Rationally-engineered reproductive barriers using CRISPR & CRISPRa: an evaluation of the synthetic species concept in *Drosophila melanogaster*

Andrew J. Waters¹, Paolo Capriotti³, David C. A. Gaboriau², Philopoulos Aris Papathanos³ & Nikolai Windbichler¹

The ability to erect rationally-engineered reproductive barriers in animal or plant species promises to enable a number of biotechnological applications such as the creation of genetic firewalls, the containment of gene drives or novel population replacement and suppression strategies for genetic control. However, to date no experimental data exist that explores this concept in a multicellular organism. Here we examine the requirements for building artificial reproductive barriers in the metazoan model *Drosophila melanogaster* by combining CRISPR-based genome editing and transcriptional transactivation (CRISPRa) of the same loci. We directed 13 single guide RNAs (sgRNAs) to the promoters of 7 evolutionary conserved genes and used 11 drivers to conduct a misactivation screen. We identify dominant-lethal activators of the *eve* locus and find that they disrupt development by strongly activating *eve* outside its native spatio-temporal context. We employ the same set of sgRNAs to isolate, by genome editing, protective INDELs that render these loci resistant to transactivation without interfering with target gene function. When these sets of genetic components are combined we find that complete synthetic lethality, a prerequisite for most applications, is achievable using this approach. However, our results suggest a steep trade-off between the level and scope of dCas9 expression, the degree of genetic isolation achievable and the resulting impact on fly fitness. The genetic engineering strategy we present here allows the creation of single or multiple reproductive barriers and could be applied to other multicellular organisms such as disease vectors or transgenic organisms of economic importance.

The advent of the CRISPR/Cas9 technology has provided the means to modify genomes with unparalleled specificity and accuracy in a wide variety of model and non-model organisms¹–⁹. These advances in genome engineering, besides enabling a wave of basic research, have also been applied with the aim of distorting inheritance in insect disease vectors. CRISPR gene drives¹,¹⁰ and sex-distorters¹¹ are currently being considered for their enormous potential to control harmful organisms such as insect vectors of human malaria. The CRISPR machinery now broadly employed for gene editing in many organisms, has recently been modified to manipulate the transcriptome; namely targeted gene transactivation and repression. CRISPR gene activation (CRISPRa) uses a nuclease-deficient, deactivated Cas9 (dCas9) protein, fused to a transactivation domain which recruits the basal transcriptional machinery to the site of sgRNA complementarity¹²–¹⁷. This expanded CRISPR toolset now allows for the exploration of more radical genetic engineering concepts one of which is the design of artificial reproductive isolation and the generation of synthetic species. While the terms artificial hybrid incompatibilities or

¹Department of Life Sciences, Sir Alexander Fleming Building, Imperial College London, South Kensington, London, SW7 2AZ, UK. ²Facility for Imaging by Light Microscopy, NHLI, Imperial College London, South Kensington, London, SW7 2AZ, UK. ³Section of Genomics and Genetics, Department of Experimental Medicine, University of Perugia, 06132, Perugia, Italy. Correspondence and requests for materials should be addressed to N.W. (email: nikolai.windbichler@imperial.ac.uk)
artificial reproductive isolation may be more precise (there are many examples of natural hybrid incompatibilities within species), we maintain herein the term synthetic species as it connects to the existing literature and better illustrates the potential use for such a technology. There are multiple potential applications of synthetic species\(^{18}\). Briefly, the prevention of undesired flow of genetic information is a major concern for the field of biotechnology. The introgression of transgenes from Genetically Modified Organisms (GMOs) into the gene-pool of their native counterparts, or the escape of gene drives could be counteracted by this technology. Alternatively, since the synthetic species and its parent species are assumed to be largely identical one can replace the other in a process that is aided by the engineered incompatibility of hybrids. In theory synthetic species released into wild populations would first predominately mate with wild-type individuals, with hybrid incompatibility reducing the number of productive mating events, leading to population suppression of the wild-type. The successive release of additional synthetic species individuals into an increasingly wild-type suppressed population would result in more and more mating between synthetic species individuals, producing viable offspring, eventually leading to population replacement. Synthetic species of insect disease vectors such as Anopheles gambiae could also be further engineered to include a pathogen suppressing transgenic ‘payload’, providing a novel genetic approach to reducing the incidence of malaria. The combination of population suppression and replacement has attractive features in that the release strain requires no special rearing conditions or sterilization. Engineered underdominance, a related approach, works by heterozygotes being less healthy than either wild-type or homozygotes of each transgene. Such systems allowing population replacement by driving of an allele with linked lethal and rescuing elements present at high frequency in a population have recently been demonstrated\(^{13,26}\). Unlike engineered underdominance the synthetic species approach causes no loss of reproductive output due to the segregation of toxin/antidote traits at every generation and can be constructed through non-linked elements, however it also does not allow the introgression of a transgene into an existing population. At an even larger scope, there are possible applications for synthetic species in ecosystem engineering\(^{21}\), e.g. the ability to close and open the reproductive barriers between closely related mating groups and the ability to introduce newly designed species are powerful concepts for environmental management.

Constructing an artificial reproductive barrier requires first the identification of an upstream enhancer or promoter region that allows for lethal, ectopic transactivation of an endogenous gene when targeted by a synthetic transcription factor. It also requires a second modification, the creation of an analogous refractory enhancer region, designed to prevent synthetic transcription factor binding. In this way synthetic lethality is triggered by the transgene in hybrids that result from a cross between modified individuals and those of the naive genetic background. This approach is in principle generalizable and could work in any tractable sexually reproducing organism, because it circumvents the need to research and employ species-specific modes of incompatibility or interfere with endogenous regulatory pathways in order to engineer isolation. This concept has recently been tested in yeast cells using the ACT1 gene\(^{18}\). In this study, the dCas9 Synergistic Activation Mediator (SAM) system was used to over-express Actin, which resulted in loss of cellular integrity when dCas9 strains were mated to wild-type\(^{18}\), although cells that escape synthetic lethality were observed. This work also highlights that the majority of experimental work utilizing dCas9 with a view to application remains cell-based\(^{22-26}\) with its potential in complex systems largely unexplored.

In the present study we combined CRISPR gene editing and transactivation, to explore the design of engineered reproductive barriers in a multicellular organism and to study their properties. Using Drosophila melanogaster as a model, we sought to achieve synthetic lethality in crosses with the wild-type by targeted misexpression of genes known to be essential for fly embryo development (Fig. 1). We reasoned that temporal and spatial perturbations of precisely orchestrated wild-type expression patterns during embryogenesis could result in developmental arrest and lethality. We sought to design sgRNAs targeting genes that function during embryogenesis, and couple these with dCas9 transcriptional activators to trigger embryonic lethality. By coupling these same sgRNA with Cas9 expressed in the germline we sought to isolate protective mutations that suppress activation, and then combine these genetic elements into engineered reproductive barriers.

**Results**

**Design of the sgRNA panel and dCas9 activation strategy.** We first constructed and tested components for ectopic transactivation of target genes in Drosophila, and evaluated the degree of transactivation. We designed a panel of 13 sgRNAs (18–20 bp) targeted to candidate upstream promoter and enhancer regions of 7 developmental genes in the Drosophila genome, namely dpp, engrailed, eve, hairy, hid, rad51 and reaper (Fig. 2A). Most sgRNAs were designed to bind close to the Transcriptional Start Site (TSS) within a window of 150 bp upstream to 48 bp downstream of the TSS, with some sgRNAs targeting known intronic enhancers. We combined these sgRNAs — expressed from U6:3 regulatory regions — with three distinct dCas9 activator domain fusions (Supplementary Fig. 1B): the dCas9-VPR fusion, which has been reported to give robust transactivation in Drosophila cells\(^{14}\) and tissue\(^{15,25}\). The dCas9-P300 Core, a histone acetyltransferase domain, which is capable of activating from promoters and distal enhancers to levels greater than VP64 in HEK cells, but had not yet been tested in Drosophila\(^{26}\). Finally, we also used a 634AA region of the P300 Drosophila orthologue Nejire, which has 77% sequence identity (Supplementary Fig. 1A) with P300 Core, as a C-terminal fusion to Human codon optimized S. pyogenes dCas9 (containing nuclease-inactivating mutations D10A and H840A) (Supplementary Fig. 1B).

**dCas9 transactivation by sgRNAs in the Drosophila eye.** First, we benchmarked the activation potential of our dCas9-VPR sgRNA combinations using the UAS/GAL4 system\(^{27}\) in the Drosophila eye (Fig. 2B, Supplementary Technical Cross 1A), where our target genes do not exhibit a significant level of expression. Testing the entire panel of sgRNAs in combination with dCas9-VPR (driven by GMR-GAL4), we found a 111 fold increase in eve expression, using a sgRNA directed to the eve promoter (eve-sgRNA). The eve gene is a pair
rule patterning factor expressed during embryogenesis and in nerve cells in the fly brain, and it is not expressed in wild-type eyes\(^{28-31}\). By comparison, the sgRNAs for dpp4, dpp5, engrailed, hairy, hid1, hid2, rad51, reaper2 increased the expression of their target genes at more moderate levels (Fig. 2B). The sgRNAs dpp1, dpp2, dpp3 and reaper1 were found to suppress gene expression compared to controls, possibly due to binding competition and steric-hindrance with cis-regulatory factors at the site of transcription (dpp\(^{12}\) and reaper\(^{13,14}\) are both known to play a developmental role in wild-type eyes). We next tested activation strength of dCas9 Histone Acetyltransferase domain fusions in complex with the panel of sgRNAs by qRT-PCR (Fig. 2C). While dCas9-P300 Core yielded no detectable increase in transcripts in the eye, dCas9-Nejire Core triggered activation with all sgRNAs (Fig. 2C), to a stronger degree than dCas9-VPR and also with sgRNAs that did not activate in combination with dCas9-VPR.

The eve-sgRNA gave a 863 fold increase with Nejire Core, a significantly stronger induction than the already pronounced effect of dCas9-VPR. Subtle increases in the expression of developmental genes induced through dCas9-VPR are known to have pronounced developmental phenotypes when expressed in Drosophila tissues\(^{32}\). The eve-sgRNA in complex with dCas9-VPR did induce a moderately aberrant eye phenotype (Fig. 2D), however no pronounced aberrant eye phenotype was observed with dCas9-VPR and the sgRNAs targeting the other genes (Supplementary Fig. 1D). As expected, the combination of dCas9-Nejire Core with the eve-sgRNA resulted in a severe eye-developmental mutant phenotype (Fig. 2D). The remaining sgRNAs assayed for activation potential were crossed to dCas9-VPR in eyes showed abnormal developmental phenotypes (Supplementary Fig. 1C). As expected, dCas9-P300 Core gave no aberrant eye phenotypes (Supplementary Fig. 1E). These results indicate that dCas9-Nejire Core is a powerful tool to over-express genes in flies using sgRNAs. Even those sgRNAs which did not increase gene expression in conjunction with VPR showed developmental phenotypes consistent with over-expression by Nejire Core. For example three sgRNAs were targeted downstream of the TSS in the dpp intron; dpp3-sgRNA and dpp4-sgRNA target 5' of the Dorsal-bound Ventral Repressive Element (VRE) 'S3', dpp5-sgRNA binds 5' of VRE 'S4'. Dorsal binds to VREs in the ventral side of the developing embryo\(^{33}\) and recruits co-factor Groucho\(^{36}\) to repress dpp at the promoter. Nejire Core induces developmental phenotypes when targeted to dpp2, 3, 4-sgRNA binding sites. This is consistent with studies with P300 Core in HEK cells which show that P300 Core is capable of activating transcription from distal enhancers whereas as VP64 cannot\(^{37}\). However, because dCas9-Nejire Core exhibited a subtle but reproducible eye phenotype even in the absence of sgRNAs we decided, in the context of this study, to focus on dCas9-VPR for the remaining experiments.

**Misexpression screen during early fly development using dCas9.** We next tested our sgRNAs against a panel of GAL4 drivers expressing dCas9-VPR, in a screen for developmental arrest/lethality during early stages of Drosophila development (Fig. 3A). The sgRNAs assayed for activation potential were crossed to a panel of GAL4 driver lines that exhibit diverse temporal/spatial, embryonic, larval or ubiquitous expression patterns. GAL4 lines were maintained over dominant balancers, the inheritance of which was scored in the F1. This allowed for the identification of GAL4-sgRNA combinations—in the presence of UAS:dCas9-VPR—giving full or partial lethality during embryogenesis and/or larval development (Supplementary Technical Cross 2A–E, Supplementary Data 1–3). Ubiquitous expression throughout tissues and at all developmental time-points with pCM Tubulin-84b-GAL4 driving dCas9-VPR gave complete embryonic lethality with sgRNAs eve, hid1, hid2 at

![Artificial reproductive barrier](image1.png)

**Figure 1.** Design of artificial reproductive barriers. An artificial barrier combines dCas9 fused to an Activation Domain (AD) and an sgRNA targeting the promoter or enhancer region of a developmental gene within a protected genome. Protection is achieved by an INDEL mutation at the sgRNA binding site (red triangle) that prevents specific dCas9 binding but maintains target gene function. When a transgenic is crossed to wild-type flies the transgene cannot be inherited as synthetic lethality is triggered. Lethality is caused in hybrids when the native target gene is misexpressed ectopically by CRISPR transactivation.
Figure 2. Design and testing of sgRNAs and dCas9 activators. (A) Gene architecture of 7 developmental target genes showing the position and orientation of the sgRNA binding sites and sequence (red) and the Protospacer Adjacent Motif (PAM, blue). Numbers flanking the target sequence indicate the sgRNA-strain. TSS indicated by 90° arrow. Base pair distance to the TSS is indicated (+ upstream, - downstream). For dpp additional sgRNAs 3, 4 and 5 were designed to target close to S3 and S4 Dorsal binding sites (green). The hid intron 1 has been truncated (slashed lines). (B) qRT-PCR analysis of target gene expression levels when the sgRNA panel was tested in a UAS::dCas9-VPR/GMR-GAL4 genetic background (Supplementary Technical Cross 1A). The control indicates eve expression in the absence of sgRNA. Two biological replicates were measured for each condition with the mean shown in red. Error bars show 95% confidence limits, calculated from three technical replicates. (C) Testing of transactivation potential of Histone Acetyltransferase domain dCas9 fusions in eyes (Supplementary Technical Crosses 1B and 1C). qRT-PCR analysis of target gene expression levels in the head using dCas9-P300 Core (mean of 2 biological replicates shown as blue diamonds) and dCas9-Nejire Core (mean of two biological replicates shown as red diamonds) under GMR-GAL4 control in eyes and sgRNAs. The control indicates eve expression in the absence of sgRNA. Error bars as in B. (D) Exemplary eye phenotypes of dCas9 activator fusions expressed from GMR-GAL4 in absence or presence of eve-sgRNA. A severe, aberrant phenotype is seen with eve-sgRNA and dCas9-Nejire Core. Moderate developmental eye phenotypes are seen with dCas9-Nejire Core in the absence of sgRNA and when dCas9-VPR is in complex with eve-sgRNA.
room temperature; dpp2, 3, 4, 5, engrailed, eve, hid1 and hid2 at 25 °C and all sgRNAs except hairy and reaper1 at 29 °C (Fig. 3A). Temperature-dependent increases in lethality were seen with dpp1 and rad51 from 25 °C to 29 °C and dpp2, 3, 4, 5 between room-temperature and 25 °C (and 29 °C). As it is known that GAL4 activity increases with temperature 27, we also performed control crosses to lines lacking sgRNAs and showed that elevated temperatures alone do not increase background lethality (Supplementary Data 3). This suggests that the activation potential of certain sgRNAs may be limited by the level of dCas9-VPR, even when dCas9-VPR is ubiquitously expressed. It is likely that increased genome binding events would lead to increased lethality, rather than increased activity of the VPR domain. This is suggested by the observation that dpp1, 2, 3 and reaper1 show temperature dependent increases in lethality (Fig. 3B) yet were found to reduce target gene expression in the eye (Fig. 2B). The eve-sgRNA was found to give complete lethality in all driver conditions except in combination with 3.1isp2, pannier and spalt. This correlates with the strong eve over-expression exhibited by eve-sgRNA when measured in eyes (Fig. 2B).

Analysis of eve misexpression during embryo development. To link this set of observations we analyzed eve mRNA and protein expression in the context of the developing embryo (Supplementary Technical Cross 3). We first quantified embryonic eve and hid over-expression by qRT-PCR (Fig. 3C) and found that, the presence of eve-sgRNA results in an 80 fold increase and hid1-sgRNA a 2.75 fold increase in early development, levels that correlate well with the expression in the eye (Fig. 2B). In the presence of eve-sgRNA, antibody staining...
showed pervasive Eve over-expression in the early embryo (stage 6–7) outside its native spatiotemporal range as a pair-rule regulator (Fig. 4B). In later stage wild-type embryos (stage 14–15) Eve protein is found in the posterior (anal pad) region and in the mesoderm along with a subset of neuronal nuclei. Lines which express the eve-sgRNA with dCas9-VPR also exhibit ectopic Eve expression throughout the embryo at this stage (Fig. 4F). At this later stage Eve misexpression is associated with a developmental delay and signs of disorganization of the embryos (Fig. 4J), which subsequently fail to hatch.

**Generation and analysis of protective INDELs using Cas9.** In parallel we performed CRISPR/Cas9 mutagenesis to identify INDELs in the targeted gene regions that would abolish sgRNA binding at the target sites, but which would be tolerated in the context of target gene function (Fig. 5A). One sgRNA per gene from the panel of sgRNA flies were crossed to pvasa::Cas9 expressing lines. Progeny were scored to appropriate balancer lines and screened for INDEL mutations at the sgRNA binding site by PCR (Supplementary Technical Cross 3A, B). F2 progeny were then tested for homozygote mutant viability, fitness and fertility (Fig. 5B and Supplementary Table 1). We found that all sgRNAs tested resulted in the production of INDEL mutations. This confirms that the
lack of significant transactivation observed in certain conditions with dCas9-VPR can’t be attributed to sgRNA
function, a point further consolidated by the potential of dCas9-Nejire Core to activate with all sgRNAs tested.
We found that mutations dpp2-1, dpp2-4 and eve-16 resulted in a significant reduction in fitness and the com-
plete loss of female fertility. Only a single mutation reaper1–9 was found to cause homozygous lethality. All other
INDEL mutations were viable as homozygotes with no obvious fitness cost observed. Although our sgRNA panel
was designed to avoid known regulatory elements within the target loci, it is possible that dpp2-1, dpp2-4 and eve-
16 mutations disrupt cis-regulatory sequences needed for germline development in female flies and an essential
developmental process in the case of reaper1–9.

Combining transactivators with protective INDELS. Next we incorporated lethal components iden-
tified in the CRISPRa screen into single expression plasmids containing dCas9-VPR under the direct control
of four different promoters: ptwist (TW), phow (H), poTubulin-84b-Long (LT) and poTubulin-84b-Short (ST)
(Supplementary Fig. 2). The plasmids also contained the eve-sgRNA expressed ubiquitously under the control of the
U6::3 regulatory regions. These constructs therefore target a single locus, in this case eve and are designed to
form a single barrier (SB) to hybridization. Embryos homozygous for the eveΔ11 mutation, which had no impact

Figure 5. CRISPR/Cas9 mutagenesis screen to isolate protective INDELS. (A) Experimental scheme detailing
mutation screen. Female virgin flies expressing Cas9 in the germ-line under the vas promoter were crossed
to 7 sgRNA lines to induce INDEL mutations within the target gene promoter regions. INDEL lines were
assessed for viability as homozygous mutants (Supplementary Technical Cross 3A,B). (B) INDEL lines obtained
with the fitness, viability and fertility of homozygous mutant flies indicated (green indicates no observable
negative effect in homozygotes). Fitness measure derived from Chi-Squared test (see Supplementary Table 1).
Wild-type sgRNA binding site sequences are shown above show INDEL mutants for each sgRNA tested. Some
independent isolated mutant lines screened were found to contain identical lesions; this is the case for dpp2-
3/5, engrailed-1/2/3, engrailed-5/17/99, eve-2/3/11, eve-13/14 and hairy-3/10. PAM sequence is shown in blue,
sgRNA binding site shown in red, deletions by black hyphen, insertions shown in orange.
on gene function (Fig. 5B and Supplementary Table 1) were injected with the expectation that this mutation would render them resistant to the effects of the lethal components. Constructs were either integrated at genomic AttP docking sites known to exhibit high expression from integrated transgenes 37–39, or by using P-element random integration and positive transformants were back-crossed to obtain, if possible, homozygous inserts. Using the Mini-White dominant marker we scored for synthetic lethality by crossing to white-eyed w1118 flies with an unmodified eve target locus (which we refer to as wild-type in this context). Figure 6A summarizes the outcome of these experiments (Fig. 6).

Table: Synthetic lethality

| Transgenic Strain | sgRNA Target | Integration | Homozygous | Progeny (transgenic / total) | Synthetic Lethality (%) |
|-------------------|--------------|-------------|------------|-----------------------------|------------------------|
| SB-H   | eve          | Yes         | wild-type  | 114/228                     | 0.0                    |
| SB-LT  | eve          | Yes         | eve Δ11    | 46/93                       | 1.1                    |
| SB-LT  | eve          | No          | wild-type  | 2/955                       | 100.0                  |
| SB-ST  | eve          | Yes         | wild-type  | 242/487                     | 1.4                    |
| SB-ST  | Eve          | Yes         | eve Δ11    | 70/295                      | 52.5                   |
| SB-ST  | Eve          | Yes         | eve Δ11    | 53/105                      | 0.0                    |
| SB-TW  | Eve          | Yes         | wild-type  | 63/222                      | 43.2                   |
| SB-TW  | Eve          | Yes         | eve Δ11    | 77/256                      | 1.3                    |
| SB-LT 1-4 | eve         | No          | wild-type  | 50/356                      | 71.9                   |
| SB-LT 1-4 | eve         | No          | eve Δ11    | 72/246                      | 3.4                    |
| SB-LT 1-2 | Eve         | No          | wild-type  | 2/348                       | 97.3                   |
| SB-LT 1-2 | Eve         | No          | eve Δ11    | 77/155                      | 0.6                    |
| SB-LT 1-3 | Eve         | No          | wild-type  | 40/211                      | 63.8                   |
| SB-LT 1-3 | Eve         | No          | eve Δ11    | 92/187                      | 1.6                    |
| SB-LT 1-1 | Eve         | No          | wild-type  | 9/310                       | 94.2                   |
| SB-LT 2-1 | Eve         | No          | wild-type  | 44/90                       | 2.2                    |
| SB-LT 2-1 | Eve         | No          | wild-type  | 138/233                     | 0.4                    |
| SB-LT 2-1 | Eve         | No          | eve Δ31    | 64/130                      | 1.5                    |
| SB-LT 2-1 | Eve         | No          | wild-type  | 138/230                     | 0.0                    |
| SB-LT 2-2 | Eve         | No          | eve Δ11    | 64/126                      | 0.0                    |
| SB-LT 2-1 | Eve         | No          | wild-type  | 74/296                      | 50.0                   |
| SB-LT 2-1 | Eve         | No          | wild-type  | 57/116                      | 1.7                    |
| SB-LT 2-1 | Eve         | No          | wild-type  | 112/225                     | 0.4                    |
| SB-LT 2-1 | Eve         | No          | wild-type  | 62/122                      | 0.0                    |
| DB-LT 1-1 | Eve         | Yes         | wild-type  | 78/310                      | 49.7                   |
| DB-LT 1-1 | Eve         | Yes         | eve Δ11    | 125/273                     | 8.5                    |
| DB-LT 1-1 | Eve         | Yes         | eve Δ11 Δ13 | 92/238                     | 22.7                   |
| DB-LT 1-1 | Eve         | Yes         | eve Δ11Δ13 Δ13 | 103/296              | 2.0                    |
| DB-LT 1-1 | Eve         | Yes         | wild-type  | 93/383                      | 3.6                    |
| DB-LT 1-1 | Eve         | Yes         | wild-type  | 95/204                      | 6.9                    |
| DB-LT 1-1 | Eve         | Yes         | wild-type  | 114/237                     | 3.8                    |
| DB-LT 1-1 | Eve         | Yes         | wild-type  | 122/277                     | 10.1                   |

Figure 6. Evaluating single construct synthetic lethality within protected genomes. (A) Single vectors containing either one (eve) or two (eve & hid1) sgRNA transgenes (for erecting a single barrier or a double barrier) as well as dCas9-VPR driven by different promoters were integrated into the Drosophila melanogaster genome either at AttP9-A (VK00027) on chromosome 3 or by random P-element integration. The target genetic background is homozygous for eveΔ11 in the case of a single barrier or double-homozygous for eveΔ11;hid1-Δ13 in the case of the double barrier experiments. The Mini-White selective marker (not shown) was used to select T0 transgenics in all cases. T1 transgenic lines were tested for associated lethality by scoring for associated red eye marker presence in F1 progeny in crosses to ‘wild-type’ (w1118) virgin females. Shown are the counts of transgenic/total flies from which we calculated the percentage of synthetic lethality of the transgene. 100% indicates the transgene was not inherited at all, 0% indicates that the transgene was inherited at a ratio consistent with no associated lethality. Single barrier (SB), Double barrier using tandem sgRNAs (DB1), Double barrier using sgRNA with a tRNA link (DB2). Promoters used: phow (-H), pαTubulin-84b-Long (-LT), pαTubulin-84b-Short (-ST), pwist (-TW). (B) qRT-PCR measuring eve expression in embryos from crosses of two P-element lines to w1118 (wild-type) and eveΔ11. Two biological replicates were measured for each condition (black bars) and the mean is shown in red. Error bars show 95% confidence limits, calculated from three technical replicates (See Supplementary Technical Cross 6).
of these experiments and the genetic crossing strategy is detailed in Supplementary Technical Cross 3A–C. Transgenic strains with dCas9-VPR driven by \textit{pc1Tubulin-84b-Long} when crossed to wild-type display the full range of possible phenotypes ranging from no observable effect on viability to complete synthetic lethality in the case of line SB-LT. Synthetic lethality of line SB-LT is completely rescued in crosses to homozygous eve\_Δ11 mutants. Furthermore the various degrees of lethality associated with SB-LT P-element strains are rescued when strains are crossed to eve\_Δ11 mutants (Fig. 6A). Besides lethality, no other developmental phenotypes were observed in lines with partial lethality.

SB-LT P-element strains were confirmed to be integrated at diverse loci through Inverse PCR (Supplementary Table 3). In order to test for a correlation between degree of lethality and degree of eve over-expression in P-element lines we analysed eve over-expression in embryos from tester crosses in two P-element lines with differing % lethality (Fig. 6B, Supplementary Technical Cross 6). Line SB-LT-8-1 and SB-LT-4-2 show 50% and 97.3% lethality respectively when crossed to wild-type tester stocks (Fig. 6A), with crosses to eve\_Δ11 rescuing lethality (0.6% for SB-LT-8-1 and 1.7% for SB-LT-4-2). Through qRT-PCR on F1 embryos we found SB-LT-8-1 to induce a 6.4 fold increase in eve expression and SB-LT-4-2 to induce a 14 fold increase (Fig. 6B), this is consistent with higher lethality seen in line SB-LT-4-2. It is notable that the degree of over-expression for eve in embryos is much less with SB-LT strains than seen when using UAS::dCas9-VPR::\textit{Tubulin-84b-GAL4} and eve-sgRNA (CF, Figs 3C and 6B), likely due to the absence of UAS/GAL4 amplification loopings of dCas9-VPR transcription in the SB-LT vector. Similar degrees of lethality and rescue were seen in reciprocal crosses between female P-element transgenic strains and male tester stocks (See Supplementary Table 2), showing that isolation elements are functional when either male or female synthetic strain individuals are used.

The SB-TW line conferred ~43% lethality whereas the SB-H line was found to confer no significant lethality. We found that few \textit{pTubulin-84b-Long} strains were lethal or parthenogeny viable. This is likely because of a significant fitness costs associated with ubiquitous dCas9-VPR tissue expression. For other lines such as SB-H, SB-TW and SB-ST homozygous expression of the activator was possible, but they induced more moderate levels of genetic isolation. In order to determine fitness effects of transgene expression in homozygous viable lines we assessed % survival of embryos progressing to larval stages of development in the lines SH-TW and SB-ST and also in stocks w1118 and eve\_Δ11. We found that background lethality in w1118 and eve\_Δ11 was similar with slightly higher background lethality seen with eve\_Δ11, suggesting there may be some subtle fitness cost with the eve\_Δ11 INDEL (Supplementary Table 4), although segregation analysis and a Chi-Squared test suggests that this does not significantly affect heritability of the lesion (Supplementary Table 1). SB-TW and SB-ST exhibit slightly higher % lethality than eve\_Δ11 background, suggesting that there are some subtle costs of transgene expression in the lines. SB-LT shows appreciably higher background % lethality than SB-TW consistent with ubiquitous expression of dCas9-VPR from a Tubulin promoter negatively effecting fitness (Supplementary Table 4).

Our genetic strategy also allows for the stacking of genetic barriers. To attempt this we chose to combine eve and hid1-sgRNAs into a single vector. We combined mutants eve\_Δ11 with hid1\_Δ13 into a genetic background with double-refractory promoter regions in homozygosity. We then inserted into this background — using P-element integration — a vector expressing dCas9-VPR from \textit{pTubulin-84b-Long} along with eve and hid1-sgRNAs expressed from \textit{U6::3}. We attempted this with \textit{U6::3} regulatory regions flanking each sgRNA in tandem (DB1 constructs) and also with a tRNA separating sgrRNAs to allow for post-transcriptional cleavage of sgrRNAs (DB2 constructs) (Fig. 6, Supplementary Fig. 2). We observed 50% lethality with the tandem DB1 construct when crossed as a heterozygote to a ‘wild-type’ background. We then performed separate crosses to the eve\_Δ11 and also hid1\_Δ13 homozygous backgrounds as well as the double-homozygous eve\_Δ11 & hid1\_Δ13 stock and confirmed that synthetic lethality induced by both sgRNAs (Fig. 6), albeit moderate overall, is synergistic.

**Discussion**

The concept of artificial reproductive isolation or “synthetic species” has been investigated previously in \textit{Drosophila}. However in the pre-CRISPR era, it required the exploitation of complex fly genetics and knowledge of unique biological aspects of the \textit{Drosophila glass} gene. This strategy therefore could not easily be applied to other species or even other loci. One of the principle attractions of the approach we present here, is that it achieves synthetic lethality through minimal modifications of evolutionary conserved loci using the expanded CRISPR toolset that is now available in a range of organisms. This strategy should facilitate the transfer of this technology to diverse organisms and the rational design of genetic isolation. Recently, a similar strategy was tested in a proof-of-principle in the lower unicellular eukaryote \textit{S. cerevisiae}; yeast however requires comparatively fewer and less complex genetic engineering steps than higher eukaryotes.

Our survey of protective mutations and lethal CRISPRa elements suggests that protecting genomes from transactivation can be achieved at most loci, as INDELS rarely seem to affect nearby cis-elements in the promoter or enhancer regions. Isolating a range of INDELS at a target site is straightforward and allows the selection of those that have no negative effects. Achieving synthetic lethality through transactivation by CRISPRa was more challenging, and whilst all sgrRNAs used for gene editing yielded INDEL mutations, not all drive strong gene activation with dCas9-VPR. In this study we directed single sgrRNAs to enhance regions and found in most cases a less than 2-fold increase in mRNA levels with one notable expression being the eve-sgRNA which to our knowledge leads to the highest increase in gene activation for a single sgRNA tested in vivo. This highlights that whilst there are some parameters for the design of effective sgrRNAs for DNA cleavage, knowledge of what makes a sgRNA particularly effective at transactivation with widely used activators such as dCas9-VPR, is still lacking. It has been reported that targeting sgrRNAs within 400 bp upstream of the transcriptional start site is effective in some contexts, but clearly such positioning alone is insufficient to guarantee strong activation. It has been shown that when multiple sgrRNAs are targeted to a promoter the probability of achieving biologically relevant transactivation is increased. However, it has also been shown that when multiple sgrRNAs are targeted to a promoter, the effect on transactivation is not synergistic, and can be mostly attributed to a single highly
active sgRNA. On the other hand it has also been demonstrated that a significant number of sgRNA pairs that target within this window generate observable phenotypes with dCas9-VPR despite relatively modest increases in gene product in some cases. Consistent with this, we observed lethality with most sgRNAs when dCas9-VPR is expressed by a ubiquitous GAL4 driver. It is likely, that assaying mRNA expression levels on a tissue aggregate level may overlook more pronounced transactivation in particularly receptive cell types. For example, it has been reported that chromatin accessibility may prevent the binding of Cas9 and the binding of synthetic transcription factors at transcriptionally inactive loci. Therefore targeting genes for transactivation in tissues which show low basal levels of expression may be a more effective approach to obtaining sgRNAs capable of strong activation with VPR. It also suggests a role for directed chromatin modifiers which could yield activation for some loci that are refractory to transactivation by more traditional activation domains such as VPR. We found that all sgRNAs tested with dCas9-Nejire Core led to over-expression of targets and morphological phenotypes when the trans- activators were expressed in the eye, which was not the case with dCas9-VPR. As expression of dCas9-Nejire Core had subtle developmental phenotypes in the absence of sgRNAs it likely triggered non-specific effects that are too strong to allow it to be used in the context of generating reproductive barriers at present. However, for other applications, and possibly with further modifications to modulate its strength, it could become a powerful building block for the in vivo CRISPRa toolbox.

We tested the synthetic lethality of single-vector CRISPRa constructs established in a genetic background homozygous for protective mutations. To obtain lines harboring elements which could not be inherited in crosses to wild-type required a ubiquitous expression pattern of dCas9-VPR using repo-tubulin. However, ubiquitous expression by repo-tubulin when associated with strong synthetic lethality also precluded the generation of viable homozygotes and thus full genetic isolation of the strain. We achieved a medium level of synthetic lethality in combination with homozygous viability with ptwist driven dCas9-VPR. In our misexpression screen ptwist had previously triggered complete lethality with eve-sgRNA, and it is likely that the lack of the GAL4 amplification loop accounts for this difference. As expected, we also detected varying levels of synthetic lethality with identical constructs depending on the insertion site, suggesting that the genetic context may modulate CRISPRa penetrance.

Whilst we didn’t explore the use of multiple stacked genetic barriers in great detail beyond demonstrating a proof-of-principle, their use will likely be a necessity for any realistic application of engineered reproductive isolation. Genetic diversity in large wild-type populations means that circulating rare variants at the sgRNA target sites could abrogate synthetic lethality and lead to the generation of escapers and the eventual breakdown of genetic isolation. By combining two sgRNA activators with their corresponding protective mutations, genetic isolation could be fortified to safeguard against failure through rare recombination, random mutations within the constructs or transgene silencing events. As a proof of principle we demonstrate that using eve and hid1-sgRNAs allows for stacking of single transactivation elements to induce synergistic effects on synthetic lethality.

We suggest that fine tuning the approach presented here will likely require the refinement of the expression and strength of transactivators, to increase lethality and at the same time reduce fitness costs associated with unacceptably broad expression of CRISPRa transgenic elements. The testing of early tissue-specific promoters with GAL4-UAS amplification to drive transactivator expression from a range of known AtTP integration sites with varying expressivity capacity presents itself as a logical next step to fine tune the system. Real-world robustness of genetic barriers could then be achieved by combining two separately tuned barriers. Application of the system in non-model organisms with fewer transgenic tools than that of Drosophila may complicate a fine-tuning approach. An approach to mitigate limitations in such organisms may be to attempt multiple combinatorial stacking of less finely tuned barriers targeting diverse loci.

In summary we have generated protective genomes and benchmarked sgRNA targets and activator domains for use in the construction of synthetic species-barriers. With modulation of the expression strength of synthetic lethal elements, complete genetic isolation in the absence of a significant effect on fitness should be achievable in the Drosophila model. More importantly the candidate genes that we have analyzed here have orthologs in other insect species which should greatly aid the transition of this approach to medically or agriculturally relevant insects.

Materials and Methods
sgRNA design and cloning. The flyCRISPR Target Finder tool (http://tools.flycrispr.molbio.wisc.edu/ta getFinder/) was used to identify 18-20nt sgRNA binding sites in genomic regions of interest. DNA sequences were obtained from FlyBase. All sgRNA candidates which overlapped with known transcription factor binding sites (annotated by the modenCODE and RedFly projects) were disregarded to avoid interfering with endogenous transcriptional control. sgRNAs were cloned into the pCFD3 vector, under the control of U6::3 regulatory regions (as reported by Port et al.). pCFD3 vectors were incorporated at AtTP-9A sites (VK00027) on the third chromosome.

GAL4 transactivation experiments. The how-GAL4 and GMR-GAL4 UAS::dCas9-VPR driver line was created by crossing UAS::dCas9-VPR lines (integrated into attP40 (BL25709) flies) with BL#1767 (how) and BL#8121 (GMR) using the double balancer line BL#33821. Other UAS::dCas9-VPR GAL4 drivers were a gift from Norbert Perrimon. All lines were maintained over balancers. The aTubulin-84b-GAL4 line is maintained over a fused 2nd and 3rd chromosome balancer UAS::dCas9-VPR; aTubulin-84b-GAL4/SM5=TM6b. Lines homozygous for pCFD3 integration were crossed to balanced UAS::dCas9-VPR; driver-GAL4 lines and F1 adult progeny analyzed by observing Curly and Humeral dominant markers on CyO and TM6b balancers. The genotype of the female activator parent used in crosses differed between lines, the genotype of the female parent used in each cross is listed in Supplementary Data 1. Technical Cross 2 A-E lists the crossing strategy for each possible female parent genotype used in the CRISPRa screen. F1 adult flies were scored for segregating marker phenotypes; to
removed and plates incubated at 25 °C for 3 hours or 12 hours to collect stage 6–7 (early embryogenesis) and stage 14–15 (late embryogenesis) embryos respectively. Embryos were washed in ddH₂O, dechorionated in 1 hr on fresh plates. Plates were replaced and flies allowed to deposit embryos for 1 hr. Plates were removed and allowed to age for 3 hours at 25 °C. 50 embryos were collected with a paintbrush, washed in ddH₂O and frozen.

Generation of INDEL mutations. To induce INDELS at sgRNA target sites in the genome (see Supplementary Technical Cross 3A,B), sgRNA lines were crossed as males to pvasa::Cas9 virgin females (BL#51324) to give F1 flies expressing Cas9 and sgRNA in the germ cells. F1 adult females were crossed to appropriate balancer lines to maintain putative INDELS. F2 males were back-crossed as individuals to the balancer line, after successful mating determined by egg-laying the founder F2 parent fly was macerated and DNA extracted by Phenol-Chloroform purification. PCR amplification of the genomic region of interest (~1 kb region around sgRNA site) was performed using loci-specific primers (See Supplementary Primers). PCR amplicons were incubated with T7 endonuclease I (NEB) to identify heteroduplex PCR products indicative of INDEL containing amplicons. PCR amplicons from T7 positive samples were sequenced and heterogeneous sample reads confirmed from the point of DNA mismatch (and INDEL induction). Mutations induced on the 3rd Chromosome required additional screening in order to remove the pvasa::Cas9 construct which is marked by PAX::GFP. This required recombination in the female germ-line (See Supplementary Technical Cross 4B). Where possible, flies homozygous for INDELS were selected in the F3 generation and the nature of the lesion confirmed by additional sequencing. The viability of INDEL lines was confirmed through the scoring of dominant balancer markers in the F3 generation. In an inter-sibling cross between two heterozygous balanced individuals, each chromosome (INDEL and balancer) would have an inheritance ratio of roughly 1:2 (Homozygous:Balancer) if no fitness cost was associated with the INDEL. If complete non-absence of balanced was observed then homozygous lethality was assumed. Progeny were counted from individual crosses between two balanced individuals and the Chi-squared test used to ascertain if the incidence of non-balanced F1 individuals were significantly less than the expected ratio of 1:2 (See Supplementary Table 1). For lines with reduced fitness, homozygous INDEL flies were crossed to balancers in a reciprocal manner to assess male/female sterility.

RNA Expression analysis. The transactivation potential of sgRNAs and activator domains were assessed in the fly eye using the driver lines UAS::dCas9-VPR;GMR-GAL4, UAS::dCas9-P300 Core; GMR-GAL4 and UAS::Cas9-Nejire Core;GMR-GAL4. sgRNA expressing males were crossed to driver line virgin females, crosses were incubated at 29 °C for maximal GAL4 induction (See Supplementary Technical Cross 1A–C). The heads of 30 F1 progeny were collected and flash-frozen in liquid nitrogen and RNA extracted. eve and hid sgRNAs were also assessed for transactivation potential in the embryo: Virgin female UAS::dCas9-VPR; α Tubulin-84b-GAL4/SM5 = TM6b flies were crossed to each sgRNA in collection cages topped with yeast-smeared, apple juice plates and incubated for 3 days at 25 °C (See Supplementary Technical Cross 3A). Flies were allowed to pre-lay for 1 hr on fresh plates. Plates were replaced and flies allowed to deposit embryos for 1 hr. Plates were removed and allowed to age for 3 hours at 25 °C. 50 embryos were collected with a paintbrush, washed in ddH₂O and frozen in liquid nitrogen. In all cases total RNA was extracted in 500µl of Trizol Reagent and aqueous layers purified with Qiagen Micro-RNAEasy kits. Equal amounts of RNA were used to make cDNA from control and activation crosses using Qiagen QuantiTect cDNA synthesis kit as per manufacturer's instruction. 1/10 diluted cDNA was assayed in SYBR green qRT-PCR (Applied Biosystems) mix with gene specific primers and normalized using the comparative ΔΔCt method against actin expression. Where available, previously confirmed (DRSC FlyPrimerBank)/intron-spanning primers were used (See Supplementary Primers for gene specific primers used). Samples were assessed using biological duplicates and technical triplicates. Error bars were calculated using Applied Biosystems StepOne software and represent variance across technical replicates using the same cDNA template to a 95% confidence level (i.e. there is a 95% confidence that actual measured expression level is represented within the error range).

Protein expression analysis. Virgin female UAS::dCas9-VPR; α Tubulin-84b-GAL4/SM5 = TM6b flies were mated to male eve-sgRNA expressing males (See Supplementary Technical Crosses 3A). Adults were removed and plates incubated at 25 °C for 3 hours or 12 hours to collect stage 6–7 (early embryogenesis) and stage 14–15 (late embryogenesis) embryos respectively. Embryos were washed in ddH₂O dechorionated in 50% bleach and fixed for 20 minutes in 4% paraformaldehyde following standard protocols. Fixed embryos were incubated overnight at 4 °C with primary antibodies for Eve (3C10-DSHB, from mouse), Thermofisher). Embryos were mounted in 50% glycerol containing DAPI counterstain. Images were acquired on a Zeiss LSM 510 inverted confocal microscope, using a Plan-Apochromat 20× 0.8 NA objective, with a x,y pixel size of 0.88 µm and a z-step of 1.0 µm for Fig. 4A,B, and x,y pixel size of 0.83 µm and a z-step of 1.8 µm for Fig. 4C–J. Excitation and emission for DAPI, Eve and FLAG were 405 nm (BP 450–480 nm), 488 nm (BP 505–550 nm) and 543 nm (LP 560 nm), respectively. Acquisition and display parameters were kept identical for all samples. Data shown are maximum intensity projections, prepared with FIJI.

Construction of dCas9-P300 Core and dCas9-Nejire Core. UAS::dCas9-VPR in pWalium20 (Gift from Norbert Perrimon) was digested with BstAPI and EcoRI to remove the VPR domain. A C terminal section of dCas9 is also removed by this digestion. Gibson Assembly primers were used to amplify the fragment of dCas9 removed by cleavage at the EcoRI site by PCR using pWalium20 as a template (Primers ‘GIB–Nejire Frag 1 F + R’ and ‘GIB–P300 Frag 1 F + R; Supplementary Primers). Gibson Assembly primers (‘GIB–P300 Frag 2 F + R’) were used to amplify P300 Core from Addgene vector 61357 as template. Gibson Primers (‘GIB–Nejire Frag 2 F + R’).
were used to amplify the identified Nejire Core region from a *Drosophila melanogaster* whole body cDNA library of w1118 flies. Digested and gel-purified *pWalium20* was incubated with appropriate Gibson Assembly PCR products at a molar ratio of 1:3 (Vector:Insert) using 50 ng of vector DNA. Reactions were performed using the Gibson Assembly Cloning Kit (NEB). See Supplementary Primers for sequence information.

**Construction of transactivation vectors.** A combination of restriction enzyme digestion, PCR, ligation, Gibson Assembly and Gateway recombination were used to construct transactivation vectors. UAS::dCas9-VPR in *pWalium20* was digested with *NheI* and *Spel* to remove the 10xUAS sequence, the backbone was gel purified and blunt-ended with dNTPs and T4 DNA polymerase (NEB). The blunt backbone was ligated to Gateway Destination vector conversion fragment reading frame B (Thermo-Fisher) to replace UAS with an AttR1-AttR2 *ccdB* containing cassette. *pWalium20* contains an AttB sequence for integration at AttP docking-sites in *Drosophila*. To enable P-element integration at various genomic locations 5′ and 3′ P-element termini along with Ori and Ampicillin Resistance regions were amplified from *pGMR-mKate* using Gibson Assembly primers (‘P-ELEMENT-GIB F + R’). *AttR1-AttR2::dCas9-VPR* in *pWalium20* was digested with *ApaI* and *SapI* and the *dCas9-VPR* containing fragment gel purified and assembled with the Gibson ampiclon from *pGMR-mKate* to create *pAJW1* (See Supplementary Fig. 2). Assembled plasmids were propagated in One Shot *ccdB* Survival 2 T1R competent cells (Thermo-Fisher). Gibson primers (‘*EVE*-gRNA-GIB F + R’) were used to amplify eve-sgRNA sequence flanked by U6:3 regulatory regions (pU6-3::*eve*-gRNA-U6-3′-3′UTR) from *pCFD3*-eve plasmid (made for use in the transactivation lethality screen); *pAJW1* was digested with *SapI*—which cleaves once 3′ of the SV40 terminator 3′ of *dCas9-VPR*—and gel-purified, the Gibson ampiclon containing eve-sgRNA was ligated to the backbone to create plasmid *pAJW2* (in summary, *pAJW2 = [5′-ATTB-AttR1-AttR2::dCas9-VPR-SV4 0-pU6-3::*eve*-gRNA-3′-3′UTR]*). In order to convert *pAJW2* into an expression vector promoter regions were inserted into the Gateway R2 site (at the AttR1-R2 site). This was achieved by cloning promoter regions, PCR amplification from w1118 fly genomic DNA (except *ptubulin-84b-Long* which was amplified from *pCasper-tubulin-GAL80*, Addgene#24352), into *d-TOPO/pENTR* plasmids and performing subsequent Gateway LR-reactions with *pAJW2* as the destination vector. Primers used to amplify genomic regions are listed in Supplementary Primers. This results in a vector with *dCas9-VPR* expression driven by the inserted promoter. The following promoter regions were used to create expression plasmids with *pAJW2: ptubulin-84b-Long* (giving SB-LT), *ptubulin-84b-Short* (giving SB-ST), *ptwist* (giving SB-TW) and *phow* (giving SB-H). Correct construction was confirmed by Sanger sequencing throughout. Plasmid DNA was prepared for microinjection using the Qiagen Maxiprep kit. Plasmids were injected into a w1118 (white eyed) line homozygous for eveΔ11 on 2nd chromosome and homozygous for the AttP-9A site VK00027 on the 3rd Chromosome with *Φc31-Integrase* expressed from the X chromosome (AttP site and integrase were derived from Bloomington stock #35569). Injections were performed by The University of Cambridge Fly Injection Facility. Transgenic individuals were selected based on *Mini-White* marker selection and inter-sibling crosses performed to create stocks. In addition the *ptubulin-84b-Long* expression vector was integrated into w1118 eveΔ11 homozygous flies using P-element integration methodology, positive transgenic individuals were selected by *Mini-White* selection (See Fig. 6).

**Construction of DB1 and DB2 vectors: Gibson assembly compatible primers (‘DB1-GIB F + R’) were used to amplify hid1-sgRNA and flanking U6:3 regulatory sequences from hid1 in *pCFD3* (cloned for transactivation lethality screen). This ampiclon was then ligated to *SapI* digested *pAJW2* plasmid. This creates a tandem sgRNA region [pU6-3::hid1-sgRNA-U6-3′-pU6-3::*eve*-gRNA-U6-3] this vector *pAJW3* was recombined in a Gateway reaction with *pENTR: ptubulin-84b-Long* and *pENTR: ptubulin-84b-Short* to create vector DB1-LT and DB1-ST respectively. DB2-LT was created by cloning hid1-sgRNA and eve-sgRNA sequences into *pCFD5* (as previously described by Simon Bullock) which has one set of U6:3 regulatory sequences with sgRNAs separated by a tRNA sequence. Gibson Assembly primers (‘DB2-gRNA F + R’) were used to clone sgRNA sequences in BbsI digested *pCFD5*. Primers ‘DB2-GIB F + R’ were used to amplify an ampiclon with eve-tRNA-hid1-sgRNA in *pCFD5* as a template and the ampiclon was then ligated to *SapI* digested *pAJW1* plasmid to create *pAJW4*, a Gateway reaction with *pENTR: ptubulin-84b-Long* was performed to yield DB2-LT. DB1-LT, DB1-ST and DB2-LT were injected into a w1118 strain homozygous for eveΔ11 on the 2nd chromosome and for hid1-Δ3 on the 3rd chromosome, P-element integration was used. Positive transfectants were selected based on their eye colour (red).

**Generation and analysis of single-vector transgenics.** Flies harbouring P-element or AttB/AttP integrated transgenes were maintained in a genetic background containing appropriate promoter mutations; mostly homozygous eveΔ11, but also eveΔ11; hid1-Δ13 in the case of DB1 and DB2 transgenics. Red eyed flies were crossed as heterozygous males to virgin female w1118 white eyed flies. Red and white eye phenotypes were scored in the F1 progeny of crosses to establish the lethality associated with inheritance of the transgene (See Supplementary Technical Cross 5A–C). Percentage survival of the transgene in the F1 was calculated by multiplying the percentage incidence of red eyed flies by 2 to account for the two genotype categories. Collection protocol, embryo ageing times, tissue extraction and expression analysis of *eve* in embryos of transgenics crossed to tester stocks were carried out as previously stated for embryo qRT-PCR (See Supplementary Technical Cross 6). Fitness costs of homozygous stocks were assessed by mating 50 virgin females and 25 males in vials and then transferring to apple-juice-agar petri dishes smeared with yeast paste. Flies were allowed to pre-lay for 1 hour, petri dishes were replaced with fresh dishes and embryos after a 6 hour lay scored and flies removed. Covered petri dishes were incubated at room temperature for 4 days and surviving larvae/dead embryos scored.

**P-Element insertion loci analysis.** Genomic DNA from 5 transgenic flies was extracted using the Qiagen Blood and Tissue Extraction kit, following manufacturer’s protocol. 10 μg DNA in ddH2O was digested with 1 Unit of *Hinpol* (NEB) for 2.5 hours and heat inactivated at 65 °C for 20 mins. Samples were ligated with 0.25 Units.
of T4 DNA Ligase (NEB) at 4°C overnight then heat-inactivated at 95°C for 3 mins. 5 µl was used as template in Phusion Polymerase (NEB) PCR reactions using primers ‘5’P Element F’ and ‘5’P Element R’ with the cycling conditions: 95°C 5 min, 35 × (95°C 30 sec, 60°C 1 min, 68°C 2 mins), 72°C 10 min, 4°C hold. Products were analyzed on an agarose gel to confirm single bands. Reactions were purified using Qagen PCR purification kit (following instructions) and sequenced using primer ‘5’P Element Seq. For primer sequences see Supplementary Primers.

Data Availability
All data generated or analysed during this study are included in the manuscript and the supplement.

References
1. Hammond, A. et al. A CRISPR-Cas9 gene drive system targeting female reproduction in the malaria vector Anopheles gambiae. Nat. Biotechnol. 34, 78–83 (2016).
2. Moreno, E. et al. CRISPR-mediated direct mutation of cancer genes in the mouse liver. Nature 514, 380–384 (2014).
3. Bräu, A. et al. CRISPR/Cas-mediated genome editing in the rat via direct injection of one-cell embryos. Nat. Protoc. 9, 2493–2512 (2014).
4. Niu, D. et al. Inactivation of porcine endogenous retrovirus in pigs using CRISPR-Cas9. Science 359, eaan4187 (2017).
5. Chang, N. et al. Genome editing with RNA-guided Cas9 nucleases in Zebrafish embryos. Cell Res. 23, 465–472 (2013).
6. Dickinson, D. J., Ward, J. D., Reiner, D. J. & Goldstein, B. Engineering the Caenorhabditis elegans genome using Cas9-triggered homologous recombination. Nat. Methods 10, 1028–1034 (2013).
7. Bassett, A. R., Tibbon, C., Ponting, C. P. & Liu, J. I. Highly Efficient Targeted Mutagenesis of Drosophila with the CRISPR/Cas9 System. Cell Rep. 4, 220–228 (2013).
8. Port, F., Chen, H.-M., Lee, T. & Bullock, S. L. Optimized CRISPR/Cas tools for efficient germline and somatic genome engineering in Drosophila. Proc. Natl. Acad. Sci. 111, E2967–E2976 (2014).
9. Feng, Z. et al. Efficient genome editing in plants using a CRISPR/Cas system. Cell Res. 23, 1229–1232 (2013).
10. Gantz, V. M. et al. Highly efficient Cas9-mediated gene drive for population modification of the malaria vector mosquito Anopheles stephensi. https://doi.org/10.1073/pnas.1521077112 (2015).
11. Galizia, R. et al. A CRISPR-Cas9 sex-ratio distortion system for genetic control. Sci. Rep. 6, 2–6 (2016).
12. Chavez, A. et al. Highly efficient Cas9-mediated transcriptional programmimg. Nat. Methods 12, 326–328 (2015).
13. Jin, M. et al. A Programmable Dual-RNA − Guided. 337, 816–822 (2012).
14. Mali, P. et al. CAS9 transcriptional activators for target specificity screening and paired nickases for cooperative genome engineering. Nat. Biotechnol. 31, 833–838 (2013).
15. Maeder, M. L. et al. CRISPR RNA-guided activation of endogenous human genes. Nat. Methods 10, 977–979 (2013).
16. Perez-pineda, P. et al. Transcription Factors. Nat. Methods 10, 973–976 (2013).
17. Konermann, S. et al. Genome-scale transcriptional activation by an engineered CRISPR-Cas9 complex. Nature 517, 583–588 (2015).
18. Maselko, M., Heinsch, S. C., Chacon, J. M., Harcombe, W. R. & Smanski, M. J. Engineering species-like barriers to sexual reproduction. Nat. Commun. 8, 1–7 (2017).
19. Reed, F. A. First steps towards underdominant genetic transformation of insect populations. PLoS One 9, e97557 (2014).
20. Akbari, O. S. et al. Report A Synthetic Gene Drive System for Local, Reversible Modification and Suppression of Insect Populations. Curr. Biol. 23, 671–677 (2013).
21. Mitsch, W. J. et al. Ecological engineering: an introduction to ecotechnology. (Wiley, 1989).
22. Dominguez, A. A., Lim, W. A. & Qi, L. S. Beyond editing: Repurposing CRISPR-Cas9 for precision genome regulation and interrogation. Nat. Rev. Mol. Cell Biol. 17, 5–15 (2016).
23. Chavez, A. et al. Comparison of Cas9 activators in multiple species. Nat. Methods 13, 563–567 (2016).
24. Lin, S., Ewen-Campen, B., Ni, X., Housden, B. E. & Perrimon, N. In vivo transcriptional activation using CRISPR/Cas9 in Drosophila. Genetics 201, 433–442 (2015).
25. Ewen-Campen, B. et al. Optimized strategy for in vivo Cas9-activation in Drosophila. Proc. Natl. Acad. Sci. 114, 201707635 (2017).
26. Hilton, I. B. et al. Epigenome editing by a CRISPR-Cas9-based acetyltransferase activates genes from promoters and enhancers. Nat. Biotechnol. 33, 510–517 (2015).
27. Brand, A. H. & Perrimon, N. Targeted gene expression as a means of altering cell fates and generating dominant phenotypes. Development 118, 401–415 (1993).
28. Frasch, M., Hoey, T., Rushlow, C., Doyle, H. & Levine, M. Characterization and localization of the even-skipped protein of Drosophila. in Drosophila eye. Development 1262–1267 (2014).
29. Perez-pinera, P. et al. A CRISPR-Cas9 sex-ratio distortion system for genetic control. Proc. Natl. Acad. Sci. 112, 3312–3317 (2015).
30. Mohr, S. E. et al. CRISPR guide RNA design for research applications. FEBS J. 283, 3232–3238 (2016).
43. Caudle, A. S., Yang, W. T., Mittendorf, E. A. & Kuerer, H. M. The Impact of Chromatin Dynamics on Cas9-Mediated Genome Editing in Human Cells. *ACS Synth Biol* **150**, 137–143 (2017).
44. Crocker, J. & Stern, D. L. TALE-mediated modulation of transcriptional enhancers in vivo. *Nat. Methods* **10**, 762–767 (2013).
45. Wefers, B., Ortiz, O., Wurst, W. & Kühn, R. Generation of targeted mouse mutants by embryo microinjection of TALENs. *Methods* **69**, 94–101 (2014).
46. José A, Campos-Ortega & Hartenstein, V. *The Embryonic Development of Drosophila melanogaster*. (Springer, Berlin, Heidelberg, 1985).
47. Schindelin, J. *et al*. Fiji: an open-source platform for biological-image analysis. *Nat. Methods* **9**, 676–682 (2012).
48. Port, F. & Bullock, S. L. Augmenting CRISPR applications in Drosophila with tRNA-flanked sgRNAs. *Nat. Methods* **13**, 852–854 (2016).

**Acknowledgements**

We would like to thank Joshua Sealy, Yuxiang Zhang, Yew Quek, Manon Ricard, Jelle van der Hilst, Max Addison, Giles Wane and Jonathan Chan. We thank Kadri Oras at the University of Cambridge fly injection facility, DSHB, Bloomington Stock center and Addgene. We would also like to thank the Perrimon lab for providing many of the GAL4/UAS::dCas9-VPR lines. This study was funded by the European Research Council under the European Union’s Seventh Framework Programme ERC grant no. 335724 awarded to N.W.

**Author Contributions**

N.W. and A.J.W. planned the experiments. A.J.W., P.C., D.G. performed the experiments. N.W., P.A.P. and A.J.W. wrote the paper.

**Additional Information**

Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-018-31433-2.

**Competing Interests:** The authors declare no competing interests.

**Publisher’s note:** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2018