Rapid Screening for CRISPR-Directed Editing of the Drosophila Genome Using white Coconversion

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ABSTRACT Adoption of a streamlined version of the bacterial clustered regular interspersed palindromic repeat (CRISPR)/Cas9 defense system has accelerated targeted genome engineering. The Streptococcus pyogenes Cas9 protein, directed by a simplified, CRISPR-like single-guide RNA, catalyzes a double-stranded DNA break at a specific genomic site; subsequent repair by end joining can introduce mutagenic insertions or deletions, while repair by homologous recombination using an exogenous DNA template can incorporate new sequences at the target locus. However, the efficiency of Cas9-directed mutagenesis is low in Drosophila melanogaster. Here, we describe a strategy that reduces the time and effort required to identify flies with targeted genomic changes. The strategy uses editing of the white gene, evidenced by altered eye color, to predict successful editing of an unrelated gene-of-interest. The red eyes of wild-type flies are readily distinguished from white-eyed (end-joining-mediated loss of White function) or brown-eyed (recombination-mediated conversion to the white coffee allele) mutant flies. When single injected G0 flies produce individual G1 broods, flies carrying edits at a gene-of-interest were readily found in broods in which all G1 offspring carried white mutations. Thus, visual assessment of eye color substitutes for wholesale PCR screening of large numbers of G1 offspring. We find that end-joining-mediated mutations often show signatures of microhomology-mediated repair and that recombination-based mutations frequently involve donor plasmid integration at the target locus. Finally, we show that gap repair induced by two guide RNAs more reliably converts the intervening target sequence, whereas the use of Lig4169 mutants to suppress end joining does not improve recombination efficacy.

The ability to make targeted changes in the genome of virtually any organism is transforming biological research. Early genome editing strategies used zinc-finger nucleases (Kim et al. 1996; Smith et al. 1999; Bibikova et al. 2001) or transcription activator-like effector nucleases (Boch et al. 2009; Moscou and Bogdanove 2009; Christian et al. 2010) that required the construction of unique proteins for each target site. In contrast, the discovery that a chimeric single-guide RNA (sgRNA) can direct the Streptococcus pyogenes type II clustered regular interspersed short palindromic repeat (CRISPR)-associated protein 9 (Cas9) to catalyze site-specific double-stranded DNA breaks (DSBs) has eliminated laborious protein construction (Jinek et al. 2012; Qi et al. 2013). To date, Cas9 is active in all tested organisms including bacteria, plants, fungi, and animals (for reviews see Hsu et al. 2014; Sander and Joung 2014; Sternberg and Doudna 2015; Govindan and Ramalingam 2016).

DSBs induced by sgRNA-guided Cas9 stimulate host DNA repair pathways. In many cases the breaks are perfectly rejoined, recreating the original target site, which can be cut again. Occasionally, error-prone end joining inserts or deletes nucleotides at the target site thereby preventing recutting. Such insertions, deletions, and substitutions, collectively called indels, can disrupt a protein-coding sequence. When a DNA donor is supplied exogenously, the DSB can be repaired by homologous recombination (HR), allowing the incorporation of novel sequences at the target site. Unlike sequences incorporated via transgenes, modifying an endogenous gene preserves the chromatin context,
enhancers, promoters, introns, and post-transcriptional regulatory elements of the wild-type locus.

Cas9-mediated genome editing requires just three components: (1) Cas9, which can be provided as a purified protein, mRNA, or gene; (2) sgRNA, which can be provided as an RNA or transcribed in vivo from a DNA template; and (3) a DNA donor bearing the target sequence containing indels or novel sequences to be incorporated. In Drosophila, providing Cas9, sgRNA, and donor DNA transgenes efficiently triggers editing, but establishing the requisite fly stocks takes over a month (Kondo and Ueda 2013; Port et al. 2014, 2015; Chen et al. 2015). Injecting sgRNA and donor DNA into Cas9-expressing embryos requires far less time but is also less efficient, making it necessary to screen large numbers of animals. Cointegrating a visible marker such as GFP into the target locus can speed the identification of recombinants (Baena-Lopez et al. 2013; Gratz et al. 2014; Port et al. 2014, 2015; Ren et al. 2014a,b; Yu et al. 2014; Zhang et al. 2014b; Chen et al. 2015). However, removing the GFP marker by site-specific recombination (e.g., Cre-LoxP) takes multiple generations, negating the time advantage of injection and leaving a “scar” sequence (e.g., LoxP) at the target site. Indels, of course, must be identified molecularly or through complementation analysis.

In Caenorhabditis elegans, coconversion strategies targeting a marker gene together with the gene-of-interest speed the screening for indels and recombinants and avoid introducing an exogenous marker gene at the target locus (Arribere et al. 2014; Kim et al. 2014; Ward 2015). The coconversion strategy restricts molecular screening to marker-positive animals, substantially reducing the work required to find mutant or recombinant animals. In theory, a similar coconversion system should speed genome editing in Drosophila melanogaster.

Here, we describe a strategy in which cotargeting the eye-color gene white (w) speeds identification of both mutants and recombinants at the gene-of-interest. In our strategy, indels generate loss-of-function w mutants whose eyes are white, instead of the wild-type red. In contrast, recombination with the exogenous w’fixc (w’) donor DNA produces flies with reddish brown eyes. Mating the injected animals to w1118 null flies and examining the eye color of their offspring allows rapid identification of parents that produce only w or w’ gametes. These flies have an enhanced frequency of indels or recombination at the cotargeted gene-of-interest.

While developing this coconversion strategy for fly genome editing, we also discovered that Cas9-induced recombinants frequently harbor undesirable integration of the entire donor plasmid at the target locus. We find that inducing gap repair with a pair of sgRNAs increases the likelihood of conversion of the intervening target region. Moreover, when DSBS are repaired by end joining, the junction site frequently contains indels or novel sequences to be incorporated. In Drosophila, providing Cas9, sgRNA, and donor DNA transgenes efficiently triggers editing, but establishing the requisite fly stocks takes over a month (Kondo and Ueda 2013; Port et al. 2014, 2015; Chen et al. 2015). Injecting sgRNA and donor DNA into Cas9-expressing embryos requires far less time but is also less efficient, making it necessary to screen large numbers of animals. Cointegrating a visible marker such as GFP into the target locus can speed the identification of recombinants (Baena-Lopez et al. 2013; Gratz et al. 2014; Port et al. 2014, 2015; Ren et al. 2014a,b; Yu et al. 2014; Zhang et al. 2014b; Chen et al. 2015). However, removing the GFP marker by site-specific recombination (e.g., Cre-LoxP) takes multiple generations, negating the time advantage of injection and leaving a “scar” sequence (e.g., LoxP) at the target site. Indels, of course, must be identified molecularly or through complementation analysis.

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MATERIALS AND METHODS

Fly stocks

vas-Cas9 (y¹, M[vas-Cas9]ZH-2A) was generated by recombining y¹, M[vas-Cas9]ZH-2A, w¹¹¹⁸ (Bloomington #51323; Gratz et al. 2014) with Oregon-R. vas-Cas9, Lig4(¹9) (y¹, M[vas-Cas9]ZH-2A, Lig4(¹9) was generated by recombining y¹, M[vas-Cas9]ZH-2A with w¹¹¹⁸, Lig4(¹⁹) (Bloomington #28877; McVey et al. 2004b). Rainbow Transgenic Flies, Inc. (Camarillo, CA) performed injections.

sgRNA-expressing plasmid construction

sgRNA design: Target loci of the injection strains were sequenced before sgRNAs designed using crispr.mit.edu (Hsu et al. 2013). Guides were preferred if nucleotides 19 and 20 were purines (Farboud and Meyer 2015); positions 15–20, the protospacer-adjacent motif-proximal nucleotides, were ≥33% GC (Ren et al. 2014b); and the sequence placed the guide close to the site of modification. Supplemental Material, Table S7 in File S1 lists sgRNA sequences.

sgRNA cloning: pCFD4, which expresses one sgRNA from a U6:3 promoter and another sgRNA from a U6:1 promoter (Addgene #49411; Port et al. 2014), was modified to remove vermillion and attB (pCFD4d). Sequence- and ligation-independent cloning (Jeong et al. 2012) was used to clone two guide into BbsI-digested pCFD4d following a PCR incorporating one guide after the U6:1 promoter, and the other after the U6:3 promoter (Port et al. 2014). Table S8 in File S1 lists the PCR primers. The 20 nt sgRNA-2 template was inserted into the BbsI sites of pDCC6, which expresses sgRNA from a U6:2 promoter and Cas9 mRNA from the hp708Bb promoter (Gokcezade et al. 2014). Plasmids were purified (Plasmid Midi Kit; QIAGEN, Hilden, Germany) and dissolved in water.

Donor template construction

pUC-w: A 2080 bp fragment, spanning genomic nucleotides X:2,792,206–2,790,141 (D. melanogaster genome release r6.07), was amplified by PCR from w⁰ genomic DNA, sequenced to confirm the w⁰ point mutation and identify natural polymorphisms, and inserted into pUC57 between the SacI and SpIh sites to produce pUC-w. Site-directed mutagenesis was used to mutate the sites targeted by w sgRNAs-1, -2, -3, and -4.

pUC-armi: A 2280 bp DNA (synthesized at GenScript, Inc., Piscataway, NJ) spanning genomic nucleotides 31L:3,464,383–3,466,434 was inserted into pUC57 between the SacI and SpIh sites. The sequence included silent mutations, a naturally occurring nine-nucleotide deletion polymorphism in armi exon 8 that disrupts the armi sgRNA-1 target site, a naturally occurring 12-nucleotide deletion polymorphism in the armi 3’ UTR, and a 36 nt C-terminal Strep-tag II peptide tag.

pCR-zuc: A 2120 bp PCR fragment spanning genomic nucleotides 2L:11,990,382–11,988,263 was inserted into pCR-Blunt II-TOPO to make pCR-zucWT. A 991 bp fragment containing a 3xFLAG peptide tag before the stop codon of zuc and silent mutations disrupting four potential sgRNA binding sites were synthesized as a gblock (Integrated DNA Technologies, Coralville, IA), digested with Ndel and PacI, and inserted into pCR-zucWT between the Ndel and PacI sites to produce pCR-zuc.

Screening for mutations at w

For armi targeting, individual injected G0 adults were mated with two w¹¹¹⁸; D; Dr/TM3, Sb males or virgin females. For zuc targeting, w¹¹¹⁸; Sb/CyO, + ; Dr/TM3, Sb males and wild-type females were used. For zuc targeting, w¹¹¹⁸; Sb/CyO; + ; Dr/TM3, Sb. Three to five-day-old G1 progeny (25°) were assessed by light microscopy (MZ6 Stereomicroscope, Leica Microsystems GmbH, Wetzlar, Germany).

Screening for mutations at the gene-of-interest

Due to the large number of all-red, white-and-red, and coffee-and-red broods, and their lower chance of harboring gene-of-interest conversion,
not all G1 broods were PCR screened. Instead, 44 all-red (37% of total), 46 white-and-red (59%), 8 all-white (100%), 29 coffee-and-red (78%), 11 coffee-and-white (92%), and 15 all-coffee broods (88%) were picked for genotyping. Anesthetized G1 males were deposited on a CO2 pad, and the 9–10 flies closest to the front edge of the pad were individually mated to corresponding balancer virgin females to generate stocks. After 5 d, the G1 males were removed from the crosses, and 1–3 flies from the same brood were homogenized (Gloor et al. 1993) in 30 μl per fly “squishing buffer” [10 mM Tris-Cl, pH 8.0, 1 mM EDTA, 25 mM NaCl, 200 μg/ml freshly diluted Proteinase K solution (AM2546; Thermo Fisher Scientific)] with a plastic pestle (Kimble-Chase Kontes, Vineland, NJ) in 1.7 ml microcentrifuge tubes, incubated at 37°C for 30 min, and then the Proteinase K inactivated at 95°C for 5 min. PCR was used to amplify 505–1225 bp amplicons spanning the target loci from 1 μl homogenate (15 μl final reaction volume; MeanGreen 2× Taq Master Mix, Empirical Bioscience, Inc., Grand Rapids, MI). We note that using this experimental setup, PCR efficiency drops for amplicons longer than 1 kb. Because different sgRNAs targeted different regions of armi or zuc, different PCR primers were designed for each target locus (Table S8 in File S1). Whenever possible, one of the two primers bound only to the genome and not the donor, to avoid amplifying extrachromosomal or ectopically inserted donor DNA. When screening for recombinants with novel sequences knocked-in at the target locus, PCR with one primer bound to the novel sequence (e.g., 3×FLAG) and another primer bound only to the genome and not the donor (Table S8 in File S1) can quickly identify the positive recombinaants. When scanning for indels or recombinants with point mutations at the target loci, we used the following strategies to identify PCR products that contained such mutations.

Restriction enzyme digestion: Because G1 flies inherit one chromosome from the injected G0 embryo and the other from the balancer fly, at least half of the PCR products were amplified from the wild-type gene. We digested the PCR reaction with a restriction enzyme that cleaves adjacent to the predicted DSB in the wild-type ampiclon: PCR products resistant to the restriction digestion should harbor mutations at the recognition site. The uncut PCR product was then gel isolated (QIAquick Gel Extraction Kit, QIAGEN) and sequenced to identify the underlying mutation. This approach ensures that the wild-type PCR products do not confound the sequencing trace and allows the detection of one mutant allele among ≥6 alleles, allowing multiple G1 flies to be pooled in the same PCR. In addition to indels, HR can also be detected by this method, as long as the HR donors are engineered to contain silent mutations that disrupt the restriction enzyme site. A drawback is that the deletion or HR must affect the restriction enzyme recognition sequence; those that do not will not be undetected. The following restriction digestions were used:

armi sgRNA-1 DSB: an AvaII site 6 bp away; 5 μl of PCR digested with AvaII [0.2 U/μl final concentration (f.c.)] in 0.5× CutSmart Buffer (New England Biolabs, Inc., Ipswich, MA) in 10 μl final volume (f.v.) at 37°C for 2 hr;

armi sgRNA-2/3 DSBs: a BstNI site 1 bp (sgRNA-2) or 1 bp (sgRNA-3) away; 5 μl of PCR with BstNI (0.5 U/μl f.c.) in 1× NEBuffer 3.1 (NEB) in 10.5 μl f.v. at 60°C for 1 hr;

armi sgRNA-4 DSB: no restriction enzyme site nearby; digested with T7E1 as described below;

armi sgRNA-5/6 DSBs: a PmII site 17 bp (sgRNA-5) or 11 bp (sgRNA-6) away; 10 μl PCR with Eco72I (0.5 U/μl f.c., Thermo Fisher) in 12.5 μl f.v. at room temperature for 1 hr;

zuc sgRNA-1 DSB: a BclI site 9 bp away; 5 μl of PCR with BclI (0.5 U/μl f.c.) in 0.5× CutSmart Buffer in 10 μl f.v. at 37°C for 1 hr;

zuc sgRNA-2 DSB: a HpyCH4III site 7 bp away; 5 μl of PCR with HpyCH4III (0.25 U/μl f.c.) in 0.5× CutSmart Buffer in 10 μl f.v. at 37°C for 2 hr.

T7 endonuclease I (T7E1) digestion: To complement the restriction enzyme digestion, the same PCR products were denatured, reannealed to form heteroduplex, and digested with the mismatch-specific, sequence-independent T7E1. In G1 single-fly PCR, either 0% (both alleles are wild-type) or 50% (one allele is mutant) of reannealed products will be substrates for T7E1. The drawback of this approach is: (1) some small sequence changes may escape T7E1 digestion; (2) lower sensitivity and higher background prevents the pooling of G1 flies in the same PCR; and (3) as the wild-type PCR products cannot be specifically destroyed, the sequencing trace has to be manually inspected to detect a mutation. To digest with T7E1, 5 μl PCR product was denatured at 95°C for 5 min, reannealed by reducing the temperature 0.1°C/sec to 25°C to allow heteroduplex to form, and then digested with T7E1.
Plasmids and fly strains are available upon request. The authors state that all data necessary for confirming the conclusions presented in the article are represented fully within the article.

**RESULTS**

**w coconversion facilitates screening for both indels and recombinants**

Changes in eye color are among the most readily identified phenotypes in *Drosophila*. Wild-type eyes are bright red with an obvious pseudopupil. Mutations in *w* generate eye colors ranging from brown to yellow for hypomorphic alleles and white for null alleles. Among the alleles of *w* that are caused by point mutations, *w*coffee (w^co^) (Zachar and Bingham 1982) was chosen as the coconversion marker because of its easy-to-screen, reddish brown eyes lacking a pseudopupil. We designed a *w*sgRNA (Figure 1A). HR-mediated coconversion facilitates screening for both indels and recombinants. (See Table 1 for statistics).

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gene-of-interest were coinjected into Lig4+ or Lig4− preblastoderm embryos that express S. pyogenes Cas9 (vas-Cas9) (Gratz et al. 2014). The adult flies that developed from the injected embryos were mated with w1118 flies; the eye colors of the resulting G1 offspring revealed the w genotype of the germline stem cells of the G0 parent. The G1 progeny included coffee-, white-, and red-eyed flies (Figure 1B). Sequencing white and coffee G1 flies confirmed that white-eyed flies (n = 10/10) had indels at the target site in w, whereas flies with coffee-colored eyes contained the G1766A, C1767A w/fl mutation (n = 6/6). Thus, eye color provides an effective reporter for w sgRNA-directed mutagenesis in the fly germline.

Some G0 produced broods with uniformly red-, white-, or coffee-eyed flies, while others produced broods comprising flies of all possible combinations of the three eye colors. Editing of w can occur early in any of the dozens of pole cells that form at the posterior pole of the syncytial blastoderm embryo or later in the descendants of these germ cell progenitors. Because individual G0 pole cells may incorporate different amounts of the injected plasmids, the frequency of DNA cleavage by sgRNA-guided Cas9 and the choice of repair pathways will differ among germ cells, generating variation in the ratio of red-, white-, and coffee-eyed G1 flies. The percentage of nonred G1 flies should reflect the allele frequency of mutant chromosomes in G0 germline stem cells, which in turn reflects the overall targeting efficiency.

To test this idea, we assigned each fertile G0 to one of six groups according to the eye color composition of its G1 brood: (1) all red; (2) white and red; (3) all white; (4) coffee and red or coffee, white, and red; (5) coffee and white; and (6) all coffee (Table 1). Six independent experiments cotargeted w and armi (armi), a third chromosome gene; one experiment cotargeted w and zuc (zuc), a second chromosome gene. Representative numbers of broods across the six eye color groups were screened by genotyping 9–10 G1 flies from each brood for sequence changes at the gene-of-interest (i.e., armi or zuc; Table 1). For simplicity, we combined the three groups containing no red-eyed progeny into a single category, “no red in broods,” and the three groups containing at least some red-eyed flies into a single category, “with red in broods.” The fraction of broods that yielded indels or recombinants was 21% ± 19% in the “with red” category, and 65% ± 34% (mean ± SD) in the “no red” category (Figure 2). Therefore, screening for mutations at a gene-of-interest can be restricted to the “no red” broods, which account for 63–21% of all broods (mean ± SD = 14% ± 6%, Table 1). For these seven experiments, w coconversion would have successfully identified mutants in the gene-of-interest by screening just the 37 “no red” broods (14% of the total 272) using a simple genetic scheme (Figure S1 and Materials and Methods).

**Microhomology-mediated end joining is frequent**

We identified 82 independent indels at seven sgRNA target sites (Figure 3B and Tables S1–S6 in File S1), and grouped them by ligation junction signatures. Two types of deletions were observed: 13 events showed a pair of ≥2 nt long, identical sequences (microhomology) being reduced...
A circular plasmid donor frequently integrates at the target locus

HR in the gene-of-interest was identified by PCR screening using a primer that binds within both the donor and the genomic locus and a primer that binds exclusively to the genomic sequence. This primer pair can amplify the original or the edited genomic locus, but not donor DNA present extrachromosomally or integrated at an ectopic location. As previously reported (Yu et al. 2014), some of the recombinants identified by this strategy corresponded to genomic integration at the gene-of-interest of the entire donor, including the plasmid backbone. In addition to converting the genomic locus to the donor sequence, these recombination events also duplicate the genomic sequence present in the donor (Figure 4A). To distinguish between gene conversion and plasmid integration, we repeated the PCR using primers binding only to the genome and not to sequence present in the HR donor. This strategy readily identified plasmid integration events by their lack of a PCR product or the amplification of a larger-than-expected product. Of the 16 independent HR events identified at armi, seven reflected gene conversion while nine integrated the plasmid, a 56% false-positive rate; of the 12 independent HR events identified at zuc, 10 underwent gene conversion while 2 integrated the plasmid, a 17% false-positive rate (Figure 4B).

Gap repair reliably converts the intervening sequence

When gene conversion occurs, the genomic sequence replaced by donor sequence is termed the “conversion tract.” If the conversion tract is short, mutations can only be introduced near the DSB. On the other hand, long conversion tracts allow a single HR event to introduce multiple mutations that are distant from the sgRNA-complementary site. Given that each gene-targeting experiment in Drosophila takes 2 to 3 months to accomplish, the ability to introduce two or more edits via a long conversion tract is advantageous. We therefore determined the length of conversion tracts in our experiments.

To introduce a peptide tag at the carboxy terminus of the ArmI protein, we assembled a donor plasmid harboring 2280 bp of sequence from the endogenous armi locus and introducing a Strep-tag II peptide tag before the stop codon (Figure 5). The donor harbored 19 sites different in sequence from the injected strain, allowing measurement of the length of the conversion tract. We first designed armi sgRNA-1
of the second, and templated one gene conversion event (Table 1). As expected when both guides direct Cas9 to cleave the genome, the 454 bp interval between the two DSBs was fully replaced with the sequence contained in the HR template plasmid (Figure 5).

We repeated the same strategy with three sgRNAs whose target sites were separated by 280 bp (sgRNA-1, sgRNA-2, and sgRNA-3; sgRNA-2 and -3 had predicted cleavage sites separated only by 7 bp therefore can be considered as a single target site). The donor included 1530 bp upstream of the first target site and 484 bp downstream of the second, and templated three gene conversion events (Table 1). The first tract reliably replaced the 280 bp gap with that of the donor; the second tract converted between 1117 and 1525 bp upstream of the first target site in addition to a full replacement of the 280 bp gap. The third tract lacked gap repair: the first target site harbored a 2 bp insertion after an 11 bp deletion (Table S4 in File S1); the second site harbored a ≥77 bp conversion tract downstream of the DSB. The 280 bp gap was not converted, suggesting separate repair events at the two target sites.

We observed a similar gap repair phenomenon when introducing sequence encoding a carboxy terminal 3×FLAG peptide tag into the zucchini genomic locus (Table 1 and Figure 6). The two guides, zuc sgRNA-1 and -2, targeted sites 395 bp apart. The zucchini HR template included 970 bp upstream of the first target site and 760 bp downstream of the second and templated 18 gene conversion events. Of the two gap repair events, one reliably converted the predicted gap, and the other converted ≥720 bp upstream of the first target site in addition to fully replacing the 395 bp gap. The remaining 16 gene conversion events lacked gap repair: only markers near the zuc sgRNA-1 target site were converted. At the zuc sgRNA-2 target site, six contained indels, and ten had wild-type sequence, suggesting separate repair events at the two target sites.

**Ligase 4 mutation does not inhibit end joining or improve HR**

In flies, mutation of Ligase 4 (Lig4169), a key enzyme in the canonical nonhomologous end-joining pathway, has been proposed to promote HR by suppressing end joining. Zinc-finger nuclease-catalyzed DSBs yield a greater proportion of recombinants in Lig4169 null mutant embryos than in wild-type, but at the cost of decreased fitness of the injected animals (Beumer et al. 2008, 2013; Bozas et al. 2009). Inhibition of Ligase 4 using RNA interference or small molecule protein inhibitors similarly increased HR efficiency in mosquitos (Basu et al. 2015), mice
To test whether Lig4169 null mutants increased the yield of recombinants, we coinjected sgRNA-expressing and HR donor plasmids targeting white into vas-Cas9, Lig4169 or vas-Cas9, Lig4+ embryos. We used the fraction of coffee-producing broods and percentage of coffee-eyed G1 in such broods to score for HR efficiency (Table 2 and Figure 7). Three independent comparisons were conducted, each with a unique sgRNA targeting white. sgRNA-1 and sgRNA-3 were provided on the pCFD4d vector together with armi sgRNA-1, and pDCC6 also carries a Cas9 gene expression unit.

Mothers homozygous for vas-Cas9 and either Lig4+ or Lig4169 produce the expected 1:1 Mendelian ratio of red/coffee-eyed or red/white-eyed siblings, excluding the formal possibility that the Cas9-expressing, Lig4169 background affects the recovery of white mutant flies. We conclude that the use of Lig4169 embryos does not reduce the recovery of Cas9-induced indels or increase the rate of HR.

**DISCUSSION**

Our data demonstrate that the coconversion strategy previously used in C. elegans (Arribere et al. 2014; Kim et al. 2014; Ward 2015) can be successfully applied to Drosophila, reducing the burden of screening for mutations at the gene-of-interest. The coconversion strategy worked equally well for the generation of indels or recombinants: both types of mutations were enriched in the broods that had no red-eyed progeny (Figure 2 and Table 1). The absence of red-eyed G1 flies in a brood indicates that all germline alleles in the G0 animal underwent targeted genome modification at white, reflecting efficient delivery of the guide plasmid to all the pole cells after injection. Our data suggest that when this happens, regardless of the choice of repair pathway, the cotargeted gene-of-interest is more likely to be modified. It is worth noting that Cas9-catalyzed DSBs at white and the gene-of-interest were correlated, but we did not observe a correlation between the repair pathways used at white and at the gene-of-interest: broods with HR at white did not necessarily produce recombinants at the gene-of-interest.

**Table 2 Targeting white in Lig4+ or Lig4169, vas-Cas9 G0 embryos**

<table>
<thead>
<tr>
<th>w sgRNA</th>
<th>G0 Lig4 Genotype</th>
<th>Fertile G0 (n)</th>
<th>Percent of Fertile G0 Whose G1 Offspring Had Eyes That Were:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>All Red</td>
</tr>
<tr>
<td>pCFD4d-1 (26 nM)</td>
<td>Lig4+</td>
<td>23% (255)</td>
<td>48</td>
</tr>
<tr>
<td>pCFD4d-3 (26 nM)</td>
<td>Lig4+</td>
<td>14% (310)</td>
<td>76</td>
</tr>
<tr>
<td>pDCC6-2 (26 nM)</td>
<td>Lig4+</td>
<td>28% (240)</td>
<td>42</td>
</tr>
<tr>
<td>pDCC6-2 (26 nM)</td>
<td>Lig4+</td>
<td>6.3% (240)</td>
<td>73</td>
</tr>
</tbody>
</table>

sgRNA templates were coinjected with 33 nM pUC-white HR donor plasmid DNA. n, the number of G0 embryos injected, irrespective of fertility or survival. The coffee & red/white group includes G0 with coffee- and red-eyed, or with coffee-, white-, and red-eyed G1 broods. pCFD4d also carries armi sgRNA-1, and pDCC6 also carries a Cas9 gene expression unit.
We frequently recovered more than one type of mutation at the gene-of-interest from a single G1 brood, evidence that independent repair events occurred among the dozens of germline stem cells of the G0 founder parent. In other words, the G0 germline is frequently mosaic. As an extreme example, five different indels and three different HR events at zinc were identified in the ten G1 flies we genotyped from a brood consisting of 15% white-eyed and 85% coffee-eyed offspring.

At the seven sgRNA target sites we tested, 39% of the 82 independent indels had junctional microhomologies or templated insertions (Figure 3 and Tables S1–S6 in File S1), signatures of the Lig4-independent, microhomology-dependent end-joining pathway (Yu and McVey 2010; Steir and Symington 2015). We recovered many indels containing such signatures from Lig4+ embryos (Tables S1, S2, S3 and S6 in File S1), suggesting that the microhomology-mediated end-joining pathway normally operates even in the presence of Ligase 4. In fact, Lig4+ mutant embryos produced no fewer indels than Lig4− embryos (Figure 7), suggesting that a Ligase 4-independent end-joining pathway predominates at generating indels. In C. elegans, polymerase 0, but not Lig4, is used to repair Cas9-induced DSBs (van Schendel et al. 2015). As in worms, the Drosophila polymerase 0 (mus308) is important for Lig4-independent end joining (Chan et al. 2010). Future experiments to test whether inactivation of mus308, alone or together with Lig4, reduces indel mutations in flies are clearly needed.

Eliminating donor integration, in which the plasmid integrates into the target locus instead of promoting the desired gene conversion, remains a challenge for Cas9-targeted HR: in our experiments, such integration accounted for 17–67% (median, 50%) of all HR events (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4). Plasmid integration has been reported to account for 70% indel mutations in whether inactivation of mus308 (Figure 4).

Variability in conversion tract length was observed in regions flanking a single DSB or flanking the gap deleted by two concomitant DSBs. Measured from the breaks, some tracts were ~1000 bp, while others were <50 bp (Figure 5); some were even <7 bp (Figure 6). The conversion of the region flanking the DSB(s) is therefore unpredictable. In contrast, when a pair of sgRNAs was used to direct two DSBs, the intervening sequence was reliably replaced with that of the donor (Figure 5 and Figure 6). Pairs of sgRNAs have been used to change or insert 1–3 kbp of novel sequence into a gene in Drosophila, presumably through the same gap repair mechanism (Gratz et al. 2014; Ren et al. 2014a,b; Yu et al. 2014; Zhang et al. 2014b; Port et al. 2015). Using the sister chromatid as a repair template, gap repair readily restores a 9 kbp gap following P element excision (McVey et al. 2004a). Alternatively, the conversion of intervening sequence between two DSBs may result from two convergent HR events initiated from each DSB separately. In this scenario, the two DSBs do not have to be created concomitantly. It is worth noting that gap repair does not always happen when two sgRNAs were coinjected, as we frequently observed gene conversion at one target site and either an indel or wild-type sequence at the other (Figure 5 and Figure 6). One possibility is that one of the two sgRNAs was more active than the other, reducing the chance of generating two DSBs at the same time—a prerequisite of gap repair. Thus, it may be prudent to carry out two experiments each using a unique pair of sgRNAs to ensure successful gap repair, which also offers the opportunity to generate two independent recombinants with nonoverlapping potential off-target mutations.

Previous studies with zinc-finger nucleases suggested that Lig4+ mutant embryos promote HR (Beummer et al. 2008, 2013; Bozas et al. 2009). Surprisingly, the use of Lig4+ embryos did not increase HR efficiency in our experiments (Figure 7), perhaps because Cas9, unlike zinc-finger nucleases, leaves blunt ends (Kim et al. 1996; Jinek et al. 2012).

In conclusion, cotargeting the w gene in Drosophila using Cas9 to alter the fly genome substantially reduces the time and effort required for the molecular identification of mutations in the gene-of-interest. Other organisms with available endogenous or transgenic marker genes should be able to adopt a similar coconversion strategy.

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LITERATURE CITED


