Conditional targeted genome editing using somatically expressed TALENs in *C. elegans*

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We have developed a method for the generation of conditional knockouts in *Caenorhabditis elegans* by expressing transcription activator–like effector nucleases (TALENs) in somatic cells. Using germline transformation with plasmids encoding TALENs under the control of an inducible or tissue-specific promoter, we observed effective gene modifications and resulting phenotypes in specific developmental stages and tissues. We further used this method to bypass the embryonic requirement of *cor-1*, which encodes the homolog of human severe combined immunodeficiency (SCID) protein coronin, and we determined its essential role in cell migration in larval Q-cell lineages. Our results show that TALENs expressed in the somatic cells of model organisms provide a versatile tool for functional genomics.

The nematode *C. elegans* has been a popular model system to study basic biology and human diseases for decades, and many genetic tools and resources are available for this organism^{1–7}. Among the 20,377 predicted protein-coding genes, 6,764 currently have deletion or null mutations, which were generated primarily by random chemical mutagenesis or Mos1 transposon–based targeted gene deletion^{1,3}. However, the inability to conditionally edit the wild-type *C. elegans* genome in a targeted fashion has limited the biological questions that can be addressed with current technology.

TALEN technology has recently been developed to generate locusspecific mutations in the genome^{8,9}. TALENs comprise a nonspecific FokI nuclease domain fused to a customizable repeat domain that recognizes a predictable DNA sequence. The specific DNA recognition domain directs the nuclease to introduce DNA double-strand breaks at the target site, and the erroneous repair by nonhomologous end-joining often induces a mutagenic deletion or insertion at the breakpoint^{8,9}. Although TALENs have been used to edit the genome in a wide range of organisms, their use has been limited to cultured cells or embryos, as well as the germ line in *C. elegans*^{7–10}. So far it has been unknown whether TALENs can be directly applied to somatic cells of a multicellular organism. In this study, we report that a conditional knockout can be achieved in somatic lineages of *C. elegans* by expressing TALEN constructs with an inducible or tissue-specific promoter (hereafter referred to as somatic TALENs).

As a proof-of-principle experiment, we examined whether somatic TALENs could conditionally disrupt the dpy-5 gene in the C. elegans genome. *dpy-5* encodes a cuticle collagen that affects body length¹¹, and its mutation causes a dumpy, short phenotype that can be easily scored. We first generated a TALEN pair targeting dpy-5 by selecting a 52-base-pair (bp) region in the dpy-5 coding sequence, which includes the left and right binding sites and a restriction site for SacI (Fig. 1a). We chose to use disruption by TALENs of a SacI site in the dpy-5 sequence as a molecular assay. We used the 'unit assembly' method to construct the transcription activator-like effector repeats¹⁰. To achieve temporally controlled mutation of *dpy-5*, we used the promoter of the heat shock gene *hsp-16.2* (Phsp)¹² to express *dpy-5* TALEN constructs (Fig. 1a). We generated transgenic *C. elegans* by germline injection of TALEN plasmids expressing both TALEN left and right constructs and a selection marker¹³. After heat shock treatment of transgenic animals at the first larval L1 stage, we detected the DPY phenotype in 93 \pm 4% of all adult animals (**Fig. 1b**,c; *n* = 267 from three generations). The DPY phenotype was not observed in the next generation of these animals because the heat shock promoter is not active in C. elegans germline cells¹².

We found that the DPY penetrance depended on the developmental stage during which the expression of the *dpy-5* TALENs was induced. Penetrance increased in the late embryos as the heat shock promoter (*Phsp*) was first activated¹² and reached its maximum in L1 larvae, which is consistent with previous work showing that *dpy-5* is expressed before secretion of new cuticle from the L1 larval stage¹¹. We did not observe the DPY phenotype in transgenic animals without heat shock (**Fig. 1b,c**; *n* > 100) or in animals expressing only the left or the right portion of *dpy-5* TALENs (*n* = 81 or 90) and subjected to heat shock.

We examined the molecular lesions generated by *dpy-5* TALENs by detecting the disruption of the SacI site in the *dpy-5* sequence (**Fig. 1a**). We PCR-amplified a 402-bp genomic DNA fragment containing the target site from transgenic animals and digested the amplified DNA with SacI. In transgenic animals without heat shock, we showed that the PCR product was completely digested by SacI to a 239-bp and a 163-bp fragment (**Fig. 1d**; left). However, we detected intact DNA fragments after digestion in transgenic animals with heat shock, indicating mutations of the Sac I site were caused by *dpy-5* TALENs (**Fig. 1d**; right). We noticed that the molecular knockout

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Figure 1 dpy-5 was conditionally edited by spatially or temporally controlled somatic TALENs in C. elegans. (a) Schematic representation of TALEN-mediated temporal regulation of *dpy-5* function in *C. elegans*. Two plasmids that express the left and right recognition sites of dpy-5 (green) and FokI (red) under the control of the heat shock promoter (gray, Phsp) transformed C. elegans by germline injection. Upon heat shock, Fokl enzymes, fused with the *dpy-5* TALENs binding domain, were generated and mutated the dyp-5 locus (right). The sequence cut by Fokl contains a recognition site for the Sacl restriction enzyme (blue). (b) Animals carrying Phsp::/dpy-5-TALENs (TLR) constructs without (left) or with (right) heat shock treatment. Animals express both the left and right TALEN constructs. Scale bar, 50 µm. (c) Left; quantification of DPY animals carrying Phsp::dpy-5-TALENs (TLR) constructs without (red) or with (green) heat shock treatment at the indicated developmental stages. Right; the penetrance of the DPY phenotype in animals expressing Punc-119::dpy-5-TLR (blue) or Pdpy-5::dpy-5-TLR (purple). N = 92-267 from three independent experiments. (d) A representative gel of the Sacl restriction enzyme assay of dpy-5 PCR products amplified from the genomic DNA in animals carrying Phsp::/dpy-5-TALENs before (left) or after (right) heat shock. The full-length gel is presented in Supplementary Figure 1c. (e) DNA sequence of the *dpy-5* locus from animals carrying Phsp::/dpy-5-TALEN constructs after heat shock. Dashed lines, deleted nucleotides. Red shows inserted nucleotides. Asterisks, the deletion or insertion causes a frameshift of the dpy-5 open reading frame. Underlined sequences, binding sites of dpy-5 TALENs.

frequency is low in the transgenic animals, which is probably because TALENs are only expressed in a subset of *C. elegans* tissues. Sequencing of intact fragments confirmed that different insertions and deletions had occurred at the target site (**Fig. 1e**).

To achieve spatially controlled mutation of dpy-5, we expressed dpy-5 TALENs using tissue-specific promoters. Expression of dpy-5 TALENs by either the dpy-5 endogenous promoter (Pdpy-5) or a neuronal promoter (Punc-119) caused mutations in the dpy-5 locus (**Supplementary Fig. 1a**). However, only dpy-5 TALENs expressed by the Pdpy-5 promoter, but not the Punc-119 promoter, produced the DPY phenotype (83% DPY, n = 76 for Pdpy-5; 0% DPY, n = 82 for Punc-119; **Fig. 1c**, right), consistent with the autonomous function of dpy-5 in hypodermal cells¹¹. We noticed that the expression of the dpy-5 TALENs by the heat shock promoter generated a more pronounced DPY phenotype than expression by the dpy-5 promoter, which is likely owing to the mutation of dpy-5 in the hypodermal precursor cells by Phsp but not by Pdpy-5.

We further examined whether somatic TALENs targeting *lon-2*, a member of the glypican family of heparan sulfate proteoglycans¹⁴, could increase *C. elegans* body length. *lon-2* mutation causes the long phenotype¹⁴. We generated transgenic animals expressing *lon-2* TALENs under the control of *Phsp*. After the heat shock treatment, we found that 63% of animals expressing the *lon-2* TALENs (n = 200) developed the long phenotype (**Supplementary Fig. 2**), indicating that TALENs may be generally used to edit genes in *C. elegans* somatic tissues.

We next investigated whether somatic TALENs can disrupt multiple copies of a transgene in the genome. We used the Pgcy-32 promoter to express the *gfp* gene in oxygen-sensory URX, and AQR and PQR (A/PQR) neurons, and Q-cell asymmetric divisions generate A/PQR neurons. (**Supplementary Fig. 3a,b**). Transgenic animals were generated by standard germline injection and subsequent integration of extra-chromosomal arrays, which normally occurs with the insertion of multiple copies of transgenes into the genome¹³. We detected the GFP fluorescence from Pgcy-32::gfp in the expected cells at low magnification (**Supplementary Fig. 3b,c**). We then expressed gfp TALENs in these animals using the *C. elegans* Q cell–specific Pegl-17 promoter. Because A/PQR neurons but not URX neurons are Q-cell



descendants (**Supplementary Fig. 3a**), Pegl-17::gfp-TALENs should mutate gfp in A/PQR neurons only (**Fig. 2a**). Indeed, we found that 57 \pm 5% of the animals (n = 85 from three generations) lost GFP fluorescence in A/PQR but not in URX neurons. It is possible that TALENs cut the transgenes in multiple sites, which led to the instability or the loss of the transgene. None of the transgenic animals lost red fluorescence from Pgcy-32::mCherry in any neurons (n = 85, **Fig. 2b** and **Supplementary Figs. 1b** and **3b**,c). These data not only reinforce our finding that the conditional editing by somatic TALENs can be achieved using cell lineage–specific promoters but also demonstrate that somatic TALENs can efficiently knock out multiple copies of a transgene.

We applied somatic TALENs to address the function of a *C. elegans* embryonic lethal gene during larval development. We chose to study *cor-1*, a worm homolog of the SCID gene coronin¹⁵. Coronins encode a conserved family of actin-binding proteins¹⁶, and mouse genetics indicates that coronin 1 is essential for T-cell migration¹⁷. However, recent studies suggested that coronin 1 regulates T-cell function not through the actin cytoskeleton but by affecting cell viability¹⁸. *cor-1* has—to the best of our knowledge—not been studied in *C. elegans* previously. Q-cell development in L1 larvae can be a useful model to dissect the function of COR-1 because Q-cell migration, asymmetric division and apoptosis are actin-dependent processes and these events can be followed at single-cell resolution¹⁹. However, the embryonic lethality of *cor-1* in larvae.

To visualize the final position of Q-cell progenies after migration, we used Pmec-4::gfp and Pgcy-32::mCherry to mark the mechanosensory AVM and PVM (A/PVM) neurons, and A/PQR cells, respectively. In wild-type (WT) animals, the bilateral Q neuroblasts

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Figure 2 Somatic TALENs mutated gfp in C. elegans Q-cell lineages. (a) GFP (green) and mCherry (red) fluorescence in URX and AQR/PQR neurons due to the expression of Pgcy-32::gfp and Pgcy-32::mCherry in WT animals (upper) or worms expressing Q cell-specific gfp TALENs constructs driven by the egl-17 gene promoter (Pegl-17, lower). Still images are shown in Supplementary Figure 3b,c. (b) Quantification of GFP- and mCherry-positive AQR/PQR and URX neurons in WT (purple) or animals expressing Pegl-17::gfp-TLR (blue). N = 88–121.

on the left (QL) and right side (QR) of the animal produce cells that migrate in opposite directions along the anterioposterior body axis¹⁹. QR descendants (QR.x) migrate anteriorly; AQR cells reach their final destination close to the posterior of URX cells, whereas AVM cells move past ALM cells (Fig. 3a,b). The QL-descendant PQR cells migrate posteriorly near PLMs, whereas PVM cells stay in the birthplace (Fig. 3a,b).

We first examined whether somatic TALENs of a gene, for which the null phenotype in Q-cell migration is already known and is not lethal, could reproduce the migration defects. mab-5 encodes a homeodomain transcription factor, and with a *mab-5(e2088)* null allele, QL.x switches its posterior migration to the anterior whereas QR.x anterior migration is not affected (Fig. 3c)²⁰. We generated transgenic animals expressing somatic mab-5 TALEN constructs in Q cells using the Pegl-17 promoter. We found that QL.x (PQR and PVM cells) switched their posterior migration to the anterior in 61% of the *mab*-5-TALEN animals (n = 71) and that QR.x (AQR and AVM cells) migrate normally to the anterior in these animals (Fig. 3d and Supplementary Fig. 4). Our data demonstrate that somatic TALENs of *mab-5* can specifically recapitulate the QL.x migration phenotype in *mab-5* null allele.

We next used somatic TALENs to mutate cor-1 within Q-cell lineages and then analyzed migration and other events in Q-cell descendants. In transgenic animals expressing Pegl-17::cor-1-TALENs in Q-cell lineages, Q-cell descendants moved in the correct directions; however, their migration distances were substantially reduced (Fig. 3e,f). AQR cells went further in the posterior direction than URX cells, whereas AVM cells were in the posterior of ALM cells and PQR cells

Figure 3 Somatic TALENs conditionally mutated mab-5 and cor-1 and revealed the function of COR-1 in C. elegans Q-cell migration. (a) The final position of mechanosensory neurons (green, marked by Pmec-4::gfp) and oxygen sensory neurons (red, marked by Pgcy-32::mCherry) in WT animals. Asterisks, Q-cell birthplace. AQR/PQR and AVM/PVM are derived from Q-cell lineages. (b-e) Still images show the final position of Q-cell progenies in WT animals (b) or mab-5(e2088) mutants (c) or in worms expressing Q cell-specific mab-5 (d) or cor-1 (e) TALENs constructs driven by Pegl-17. Yellow arrows point to the position of AQR, AVM, PVM and PQR. URX and PLM were used as fiducial markers to quantify their positions. AQR and AVM are on the right side of the animal whereas PQR and PVM are on the left. In mab-5 mutants or mab-5 TALEN animals, PVM is always more posterior than AVM. Dashed lines show the animal periphery. +, vulva. Scale bar, 50 µm. (f) Quantification of AQR (left), AVM (middle) and PQR (right) positions in WT (red; n = 25) and animals with *cor-1* TALENs in Q cells (green; n = 40).

were close to the nonmigratory PVM cells (Fig. 3e,f). We showed that deletion/insertion mutations by cor-1 TALENs caused the frameshift in the cor-1 open reading frame (Supplementary Fig. 5). These data demonstrate that the disruption of cor-1 in Q-cell lineages reduces Q-cell migration (Fig. 3a). Defects in Q-cell asymmetric division or apoptosis produce ectopic neurons, whereas the failure of Q-cell survival should cause a loss of neurons. Quantifying the number of A/PQR or A/PVM cells in these animals (n = 83), we did not find any gain or loss of these neurons. Our data demonstrate that COR-1 is only essential for cell migration and is not involved in cell proliferation or survival, at least, in *C. elegans* Q-cell lineages.

Compared to current C. elegans conditional gene-inactivating techniques or tools for reverse genetics, somatic TALENs have several advantages. First, the technique conditionally edits the WT C. elegans genome and can generate mutations in the specific somatic cell lineages or developmental stages, providing a versatile tool to address gene function. Alternative techniques such as the Cre/LoxP system⁴ and FLP-FRT conditional system² were applied in C. elegans, but a null allele mutant of the target gene, which is not available for every *C. elegans* gene, is required to start with. Somatic TALENs can work on any genome, which will greatly facilitate functional genomics. Second, somatic TALENs can produce robust phenotypes with low variability, which can be explained by the same type of mutation (e.g., frameshift) that is generated during DNA repair in most cells (Fig. 1e and Supplementary Figs. 1b and 5). A conditional RNA interference (RNAi) technique was developed using neuron-specific expression of the membrane protein SID-1 in the *sid-1* mutant background²¹. However, RNAi is sensitive to experimental conditions and its use can cause residual gene expression⁶, causing high variability or low penetrance of the phenotype. Furthermore, C. elegans neurons are wellknown for being refractory to RNAi⁶. Indeed, our attempts to knock down gfp in transgenic animals expressing Pgcy-32::gfp, including the use of various RNAi-sensitive mutants or the direct expression



of double-stranded RNA in target neurons, have not been successful (data not shown). By contrast, somatic TALENs can efficiently eliminate GFP fluorescence from the *Pgcy-32::gfp* transgene (**Fig. 2a,b**, and **Supplementary Figs. 1b** and **3b,c**). Temperature-sensitive alleles can be utilized to conditionally inactivate genes, but the penetrance is often limited because of only partial loss of function of the gene⁵. Moreover, producing a specific temperature-sensitive allele of every gene in *C. elegans* does not seem feasible. A third potential advantage is that the somatic TALENs technique is relatively fast and efficient. Starting from the experimental design, it might take only 3 weeks or less to obtain edited animals for phenotypic analysis.

This study focused on the local genome-editing changes induced by somatic TALENs, and more complete analyses will be required to assess potential off-target effects of TALENs. Prior deep sequencing studies on germ line–transmitting fish or human pluripotent cells did not uncover off-target effects caused by TALENs^{10,22}. Our somatic TALENs used obligate heterodimer-based nuclease fusions, which can considerably increase the specificity of the gene-editing²³. In our studies of *dpy-5*, *lon-2*, *mab-5*, *cor-1* and *gfp*, we did not notice additional morphological defects. Moreover, somatic TALENs of *cor-1* revealed its essential role in cell migration but not in other events, suggesting that somatic TALENs may not generally result in off-target effects.

In conclusion, somatic TALENs offer a useful tool for the *C. elegans* community. In principle, somatic TALENs should be useful in a variety of other organisms, including those in which conditional knockout techniques do not exist or are time consuming to implement. If the issue of off-target mutations can be addressed in the future, somatic TALENs may also provide an alternative strategy for gene therapy.

METHODS

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

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

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AUTHOR CONTRIBUTIONS

Z.C., P.Y., W.L. and G.O. designed experiments; Z.C., P.Y., X.W., Y.C., G.F., Y.Y., X.L., P.Y., Z.Z. and W.L. performed experiments; G.O. wrote the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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C. elegans strains, genetics and DNA manipulations. *C. elegans* strains are listed in Supplementary Table 1. All strains were raised at 20 °C on nematode growth medium (NGM) plates seeded with the *Escherichia coli* strain OP50. All PCR reactions were done with Phusion DNA polymerase (New England Biolabs). PCR templates, primers and plasmid constructs are listed in Supplementary Table 2. Transgenic *C. elegans* were created by germline transformation. PCR products and DNA plasmids at 10–30 ng/µl were injected into N2 hermaphrodites with a selection marker Podr-1::dsRed, pRF4 or Pegl-17::mCherry::TEV-S::his-24.

TALEN design. TALENs were designed based on three criteria. First, the binding sites were selected to be 16–18 bases in length to ensure specificity. Second, the last base of each binding site was fixed as thymine. Third, the spacer regions were selected to be 14–18 base pairs (bp) in length. BLAST (NCBI) was run on TALEN binding sites to check potential off-target sites. For example, the *dpy-5*-TALEN recognition sequences are: left 5'-GACATGTATGATGAT GT-3' and right 5'-CAGAGATATCTCTGAAT-3'. The spacer region contains a SacI restriction site (underlined): 5'-GATGGGA<u>GAGGCTC</u>GGAGG-3'. The information on TALEN recognition sequences of *lon-2, gfp, mab-5* and *cor-1* is available in **Supplementary Table 3**.

TALEN constructs. TALEN somatic expression backbones were modified from pCS2-PEAS and pCS2-PERR¹⁰. TALEN coding sequences, referred to as PEAS and PERR in the original plasmids, were cloned into *C. elegans* expression vector pPD95.77. The *Pegl-17*, *Pdpy-5*, *Punc-119* or *Phsp-16.2* promoter was then inserted for Q-cell, hypodermal, neuronal or inducible expression, respectively. All the RVD-containing repeats in this study were constructed using a unit assembly approach¹⁰. The fulllength repeats were cut down from TALEN unit assembly plasmid by SpeI and NheI and then cloned into somatic expression backbones by means of In-Fusion Advantage PCR cloning kit (Clontech, cat. no. 639621). To reduce the potential off-targeting effects, we used a sharkey form with the obligate heterodimeric AS:RR pair of FokI cleavage domain in TALEN expression vectors²³. *C. elegans* heat shock treatment. We first synchronized the culture by allowing 50–100 adult worms to lay eggs for 2 h in a seeded NGM plate. The eggs were raised at 20 °C before heat shock. At the desired stage, we treated worms at 33 °C for 1 h. Worms were then raised at 20 °C. The DPY or LON phenotype or the loss of GFP was quantified at the adult stage.

Molecular analysis of *dpy-5* mutations caused by somatic TALENs. Genome DNA was extracted from worms at the L3 or L4 stage. A 402-bp DNA fragment was PCR amplified, concentrated by QIAquick PCR Purification Kit (Qiagen, cat. no. 28104), and purified by MinElute gel extraction kit (Qiagen, cat. no. 28606). DNA concentration was normalized to 60 ng/µl and digested by SacI overnight. 10 µl of digestion product was loaded to 2% (wt/vol) agarose gel in 1× TAE electrophoresis buffer with Golden View dye and then imaged by gel imaging system. The intact 402-bp bands were excised and DNA was extracted using the MinElute gel extraction kit (Qiagen, cat. no. 28606). To sequence the mutations, the intact DNA was cloned into vectors by In-Fusion cloning and used to transform Trans5 α Chemically Competent Cell (Transgen, cat. no. CD201-01). Single colonies were sequenced by Sanger sequencing.

C. elegans imaging. C. elegans adults were anesthetized with 0.1 mmol/L levamisole in M9 buffer, then mounted on 2% agar pads and maintained at room temperature (20 °C). Our imaging system includes an Axio Observer Z1 microscope (Carl Zeiss MicroImaging, Inc.) equipped with a 10× objective, an EM CCD camera (Andor iXon+ DU-897D-C00-#BV-500), and the 488-nm and 568-nm lines of a Sapphire CW CDRH USB Laser System attached to a spinning disk confocal scan head (Yokogawa CSU-X1 Spinning Disk Unit). Images were acquired with exposure time of 300 msec with μ Manager (http://valelab.ucsf.edu/~MM/MMwiki/). ImageJ software (http://rsbweb.nih.gov/ij/) was used to process the images.

Quantification of Q-cell final position. We quantified the final positions of Q-cell progenies in adult worms. The nonmotile cells, URX, PLM and PVM, which were labeled by either Pgcy-32::mCherry or Pmec-4::gfp, were chosen as fiducial markers. For instance, the relative position of AQR is calculated as the distance between URX and AQR divided by the distance between URX and PLM.