

# Targeted gene deletions in *C. elegans* using transposon excision

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**We developed a method, MosDEL, to generate targeted knockouts of genes in *Caenorhabditis elegans* by injection. We generated a double-strand break by mobilizing a *Mos1* transposon adjacent to the region to be deleted; the double-stranded break is repaired using injected DNA as a template. Repair can delete up to 25 kb of DNA and simultaneously insert a positive selection marker.**

Gene-specific knockouts are a defining technology of reverse genetics, allowing phenotypes to be assigned to any of the thousands of genes identified in genome sequencing projects. In *Caenorhabditis elegans*, reverse genetics has mainly relied on random chemical mutagenesis to generate loss-of-function mutants<sup>1</sup> or, more recently, random deletions downstream of guanine-quadruplex DNA<sup>2</sup>. In both cases, mutagenized populations are screened by PCR with gene-specific primers for random deletions in target genes. This approach has been used to generate putative loss-of-function alleles in more than 5,000 genes, mainly through the efforts of the *C. elegans* Gene Knockout Consortium in the United States and Canada and the National BioResource Project in Japan<sup>1,3</sup>. Random deletions have a few limitations. First, deletions are typically small and not necessarily molecularly null. Second, chemical mutagenesis invariably leads to background mutations. And third, some deletions involve complex rearrangements<sup>1</sup>.

In fruit flies, large deficiencies can be generated by recombination between *FRT* sites in *P* elements<sup>4</sup>. In other genetic model organisms (for example, yeast and mice) transgenic DNA fragments and homologous recombination are used to generate targeted deletions. Bombardment with DNA-coated gold particles can lead to gene replacement in *C. elegans* as well<sup>5</sup>. Unfortunately, the frequency of homologous recombination is low and the technique has not been widely adopted.

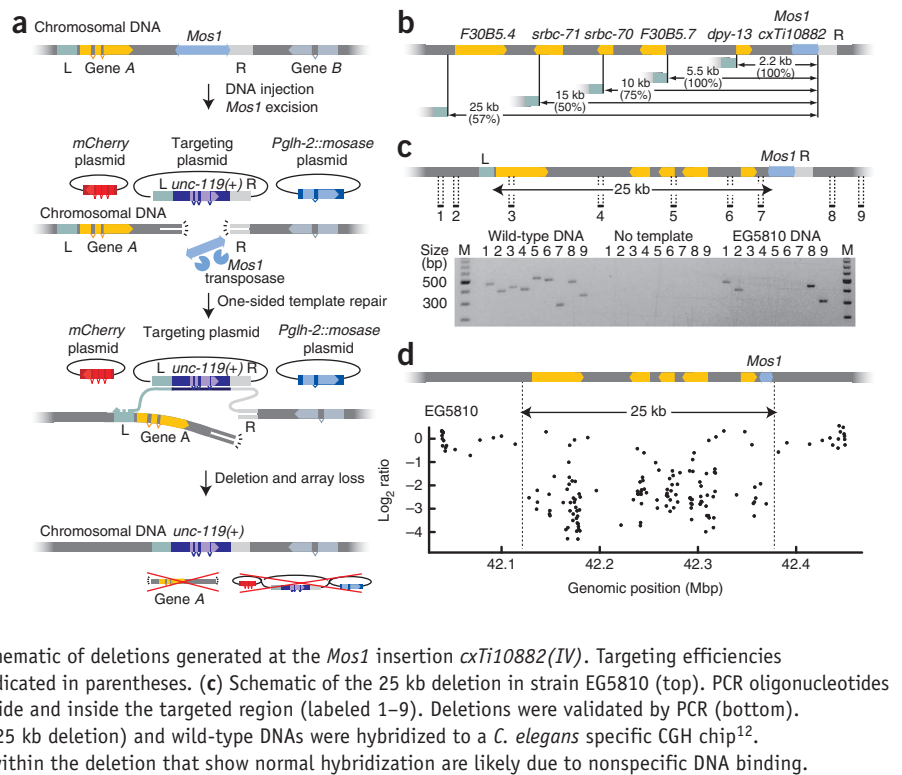
Endogenous Tc1 transposons have been used in *C. elegans* to inactivate genes by causing random deletions after excision<sup>6</sup>. More recently the *Drosophila melanogaster* transposon *Mos1* has been adapted for gene targeting<sup>7</sup>. To facilitate the use of *Mos1* elements, approximately 14,000 molecularly mapped transposon inserts have been generated by the NemaGENETAG consortium<sup>8</sup>. Mobilization of transposons generates double-stranded DNA breaks; repair of the breakpoint can lead to targeted modifications when a repair template is present<sup>9</sup>. This repair mechanism has been used to develop *Mos1* excision-induced transgene-instructed gene conversion (*mosTIC*)<sup>7</sup>, a technique that can be used to reliably modify DNA within one kilobase of a *Mos1* insert. Building on these efforts, we had developed *Mos1*-mediated single copy insertion (*MosSCI*), a technique to insert transgenes into well-defined genomic loci<sup>10</sup>. *Mos1* excision is induced using a simple injection-based method and successful insertions are identified using an inserted selectable marker<sup>10</sup>. Here we demonstrate that *Mos1* excision can be used to generate targeted deletions of up to 25 kb. We call this technique *Mos1*-mediated deletion (*MosDEL*).

We generated deletions in a strain with a *Mos1* element adjacent to the gene of interest (**Fig. 1a**). The *Mos1* element was excised in the germline by the *Mos1* transposase encoded on an injected helper plasmid. Excision resulted in a double-stranded DNA break, which was repaired using the co-injected repair template. Previously we had demonstrated that single-copy DNA can be incorporated into the genome by flanking the DNA with homology regions from both sides of the double-strand break. Under these conditions, both free ends of the chromosome break have homologous DNA in the repair template to initiate repair. Here we used targeting constructs capable of only one-sided repair, such that one of the broken ends has no homologous sequence to invade (**Fig. 1a**). Repair is initiated from one side by strand invasion of the homology arm on the template (right homology region; typically 2 kb). DNA is then copied from the extrachromosomal array, including sequence from a distal homology arm (left homology region; typically 3 kb). The 3' end can then invade the other half of the broken chromosome at a distance from the break. A wild-type copy of the *unc-119* gene is simultaneously inserted to provide a positive selection marker for deletions. Red fluorescent markers encoded on a co-injected plasmid are used to visually identify worms rescued by extrachromosomal arrays so that they can be disregarded.

To test the targeting strategy, we selected a *Mos1* element (*cxTi10882*) located 1.1 kb 3' from *dpy-13* (**Fig. 1b**). *dpy-13* mutants are viable and can be identified based on the dumpy phenotype. We injected 83 *unc-119 cxTi10882* worms with

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**Figure 1** | Using *Mos1* transposons to create targeted deletions. (a) Schematic of MosDEL. A *Mos1* transposon near the gene of interest is placed into an *unc-119(ed3)* strain. Injection of a plasmid encoding *Mos1* transposase (*Pglh-2::mosase*) results in excision of the transposon. The resulting double-strand DNA break is repaired by synthesis-dependent strand annealing at the right homology region (labeled “R”) on the extrachromosomal targeting plasmid, incorporating the positive selection marker *unc-119(+)* into the chromosomal locus. Genomic DNA between the right homology region and left homology region (labeled “L”) is deleted when the nascent repair strand re-invades the genomic DNA and resolves the break by homologous recombination. Red fluorescence (from co-injected plasmids encoding mCherry) marks the extra-chromosomal array; deletion mutants are isolated by screening for *unc-119* rescued worms lacking red fluorescence. *Pglh-2::mosase*, mCherry plasmid and the targeting plasmid are injected as separate plasmids; extrachromosomal arrays are genetically unstable and therefore lost at high frequency. (b) Schematic of deletions generated at the *Mos1* insertion *cxTi10882(IV)*. Targeting efficiencies (percentage of deletions per *unc-119* insertion) are indicated in parentheses. (c) Schematic of the 25 kb deletion in strain EG5810 (top). PCR oligonucleotides were designed to amplify 200–500 bp fragments outside and inside the targeted region (labeled 1–9). Deletions were validated by PCR (bottom). M, 100 bp marker. (d) For CGH verification, EG5810 (25 kb deletion) and wild-type DNAs were hybridized to a *C. elegans* specific CGH chip<sup>12</sup>. A  $\log_2$  ratio below  $-2$  indicates deleted DNA. Points within the deletion that show normal hybridization are likely due to nonspecific DNA binding.



a targeting plasmid that deleted the full *dpy-13* coding region (2.2 kb deletion), and inserted a GFP marker expressed in coelomocytes and the *unc-119(+)* selection marker. After two generations (one week), we visually screened plates for the presence of *unc-119(+)* worms that lacked the co-injection markers. We identified three putative deletion strains (3 of 83 injected worms = 3.7%). Each strain was homozygous for *dpy-13* mutations, the *unc-119(+)* marker and expressed GFP in coelomocytes; we confirmed the 2.2 kb deletion by PCR analysis (Supplementary Fig. 1 and Supplementary Table 1).

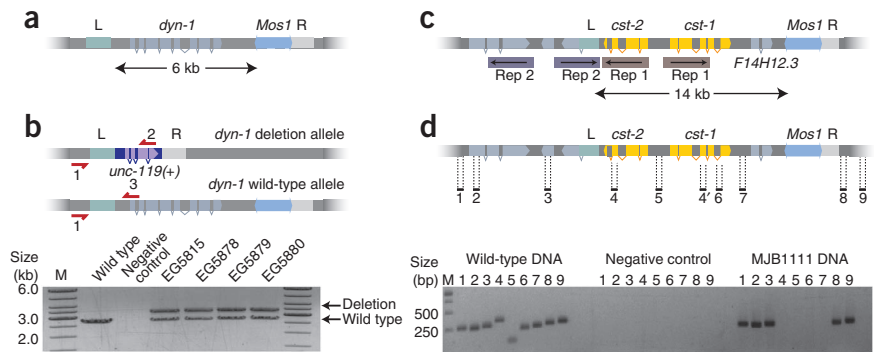
In most organisms, the efficiency of transgene-instructed gene conversion drops rapidly with distance from the DNA breakpoint; only 25% of conversions extend 1 kb from the breakpoint<sup>7,11</sup>. To determine how much DNA could be feasibly deleted using MosDEL, we injected additional constructs coding for 5, 10, 15, 25, 35 and 50 kb deletions (Fig. 1b). The frequency of *unc-119(+)* insertion was largely independent of the targeting construct: we found rescued strains in the progeny of about 5% of the injected worms (Supplementary Table 1). For example, we isolated seven stable lines with *unc-119(+)* insertions from 81 injected worms (9%) using the 25 kb construct. Two of the lines did not exhibit a Dpy phenotype and thus were simply insertions of the *unc-119(+)* transgene. Five strains exhibited a Dpy phenotype. One of these five strains did not have a deletion of the entire 25 kb stretch, whereas the other four had deletions of the entire 25 kb region (4/7 lines = 57% deletions correct) based on PCR analysis (Fig. 1c). However, the generation of deletions decreased sharply for events larger than 25 kb: we did not recover full-length deletions using the 35 kb and 50 kb templates (Supplementary Table 1). For these larger deletions, all *unc-119(+)* insertions (13 strains) were incomplete deletions (data not shown).

We used comparative genome hybridization (CGH) to verify deletion endpoints and confirm PCR results. In CGH, fluorescently labeled mutant DNA is compared to binding of wild-type DNA on a high-density array of gene-specific oligonucleotides<sup>12</sup>. CGH analysis verified the deletion of the targeted genomic regions in the three strains tested: a 25 kb deletion (Fig. 1d) and two 15 kb deletions (data not shown).

Deletion of essential genes is lethal when homozygous. Such deletions must be maintained as heterozygotes over a balancer chromosome; the balancer chromosome usually contains a marker that identifies loss of heterozygosity. Balanced strains often degenerate because of recombination between the marker and deletion on the homologous chromosomes. MosDEL leads to perfectly balanced lethal chromosomes by inserting a selectable marker (*unc-119(+)*) at the site of the deletion. Rescue of the *unc-119(-)* uncoordinated phenotype identifies the presence of the deletion in heterozygotes. Because *unc-119* worms are subviable, the homozygous balancer worms are selected against, and the deletion chromosome is maintained as a heterozygote. To test the utility of this feature, we deleted the entire coding region of the *C. elegans* dynamin ortholog *dyn-1* (Fig. 2a). From 100 injected worms, we obtained 17 strains that were *unc-119(+)* and did not contain extrachromosomal arrays based on the absence of red fluorescent markers. As expected from a lethal mutation balanced by the insertion of *unc-119(+)*, we could not generate homozygous *unc-119(+)* rescued worms for 11 of 17 strains (65%). These putative deletion strains segregated as 50% wild-type heterozygotes, 25% *unc-119* worms and 25% dead *dyn-1* embryos. We selected five of these 11 strains for verification. PCR analysis confirmed that four of five strains contained the full targeted deletion (Fig. 2b), and lethality could be rescued by a *dyn-1(+)* transgene.

**Figure 2** | Deleting lethal genes and multiple similar genes. (a) Schematic for the deletion of the essential gene *dynamain dyn-1*.

(b) Schematic showing the balanced heterozygous deletion mutant. The *dyn-1* deletion allele is balanced with the insertion of *unc-119(+)*. The gel shows PCR verification of deletions in heterozygous *dyn-1* strains (EG5815, EG5878, EG5879 and EG5880) with the three oligos (1, 2 and 3) shown in the schematic. M, 1 kb DNA marker. (c) Schematic showing the *cst-1* and *cst-2* genomic region. *cst-1* and *cst-2* are 100% identical in the inverted repeats ("Rep 1"). An adjacent region contains a second inverted repeat ("Rep 2"), which overlaps the left homology region ("L"). The targeting construct removes 14 kb of genomic DNA, including *cst-1*, *cst-2* and *F14H12.3*. (d) PCR verification of the strain MJB1111 with deletions of *cst-1*, *cst-2* and *F14H12.3*. Oligos were designed to amplify 200–300 bp fragments inside and outside the deletion. M, 1 kb DNA marker.



Occasionally, the genome contains tandem gene duplications that provide redundant function or operons encoding genes with related functions. The loss-of-function phenotype of such loci requires the deletion of both genes, which can be accomplished using MosDEL. To test our system for this application, we targeted protein kinase genes *cst-1* and *cst-2* (Fig. 2c). These genes are adjacent to each other as two identical inverted repeats in a complex genomic region. We designed a deletion template for these two genes and isolated two strains containing *unc-119(+)* insertions; of these, one strain contained the correct deletion (Fig. 2d).

These results show that MosDEL can be used to target a gene if there is a *Mos1* insertion within 25 kb of the gene. Our method has several advantages relative to current knockout techniques. First, the technique is relatively fast and efficient. Second, the endpoints of deletions can be specified to completely eliminate the gene so that no partial gene products are generated. Third, lethal mutations are balanced by the positive insertion marker. In cases in which alleles do not have an obvious phenotype, a fluorescent protein marker can be inserted to follow the mutation in crosses. Finally, several adjacent genes can be deleted. This is particularly useful when genes with similar function are grouped.

The technique relies on the prevalence of *Mos1* insertions in the genome. We analyzed the distribution of the 14,305 *Mos1* elements relative to all 20,160 genes in *C. elegans*. Of these, 20,043 genes (99.4%) fell within 25 kb of at least one *Mos1* element (median distance to nearest *Mos1* element was 3.1 kb), so essentially all *C. elegans* genes can be targeted by MosDEL (Supplementary Fig. 2 and Supplementary Table 2). Thus, 10,069 of 10,154 genes presently lacking a deletion allele can be deleted by MosDEL. Moreover, 8,803 genes have no other genes between them and the *Mos1* element so these genes can be removed as a single-gene deletion.

Plasmid-mediated repair of double-strand breaks is not limited to *C. elegans*. For example, plasmid-driven repair has been described in *Drosophila*<sup>13</sup>, and MosDEL could be adapted to make deletions in fruit flies. This technique should be a useful tool for the *C. elegans* research community and possibly for other genetic model organisms.

## METHODS

Methods and any associated references are available in the online version of the paper at <http://www.nature.com/naturemethods/>.

Note: Supplementary information is available on the Nature Methods website.

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## AUTHOR CONTRIBUTIONS

C.F.-J., M.W.D., G.H. and E.M.J. conceived and designed experiments; C.F.-J., G.H., J.T., P.N., R.L. and M.P.-D. performed experiments; T.W.H. performed the bioinformatic analysis of *Mos1* distribution; M.B., D.G.M. and E.M.J. provided supervision and funding; C.F.-J. and E.M.J. wrote the manuscript.

## COMPETING FINANCIAL INTERESTS

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## ONLINE METHODS

**Nematode strains.** *Mos1* alleles were selected by visual screening in Wormbase (<http://wormbase.org/db/gene/variation?name=cxti10882>) for appropriately located transposon insertions and provided by members of the NemaGENETAG consortium. *Mos1* insertions were made homozygous and analyzed in crosses by PCR. Strains were maintained on nematode growth medium (NGM) plates seeded with OP50 or HB101 bacteria. Strains with *Mos1* elements and all deletions generated are listed in **Supplementary Table 3**.

**Deletion protocol.** *Mos1* alleles were identified in Wormbase and requested from the NemaGENETAG consortium (<http://elegans.imbb.forth.gr/nemagenetag/>). The presence of the *Mos1* insertion was verified with gene-specific primers, annealing inside and outside of the *Mos1* element. The *Mos1* element was crossed into the *unc-119(ed3)* background to make an injection strain and verified by PCR analysis.

Targeting constructs were designed to contain (i) a right homology region (adjacent to the *Mos1* element), (ii) a left homology region (distant from the *Mos1* element) and (iii) a *Caenorhabditis briggsae unc-119(+)* rescue region. The right homology regions comprised approximately 2 kb of homologous DNA immediately adjacent to the *Mos1* insertion site. The left homology regions comprised 2–3 kb of homologous DNA, which specifies the endpoint of the targeted deletion. These regions were selected to avoid repetitive DNA sequences, in particular short inverted repeats, which are likely to anneal and reduce the frequency of correct deletions. DNA between the right and left homology region is deleted; DNA contained in the left region is retained in the deletion strain. A *C. briggsae unc-119(+)* rescue fragment was chosen because it is smaller than the *C. elegans unc-119* gene.

An injection mix was made that contained the targeting plasmid (50 ng  $\mu\text{l}^{-1}$ ), *Mos1* transposase helper plasmid pJL43.1 (ref. 14) (50 ng  $\mu\text{l}^{-1}$ ), plasmids encoding co-injected mCherry markers pGH8 (ref. 10) (*Prab-3::mCherry*, 10 ng  $\mu\text{l}^{-1}$ ), pCFJ90 (ref. 10) (*Pmyo-2::mCherry*, 2.5 ng  $\mu\text{l}^{-1}$ ) and pCFJ104 (ref. 10) (*Pmyo-3::mCherry*, 5 ng  $\mu\text{l}^{-1}$ ). The injection strain was maintained at 15 °C on HB101 bacteria. Young adult worms were mounted on an agarose pad under halocarbon oil and injected following standard techniques. Injected worms were left to recover at 15 °C for several hours and then transferred three at a time to HB101- or OP50-seeded NGM plates and placed at 25 °C. In our hands, approximately 70% of all injected worms resulted in transgenic progeny. We found that growing worms on HB101 bacteria at 15 °C considerably improved the health of *unc-119* worms.

After approximately 7 days each plate was screened for deletion mutants. Screening was greatly facilitated by allowing the plate to starve out completely because *unc-119* worms cannot form dauers and are therefore selected against. Strains with an *unc-119(+)* insertion were identified on a fluorescence dissection microscope as first stage larval (L1) worms that move like wild-type worms but have none of the fluorescent co-injection markers. A single rescued, nonfluorescent worm was picked to a new plate and allowed to propagate for one generation. In the case of obvious phenotypes (for example, *Dpy-13*) a single mutant worm was picked from the progeny; in cases in which the phenotype was wild type, ten worms were picked to individual plates and tested for homozygosity.

It took approximately 2 weeks from injecting the strain to recovering a homozygous deletion worm.

**Analysis of *Mos1* distribution.** We calculated the distance of every protein-coding gene in the WS205 referential release of WormBase (<http://ws205.wormbase.org/>)<sup>15</sup> to all current *Mos1* insertion alleles using a state machine algorithm written in Perl. The closest *Mos1* element was defined as the distance from the insertion site to the ATG start codon. The number of *Mos1* elements in the vicinity of each gene was determined by extracting a sequence segment upstream and downstream of the ATG and summing the number of elements contained within that span. The number of intervening genes between a given gene and its nearest *Mos1* element was determined by extracting the segment ranging from the ATG to the insertion site and tallying the number of genes present, including genes that partially reside within the interval. The analysis was repeated against all genes lacking a deletion allele from either the *C. elegans* Gene Knockout Consortium in the US and Canada (*ok* alleles) and the National BioResource Project in Japan (*tm* alleles).

**Comparative genome hybridization.** Genomic DNA from worms was isolated with the Genra Puregene Tissue kit (Qiagen) following the manufacturer's supplementary protocol, "Purification of archive-quality DNA from nematodes or nematode suspensions using the Genra Puregene Tissue Kit". DNA labeling, sample hybridization, image acquisition and determination of fluorescence were all performed as previously described<sup>12</sup>. We used a 3× high-density (HD) chip divided into three 720,000 whole-genome sections for all experiments. The chip design was based on our original 385,000 whole genome chip<sup>12</sup>. All microarrays were manufactured by Roche-NimbleGen with oligonucleotides synthesized at random positions on the arrays. Chip design name is 90420\_Cele\_RZ\_CGH\_HX3. For all experiments, normalization of intensity ratios were performed with a local scatterplot smoothing (LOESS) regression as previously described<sup>12</sup>. Three strains, EG5810 (25 kb deletion), EG5620 (15 kb) and EG5621 (15 kb), were tested against wild-type DNA. All strains had the targeted deletions. All samples also had two identical untargeted deletions: an approximately 8 kb deletion of *pgp-6* and *pgp-7* on chromosome X and an approximately 4 kb deletion of the telomeric region *cTelX3.1* at the left end of chromosome V. We verified by CGH that these deletions were present in the parent strain (EG5003), and the deletions therefore do not represent second site mutations caused by the MosDEL technique.

**Molecular biology.** Targeting vectors typically consist of three distinct fragments: a right homology region, a positive selection marker (*cb-unc-119(+)*) and a left homology region. In some cases the positive selection marker was flanked by the fluorescent marker *Punc-122::GFP*, which is dimly expressed in the coelomocytes. See **Figure 1a** for a schematic overview of the components of the targeting vector. All targeting vectors were made using the MultiSite Gateway Three-Fragment Vector Construction kit (Invitrogen).

To verify deletions by PCR, we designed oligos that would amplify short genomic DNA fragments inside and outside the targeted regions. These reactions were reproducible and robust; PCR amplification was successful on crude genomic lysates

from five to ten worms or from high quality DNA samples prepared with a Gentra Puregene kit (Qiagen).

***dyn-1* heterozygous verification primers.** Complete *dyn-1* deletions were verified at the 5' end by PCR amplification with the three primers: oGH154, oGH133 and oCF125. Oligos oGH154 and oGH133 gave a 2.7 kb PCR product when the wild-type copy of *dyn-1* is present. Oligos oGH154 and oCF125 gave a 3.4 kb PCR product when the *dyn-1* locus is deleted. The deletion was verified at the 3' end by PCR with oligos oGH155, oGH156 and oCF400. Oligos oGH155 and oGH156

produce a 2.5 kb PCR product when the wild-type *dyn-1* locus and the *ttTi14024 Mos1* element are present. Oligos oGH155 + oCF400 produce a 3.0 kb PCR product when the *dyn-1* gene is deleted.

All molecular biology analysis and design was done with the program ApE, a plasmid editor, that is freely available at <http://www.biology.utah.edu/jorgensen/wayned/apel/>.

All oligos and plasmids are listed in **Supplementary Table 3**.

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