developing such an effective procedure and for sharing advice and reagents.

- Obtain worms of an appropriate strain for injection. Although we used strain DP38 (unc119 (ed3) III) for initial experiments, we have subsequently switched to the outcrossed derivative HT1593. HT1593 has a slightly weaker phenotype than DP38 (HT1593 animals are less Dpy and more active), but this has not affected our ability to recognize plates with rescued animals or isolate recombinants. Therefore we recommend that others use HT1593 for unc-119 selection experiments. In principle, any strain compatible with your selectable marker can be substituted. Our selection procedure is also optimized for unc-119 selection, and the use of other selectable markers may require subtle changes. If using unc-119 selection, note that unc-119 worms are healthier and easier to inject when grown on HB101 bacteria at $15^{\circ} \mathrm{C}$.
- The night before injection, pick $\sim 80$ L4 animals to a fresh plate and allow them to mature into adults at $15^{\circ} \mathrm{C}$ overnight.


## Day 0: Injection

- Prepare an injection mix containing the following plasmids:
- $10 \mathrm{ng} / \mu \mathrm{L}$ homologous repair template
- $50 \mathrm{ng} / \mu \mathrm{L}$ Cas $9-\mathrm{sg}$ RNA construct with your targeting sequence
- $10 \mathrm{ng} / \mu \mathrm{L}$ pMA122 (heat-shock driven PEEL-1 negative selection; Addgene \#34873)
- $10 \mathrm{ng} / \mu \mathrm{L}$ pGH8 (Prab-8::mCherry neuronal co-injection marker; Addgene \#19359)
- $5 \mathrm{ng} / \mu \mathrm{L}$ pCFJ104 (Pmyo-3::mCherry body wall muscle co-injection marker; Addgene \#19328)
- $2.5 \mathrm{ng} / \mu \mathrm{L}$ pCFJ90 (Pmyo-2::mCherry pharyngeal co-injection marker; Addgene \#19327) Miniprep-quality DNA is sufficient, and we do not linearize plasmids prior to injection.
- Inject the mixture into the gonads of the young adult worms.
- Transfer the injected worms to new seeded plates (three animals per plate works well in our hands).
- Put the plates at $25^{\circ} \mathrm{C}$ until the food bacteria are consumed and the Unc worms make piles (these are visible to the naked eye as big clumps of worms on the plate, where the edge of the bacterial lawn used to be).


## Day 7-9: Heat Shock

- Examine the plates and identify those that contain unc-119 rescued (i.e. normally moving) animals. In our experience, plates with rescues are obvious because they contain many crawling worms. You should not have to spend a lot of time searching to spot rescued worms. The number of plates with rescued animals will vary depending on your injection experience, but is usually greater than half in our hands.
- Heat shock rescued plates at $34^{\circ} \mathrm{C}$ for 4 hours in an air incubator. This activates expression of the PEEL-1 toxin, which kills animals that carry extrachromosomal arrays. After heat shock, return the plates to $25^{\circ} \mathrm{C}$ overnight to allow the array-carrying animals to die.


## Day 8-10: Pick Candidate knock-in animals

- The day after heat shock, examine the plates for normally moving animals that survived heat shock. Such plates are candidates for a knock-in event. Again, these plates are usually obvious in our hands - intensive searching for survivors should not be necessary. Efficiency can vary, but we routinely recover between two and six independent candidate knock-ins lines from each set of injections.
- Using a fluorescence dissecting scope, verify that the worms that survived heat shock also lack expression of the red fluorescent co-injection markers. From each candidate knock-in plate, single 8-10 non-Unc animals lacking co-injection marker fluorescence to new plates (some of these worms will be sterile after the heat shock, so don't try to get away with picking fewer).
- After these worms produce progeny, look at the plates and identify those that did not segregate Unc progeny. These are likely to be homozygous knock-in strains. Failure to identify any homozygotes may indicate that your genomic modification is homozygous lethal, or it may be a sign that you don't have a true knock-in. Use PCR to distinguish between these two possibilities. If you are making a GFP knock-in, you can also look for fluorescence at this stage to establish whether you have an integrant.


## PCR verification

- Design primers that anneal inside and outside the homologous repair template (see Fig. 3a for an example).
- Isolate genomic DNA from the strain to be characterized, using detergent and proteinase K digestion followed by phenol-chloroform extraction (for a detailed protocol, see http://thalamus.wustl.edu/nonetlab/ResourcesF/worm\ genomic\ DNA.pdf). A crowded 5 cm plate yields more than enough DNA. Also isolate DNA from your parental strain as a control.
- Set up PCR reactions with primer pairs similar to those shown in Fig. 3a-b. We have found that LongAmp Taq DNA polymerase from NEB gives particularly robust amplification from genomic DNA.
- Note: the primer pairs outside the homologous repair template (flanking the insertion site) are particularly useful since you should see a band shift for a knock-in, and the disappearance of the wild-type band indicates that your strain is homozygous.
- If desired, sequence the PCR products obtained in this step to verify a mutant allele.


## Removal of the selectable marker with Cre recombinase

These instructions assume you are using unc-119 selection, but the same basic approach should be applicable to other selectable markers.

- The night before injection, pick 25-30 L4 knock-in worms to a fresh plate and allow them to mature into young adults overnight.
- Prepare an injection mix containing the following:
- $50 \mathrm{ng} / \mu \mathrm{L}$ pDD104 (Peft-3.::Cre::tbb-2 $3^{3}$ UTR; Addgene \#47551)
- $2.5 \mathrm{ng} / \mu \mathrm{L}$ pCFJ90 (Pmyo-2::mCherry pharyngeal co-injection marker) or any other fluorescent marker.
- Inject the mixture into the gonads of young adult worms. Place the injected worms at $25^{\circ} \mathrm{C}$, three worms per plate.
- Two days after injection, single 10-20 F1 progeny that express the fluorescent marker to new plates. These animals are progeny resulting from successful injections and can segregate Uncs in the next generation. Do not put more than one animal per plate at this stage - more animals will cause the plates to be overcrowded and make screening difficult.
- When the F2 progeny have reached adulthood, pick Unc animals to new plates to establish lines. Timing is important here: at $25^{\circ} \mathrm{C}$, unc-119 animals are slightly egg laying-defective (Egl), and therefore are easiest to spot when they are old enough to have accumulated some eggs. However, it is important not to wait too long, as screening becomes more difficult once the plates become crowded.
- Note: because our Cre expression construct also carries unc-119(+), only animals that have 1) Excised both genomic copies of the unc-119(+) cassette and 2) Lost the extrachromosomal array generated by injecting the Cre expression construct will have an Unc phenotype. Loss of the array can be verified by loss of the fluorescent co-injection marker.
- If desired, confirm removal of the selectable marker by PCR or sequencing. At this stage, the Unc animals resulting from this procedure can be used in another round of homologous recombination to produce more complicated modifications, or outcrossed to remove the unc-119 mutation.


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