A home and rescue gene drive forces its inheritance stably persisting in populations

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Abstract

Homing based gene drives, engineered using CRISPR/Cas9, have been proposed to spread desirable genes into target populations. However, spread of such drives can be hindered by the accumulation of resistance alleles. To overcome this significant obstacle, we engineer a population modification Home-and-Rescue (HomeR) drive in *Drosophila melanogaster* that, by creative design, limits the accumulation of such alleles. We demonstrate that HomeR can achieve nearly ~100% transmission enabling it to persist at genotypic fixation in several multi-generational population cage experiments, underscoring its long term stability. Finally, we conduct mathematical modeling determining HomeR can outperform contemporary gene drive architectures for population modification over wide ranges of fitness and transmission rates. Given its straightforward design, HomeR could be universally adapted to a wide range of species.

Keywords: Localized, Homing, Gene Drive, Population Modification, Stable, Haplosifficient, Resistance.

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Introduction

Effective insect control strategies are necessary for preventing human diseases, such as malaria and dengue virus, and protecting crops from pests. These challenges have fostered the development of innovative population control technologies such as Cas9/guideRNA (Cas9/gRNA) homing-based gene drives (HGDs) (Champer et al., 2016; Esvelt et al., 2014) which have been tested in the laboratory for either population modification (Adolfi et al., 2020; Gantz et al., 2015; Li et al., 2020; Pham et al., 2019) to spread desirable traits that can impair the mosquitoes ability to transmit pathogens (Buchman et al., 2020, 2019; Hoermann et al., 2020; Isaacs et al., 2012; Marshall et al., 2019), or population suppression (Hammond et al., 2016; Kyrou et al., 2018; Simoni et al., 2020) to reduce and eliminate wild disease transmitting populations of mosquitoes. Despite significant progress, HGDs are still an emerging technology that can suffer from the rapid formation of resistance alleles, hindering their efficacy.

In CRISPR/Cas9, the Cas9 endonuclease cuts a programmed DNA sequence complementary to a user defined short guide RNA molecule (gRNA). To engineer a HGD, CRISPR components are integrated at the cut site in the genome so that when they cut the recipient wildtype (wt) allele it is repaired via homology-directed repair (HDR) in heterozygotes, using the donor allele (i.e. allele harboring the HGD) as a reference for DNA repair. This enables the HGD to home, or copy, itself into the recipient allele (Champer et al., 2016; Esvelt et al., 2014; Kandul et al., 2019a) (referred to as homing from hereon). This general design for HGD was quickly adopted, and many HGDs were developed in several insect species (Esvelt et al., 2014; Gantz et al., 2015; Hammond et al., 2016; Kandul et al., 2019a; Kyrou et al., 2018; Li et al., 2020; Simoni et al., 2020). However, it soon became apparent that HGDs unintentionally promote the formation of resistance alleles through mutagenic repair. When these alleles are positively selected they can hinder HGD spread in laboratory cage populations (Champer et al., 2017; Hammond et al., 2017; Kandul et al., 2019a; KaramiNejadRanjbar et al., 2018; Oberhofer et al., 2018), with one exception that targeted an ultra-conserved sex determination gene for population suppression (Kyrou et al., 2018; Simoni et al., 2020). This resistance arises from, in addition to HDR, Cas9/gRNA-directed DNA cuts are also repaired by non-homologous end joining (NHEJ), an alternative DNA repair pathway that occasionally introduces insertions or deletions (indels) at the target site. Many of these indels produce loss-of-function (LOF) alleles, which can be selected against. However, functional inframe NHEJ-induced indel alleles that survive and propagate, that are unrecognized by the same Cas9/gRNA complex, become drive resistance alleles. When resistance alleles are induced in germ cells, they are heritable and can hinder spread of HGDs (Champer et al., 2017; Hammond et al., 2017; Kandul et al., 2019a; KaramiNejadRanjbar et al., 2018; Oberhofer et al., 2018). Both induced and naturally existing resistance alleles can pose serious challenges to engineering a stable HGD capable of spreading and persisting in a population.

To overcome the accumulation of drive resistance alleles, toxin-antidote (TA) based systems, in which embryos are essentially "poisoned" and only those embryos harboring the TA genetic cassette are rescued, were described (Fig. S8 in (Kandul et al., 2019b)) and engineered (Champer et al., 2020a; Oberhofer et al., 2020a, 2020b, 2019). Generally these designs utilize a toxin consisting of a non-homing GD harboring multiple gRNAs targeting a vital gene, and an "addictive" antidote that is a re-coded, cleavage-immune version of the targeted gene. These TA based drives are Mendelianly transmitted and spread by killing progeny that fail to inherit the drive (e.g. 50% perish from heterozygous mother). Alternative HDR-based designs were also described (Champer et al., 2016; Esvelt et al., 2014), modeled (Noble et al., 2017), and recently tested in *Anopheles stephensi (Adolfi et al., 2020)* targeting a non-essential gene for viability (Adolfi et al., 2020; Gantz et al., 2015; Pham et al., 2019), and in *Drosophila melanogaster* targeting a

haploinsuffienct (i.e. a single functional copy of the target gene is <u>not sufficient</u> for normal function) gene (Champer et al., 2020b), demonstrating drive capacity.

Building upon prior work, here we describe the development of a Home-and-Rescue (HomeR) split-drive (i.e. Cas9 separated from the drive) targeting an 100% essential, haplosufficient (i.e. a single functional copy of the target gene is sufficient for normal function) gene in *Drosophila* melanogaster. We demonstrate that the accumulation of NHEJ-induced resistance alleles can be reduced by strategically (i) designing the HGD to target the conserved 3' coding sequence of a haplosufficient gene required for insect viability using a single gRNA, and (ii) encoding the rescue of its wt function into HomeR, and (iii) using an exogenous 3' UTR to prevent expected deleterious recombination events between the drive and the endogenous target gene. We demonstrate that efficient cleavage of the target sequence by HomeR and rescue of the gene's function are requisites to achieve nearly ~100% transmission, which is accomplished by homing in ~90% of wt alleles and by destroying the remaining ~10% from trans-heterozygous females. Further, we perform multi-generational population cage experiments demonstrating long term stability. Finally, we conduct comprehensive mathematical modeling to demonstrate that HomeR can outperform contemporary gene drive systems for population modification over wide ranges of fitness and transmission rates and, given the simplistic design, this system could be adapted to other species in the future.

Results

Design and testing of gRNAs targeting an essential gene.

To develop a HomeR-based drive, we first identified an essential haplosufficient gene to target. We chose Pol-Y35, required for the replication and repair of mitochondrial DNA (mtDNA) (Carrodeguas, 2000; Carrodeguas et al., 2001) and whose LOF results in lethality (lyengar et al., 2002) (Fig. 1A). Given that separate gRNAs can result in varying degrees of cleavage efficiencies (Kandul et al., 2019b), we tested two qRNAs targeting a conserved C-terminal domain of Pol-Y35 $(gRNA\#1^{Pol-\gamma35})$ and $gRNA\#2^{Pol-\gamma35})$. Both $gRNA\#1^{Pol-\gamma35}$ and $gRNA\#2^{Pol-\gamma35}$ were inserted site specifically in the *Drosophila* genome and were expressed using the *U6.3* promoter (Port et al., 2014). To genetically assess the efficiency of Cas9/gRNA-mediated cleavage induced by each gRNA, we separately crossed these established gRNA lines to two different Cas9 expressing lines: (i) a previously characterized ubiquitously expressing Cas9 line (Port et al., 2014) in the DNA Ligase 4 null genetic background (Act5C-Cas9; Lig4-/-) (Xu Zhang et al., 2014) and also to (ii) a germline-enriched Cas9 driven by the nanos promoter (nos-Cas9) (Kandul et al., 2019a, 2019b). As mutations in Lig4 gene can result in decreased activity of DNA repair by the NHEJ pathway (McVey et al., 2004) we reasoned that by testing the gRNA efficiency in a Lig4-/background, we may increase the penetrance of lethality phenotypes since we are targeting an essential gene that cannot be repaired efficiently by NHEJ which simplifies scoring of the gRNAs for efficacy. Given that the Lig4 gene is located on the X chromosome, maternal Lig4- alleles will be inherited by all male progeny, making them hemizygous Lig4- mutants, while females will be heterozygous *Lig4*–/+.

We observed that the genetic cross between either U6.3- $gRNA#1^{Pol-arrow35}$ or U6.3- $gRNA#2^{Pol-arrow35}$ homozygous males to Act5C-Cas9, Lig4—/— homozygous females was lethal for all male progeny (**Fig. 1B,C**). Interestingly, trans-heterozygous Act5C-Cas9, Lig4—/+; U6.3-

we also found that the Cas9 protein deposited by *nos-Cas9/+* females without inheritance of the *nos-Cas9* transgene, aka maternal carryover (Kandul et al., 2019b; Lin and Potter, 2016), was sufficient to ensure lethality of the F₁ progeny harboring *U6.3-gRNA#1*^{Pol-}\$\footnote{3}\$ (**Fig. 1C**), while *U6.3-gRNA#2*^{Pol-}\$\footnote{3}\$ induced lethality only in a fraction of the F₁ *U6.3-gRNA#2*^{Pol-}\$\footnote{3}\$ (**Fig. 1C**). Pooled embryo sanger sequencing of trans-heterozygotes revealed expected mutations at the *Pol-*\$\footnote{3}\$ gRNA target sites. Taken together, these results indicate that both tested gRNAs induced cleavage of the *Pol-*\$\footnote{3}\$ target sequences, though *U6.3-gRNA#1*^{Pol-}\$\footnote{3}\$ induced greater cleavage than *U6.3-gRNA#2*^{Pol-}\$\footnote{3}\$ and resulted in complete lethality of females and males with either *Act5C-Cas9* or *nos-Cas9*. Note that unlike previously described LOF mutations of *Pol-*\$\footnote{3}\$ (*Iyengar et al., 2002*), Cas9/gRNA-induced cleavage of the *Pol-*\$\footnote{3}\$ C-terminal domain resulted in embryonic lethality. In sum, these results indicate that both gRNAs are functional and could be used to generate gene drives.

Development of split HomeR drives with encoded rescue

Using these characterized gRNAs, we engineered two $Pol-\gamma35$ HomeR ($HomeR1^{Pol-\gamma35}$ and $HomeR2^{Pol-\gamma35}$) drives. Fitting with the split-GD design, neither $HomeR1^{Pol-\gamma35}$ nor $HomeR2^{Pol-\gamma35}$ include the Cas9 gene, and thus are non-autonomous gene drives. To mediate HDR, left and right homology arm sequences (LHA and RHA) matching the sequences surrounding the cut site of the corresponding gRNA were used. Additionally, we included a 3xP3-eGFP-SV40 marker gene to track the presence of the drive and a re-coded C-terminal domain incorporated into the LHA with a p10 3'UTR to support robust expression of the re-coded $Pol-\gamma35$ (Fig. 2A) to eliminate homology and prevent gene conversion between the rescue allele and the endogenous allele, which proved problematic in previous drive designs (Champer et al., 2020a, 2020b). Importantly, the recoding was designed to ensure the translation of the re-coded DNA sequence in the wildtype (wt) amino acid sequence of $Pol-\gamma35$ with respect to Drosophila codon usage bias.

In case the initial HDR-mediated transgenesis of $HomeR^{Pol-\gamma35}$ at the $Pol-\gamma35$ cut site failed, both $HomeR^{Pol-\gamma35}$ constructs were assembled in a piggyBac plasmid that supported an alternative path for genome integration (**Fig. 2B**). To distinguish between the site-specific HDR-mediated integration, tagged by the eye-specific GFP fluorescence of 3xP3-eGFP-SV40, and non-site-specific integration using piggyBac-mediated insertion, the Opie2-dsRed-SV40 marker was included outside the LHA and RHA of both $HomeR^{Pol-\gamma35}$, conferring body-specific dsRed fluorescence (**Fig. 2B**). Overall, this split-GD design (Champer et al., 2020a; Kandul et al., 2019a; Li et al., 2020) restricts the spread of $Pol-\gamma35^{HomeR}$ and serves as a safeguard against unintended spread, as the homing $Pol-\gamma35^{HomeR}$ harboring a gRNA and the non-homing Cas9 are genetically unlinked, resulting in molecular confinement (Esvelt et al., 2014; Marshall and Akbari, 2018).

To generate transgenic lines, we first injected mixtures of each $HomeR^{Pol-\gamma35}$ plasmid and a helper plasmid, expressing the pBac transposase that directs genomic integration (Handler and Harrell, 1999), into w^{1118} embryos. This established transgenic lines carrying random insertions of the $HomeR1^{Pol-\gamma35}$ and $HomeR2^{Pol-\gamma35}$ plasmids, genetically tracked by both eGFP and dsRed markers. We then used Homology Assisted CRISPR Knock-in (HACK)(Gantz and Akbari, 2018; Lin and Potter, 2016) to integrate $HomeR1^{Pol-\gamma35}$ or $HomeR2^{Pol-\gamma35}$ at the corresponding $gRNA^{Pol-\gamma35}$ cut site by crossing flies harboring random insertions of $HomeR1^{Pol-\gamma35}$ or $HomeR2^{Pol-\gamma35}$ to $HomeR2^{Pol-\gamma35}$ to $HomeR2^{Pol-\gamma35}$ to $HomeR2^{Pol-\gamma35}$ to $HomeR2^{Pol-\gamma35}$ to $HomeR2^{Pol-\gamma35}$ or $HomeR2^{Pol-\gamma35}$ to $HomeR2^{Pol-\gamma35}$ or $HomeR2^{Pol-\gamma35}$ or $HomeR2^{Pol-\gamma35}$ or $HomeR2^{Pol-\gamma35}$ plasmid directly into $HomeR2^{Pol-\gamma35}$ plasmid directly into H

For both insertion approaches described above (**Fig. 2B**), the F₁ trans-heterozygous flies harboring potential *Pol-\gamma35^{HomeR1}* or *Pol-\gamma35^{HomeR2}* and tagged by double fluorescence (GFP+ and dsRed+) were individually crossed to sna^{Sco}/CyO balancer flies to isolate $Pol-\gamma35^{HomeR1}/CyO$ or $Pol-\gamma35^{HomeR2}/CyO$ flies marked with only GFP fluorescence. Multiple independent transgenic lines of each $Pol-\gamma35^{HomeR1}$ and $Pol-\gamma35^{HomeR2}$ were isolated and balanced on the second chromosome. To confirm that $Pol-\gamma35^{HomeR1}/CyO$ or $Pol-\gamma35^{HomeR2}/CyO$ lines were indeed inserted at the corresponding cut site in $Pol-\gamma35^{HomeR1}/CyO$ or $Pol-\gamma35^{HomeR2}/CyO$ lines were indeed inserted in the presence of Cas9 in $Scale{Scale}$ in $Scale{Scale}$ and $Scale{Scale}$ in $Scale{Scale}$ in $Scale{Scale}$ and $Scale{Scale}$ in $Scale{Scale}$ and $Scale{Scale}$ in $Scale{Scale}$ and $Scale{Scale}$ in $Scale{Scale$

Assessment of germline transmission and cleavage rates

To assess the effects of gRNA-mediated cleavage efficiency on transmission rates, we compared the two nearly identical HomeRs, as they harbored two distinct gRNA sequences that differed in cleavage efficiencies. The Pol-Y35^{HomeR1} and Pol-Y35^{HomeR2} homozygous lines have U6.3gRNA#1^{Pol-}Y35 and U6.3-gRNA#2^{Pol-}Y35, respectively, and slightly different LHA and RHA corresponding to the appropriate DNA cut sites, which are only 13 bases apart (Fig. 1A). We found that Pol-\(\chi_35^{HomeR1}/+ \); nos-Cas9/+ trans-heterozygous females crossed to wt males transmitted $Pol-\sqrt{35^{HomeR1}}$ to 99.5 ± 0.6% of progeny, while $Pol-\sqrt{35^{HomeR2}}/+$; nos-Cas9/+ females transmitted the corresponding *Pol-\35^{HomeR2}* to a significantly lower fraction of F₁ progeny (68.7 ± 6.2%, two-sided Student's *t*-test with equal variances, p < 0.0001; Fig. 3A, Supplementary Data 3). Genetic crosses of either Pol-Y35^{HomeR1}/+; nos-Cas9/+ or Pol-Y35^{HomeR2}/+; nos-Cas9/+ transheterozygous males to wt females did not result in biased transmission to F₁ progeny (60.7 ± 5.3% vs 52.9 \pm 4.0% or 54.3 \pm 4.0% vs 51.5 \pm 1.8%, respectively, two-sided Student *t*-test with equal variances, p > 0.05; Fig. 3A). Maternal carryover of Cas9 protein by nos-Cas9/+ females significantly increased transmission of Pol-Y35^{HomeR1} by F₁ Pol-Y35^{HomeR1}/CyO females, 66.1 ± 0.8% vs 52.9 \pm 4.0% (two-sided Student *t*-test with equal variances, p < 0.001; Fig. 3A, Supplementary Data 3). These results suggest that the higher cleavage efficiently of U6.3 $gRNA#1^{Pol-\gamma35}$, as measured by the induced lethality in the Lig4 null genetic background (**Fig. 1B**), likely contributes to the higher homing rate of Pol-V35HomeR1 harboring U6.3-gRNA#1Pol-V35, and underscores the importance of selecting an efficient gRNA for gene drives (GDs).

Majority of *Pol-*\$\cong 35^{WT}\$ alleles are converted into *Pol-*\$\cong 35^{HomeR1}\$ alleles in trans-heterozygous females

Either homing (indicating allelic conversion) in oocytes or "destruction" of the *wt* alleles in embryos of trans-heterozygous *Pol-\gamma35^{HomeR1}/+*; *Cas9/+* females via a process termed "biallelic lethal mosaicism (BLM)" (Kandul et al., 2019b) could contribute to biased *Pol-\gamma35^{HomeR1}* transmission rates. BLM contributes to RNA-guided dominant biallelic knockouts of the target gene throughout development thereby converting recessive non-functional resistant alleles into dominant deleterious/lethal mutations that can get selected out of a population. Previously, destruction of the *wt* allele in conjunction with maternal carryover of a "toxin" was used to engineer gene drives based on an "addictive" TA approach (Champer et al., 2020a; Oberhofer et al., 2020a, 2019). In these TA drives, one half of the F₁ progeny did not inherit the TA cassette, meaning not rescued, and were killed—resulting in a rapid spread of the genetic cassette in laboratory populations.

In our experiments, the U6.3-gRNA#1^{Pol-}Y35 induced embryonic lethality in the presence of nos-Cas9 or maternal carryover of the Cas9 protein (Fig. 1C). Therefore, to explore the mechanism resulting in the super-Mendelian transmission of Pol-Y35HomeR1, and given that disruption of Pol-V35 by 6.3-qRNA#1^{Pol-Y35} in Pol-Y35^{HomeR1} results in fully penetrant embryonic lethality, we determined the egg hatching rate, as the percentage of embryonic lethality, for transheterozygous females and compared it to those of females heterozygous for Pol-Y35HomeR1 or Cas9 (**Fig. 3B**). The hatching rate of F₁ eggs generated by *Pol-Y35^{HomeR1}/+*: nos-Cas9/+ transheterozygous females crossed to wt males was reduced by 5% as compared to Pol-\35^{HomeR1}/+; +/+ heterozygous females crossed to wt males (88.8 ± 2.4% vs 94.4 ± 1.2%; two-sided Student t-test with equal variances, p < 0.004). Furthermore, the hatching rate of eggs laid by transheterozygous females was not statistically different from that laid by +/+; nos-Cas9/+ heterozygous females crossed to wt males (88.8 ± 2.4% vs 93.4 ± 3.7%; two-sided Student's ttest with equal variances, p < 0.052, Fig. 3B, Supplementary Data 4). Moreover, there was no significant difference between the larvae-adult survival rates comparing Pol-\(\chi 35^{HomeR1} / +; \ nos-Cas9/+ to Pol-\gamma35^HomeR1/+; +/+ indicating that there is no bias at these later stages. Taken together, these data indicate that from the expected 50% of Pol-Y35WT alleles transmitted by transheterozygous females, ~5% were "destroyed" via BLM—meaning mutated and not complemented by the wt allele delivered with a sperm, since it was also mutated by Cas9/gRNA maternal carryover—and the remaining ~45% were converted into *Pol-\/35*^{HomeR1}—resulting in an estimated conversion rate of ~90% (45% X 2). Therefore, these data indicate that the observed transmission rate of nearly ~100% was caused by ~90% conversion and ~10% "destruction" of the Pol-Y35WT alleles. In sum, the Pol-Y35^{HomeR1} transmission rate of nearly ~100% observed in Pol-Y35^{HomeR1}/+; nos-Cas9/+ trans-heterozygous females could not be simply explained by the "destruction" of all wt Pol-Y35 alleles, which would result in the lethality of 50% progeny as in non-homing ClvR (Oberhofer et al., 2020a, 2020b, 2019) and TARE (Champer et al., 2020a) drives, and instead is a result of both conversion and destruction of the recipient allele at the *Pol-Y35* locus (**Fig. 2A**).

Nos- and ubiq-Cas9 support the strongest female-specific transmission of Pol-¥35^{HomeR1}

The split-drive design facilitates testing of different Cas9 promoters. Therefore, we were able to estimate the transmission of Pol-Y35^{HomeR1} by trans-heterozygous females and males harboring Pol-Y35^{HomeR1} in combination with four alternative Cas9 promoters active in germ cells of both sexes. Nanos (nos) and vasa (vas) promoters were previously described as germline-specific promoters active in both sexes (Hay et al., 1988; Sano et al., 2002; Van Doren et al., 1998), though recent evidence indicates ectopic expression in somatic tissues from both nos-Cas9 and vas-Cas9 (Kandul et al., 2019a, 2019b). The Ubiquitin 63E (Ubiq) and Actic 5C (Act5C) promoters in ubig-Cas9 (Kandul et al., 2019b) and Act5C-Cas9 (Port et al., 2014) transgenic lines, respectively, support strong expression in both somatic and germ cells (Kandul et al., 2019a, 2019b; Port et al., 2014; Preston et al., 2006). To control for genome insertion effects, each Cas9 transgene was inserted at the same attP docking site on the 3rd chromosome, except for Act5C-Cas9 that was integrated on the X chromosome (Port et al., 2014). Since maternal carryover of the Cas9 protein was shown to induce a "shadow gene drive" two generations later (Guichard et al., 2019; Kandul et al., 2019a), we used trans-heterozygous flies that inherited paternal Cas9 to quantify the transmission of Pol-Y35^{HomeR1}. Trans-heterozygous females carrying Pol-Y35^{HomeR1} together with nos-Cas9, vas-Cas9, ubiq-Cas9, or Act5C-Cas9 crossed to wt males biased transmission of Pol- $\sqrt{35}^{HomeR1}$ to nearly ~100% of F₁ progeny (99.5 ± 0.6%, 97.6 ± 2.6%, 99.6 ± 0.6%, and 99.0 \pm 0.4%, respectively, vs 52.9 \pm 4.0% by Pol-Y35^{HomeR1}/Pol-Y35^{WT}; +/+ females, two-sided Student's *t*-test with equal variances, p < 0.001; **Fig. 3C**). Note that the corresponding trans-heterozygous males only modestly biased Pol-\gamma35^{HomeR1} transmission from 55.3 ± 5.0% of F_1 progeny to $60.7 \pm 5.3\%$ (p > 0.05), $63.2 \pm 6.6\%$ (p < 0.03), $66.1 \pm 4.6\%$ (p < 0.004), and 62.0

 \pm 1.7% (p < 0.017, two-sided Student's t-test with equal variances, **Fig. 3C, Supplementary Data 3**), respectively.

ExuL-Cas9 supports the strongest male-specific transmission of Pol-Y35HomeR1

To assess whether males could support robust homing similar to females, we investigated three alternative male-specific promoters. We established the *Drosophila exuperantia* (CG8994) large fragment (exuL) promoter for an early male-specific expression. The Rcd-1 related (Rcd1r, CG9573)(Chan et al., 2013) and \$\beta Tubulin 85D (\beta Tub)(Chan et al., 2011; Michiels et al., 1989) promoters support an early and late, respectively, testis-specific expression in *Drosophila* males. We found that only exuL-Cas9 induced the male-specific super-Mendelian inheritance of Pol-Y35^{HomeR1}: trans-heterozygous males, but not females, transmitted *Pol-Y*35^{HomeR1} to more than 50% of F₁ progeny (75.0 \pm 6.1% vs 55.3 \pm 5.0% in $\sqrt[3]{p}$ < 0.0001; and 50.7 \pm 3.4 % vs 52.8 \pm 4.0% in \mathcal{L} , p > 0.05, two-sided Student's *t*-test with equal variances; Fig. 3C, Supplementary Data 3). To our surprise, Rcd1r-Cas9 induced super-Mendelian inheritance of Pol-Y35HomeR1 in both trans-heterozygous males and females (68.2 \pm 3.8% vs 55.3 \pm 5.0% in β , p < 0.002; and 90.8 \pm 0.5% vs 52.8 \pm 4.0% in Ω . p > 0.0001, two-sided Student's t-test with equal variances: Fig. 3B). Finally, βTub-Cas9 did not induce changes in transmission of Pol-¥35^{HomeR1} by either trans-heterozygous males or females (55.6 \pm 5.7% vs 55.3 \pm 5.0% in $\sqrt[3]{p}$ = 0.55; and 51.5 \pm 2.1% vs 52.9 \pm 4.0% in Ω , p = 0.94, two-sided Student's t-test with equal variances; Fig. 3B, Supplementary Data 3). These results suggest that Drosophila males bias Pol-Y35^{HomeR1} transmission, however this bias is substantially lower than the nearly ~100% transmission of Pol- $\chi 35^{HomeR1}$ in females.

Functional *Pol-* $\upgamma35$ resistance alleles (*Pol-* $\upgamma35^{R1}$) did not hinder drive persistence in 10 generations

Only one Cas9-mediated HGD, targeting an ultra-conserved female-specific dsx splice variant in Anopheles gambiae, which is dispensable for male viability and fertility, has successfully avoided generation of resistance alleles in small laboratory populations (Kyrou et al., 2018; Simoni et al., 2020). We reasoned that insertion of *HomeR* into the gene required for viability could also prevent the accumulation of Pol-Y35 LOF resistance alleles (R2 type, Pol-Y35^{R2}) by exploiting BLM. However, functional resistance alleles (R1 type, Pol-\(\chi_3\)5^{R1}), either from in-frame indels or synonymous base substitutions (SBS), could still be induced by Cas9/gRNA#1^{Pol-}Y³⁵. This could be problematic if *Pol-Y35^{R1}* resistance alleles do not impose fitness costs on homozygous carriers. as they would be expected to spread at the expense of the drive. To explore this potential, we set up three laboratory populations of heterozygous Pol-Y35^{HomeR1}/+ flies in the nos-Cas9/nos-Cas9 genetic background and assessed the emergence and spread of induced resistance alleles over ten discrete generations (Fig. 4A). Although we could not distinguish homozygous flies from heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes heterozygotes with respect to the dominant marker of Pol-\(\frac{7}{35}\) heterozygotes hetero alleles were expected to spread at the expense of Pol-Y35HomeR1 alleles, which would be straightforward to score in our assay by loss of the GFP marker and would be predicted to become homozygous over multiple generations.

Out of 10 generations with three distinct populations, we found only nine flies lacking the $Pol-\gamma 35^{HomeR1}$ allele (i.e. GFP negative) at generations 2 and 3 in two out of three population lineages (drives #1 and #3, **Fig. 4A**; **Supplementary Data 5**). To rescue their viability in the absence of $Pol-\gamma 35^{HomeR1}$, these flies had to harbor either $Pol-\gamma 35^{WT}$ or $Pol-\gamma 35^{R1}$ and were analyzed to determine their genotype. To ensure that any generated $Pol-\gamma 35^{R1}$ alleles had a chance to spread and compete with $Pol-\gamma 35^{HomeR1}$ alleles, these flies were transferred among the subsequent generation and allowed to mate with other flies and lay eggs in each population lineage before

they were genetically analyzed. The analysis revealed that each fly harbored at least one $Pol-\gamma 35^{R1}$ resistance allele that rescued viability. Two different $Pol-\gamma 35^{R1}$ alleles were identified by sequencing multiple clones of amplicons from each of the nine genotyped flies. One $Pol-\gamma 35^{R1}$ allele was sampled in eight flies from two independent lineages. It had a three-base-insertion that inserted one amino acid as well as a change in one amino acid, p.T358_L360insAG (R1.1, **Fig. 4B**). The other $Pol-\gamma 35^{R1}$ allele was found in two flies, and its three-base-substitution caused one amino acid change, p.T358S (R1.2, **Fig. 4B**). Seven out of nine genotyped flies were heterozygous, harboring both functional and LOF alleles at the $Pol-\gamma 35^{R1}$ locus, one fly had two different $Pol-\gamma 35^{R1}$ alleles, and one fly could be homozygous for the R1.1 allele, since ten sequenced clones gave the same R1.1 allele (**Fig. 4B**). We did not identify any fly without the $Pol-\gamma 35^{HomeR1}$ allele after generation 3, as indicated by the eye-specific GFP expression (**Fig. 4A**, **Supplementary Data 5**), thus the identified $Pol-\gamma 35^{R1}$ alleles did not spread and we were unable to establish these as isolated strains, indicating that flies harboring these alleles were likely less competitive than those with one copy of the $Pol-\gamma 35^{HomeR1}$ allele as would be expected when targeting a haplosufficient gene.

To further explore the diversity of resistance alleles remaining after ten generations, starting with 100% heterozygous flies at the 1st generation, we performed next-generation sequencing on sixty randomly chosen flies (each fly had at least one copy of Pol-Y35 FlomeR1 since the drives were at genotypic fixation) from each drive to identify and quantify any "short" Pol-Y35 alleles, which did not harbor the large insert of HomeR1^{Pol-Y35} (~2.5 KB, Fig. 2A). Neither the Pol-Y35^{WT} nor previously identified Pol-Y35^{R1} alleles were sampled among nearly 150,000 sequence reads for the three drive experiments. Instead, we found two novel in-frame indel alleles, 18 bp and 9 bp deletions, in drives #2 and #3 (Fig. 4C). The 18 bp deletion was responsible for 80% of the resistance alleles sampled for drive #2, while the 9 bp deletion was the least abundant (5%, Fig. 4C) resistance allele in drive #3. Since both in-frame indel alleles cause the deletion of either 6 or 3 amino acids in the middle of the highly conserved c-terminal domain of Pol-Y35, they are likely deleterious recessives. The remaining eleven alleles were out-of-frame indels, ranging from a 1 bp insertion to a 23 bp deletion (Fig. 4C). Two LOF alleles, 2 bp and 4 bp deletions, were also seen in the genotyped flies at generations 2 and 3. The relative abundance of each allele can be used to extrapolate the minimum number of resistance alleles sampled in the sixty heterozygous and/or homozygous flies and thus persisted though ten generations of drive. We inferred that at least 9, 5, and 17 resistance alleles persisted for ten generations and were rescued by the Pol-V35^{HomeR1} allele in 60 sampled flies from drives #1, #2, and #3, respectively (**Fig. 4C**). The fact that these induced functional resistance alleles did not take over at the expense of HomeR1 in 10 generations (i.e. the drives were all at genotypic fixation) indicates that these alleles could not compete with the *Pol-Y35*^{HomeR1} re-coded rescue and spread at its expense.

Modeling indicates that *HomeR* is an efficacious gene drive

To compare the performance of HomeR against contemporary gene drive systems for population replacement, we modeled one- (aka. autonomous) and two-locus (aka. split-drive) versions of ClvR (Oberhofer et al., 2020a, 2020b, 2019), the one-locus TARE system from (Champer et al., 2020a) as well as a two-locus TARE configuration based on their design, a HGD targeting a non-essential gene (Gantz et al., 2015; Hammond et al., 2016), and HomeR. In each case, we first simulated population spread of each gene drive system for an ideal parameterization (see Methods for more details) and included additional simulations for HomeR under current experimentally-derived parameters (HomeR-exp, **Fig. 5A and 5C**). To gauge behavior across a range of scenarios, we performed simulations for a range of fitness costs (implemented as female fecundity reduction) and drive system transmission rates (implemented by varying the cleavage rate), providing heatmaps of the expected performance for each drive system at each parameter

combination (**Fig. 5B and 5D**). Drive efficacy, the outcome in these comparisons, is defined as the expected fraction of individuals that carry the drive (and a linked effector) allele, in either heterozygous or homozygous form, 20 generations following a 25% release of female and male heterozygotes for each drive system.

When one-locus GD systems are compared for ideal parameter values, HomeR outperforms all other GDs in terms of speed of spread, and reaches near fixation in terms of carrier frequency. as does ClvR and TARE (Fig. 5A). HGD displays a similar speed of spread to HomeR initially; however, fitness costs from the targeted gene knockout and loss-of-function (R2) alleles build up over time and progressively reduce its speed of spread and efficacy. The HomeR design overcomes this fitness reduction and R2 allele buildup by rescuing the wt function of a targeted essential gene (Fig. 6). ClvR and TARE perform similarly to each other for ideal parameter values. but reach near carrier fixation ~8-9 generations after HomeR does for ideal parameter values (Fig. 5A). When experimental parameters are used for HomeR (HomeR-Exp, in Fig. 5A), it reaches near carrier fixation ~3 generations later than for ideal parameter values; but still spreads faster than the ClvR and TARE systems with ideal parameters. HomeR also reaches near carrier fixation for the widest range of fitness and transmission rate parameter values (Fig. 5B). outperforming all alternative drives by this criterion. As a HDR, HomeR drives to high carrier frequencies provided its inheritance bias (or transmission rate) exceeds its associated fitness cost. Drive efficacy of ClvR and TARE, on the other hand, is strongly dependent on fitness cost and weakly dependent on transmission rate. Indeed, ClvR and TARE can each only tolerate fitness costs less than ~20% (Fig. 5B). This is a consequence of their design, employing a toxinantidote scheme, which induces a significant fecundity reduction (Fig. 6A-B) in addition to other fitness costs. A one-locus HGD also exhibits efficacy across a wide range of parameter combinations, but its efficacy is reduced compared to HomeR due to the build-up of R2 and R1 alleles (Fig. 6C-D), which can potentially block spread of the HGD in large populations (Fig. 5B).

In two-locus simulations, Cas9 is separated from the gRNAs in all designs and undergoes independent assortment during gametogenesis. The effects of this design change are evident (Fig. 5C). Under the same experimental conditions as one-locus simulations, there is significantly more variation in behavior of two-locus GDs, with a reduced speed of introgression into the population and slightly reduced overall efficacy. Nevertheless, HomeR still demonstrates strong performance, spreading significantly faster than ClvR and TARE. TARE performs significantly worse in a split configuration (Champer et al., 2020a). ClvR, when completely unlinked, also performs significantly worse, in-agreement with results from (Oberhofer et al., 2020a). Exploring the performance under a range of parameters, we found reduced efficacy for all drives (Fig. 5D); however, trends of between-drive performance are maintained in this new configuration. TARE and ClvR drive efficacy is still nearly independent of transmission rate, but sharply dependent on fitness costs, rarely reaching carrier fixation for the explored parameter values. HGD still exhibits efficacy across a wide range of parameter combinations, but efficacy is limited in all of them. HomeR still demonstrates higher efficacy than other drives, but for a smaller range of parameter combinations.

Discussion

We have engineered a system we term HomeR, for population modification that mitigates existing issues related to drive resistance. To limit the potential for inducing functional resistance alleles, an ultraconserved, haplosufficient gene required for insect viability was targeted. Multigenerational population cage experiments indicate that Pol-\gamma35^HomeR can persist efficiently in the presence of Cas9, and this persistence is not impacted by induced resistance alleles,

including functional resistance alleles, overcoming a major challenge for population modification HGDs.

The re-coded rescue strategy that we used to develop HomeR was also used in previous Drosophila toxin-antidote non-homing GDs (Champer et al., 2020a; Oberhofer et al., 2020a, 2020b, 2019) and a recent HGD's in both Drosophila (Champer et al., 2020b), and Anopheles stephensi (Adolfi et al., 2020), though each of these examples suffered from potential drawbacks. For example, both the haplolethal HGD (Champer et al., 2020b) and the TARE design (Champer et al., 2020a) share similar problematic design architectures that can be unstable as they are susceptible to functional resistance alleles induced via recombination between the promoter including sequences 5' of a coding sequence and 3'UTR regions, which are identical between the re-coded sequence and the wt sequence (Fig. S2C in (Champer et al., 2020a) and Fig. 2 (Champer et al., 2020b)). Moreover, the haplolethal HGD (Champer et al., 2020b) requires a strict germline specific promoter that lacks maternal carryover otherwise lethal mosaicism, either monoor bi-allelic, will result in dominant negative fitness costs to its carrier and impede drive spread. In fact, our efforts to find such promoters in Drosophila proved exceedingly difficult - with previously tested "germline specific" promoters such as nanos and vasa showing significant somatic activity at multiple insertion sites (Kandul et al., 2019a, 2019b). The recent HGD in Anopheles stephensi (Adolfi et al., 2020) was designed to target and rescue a non-essential gene for viability (i.e. the eye pigmentation kynurenine hydroxylase (kh) gene), whose knockout was pleiotropic and only partially costly to female fecundity and survival (Adolfi et al., 2020; Gantz et al., 2015; Pham et al., 2019). Notwithstanding, this drive spread efficiently in small, multigenerational laboratory population cages under several release thresholds, however, many drives did not reach nor maintain complete fixation presumably due to the viability and partial fertility of drive generated homozygous LOF resistance alleles, underscoring the critical importance of targeting a haplosufficient essential gene for such drives especially for larger releases. Comparatively, the creative design of the ClvR system is guite stable, however it can be cumbersome to engineer requiring re-coding of the essential rescue gene, including all target sequences within the coding sequence (lacking introns), and uses an exogenous promoter and 3'UTR, necessitating precise titration of expression from a distal genomic location with exogenous sequences to guarantee rescue without imposing deleterious fitness costs, a feat that may be difficult to accomplish for essential genes requiring complex regulatory elements and networks not directly adjacent to the target gene. In contrast to the aforementioned drives, (i) HomeR relies on the endogenous promoter sequence of the target gene to facilitate rescue expression which significantly simplifies the design and ensures endogenous expression of the rescue, (ii) creatively designed to target the 3' end of the essential haplosufficient gene to limit the degree of recording required for the rescue, (iii) an exogenous 3'UTR to prevent recombination, and (iv) exploits BLM (Kandul et al., 2019b) by targeting an essential haplosufficient gene to convert recessive non-functional resistant alleles into dominant deleterious/lethal mutations that can get selected out of a population, four important features that should be incorporated into future population modification drives.

Results of three independent multi-generational population cage experiments initiating with 100% heterozygous populations (I.e. initial drive allele frequency of 50%; Cas9 allele frequency of 100%) indicate that NHEJ-induced loss-of-function (R2) alleles can persist for many generations. As expected, a single copy of the HomeR inserted at a haplosufficient gene provides sufficient rescue and complements the corresponding R2 allele. However, it was unexpected that up to 28% of sampled flies potentially harbor R2 *indels* that can persist for ten generations without being selected out (**Fig. 4C**). This high frequency of R2 alleles, for a HGD with nearly ~100% transmission rate, suggest that these R2 alleles are likely induced from the paternal *wt* alleles by maternal carryover of Cas9/gRNA in zygotes harboring the maternal rescuing *Pol-\gamma35* deletes, for a HGD. The maternal carryover can be a major source of both R2 and R1 resistance alleles,

because it can possibly facilitate the cleavage and NHEJ repair of paternal wt allele before it comes into proximity with a maternal carrier allele to facilitate HDR (Adolfi et al., 2020; Champer et al., 2019; Gantz et al., 2015; Kandul et al., 2019a). Once R2 alleles are complemented by the $Pol-\chi 35^{HomeR1}$ allele, it takes several generations for R2 alleles to combine as lethal homozygotes and be selected out from a population. The elimination of R2 alleles takes especially long time by HGDs targeting non-essential genes or genes whose knockouts do not cause complete lethality or sterility of homozygous carriers (Adolfi et al., 2020; Gantz et al., 2015) underscoring the importance of targeting essential haplosufficient genes. Therefore, to assess the stability of gene drives against accumulation of functional resistance alleles, its spread must be examined for several generations after a carrier frequency has reached 100%.

Our results are congruent with previous studies (Chan et al., 2013, 2011; Windbichler et al., 2011) demonstrating reduced homing in *Drosophila* males. We tested multiple Cas9 lines supporting Cas9 expression in early and/or late germ cells with different levels of specificity, and have not achieved high levels of homing as reported in mosquito males, aka. >90% (Gantz et al., 2015; Kyrou et al., 2018). Achiasmatic meiosis in *Drosophila* males likely correlates with the weak activity of HDR pathway (Preston et al., 2006), which in turn results in inefficient homing in *Drosophila* males. Mosquito males have chiasmatic meiosis and recombination (Kitzmiller, 1976) that require active HDR machinery in primary spermatocytes, possibly contributing to efficient homing in mosquito males. Reduced homing efficacy in *Drosophila* males should be accounted for when designing HGDs in other species exhibiting achiasmatic meiosis, such as *D. suzukii*, an invasive fruit pest.

Our results indicate that functional resistance (R1) alleles can still be induced even when a conserved haplosufficient gene required for insect viability is targeted (**Fig. 4**). However, each identified in-frame, R1 allele changes at least one amino acid and thus may affect the fitness of its carrier preventing such alleles from accumulating at the expense of the drive. This incurring fitness cost likely slows down their accumulation and results in selection out of the population, in favor of the *Pol-\gamma35* alleles, over multiple generations (**Fig. 4**) again underscoring the importance of targeting an essential haplosufficient gene. Nevertheless, encoding additional gRNAs targeting the *wt* coding sequence of *Pol-\gamma35* downstream from the Cas9/gRNA cut site, which is re-coded in *Pol-\gamma35* alleles, into HomeR may further diminish the probability of inducing functional resistance alleles.

Splitting HomeR into two genetic loci (*HomeR* and *Cas9*) integrated on different chromosomes serves as a necessary molecular containment mechanism (**Fig. 5**). The *HomeR* element is able to home into *wt* alleles and bias its transmission. However, the *Cas9* element, which is inherited Mendelianly, is required for its homing. Therefore, the independent assortment of *Cas9* and *HomeR* limits the spread of the *HomeR* element and acts as a 'brake' for HomeR propagation. The spread dynamic of split-HGDs resembles that of high-threshold drives and thus requires a high introduction rate for HomeR to spread into a local population and prevents its spread into neighboring populations (**Fig. 5**). Moreover, if unintended consequences do arise, HomeR's spread can be reversed by reintroduction of insects harboring *wt* alleles of the gene targeted by split-drive. Notwithstanding, if desired, HomeR could facilely be converted into a non-localized gene drive by incorporating the Cas9 into the Homer drive cassette. Taken together, the split-design of HomeR is safe localized gene drive technology that could be safely adopted and implemented for local control, and if a non-localized drive is desired for more wide scale spread, HomeR could be converted for that purpose too.

In sum, HomeR combines promising aspects of current population-replacement drives - confineablity, high transmission of HGD's, and resilience to NHEJ generation of TA drives (Fig.

6). Modeling illustrates success of both design aspects in linked or split-drive form, demonstrating robust behavior over a range of parameter combinations (**Fig. 5**). This underscores its resilience to NHEJ alleles, overcoming a significant hurdle for current HGD designs. Given the simplicity of the HomeR design, it could be universally adapted to a wide range of species including human disease vectors in the future.

Methods

Selection of Cas9/gRNA target sites

We inserted a <u>Home</u>-and-<u>Rescue</u> (HomeR) in *DNA Polymerase y 35-kDa* (*Pol-y35* or *PolyB*, CG33650). *Pol-y35* is a haplosufficient gene required for insect viability: a lethal knockout can be rescued by a single functional copy. The highly conserved domain of *Pol-y35* is located at the end of the coding sequence, which facilitates its re-coding (**Fig. 1A**). We PCR amplified a 413-base fragment of the domain with 1073A.S1F and 1073A.S2R from multiple *Drosophila* strains (w^{1118} , Canton S, Oregon R, *nos-Cas9* (*Kandul et al., 2019b*)) and used the consensus sequence along with the tool CHOPCHOP v2 (Labun et al., 2016) to choose two gRNA targets sites that minimize off-target cleavage.

Design and assembly of genetic constructs

We used Gibson enzymatic assembly to build all genetic constructs (Gibson et al., 2009). To assemble both gRNA constructs, we used the previously described $sgRNA^{Sxl}$ plasmid (Kandul et al., 2019b) (Addgene #112688) harboring the mini-white gene and attB docking site. We removed the fragment encompassing the U6.3 promoter and gRNA scaffold by Ascl and SacII digestion, and we cloned it back as two fragments overlapping at a novel gRNA sequence (**Fig. 1A**). Both U6.3-gRNA# $1^{Pol-rac{1}{2}35}$ and U6.3-gRNA# $2^{Pol-rac{1}{2}35}$ plasmids targeting $Pol-rac{1}{2}35$ are deposited at www.addgene.org (#159774 and #159675).

We assembled two *HomeR*^{Pol-Y35} constructs using two tested gRNAs (**Fig. 2A**). Each *HomeR*^{Pol-Y35} was built around a specific gRNA, with matching LHA and RHA: *HomeR1*^{Pol-Y35} harbored *U6.3-gRNA#1*^{Pol-Y35}, and *HomeR2* had *U6.3-gRNA#2*^{Pol-Y35}. We digested the *nos-Cas9* plasmid (Kandul et al., 2019b) (Addgene #112685) with AvrII and AscI, preserving the backbone containing the *piggyBac* left and right sequences that encompass the *Opie-dsRed-SV40* marker gene. The HomeR construct was assembled between *Opie-dsRed-SV40* and *piggyBacR* in three steps. First, we cloned the *U6.3-gRNA#1* or #2 from the corresponding plasmid together with the *3xP3-eGFP-SV40* marker gene, to tag site-specific insertion of *GDe*. Then, we cloned three fragments: (1) LHA, which was amplified from the *Drosophila* genomic DNA; (2) the re-coded fragment downstream from the gRNA cut site, which was PCR amplified from the dePol-Y35 gBlock custom synthesized by IDT® (**Supplementary Table 1**); (3) the p10 3'UTR to provide robust expression (Pfeiffer et al., 2012) of the re-coded *Pol-Y35* rescue. Finally, we cloned RHA, which was PCR amplified from genomic DNA, corresponding to each specific gRNA cut site. Both *HomeR1*^{Pol-Y35} and *HomeR2*^{Pol-Y35} plasmids, targeting the *Pol-Y35* locus, are deposited at www.addgene.org (#159676 and #159677).

To assemble the three constructs for testis-specific Cas9 expression, we used a plasmid harboring the *hSpCas9-T2A-GFP*, the *Opie2-dsRed* transformation marker, and both *piggyBac* and attB-docking sites, which were previously used to establish Cas9 transgenic lines in *Aedes aegypti* (*Li et al., 2017*) and *Drosophila melanogaster* (Kandul et al., 2019a, 2019b). We removed the *Ubiquitin 63E* promoter from the *ubiq-Cas9* plasmid (Addgene #112686) (Kandul et al., 2019b) by digesting it with Swal at +27°C and then with Notl at +37°C, and cloned a promoter fragment amplified from the *Drosophila* genomic DNA. The *Drosophila exuperantia* (CG8994) 783-bp fragment (*exuL*) upstream of the *exuperantia* gene was amplified with ExuL.1F and ExuL.2R primers (**Supplementary Table 1**) and cloned to assemble the *exuL-Cas9* plasmid. The *Rcd-1*

related (Rcd1r, CG9573) (Chan et al., 2013) and β -Tubulin 85D (β Tub) (Chan et al., 2011; Michiels et al., 1989) promoters support early and late, respectively, testis-specific expression in Drosophila males. The 937-base-long fragment upstream of Rcd1r was amplified with 1095.C1F and 1095.C2R primers and cloned to assemble the Rcd1r-Cas9 plasmid. The 481-base-long fragment upstream of β Tub was amplified with β Tub.1F and β Tub.2R primers (**Supplementary Table 1**) and cloned to build the β Tub-Cas9 plasmid. Three plasmids for testis-specific Cas9 expression are deposited at www.addgene.org (#159671 – 159773).

Fly maintenance and transgenesis

Flies were maintained under standard conditions: 26°C with a 12H/12H light/dark cycle. Embryo injections were performed by Rainbow Transgenic Flies, Inc. We used φC31-mediated integration (Groth, 2004) to insert the U6.3-qRNA#1 and U6.3-qRNA#2 constructs at the P{CaryP}attP1 site on the 2nd chromosome (BDSC #8621), and the exuL-Cas9, βTub-Cas9, and Rcd1r-Cas9 constructs at the PBac{y+-attP-3B}KV00033 on the 3rd chromosome (BDSC #9750). Two methods were used to generate the site-specific insertion of HomeR1Pol-Y35 or HomeR2Pol-Y35 constructs at the *gRNA#1*^{Pol-\gamma35} or *gRNA#2*^{Pol-\gamma35} cut sites, respectively, inside the *Pol-\gamma35* gene via HDR. First, we injected the mixture of HomeR and helper phsp-pBac, carrying the piggyBac transposase (Handler and Harrell, 1999), plasmids (500 ng/µl and 250 ng/µl, respectively, in 30 µI) into w^{1118} embryos. Random insertions of HomeR1^{Pol-Y35} and HomeR2^{Pol-Y35}, assessed by double (eye-specific GFP and body-specific dsRed) fluorescence (Fig. 2B), established with this injection were genetically crossed to nos-Cas9/nos-Cas9 (BDSC #79004) (Kandul et al., 2019b) flies to "relocate" HomeR1Pol-Y35 or HomeR2Pol-Y35 to the corresponding gRNA cut site via Homology Assisted CRISPR Knock-in (HACK) (Lin and Potter, 2016). A few site-specific Pol-Y35^{HomeR1} and Pol-Y35^{HomeR2} lines tagged with only eye-specific GFP fluorescence were recovered. Second, we injected HomeR1^{Pol-Y35} or HomeR2^{Pol-Y35} plasmids directly into nos-Cas9/nos-Cas9 (BDSC #79004) (Kandul et al., 2019b) embryos, generating multiple independent, site-specific insertions for each Pol-Y35HomeR (Fig. 2B). Recovered transgenic lines were balanced on the 2nd and 3rd chromosomes using single-chromosome balancer lines (w^{1118} ; CyO/sna^{Sco} for II and w^{1118} ; TM3, $Sb^{1}/TM6B$, Tb^{1} for III) or a double-chromosome balancer line (w^{1118} ; CyO/Sp; Dr/TM6C, Sb^{1}).

We established three homozygous lines of $Pol-\gamma 35^{HomeR1}$ and $Pol-\gamma 35^{HomeR2}$ from independent insertion lines, and confirmed the precision of site-specific insertions by sequencing the borders between HomeR constructs and the Drosophila genome (**Fig. 2C**). The 1118-base-long fragment overlapping the left border was PCR amplified with 1076B.S9F and 1076B.S2R and was sequenced with 1076B.S3F and 1076B.S4R primers. The same-length fragment at the right border was amplified with 1073A.S1F and 1076B.S10R and was sequenced with 1076B.S7F and 1076B.S8R primers (**Supplementary Table 1**).

Fly genetics and imaging

Flies were examined, scored, and imaged on a Leica M165FC fluorescent stereo microscope equipped with a Leica DMC2900 camera. We assessed the transmission rate of HomeR by following its eye-specific GFP fluorescence, while the inheritance of *Cas9* was tracked via body-specific dsRed fluorescence (**Fig. 2B, Fig. 4A**). All genetic crosses were done in fly vials using groups of ten males and ten females.

RNA^{Pol-Y35} cleavage assay

To assess the cleavage efficiency of each gRNA targeting the C-terminal domain of $Pol-\slash35$, we genetically crossed ten w^{1118} ; U6.3- $gRNA\#1^{Pol-\slash35}$ or w^{1118} ; U6.3- $gRNA\#2^{Pol-\slash35}$ homozygous males to ten y^1 , Act5C-Cas9, w^{1118} , Lig4 (X. $Zhang\ et\ al.$, 2014) (BDSC #58492) homozygous females, and we scored the lethality of F_1 males (**Fig. 1B**). The F_1 males would then inherit the X chromosome from their mothers, expressing U6.3- $gRNA\#1^{Pol-\slash35}$ or U6.3- $gRNA\#2^{Pol-\slash35}$ with

Act5C-Cas9 in a Lig4-null genetic background, and this results in male lethality when a tested gRNA directs cleavage of the Pol- γ 35 locus. To assess the induced lethality in the Lig4+/+ genetic background, we crossed ten γ^1 , Act5C-Cas9, w^{1118} (BDSC #54590) (Port et al., 2014) flies to ten U6.3-gRNA#1^{Pol- γ 35} flies in both directions, and scored survival of trans-heterozygous and heterozygous F₁ progeny. To measure the Cas9/gRNA-directed cleavage of Pol- γ 35 by maternally deposited Cas9 protein in the Lig4+ background, the same homozygous males were genetically crossed to w^{1118}/w^{1118} ; nos-Cas9/CyO females (**Fig. 1C**), and the F₁ progeny, harboring U6.3-gRNA#1^{Pol- γ 35} or U6.3-gRNA#2^{Pol- γ 35}, were scored and compared to each other.

Assessment of Pol-Y35HomeR transmission rates

To compare transmission rates of Pol-Y35^{HomeR1} and Pol-Y35^{HomeR2}, we first established transheterozygous parent flies by genetically crossing Pol-\(\chi \)35^{HomeR1}/Pol-\(\chi \)35^{HomeR1}; +/+ or Pol-V35^{HomeR2}/Pol-V35^{HomeR2}; +/+ females to +/+; nos-Cas9/nos-Cas9 males. We then assessed the transmission rates by trans-heterozygous parent females and males crossed to wt flies. For controls, we estimated the transmission rates of *HomeR1*^{Pol-}¥35 and *HomeR1*^{Pol-}¥35 in the absence of Cas9, by heterozygous Pol-\gamma35^HomeR1/+ or Pol-\gamma35^HomeR2/+ females and males crossed to wt flies (Fig. 3A). To explore the effect of maternally deposited Cas9 protein on transmission of Pol-√35^{HòmeR1}(Kandul et al., 2019a), we generated heterozygous Pol-√35^{HomeR1}/CyO embryos containing Cas9 protein deposited by nos-Cas9/CyO mothers and estimated the transmission of Pol-\(\chi 35^{HomeR1} \) by females and males raised from these embryos and crossed to wt flies. We tested five different Cas9 lines—supporting germline (vas-Cas9), ubiquitous (ubiq-Cas9, Act5C-Cas9), and early (exuL-cas9, Rcd1r-Cas9) or late testes-specific expression (β Tub-Cas9)—together with the strongest HomeR, Pol-Y35HomeR1. Ten trans-heterozygous females or males, generated by crossing homozygous Pol-V35HomeR1 females to homozygous Cas9 males, were genetically crossed to wt flies and the transmission of Pol-Y35^{HomeR1} was quantified in their F₁ progeny (Fig. 3C).

Egg hatching assay to assess the homing rate of Pol-¥35^{HomeR1}

To identify the mechanism of the super-Mendelian transmission of $Pol-\gamma35^{HomeR1}$, we assessed the percentage of F_1 hatched eggs laid by trans-heterozygous $Pol-\gamma35^{HomeR1}$ /+; nos-Cas9/+ females genetically crossed to wt males and compared it to those hatched from two types of heterozygous females: $Pol-\gamma35^{HomeR1}$ /+; +/+ φ and +/+; nos-Cas9/+ φ (**Fig. 3B**). We collected virgin females and aged them for three days inside food vials supplemented with a yeast paste, then five groups of 25 virgin females of each type were transferred into vials with fresh food containing 25 wt males and allowed to mate overnight (12 H) in the dark. Then, all males were removed from the vials, while females were transferred into small embryo collection cages (Genesee Scientific 59–100) with grape juice agar plates. After 12 H of egg laying, a batch of at least 200 laid eggs was counted for each sample group and incubated for 24 H at 26°C before the number of unhatched eggs was counted. Some fraction of hatched larvae for each test group was transferred into food vials to confirm that they would finish development.

Accumulation of functional in-frame resistance alleles, Pol-Y35R1

To explore the generation and accumulation of functional resistance alleles induced by NHEJ, we initiated three drive populations by crossing 50 +/+; nos-Cas9/nos-Cas9 females and 50 $Pol-\gamma35^{HomeR1}/Pol-\gamma35^{HomeR1}$; nos-Cas9/nos-Cas9 males in 0.3 L plastic bottles (VWR® Drosophila Bottle 75813-110). Parent (P) flies were removed after six days, and their progeny were allowed to develop, eclose, and mate for 13–15 days. This established a 100% heterozygous $Pol-\gamma35^{HomeR1}/+$; nos-Cas9/+ population in the next generation (G₀) (due to ~100% transmission efficiency) , with 50% allelic and 100% genotypic frequency of $Pol-\gamma35^{HomeR1}$ in each bottle population. Each generation, around 250–350 emerged flies were anesthetized using CO₂, and their genotypes with respect to $Pol-\gamma35^{HomeR1}$ (presence or absence) were determined using the

dominant eye-specific GFP marker. Then they were transferred to a fresh bottle and allowed to lay eggs for six days before removing them, and the cycle was repeated. Three populations were maintained in this way for eleven generations, which corresponds to ten generations of gene drive. Note that any fly scored without the $Pol-\gamma35^{HomeR1}$ allele was transferred into a fresh bottle to ensure any $Pol-\gamma35$ resistance or wt alleles could be passed to the next generation. We retrieved and froze the flies for genotyping only after six days to ensure sufficient time for breeding. We expected that the gRNA#1/Cas9-induced $Pol-\gamma35^{R1}$ alleles that did not incur fitness costs would accumulate over a few generations and block the spread of the gene drive. However, as we did not find any fly without the $Pol-\gamma35^{HomeR1}$ allele after G_3 , we stopped the population drives after ten generations of $Pol-\gamma35^{HomeR1}$ homing in the heterozygous flies. We froze 60 flies after G_{10} for further sequence analysis.

Sequencing of induced resistance alleles

To analyze the molecular changes that caused functional in-frame (R1) and loss-of-function (LOF, R2) mutations in $Pol-\chi^35$, we PCR amplified the 232-base-long genomic region containing both $gRNA\#1^{Pol-\chi^35}$ and $gRNA\#2^{Pol-\chi^35}$ cut sites using 1073A.S3F and 1073A.S4R primers (**Supplementary Table 1**). For PCR genotyping from a single fly, we followed the single-fly genomic DNA prep protocol (Kandul et al., 2019b). PCR amplicons were purified using the QIAquick PCR purification kit (QIAGEN), subcloned into the pCRTM2.1-TOPO® plasmid (Thermofisher), and 5–7 clones were sequenced in both directions by Sanger sequencing at Retrogen® and/or Genewiz® to identify both alleles in a each fly. Sequence AB1 files were aligned against the corresponding wt sequence of $Pol-\chi^35$ in SnapGene® 4.

To explore the diversity of resistance alleles persisting after 10 generations of Pol-Y35^{HomeR1} in a 100% heterozygous population, we froze 60 flies (30 \circ and 30 \circ), each harboring at least one copy of the dominant marker of Pol-Y35HomeR1, from each drive population after G10. Using these flies, we quantified any resistance and wt alleles remaining in the population via Illumina sequencing of heterogeneous PCR amplicons from the Pol-Y35 locus. Note that PCR amplicons did not include the Pol-Y35HomeR1 allele due to its length (Fig. 2A). DNA was extracted using the DNeasy Blood and Tissue Kit (QIAGEN). To analyze heterogeneous PCR products, we used the Amplicon-EZ service by Genewiz[®] and followed the Genewiz[®] guidelines for sample preparation. In brief, Illumina adapters were added to the 1073A.S3F and 1073A.S4R primers to simplify the library preparation, PCR products were purified using QIAquick PCR purification kit (QIAGEN), around 50,000 one-direction reads covering the entire amplicon length were generated, and relative abundances of recovered SBS and *indel* alleles at the *qRNA#2*^{Pol-y35} cut site were inferred using Galaxy tools (Afgan et al., 2018). Amplicon-EZ data from Genewiz® were first uploaded to Galaxy.org. A quality control was performed using FASTQC. Sequence data were then paired and aligned against the *Pol-\footnote{35}^{WT}* sequence using Map with BWA-MEM under "Simple Illumina" mode". The SBS and indel alleles were detected using FreeBayes, with the parameter selection level set to "simple diploid calling".

Modeling of gene drive spread

Simulations were performed using a discrete-generation version of the Mosquito Gene Drive Explorer (MGDrivE) modeling framework (C. et al., 2020). The first generation was seeded with 200 adults, 75% wild-type and 25% heterozygous for each gene drive, split equally between sexes. At each generation, adult females mate with males, thereby obtaining a composite mated genotype (their own, and that of their mate) with mate choice following a multinomial distribution determined by adult male genotype frequencies. Egg production by mated adult females then follows a Poisson distribution, proportional to the genotype-specific lifetime fecundity of the adult female. Offspring genotype follows a multinomial distribution informed by the composite mated female genotype and the inheritance pattern of the gene drive system. Sex distribution of offspring

follows a binomial distribution, assuming equal probability for each sex. Female and male adults from each generation are then sampled equally to seed the next generation, with a sample size of 200 individuals (100 female and 100 male), following a multivariate hypergeometric distribution. 25 repetitions were run for each drive in the trace plots (**Fig. 5A and 5C**) and 100 repetitions were run for each parameter combination in the heatmaps (**Fig. 5B and 5D**).

The inheritance pattern is captured by the "inheritance cube" module of MGDrivE. ClvR and TARE constructs were implemented to match their published descriptions (Champer et al., 2020a; Oberhofer et al., 2020a, 2019). HomeR and HGD were implemented as one or two-locus systems following equivalent inheritance rules. When Cas9 and gRNAs co-occur in the same individual, wild-type alleles are cleaved at a rate $c_F(c_M)$ (female- (male-) specific cleavage), with $1-c_F(1-c_M)$ remaining wild-type. Given cleavage, successful HDR occurs at a rate $ch_F(ch_M)$, with $1-ch_F(1-ch_M)$ alleles undergoing some form of NHEJ. Of these, a proportion, $cr_F(cr_M)$, are in-frame NHEJ alleles, while the remainder, $1-cr_F(1-cr_M)$, are LOF alleles. Maternal carryover (maternal deposition, or maternal perdurance) was modeled to occur in zygotes of mothers having both Cas9 and gRNAs, impacting a proportion, d_F , of zygotes. Of the wild-type alleles in impacted zygotes, a proportion, dr_F , become in-frame NHEJ alleles, while the remainder, $1-dr_F$, become LOF alleles. These inheritance rules apply to both HomeR and HGD, with differing fitness costs.

ClvR (Oberhofer et al., 2020a, 2019) was modeled using a 99% cleavage rate in female and male germ cells, as well as in embryos from maternal carryover. For two-locus ClvR, the two loci were assumed to undergo independent assortment (>=50cM separation), as was assumed for all twolocus systems in this analysis. For both configurations, it was assumed that 0.1% of cleaved alleles were converted to functional resistance alleles (R1 type), and the rest became LOF alleles (R2 type). In addition to the 50% egg-hatching reduction due to the non-homing drive (Fig. 6A-B), an additional 5% reduction in fecundity was applied to females that harbored Cas9. For consistency, TARE, HGD, and HomeR (for ideal parameters) also used a cleavage rate of 99% in females and males, though TARE demonstrated lower maternal carryover (Champer et al., 2020a), and was modeled with 95% cleavage. HGD and HomeR (for ideal parameters), which rely on HDR, were simulated with 90% HDR rates in females and males. Cleaved alleles that did not undergo HDR were assumed to be R1 alleles with proportion 0.5%, and R2 LOF alleles the remainder of the time. TARE and HomeR were also modeled with a small (5%) fitness reduction, applied as a reduction of female fecundity. Since a HGD does not provide a rescue for a knocked out targeted gene, its carriers demonstrate higher fitness costs, and were assigned a 20% fitness reduction with the assumption that the HGD is inserting into a non-lethal gene that imposes a low/moderate fitness cost. Experimentally-derived parameters for HomeR differed from ideal parameters in two ways: i) there was no HDR in males (although cleavage remained the same), and ii) 1% of NHEJ-repaired wt alleles were converted into R1 alleles (c.f. 0.5% for the ideal case). All simulations were performed, analyzed, and plotted in R ("Website," n.d.) (R Core Team 2017). Code is available upon request.

Statistical analysis

Statistical analysis was performed in JMP 8.0.2 by SAS Institute Inc., and graphs were constructed in Prism 8.4.1 for MacOS by GraphPad Software LLC. At least three biological replicates were used to generate statistical means for comparison. *P* values were calculated using a two-sample Student's *t*-test with equal variance.

Gene Drive safety measures

All crosses using gene drive genetics were performed in accordance to a protocol approved by the Institutional Biosafety Committee at UCSD, in which full gene drive experiments were performed in a high-security ACL2 barrier facility and split-drive experiments were performed in an ACL1 insectary in plastic vials that were autoclaved prior to being discarded, in accordance with currently suggested guidelines for the laboratory confinement of gene drive systems (Akbari et al., 2015; National Academies of Sciences, Engineering, and Medicine et al., 2016).

Ethical conduct of research

We have complied with all relevant ethical regulations for animal testing and research and conformed to the UCSD institutionally approved biological use authorization protocol (BUA #R2401).

Data and Reagent Availability

All data are represented fully within the tables and figures. The U6.3- $gRNA#1^{Pol-\gamma35}$, U6.3- $gRNA#2^{Pol-\gamma35}$, $HomeR1^{Pol-\gamma35}$, $HomeR2^{Pol-\gamma35}$, exuL-Cas9, Rcd1r-Cas9, and βTub -Cas9 plasmids and corresponding fly lines are deposited at Addgene.org (159671–159677) and the Bloomington Drosophila Stock Center, respectively.

DISCLOSURES

O.S.A is a founder of Agragene, Inc., has an equity interest, and serves on the company's Scientific Advisory Board. N.P.K is a consultant for Agragene. All other authors declare no competing interests.

AUTHOR CONTRIBUTIONS

N.P.K and O.S.A. conceived the idea and designed experiments. N.P.K and J.L. engineered plasmids, and performed all molecular and genetic experiments. J.B.B and J.M.M performed mathematical modeling. All authors analyzed the data, contributed to the writing of the manuscript, and approved the final manuscript.

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SUPPLEMENTARY INFORMATION

Supplementary Table 1. Target sequence of gRNA and primers used in this study.

Supplementary Data 1. Cleavage assay of two $gRNAs^{Pol-\gamma35}$ with Act5C-Cas9 in the $Lig4\Delta$ genetic background.

Supplementary Data 2. Cleavage assay of two *gRNAs*^{Pol-y35} with *nos-Cas9*.

Supplementary Data 3. Transmission rate of *Pol-γ35*^{HomeR1} or *Pol-γ35*^{HomeR1} in conjunction with different Cas9 lines.

Supplementary Data 4. Assessment of embryonic lethality induced by *Pol-γ35*^{HomeR1}/+; nos-Cas9/+ females.

Supplementary Data 5. Spread of resistance alleles in $Pol-\gamma 35^{HomeR1}/+$; nos-Cas9/nos-Cas9 flies over ten generations.

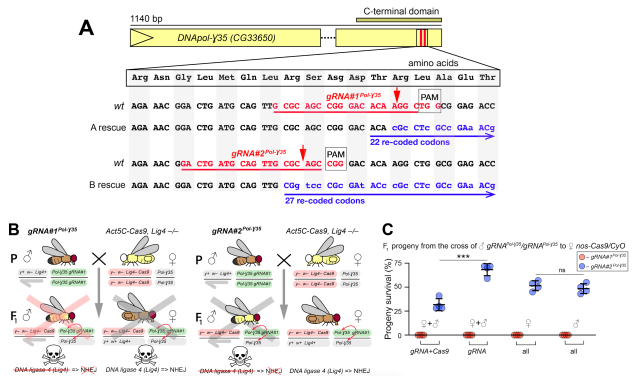


Figure 1. Assessing gRNAs targeting the *Drosophila Pol-y35* gene. (A) Schematic map of the DNA polymerase gamma 35 kb gene (Pol-y35). Two gRNAs targeting its highly conserved C-terminal domain were chosen. Both gRNA target sites (highlighted in red) are located near the 3' end of the coding sequence, facilitating re-coding for the rescue allele (highlighted in blue), resistant to Cas9/gRNA-mediated cleavage. Red arrows depict Cas9/gRNA cut sites. (**B**) Crosses to assess the cleavage efficiency of the gRNAs in the *DNA ligase 4* null genetic background (Lig4-/-), in which the Non-Homologous End Joining (NHEJ) pathway is inactive. Homozygous $gRNA\#1^{Pol-y35}$ or $gRNA\#2^{Pol-y35}$ males were crossed to homozygous Act5C-Cas9, Lig4-/- females, resulting in death of all male progeny for each gRNA. Notably, female progeny harboring $gRNA\#1^{Pol-y35}$ and Act5C-Cas9 in the Lig4+/- background also perished. (**C**) $gRNA\#1^{Pol-y35}$, but not $gRNA\#2^{Pol-y35}$, induced embryonic lethality of all F₁ progeny in conjunction with nos-Cas9 or maternal carryover of the Cas9 protein. The frequency of F₁ progeny survival presented by the genotype and/or sex. The plot shows the mean ± standard deviation (SD) over four biological replicates. Statistical significance was estimated using a two-sided Student's t test with equal variance. ($p \ge 0.05^{ns}$ and $p < 0.001^{***}$).

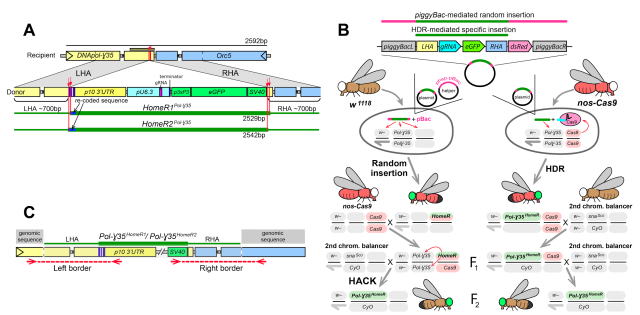


Figure 2. The Pol-y35 HomeR split-drive and its site-specific integration. (A) Schematic maps of the recipient (wildtype, wt) allele encompassing the area spanning Pol-y35 and Orc5 (CG7833) genes, and the donor allele harboring the HomeR1 or HomeR2 site-specifically integrated at gRNA cut site #1 or #2 in Pol-y35, respectively. Red arrows depict the Cas9/gRNA cut sites, which are 13 bases apart (Fig. 1A). To facilitate site-specific integration, each HomeR genetic construct is surrounded by the Left and Right Homology Arms (LHA and RHA) from the corresponding Cas9/gRNA cut site in the wt allele. The re-coded 3' end sequences of Pol-v35 (LHA) are shown in dark blue for both *HomeR1*^{Pol-γ35} and *HomeR2*^{Pol-γ35}. (**B**) Two separate approaches were used to generate transgenic lines harboring site-specific insertions of HomeR1 or HomeR2 at the Drosophila Pol-v35. In the first approach, two plasmids, one carrying the HomeR construct and the helper plasmid (phsp-pBac (Handler and Harrell, 1999)) carrying the piggyBac transposase, were injected into w^{1118} embryos to generate transgenic lines harboring a random piggyBac-mediated integration tagged by double fluorescence (GFP+ and dsRed+). Then transgenic males (GFP+, dsRed+) were crossed to nos-Cas9 females (dsRed+) to generate sitespecific Pol-v35GDe transgenic lines, tagged by GFP alone, using Homology Assisted CRISPR Knock-in (HACK) (Lin and Potter, 2016). In the second approach, the plasmid harboring the HomeR genetic construct was injected alone into nos-Cas9 embryos (dsRed+) and sitespecifically integrated via Homology Directed Repair (HDR) to generate F₁ heterozygous Polv35^{GDe}/+; nos-Cas9/+ males with double fluorescence (GFP+ and dsRed+). In both approaches, F₁ transgenic males harboring *Pol-γ35*^{HomeR} (GFP+) and *nos-cas9* (dsRed+) were crossed to the 2^{nd} chromosome balancer line, $w^{11\dot{1}8}$; CyO/sna^{Sco} , to balance and isolate the $Pol-y35^{GDe}$ insertion (GFP+). With this approach, viable homozygous lines were established. (C) Finally, to confirm site-specific integration of both *Pol-y35*^{HomeR1} and *Pol-y35*^{HomeR2}, the left and right borders between either HomeR1 or HomeR2 genetic construct and Drosophila genomic sequence around the integration site were sequenced and in three independent homozygous lines.

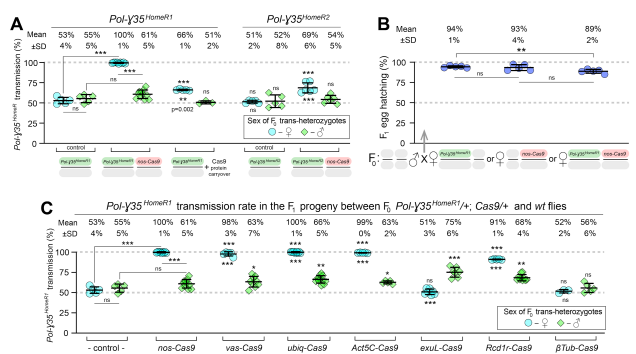


Figure 3. Transmission rates for Pol-y35HomeR1 and Pol-y35HomeR2. The HomeR element, Polv35^{HomeR1} or Pol-v35^{HomeR2}, is inactive by itself and requires Cas9 endonuclease to induce Cas9/gRNA-mediated cleavage for successful homing. This split-drive design permits genetic analysis of a single HomeR with different Cas9 lines. (A) Both Pol-y35HomeR1 and Pol-y35HomeR2 support super-Mendelian transmission in conjunction with nos-Cas9 in females, but not in males. Pol-v35^{HomeR1} induced significantly higher transmission than Pol-v35^{HomeR2}, 99.5 ± 0.6% vs 68.7 ± 6.2%, respectively. Notably, maternal carryover of Cas9 protein was sufficient to bias transmission of $Pol-\gamma 35^{HomeR1}$ by female embryos. (**B**) The hatching rate of F_1 eggs generated by Pol-y35^{HomeR1}/+; nos-Cas9/+ females mated to wildtype (wt) males was lower by 5% or 4% than that of Pol- $\frac{1}{2}$ 35^{HomeR1}/+; +/+ or +/+; nos-Cas9/+ females mated to wt males (89 ± 2% vs 94 ± 1% or 93 ± 4%). Therefore, embryonic lethality of wt Pol-\(\chi \) 35 alleles is not the sole mechanism of the nearly ~100% transmission of *Pol-γ35*^{HomeR1}. Instead, ~90% of wt *Pol-γ35* alleles were converted (i.e. homed) into Pol-y35^{HomeR1}. (C) Assessment of different Cas9 promoters to improve the transmission rate of Pol-y35^{HomeR1} in females and males. Trans-heterozygous females (♀) and males (3) harboring paternal Cas9 expressed under different promoters were mated to wt flies of the opposite sex, and F₁ progeny were scored for the GFP dominant marker of *Pol-y35*^{HomeR1}. The transmission rate was compared to that in Pol-y35HomeR1/+; +/+ flies without Cas9 (control) of the corresponding sex (statistical significance indicated above data points). In addition, the transmission rate by trans-heterozygous females was compared to that of trans-heterozygous males for each Cas9 promoter (statistical significance indicated below data points). Plots show the mean ± SD over at least three biological replicates and rounded to a whole number. Statistical significance was estimated using a two-sided Student's t test with equal variance. ($p \ge 0.05^{ns}$, p $< 0.05^*$, $p < 0.01^{**}$, and $p < 0.001^{***}$).

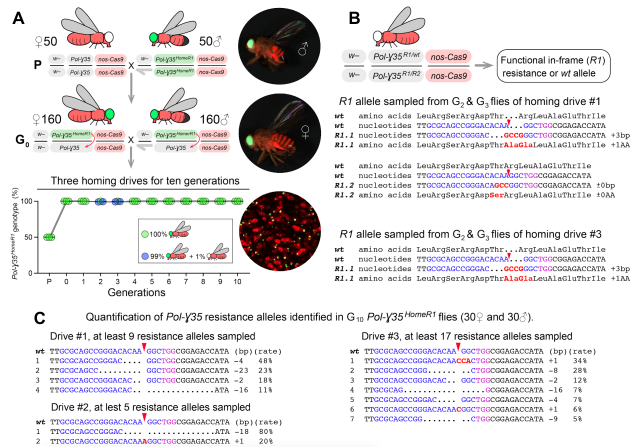


Figure 4. Functional resistance alleles (Pol-y35^{R1}) do not prevent drive persistence of Polv35^{HomeR1}. (A) To explore the fate of induced functional resistance alleles (R1), three gene population cage experiments were set up starting with Pol-y35^{HomeR1}/Pol-y35^{WT} heterozygous flies in nos-Cas9/nos-Cas9 genetic background and run for ten discrete generations. Pol-v35^{HomeR1}/+: nos-Cas9/nos-Cas9 flies were tagged by eye-specific GFP and body-specific dsRed. Images of an individual male (β), female (Ω), and a group of flies are shown. In total, nine viable flies (lacking the *Pol-y35*^{HomeR1} allele), as determined by the absence of dominant eye-specific GFP expression, were identified at generations 2 and 3 in cages #1 and #3. After these flies were allowed to mate and lay eggs for the next generation, then isolated and genotyped. (B) None of nine genotyped flies harbored the Pol-v35WT allele. Instead, each fly carried at least one functional resistance allele rescuing the viability of these flies. Both different types of R1 alleles change the amino acid sequence. Seven of nine flies were heterozygous, harboring one of the identified R1 alleles together with an out-of-frame indel allele. (C) Resistance alleles, persisting for ten generations of cleavage and homing, were sampled from sixty flies chosen randomly harboring at least one Poly35^{HomeR1} allele and were quantified using Illumina® sequencing. For each gene drive, nearly 50,000 amplicons of Pol-y35 alleles (150K total), which did not carry the 2.5 kb insert of the HomeR1, were sequenced and used to estimate the minimum number of sampled resistance alleles. Note that both R1.1 and R1.2 alleles identified at earlier generations (**B**) were not sampled. Two novel in-frame resistance alleles (-18 bp and -9 bp) resulted in deletions of 6 and/or 3 amino acids and would not be good rescues. The sequence of $gRNA#1^{Pol-y35}$ is highlighted in blue, and its PAM sequence is in purple. Red arrows depict Cas9/gRNA cut sites. Base insertions and amino acid changes are in red.

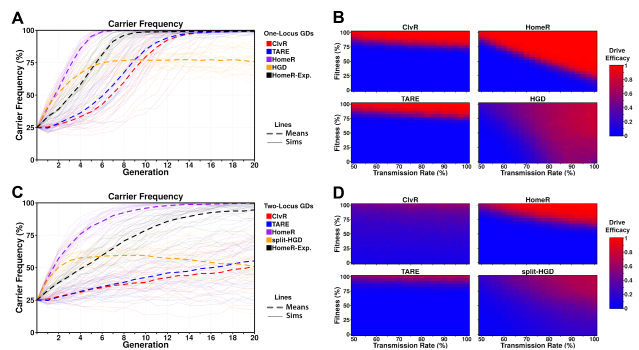


Figure 5. Performance of contemporary gene drive systems for population modification. (A) Simulations of carrier frequency trajectories (i.e. heterozygotes and homozygotes) for onelocus versions of ClvR, TARE, HomeR, and HGD for ideal parameters (see Methods), and HomeR for experimental parameters (HomeR-Exp, see Methods). 25 repetitions (lighter lines) were used to calculate the average behavior of each drive (thicker, dashed lines). Populations were initialized with 75% wildtype (+/+) adults and 25% drive heterozygotes (drive/+), equally split between females and males. (B) Heatmaps depicting drive efficacy for one-locus versions of ClvR, TARE. HomeR, and HGD for a range of fitness and transmission rate parameter values. Fitness costs were incorporated as a dominant, female-specific fecundity reduction. Transmission rate was varied based on cleavage rate, using HDR rates consistent with ideal parameters, when applicable (see Methods). Drive efficacy is defined as the average carrier frequency at generation 20 (approximately 1 year, given a generation period of two to three weeks) based on 100 stochastic simulations with the same initial conditions as A. (C) Simulations of carrier frequency trajectories for two-locus (split-drive) versions of ClvR, TARE, HomeR, and HGD for ideal parameters (see Methods), and HomeR for experimental parameters (HomeR-Exp, see Methods). 25 repetitions (lighter lines) were used to calculate the average behavior of each drive (thicker, dashed lines). Populations were initialized with 75% wildtype (+/+; +/+) adults and 25% drive heterozygotes (Cas9/Cas9; gRNA/+), equally split between females and males. (D) Heatmaps depicting drive efficacy for two-locus versions of ClvR, TARE, HomeR, and HGD for a range of fitness and transmission rate parameter values, implemented as in panel B. Drive efficacy is defined as the average carrier frequency at generation 20 (approximately 1 year, given a generation period of two to three weeks) based on 100 stochastic simulations with the same initial conditions as C.

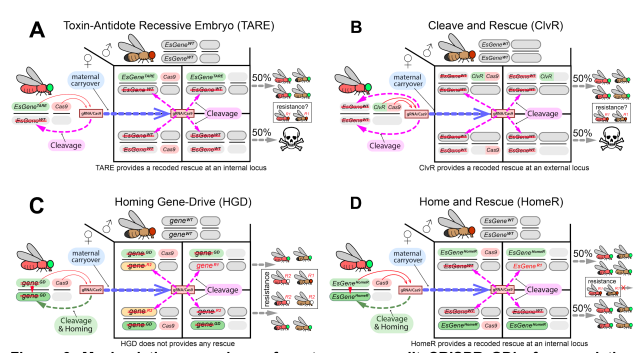


Figure 6. Mechanistic comparison of contemporary split CRISPR GD's for population replacement. Each diagram depicts the cross between females trans-heterozygous for a GD and wildtype (wt) males. Toxin-Antidote Recessive Embryo (TARE) (A) and Cleave and Rescue (ClvR) (B) are non-homing TA-based drives. TARE and ClvR force their inheritance by knocking out an essential gene (EsGeneWT) in oocytes as well as in embryos by the maternal carryover of Cas9/gRNA, and rescuing only those embryos that inherit TARE and ClvR genetic cassettes harboring a re-coded essential gene, which is resistant to Cas9/qRNA-mediated cleavage. As a result, mating between trans-heterozygous TARE and ClvR females and wt males generate 50% unviable embryos. The TARE is integrated at the target gene locus and uses its native promoter to drive expression of the re-coded rescue, hence it is referred to as EsGene^{TARE}. Panel A shows its two-locus version, in which Cas9 is expressed from a separate chromosome. Both components of a two-locus ClvR are inserted at genomic loci separate from the target gene. Panel B depicts the two-locus version of ClvR, in which a ClvR harbors a re-coded rescue with its sequencedistinct promoter and 3'UTR, and both ClvR and Cas9 are inserted into two distant loci. Since both TARE and ClvR use multiple gRNAs to target an essential gene, only very rare functional resistant alleles (R1) can survive. (C) A homing gene-drive (HGD) spreads its inheritance in heterozygous germ cells by cleaving a non-essential gene (gene) and homing, or copying itself, at the cut site (gene^{GD}). Since the knockout of a non-essential gene does not cause lethality and sterility of gene^{GD}/gene^{GD}, HGD spreads itself, though the fitness of gene^{GD}/gene^{GD} is lower than that of gene^{WT}/gene^{GD} or gene^{WT}/gene^{WT}. Maternal carryover of Cas9/gRNA knocks out paternal alleles in embryos, and NHEJ induces large levels of both R2 and R1 (gene^{R2} and gene^{R1}) resistance alleles that survive and eventually block the spread of HGD. (D) Home and Rescue (HomeR) drive as a toxin-antidote homing drive. HomeR harbors a re-coded essential gene, and its precise homing at the cut site rescues the wt function of the essential gene (EsGene^{HomeR}). Maternal carryover of HomeR's Cas9/gRNA induces cleavage of paternal EsGeneWT alleles in embryos, that are rescued by only EsGene^{HomeR} but not EsGene^{R2} maternal alleles resulting in the removal of non-rescued loss-of-function NHEJ alleles (*EsGene*^{R2}) via biallelic lethal mosaicism. HomeR targets an ultra-conserved region of an essential gene for knockout to minimize induction of functional resistance alleles (EsGene^{R1}) that are not fitness costly. Since HomeR is less costly to the female fertility than TARE and ClvR, and induces less resistance alleles that a homing GD.

HomeR outperforms contemporary gene drives (Fig. 5). Red strikethrough defines a LOF (R2) allele.

References

- Adolfi A, Gantz VM, Jasinskiene N, Lee HF, Hwang K. 2020. Efficient population modification gene-drive rescue system in the malaria mosquito Anopheles stephensi. *bioRxiv*.
- Afgan E, Baker D, Batut B, van den Beek M, Bouvier D, Cech M, Chilton J, Clements D, Coraor N, Grüning BA, Guerler A, Hillman-Jackson J, Hiltemann S, Jalili V, Rasche H, Soranzo N, Goecks J, Taylor J, Nekrutenko A, Blankenberg D. 2018. The Galaxy platform for accessible, reproducible and collaborative biomedical analyses: 2018 update. *Nucleic Acids Res* **46**:W537–W544.
- Akbari OS, Bellen HJ, Bier E, Bullock SL, Burt A, Church GM, Cook KR, Duchek P, Edwards OR, Esvelt KM, Gantz VM, Golic KG, Gratz SJ, Harrison MM, Hayes KR, James AA, Kaufman TC, Knoblich J, Malik HS, Matthews KA, O'Connor-Giles KM, Parks AL, Perrimon N, Port F, Russell S, Ueda R, Wildonger J. 2015. BIOSAFETY. Safeguarding gene drive experiments in the laboratory. *Science* **349**:927–929.
- Buchman A, Gamez S, Li M, Antoshechkin I, Lee S-H, Wang S-W, Chen C-H, Klein MJ, Duchemin J-B, Crowe JE, Paradkar PN, Akbari O. 2020. Broad Dengue Neutralization in Mosquitoes Expressing an Engineered Antibody. *SSRN Electronic Journal*. doi:10.2139/ssrn.3398490
- Buchman A, Gamez S, Li M, Antoshechkin I, Li H-H, Wang H-W, Chen C-H, Klein MJ, Duchemin J-B, Paradkar PN, Akbari OS. 2019. Engineered resistance to Zika virus in transgenic Aedes aegypti expressing a polycistronic cluster of synthetic small RNAs. *Proceedings of the National Academy of Sciences*. doi:10.1073/pnas.1810771116
- Carrodeguas JA. 2000. Protein sequences conserved in prokaryotic aminoacyl-tRNA synthetases are important for the activity of the processivity factor of human mitochondrial DNA polymerase. *Nucleic Acids Research*. doi:10.1093/nar/28.5.1237
- Carrodeguas JA, Theis K, Bogenhagen DF, Kisker C. 2001. Crystal Structure and Deletion Analysis Show that the Accessory Subunit of Mammalian DNA Polymerase γ, PolγB, Functions as a Homodimer. *Mol Cell* **7**:43–54.
- Champer J, Buchman A, Akbari OS. 2016. Cheating evolution: engineering gene drives to manipulate the fate of wild populations. *Nat Rev Genet* **17**:146–159.
- Champer J, Chung J, Lee YL, Liu C, Yang E, Wen Z, Clark AG, Messer PW. 2019. Molecular safeguarding of CRISPR gene drive experiments. *Elife* 8. doi:10.7554/eLife.41439
- Champer J, Lee E, Yang E, Liu C, Clark AG, Messer PW. 2020a. A toxin-antidote CRISPR gene drive system for regional population modification. *Nat Commun* **11**:1082.
- Champer J, Reeves R, Oh SY, Liu C, Liu J, Clark AG, Messer PW. 2017. Novel CRISPR/Cas9 gene drive constructs reveal insights into mechanisms of resistance allele formation and drive efficiency in genetically diverse populations. *PLoS Genet* **13**:e1006796.
- Champer J, Yang E, Lee YL, Liu J, Clark AG, Messer PW. 2020b. Resistance is futile: A CRISPR homing gene drive targeting a haplolethal gene. doi:10.1101/651737
- Chan Y-S, Huen DS, Glauert R, Whiteway E, Russell S. 2013. Optimising homing endonuclease gene drive performance in a semi-refractory species: the Drosophila melanogaster experience. *PLoS One* **8**:e54130.
- Chan Y-S, Naujoks DA, Huen DS, Russell S. 2011. Insect population control by homing endonuclease-based gene drive: an evaluation in Drosophila melanogaster. *Genetics* **188**:33–44.
- C. HMS, Sánchez C. HM, Wu SL, Bennett JB, Marshall JM. 2020. MGD riv E: A modular simulation framework for the spread of gene drives through spatially explicit mosquito populations. *Methods in Ecology and Evolution*. doi:10.1111/2041-210x.13318
- Esvelt KM, Smidler AL, Catteruccia F, Church GM. 2014. Concerning RNA-guided gene drives for the alteration of wild populations. *eLife*. doi:10.7554/elife.03401
- Gantz VM, Akbari OS. 2018. Gene editing technologies and applications for insects. Curr Opin

- Insect Sci 28:66-72.
- Gantz VM, Jasinskiene N, Tatarenkova O, Fazekas A, Macias VM, Bier E, James AA. 2015. Highly efficient Cas9-mediated gene drive for population modification of the malaria vector mosquito Anopheles stephensi. *Proc Natl Acad Sci U S A* **112**:E6736–43.
- Gibson DG, Young L, Chuang R-Y, Venter JC, Hutchison CA 3rd, Smith HO. 2009. Enzymatic assembly of DNA molecules up to several hundred kilobases. *Nat Methods* **6**:343–345.
- Groth AC. 2004. Construction of Transgenic Drosophila by Using the Site-Specific Integrase From Phage C31. *Genetics*. doi:10.1534/genetics.166.4.1775
- Guichard A, Haque T, Bobik M, Xu X-RS, Klanseck C, Kushwah RBS, Berni M, Kaduskar B, Gantz VM, Bier E. 2019. Efficient allelic-drive in Drosophila. *Nat Commun* **10**:1640.
- Hammond A, Galizi R, Kyrou K, Simoni A, Siniscalchi C, Katsanos D, Gribble M, Baker D, Marois E, Russell S, Burt A, Windbichler N, Crisanti A, Nolan T. 2016. A CRISPR-Cas9 gene drive system targeting female reproduction in the malaria mosquito vector Anopheles gambiae. *Nat Biotechnol* **34**:78–83.
- Hammond AM, Kyrou K, Bruttini M, North A, Galizi R, Karlsson X, Kranjc N, Carpi FM, D'Aurizio R, Crisanti A, Nolan T. 2017. The creation and selection of mutations resistant to a gene drive over multiple generations in the malaria mosquito. *PLoS Genet* **13**:e1007039.
- Handler AM, Harrell RA 2nd. 1999. Germline transformation of Drosophila melanogaster with the piggyBac transposon vector. *Insect Mol Biol* **8**:449–457.
- Hay B, Jan LY, Jan YN. 1988. A protein component of Drosophila polar granules is encoded by vasa and has extensive sequence similarity to ATP-dependent helicases. *Cell* **55**:577–587.
- Hoermann A, Tapanelli S, Capriotti P, Masters EKG, Habtewold T, Christophides GK, Windbichler N. 2020. Converting endogenous genes of the malaria mosquito into simple non-autonomous gene drives for population replacement. doi:10.1101/2020.05.09.086157
- Isaacs AT, Jasinskiene N, Tretiakov M, Thiery I, Zettor A, Bourgouin C, James AA. 2012. Transgenic Anopheles stephensi coexpressing single-chain antibodies resist Plasmodium falciparum development. *Proceedings of the National Academy of Sciences*. doi:10.1073/pnas.1207738109
- lyengar B, Luo N, Farr CL, Kaguni LS, Campos AR. 2002. The accessory subunit of DNA polymerase γ is essential for mitochondrial DNA maintenance and development in Drosophila melanogaster. *Proc Natl Acad Sci U S A* **99**:4483–4488.
- Kandul NP, Liu J, Buchman A, Gantz VM, Bier E, Akbari OS. 2019a. Assessment of a Split Homing Based Gene Drive for Efficient Knockout of Multiple Genes. *G*3. doi:10.1534/g3.119.400985
- Kandul NP, Liu J, Sanchez C HM, Wu SL, Marshall JM, Akbari OS. 2019b. Transforming insect population control with precision guided sterile males with demonstration in flies. *Nat Commun* **10**:84.
- KaramiNejadRanjbar M, Eckermann KN, Ahmed HMM, Sánchez C HM, Dippel S, Marshall JM, Wimmer EA. 2018. Consequences of resistance evolution in a Cas9-based sex conversion-suppression gene drive for insect pest management. *Proc Natl Acad Sci U S A* **115**:6189–6194.
- Kitzmiller JB. 1976. Genetics, cytogenetics, and evolution of mosquitoes. *Adv Genet* **18**:315–433.
- Kyrou K, Hammond AM, Galizi R, Kranjc N, Burt A, Beaghton AK, Nolan T, Crisanti A. 2018. A CRISPR–Cas9 gene drive targeting doublesex causes complete population suppression in caged Anopheles gambiae mosquitoes. *Nat Biotechnol* **36**:1062–1066.
- Labun K, Montague TG, Gagnon JA, Thyme SB, Valen E. 2016. CHOPCHOP v2: a web tool for the next generation of CRISPR genome engineering. *Nucleic Acids Res* **44**:W272–6.
- Li M, Bui M, Yang T, White B, Akbari O. 2017. Germline Cas9 Expression Yields Highly Efficient Genome Engineering in a Major Worldwide Disease Vector, Aedes aegypti. doi:10.1101/156778

- Li M, Yang T, Kandul NP, Bui M, Gamez S, Raban R, Bennett J, Sánchez C HM, Lanzaro GC, Schmidt H, Lee Y, Marshall JM, Akbari OS. 2020. Development of a confinable gene drive system in the human disease vector Aedes aegypti. *Elife* **9**. doi:10.7554/eLife.51701
- Lin C-C, Potter CJ. 2016. Editing Transgenic DNA Components by Inducible Gene Replacement in Drosophila melanogaster. *Genetics* **203**:1613–1628.
- Marshall JM, Akbari OS. 2018. Can CRISPR-Based Gene Drive Be Confined in the Wild? A Question for Molecular and Population Biology. *ACS Chemical Biology*. doi:10.1021/acschembio.7b00923
- Marshall JM, Raban RR, Kandul NP, Edula JR, León TM, Akbari OS. 2019. Winning the Tug-of-War Between Effector Gene Design and Pathogen Evolution in Vector Population Replacement Strategies. *Front Genet* **10**:1072.
- McVey M, Radut D, Sekelsky JJ. 2004. End-joining repair of double-strand breaks in Drosophila melanogaster is largely DNA ligase IV independent. *Genetics* **168**:2067–2076.
- Michiels F, Gasch A, Kaltschmidt B, Renkawitz-Pohl R. 1989. A 14 bp promoter element directs the testis specificity of the Drosophila beta 2 tubulin gene. *EMBO J* **8**:1559–1565.
- National Academies of Sciences, Engineering, and Medicine, Division on Earth and Life Studies, Board on Life Sciences, Committee on Gene Drive Research in Non-Human Organisms: Recommendations for Responsible Conduct. 2016. Gene Drives on the Horizon: Advancing Science, Navigating Uncertainty, and Aligning Research with Public Values. National Academies Press.
- Noble C, Olejarz J, Esvelt KM, Church GM, Nowak MA. 2017. Evolutionary dynamics of CRISPR gene drives. *Sci Adv* **3**:e1601964.
- Oberhofer G, Ivy T, Hay BA. 2020a. 2-Locus Cleave and Rescue selfish elements harness a recombination rate-dependent generational clock for self limiting gene drive. doi:10.1101/2020.07.09.196253
- Oberhofer G, Ivy T, Hay BA. 2020b. Gene drive and resilience through renewal with next generation Cleave and Rescue selfish genetic elements. doi:10.1101/2019.12.13.876169
- Oberhofer G, Ivy T, Hay BA. 2019. Cleave and Rescue, a novel selfish genetic element and general strategy for gene drive. *Proc Natl Acad Sci U S A* **116**:6250–6259.
- Oberhofer G, Ivy T, Hay BA. 2018. Behavior of homing endonuclease gene drives targeting genes required for viability or female fertility with multiplexed guide RNAs. *Proc Natl Acad Sci U S A* **115**:E9343–E9352.
- Pfeiffer BD, Truman JW, Rubin GM. 2012. Using translational enhancers to increase transgene expression in Drosophila. *Proc Natl Acad Sci U S A* **109**:6626–6631.
- Pham TB, Phong CH, Bennett JB, Hwang K, Jasinskiene N, Parker K, Stillinger D, Marshall JM, Carballar-Lejarazú R, James AA. 2019. Experimental population modification of the malaria vector mosquito, Anopheles stephensi. *PLoS Genet* **15**:e1008440.
- Port F, Chen H-M, Lee T, Bullock SL. 2014. Optimized CRISPR/Cas tools for efficient germline and somatic genome engineering in Drosophila. *Proc Natl Acad Sci U S A* **111**:E2967–76.
- Preston CR, Flores CC, Engels WR. 2006. Differential usage of alternative pathways of double-strand break repair in Drosophila. *Genetics* **172**:1055–1068.
- Sano H, Nakamura A, Kobayashi S. 2002. Identification of a transcriptional regulatory region for germline-specific expression of vasa gene in Drosophila melanogaster. *Mech Dev* **112**:129–139.
- Simoni A, Hammond AM, Beaghton AK, Galizi R, Taxiarchi C, Kyrou K, Meacci D, Gribble M, Morselli G, Burt A, Nolan T, Crisanti A. 2020. A male-biased sex-distorter gene drive for the human malaria vector Anopheles gambiae. *Nature Biotechnology*. doi:10.1038/s41587-020-0508-1
- Van Doren M, Williamson AL, Lehmann R. 1998. Regulation of zygotic gene expression in Drosophila primordial germ cells. *Curr Biol* **8**:243–246.
- Website. n.d. R Core Team (2017). R: A language and environment for statistical computing. R

- Foundation for Statistical Computing, Vienna, Austria. URL https://www.R-project.org/. Windbichler N, Menichelli M, Papathanos PA, Thyme SB, Li H, Ulge UY, Hovde BT, Baker D, Monnat RJ Jr, Burt A, Crisanti A. 2011. A synthetic homing endonuclease-based gene drive system in the human malaria mosquito. *Nature* **473**:212–215.
- Zhang X, Koolhaas WH, Schnorrer F. 2014. A Versatile Two-Step CRISPR- and RMCE-Based Strategy for Efficient Genome Engineering in Drosophila. *G3: Genes|Genomes|Genetics*. doi:10.1534/g3.114.013979
- Zhang X, Koolhaas WH, Schnorrer F. 2014. A versatile two-step CRISPR-and RMCE-based strategy for efficient genome engineering in Drosophila. G3 (Bethesda). 2014; 4: 2409-2418.