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OPEN A male-biased sex-distorter gene drive for the human malaria vector Anopheles gambiae

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Only female insects transmit diseases such as malaria, dengue and Zika; therefore, control methods that bias the sex ratio of insect offspring have long been sought. Genetic elements such as sex-chromosome drives can distort sex ratios to produce unisex populations that eventually collapse, but the underlying molecular mechanisms are unknown. We report a male-biased sex-distorter gene drive (SDGD) in the human malaria vector *Anopheles gambiae*. We induced super-Mendelian inheritance of the X-chromosome-shredding I-Ppol nuclease by coupling this to a CRISPR-based gene drive inserted into a conserved sequence of the *doublesex* (*dsx*) gene. In modeling of invasion dynamics, SDGD was predicted to have a quicker impact on female mosquito populations than previously developed gene drives targeting female fertility. The SDGD at the *dsx* locus led to a male-only population from a 2.5% starting allelic frequency in 10-14 generations, with population collapse and no selection for resistance. Our results support the use of SDGD for malaria vector control.

Sex-chromosome drivers are genetic elements that interfere with chromosome segregation during meiosis and are over-represented in progeny¹. In heterogametic sex, they cause an unbalanced male-to-female ratio among offspring, which can potentially lead to population suppression or extinction. Relatively few sex-chromosome drives have been characterized, most likely because they produce an evolutionary conflict with the rest of the genome that selects for autosomal suppressors or resistant sex chromosomes^{2,3}.

Mathematical modeling predicts that a driving sex distorter will spread in a population and, in the absence of resistance, cause eventual collapse^{4,5}. Population collapse using natural sex-chromosome drives has been reported in laboratory colonies of *Drosophila*^{6,7}. In the field, a population crash of the species *Drosophila neotestacea* was detected in Washington State due to a natural X-chromosome distorter that produced a female-only population⁸. Therefore, sex-distorter drives could conceivably be harnessed for invasive pest or vector control^{9,10}.

Although Y drives are less common than X drives, they have been described in *Aedes aegypti* and *Culex pipiens* mosquitoes^{11,12}. Y drives are particularly attractive for mosquito vector control because they can progressively reduce the number of females and hence disease transmission as they spread. In addition, Y drives are likely to be more effective than X drives because they can increase at a greater rate the fraction of heterogametic driving individuals³⁻⁵. Synthetic sex distorters have been generated in *A. gambiae* mosquitoes by using site-specific nucleases such as I-PpoI or CRISPR– Cas9, which cleave conserved repeated sequences in the mosquito ribosomal DNA gene cluster located exclusively on the X chromosome^{13,14}. These nucleases, when expressed during spermatozoa development, selectively cleave the X chromosome, thereby favoring the production of Y-bearing gametes and causing a 95% male bias in the progeny^{13,14}. However, attempts to convert synthetic sex-ratio distorters into Y-chromosome drives have been unsuccessful so far. In most insect species, including *A. gambiae*, the sex chromosomes are transcriptionally shut down during gametogenesis, a process known as meiotic sex-chromosome inactivation^{15,16}, which prevents the transcription of X-shredding nucleases if they are inserted into the Y chromosome (personal observation, A.C. and R.G.).

Recently, a gene drive that targeted the *dsx* gene reached 100% frequency in 7–11 generations and crashed a caged population of 600 mosquitoes without inducing resistance¹⁷. We hypothesized that it might be possible to circumvent meiotic sex-chromosome inactivation by developing an autosomal male-biased sex distorter and coupling sex-ratio distortion with drive. This could result in a quicker impact on disease transmission and a synergistic effect (robustness) between the sex distorter and gene-drive components. Here we report the design and validation of an SDGD to spread the X-chromosome-shredding I-PpoI endonuclease and produce a male-only insect population.

Results

Designing an SDGD. We designed an SDGD system by combining (on the same construct) a CRISPR-based gene drive that targets a haplosufficient female fertility gene with the I-PpoI endonuclease, which in turn cleaves a conserved sequence in the X-linked ribosomal gene cluster (Fig. 1a,b). We used mathematical modeling to test the likely spread of this SDGD design. Our results indicate that our SDGD could spread rapidly from a low starting frequency to produce a largely unisex male population and would also impose a fitness load by impairing female fertility, which together would eliminate the population (Fig. 1c). This SDGD design is different

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Fig. 1 | Driving a sex-distorter system in the autosome. a, Schematic overview of the construct used to build an SDGD, which contains four transcription units: the I-Ppol nuclease (variant W124L), expressed as a fusion protein with the enhanced green fluorescent protein (eGFP) visual marker, under the control of the male-specific beta2-tubulin germline promoter; the SpCas9 nuclease, regulated by a promoter that is active in the germ line of both males and females (from the vasa or zpg gene); a gRNA under the control of the ubiquitous U6 polymerase III promoter, designed for homing at previously characterized haplosufficient fertility genes; and a 3xP3::DsRed gene as a fluorescent integration marker. **b**, Mode of action of the autosomal SDGD. The sex-distorter (I-Ppol; blue square) and gene-drive (CRISPRh; red square) components are linked head to tail in the same construct that is integrated in the autosome within a fertility gene. In the germ line of a female transgenic mosquito (highlighted in red), the CRISPR^h component is active (red arrowhead), leading to super-Mendelian inheritance of the transgene by homology-directed repair. In the germ line of a male transgenic mosquito, both the gene-drive (red arrowhead) and sex-distorter (blue arrowhead) transcription units are active, leading to homing of the construct (by action of CRISPR^h) and shredding of the X chromosome (by action of I-Ppol targeting ribosomal DNA repeats; indicated by vertical lines). This results in a bias of the sex ratio toward males in the progeny and super-Mendelian inheritance of the transgene. c, Idealized predictions (discrete-generation deterministic model) of transgenic frequency for spread in a population (solid line) alongside the load on the target population (dashed line) for an SDGD construct (left; fraction of male progeny (m) = 0.95) and a gene drive (right; m = 0.50) targeting a female fertility gene. The colored shading represents the fraction of males (blue) and females (pink) in the population, with fertile females indicated by a darker color. This idealized model makes several assumptions that are likely to vary by strain, including but not limited to full fitness in males and heterozygous females (fully recessive female fertility gene); complete sterility in homozygous females; 95% SDGD transmission in male and female heterozygotes; no generation of drive-resistant mutations; no loss of function of the sex distorter; and single release of male drive heterozygotes equal to 1% of the male population.

from the previously reported CRISPR-based gene drives that target recessive female fertility genes and impose a fitness load by the generation of homozygous sterile mutants^{17,18}. The modeling predicted that this SDGD would quickly bias the population toward males and gradually reduce the abundance of biting females, which both

reduce pathogen transmission (by females only) and suppress the population (Table 1 and Supplementary Fig. 1).

We generated distinct *A. gambiae* SDGD strains targeting three haplosufficient genes (*AGAP011377*, *AGAP007280* and *AGAP005958*) with established roles in female fertility¹⁸.
 Table 1 | Comparison of performance of gene-drive and sex-distorter genetic control approaches in terms of efficacy, spread and robustness

Construct type	Construct name	Homing rate	Male-biased sex-ratio distortion	Spread in caged population	Population suppression	Development of resistance in cages	Impact of heterozygotes on population sizeª	Component redundancy	Ref.
Gene drive	<i>dsx</i> F ^{CRISPRh}	92% males; 99% females	50%	Yes	Yes	No	No	No	17
Sex-distorter gene drive	SDGD ^{dsx}	92% males; 99% females	93%	Yes	Yes	No	Yes	Yes	This study
Autosomal sex distorter	gfp111A-2	0%	95%	No	Yes (over-flooding ratio of 3x)	No	Yes	No	13
Y drive	NA	100% males; 0% females*	95%*	Yes*	NA	NA	Yes*	No	NA

Homing rate is defined as the fraction of transgenic progeny above Mendelian inheritance. An asterisk denotes values based on a hypothetical X-shredder construct inserted on the Y chromosome generating 95% male offspring, all of which inherit the transgene. NA, not applicable. *Ability of the construct to have an impact on the population size (that is, the number of females) in heterozygosity (or hemizygosity for the Y drive) as compared to constructs targeting recessive female fertility loci, which impact population size when homozygote transgenic females are generated.

We assessed the activity of three SDGD constructs (SDGD⁰¹¹³⁷⁷, SDGD⁰⁰⁷²⁸⁰ and SDGD⁰⁰⁵⁹⁵⁸) in the progeny of crosses between SDGD-heterozygous and wild-type individuals by scoring the fraction of offspring containing the drive element and the sex ratio of the progeny. SDGD⁰⁰⁷²⁸⁰ had severely reduced fertility, and we did not recover enough progeny to assess drive activity. We found average inheritance rates of 79% (±0.17% (s.d.)) for SDGD⁰¹¹³⁷⁷ and 98% (±0.08% (s.d.)) for SDGD⁰⁰⁵⁹⁵⁸ (Supplementary Fig. 2 and Supplementary Table 1). Furthermore, we observed a male bias ranging from 92% to 94% in the progeny of males heterozygous for SDGD⁰¹¹³⁷⁷ and SDGD⁰⁰⁵⁹⁵⁸. Monitoring of life history traits revealed a dramatic reduction of female fertility in females heterozygous for SDGD⁰¹¹³⁷⁷ and SDGD⁰⁰⁵⁹⁵⁸ (Supplementary Fig. 3), similarly to previous findings where in the same genes were targeted with a vasa-Cas9 gene-drive construct¹⁸. We attributed this reduction in fertility to ectopic expression of the vasa promoter and subsequent conversion to a null genotype for the target gene in somatic tissues, where the gene product is required¹⁸⁻²⁰. In addition, the vasa promoter is known to induce maternal deposition of Cas9 into the developing embryo, resulting in deleterious mutations of the paternally inherited gene copy, in addition to the null allele inherited from the mother, imposing additional fitness costs to heterozygous female offspring. We also observed a strong reduction in the fertility of heterozygous males, particularly in SDGD⁰⁰⁷²⁸⁰ and SDGD⁰⁰⁵⁹⁵⁸ (Supplementary Fig. 3). We hypothesized that male sterility in SDGD⁰⁰⁷²⁸⁰ and partial male sterility in SDGD⁰⁰⁵⁹⁵⁸ was due to locus-dependent high expression of the I-PpoI nuclease, which, if persisting in spermatozoa, shreds the maternally inherited X chromosome in the fertilized embryo, resulting in embryo lethality^{13,21}. Despite high levels of drive transmission and male bias, unintended and severe fertility costs prevented the spread of ${\rm SDGD}^{{\scriptstyle 011377}}$ and ${\rm SDGD}^{{\scriptstyle 005958}}$ into caged mosquito populations when these were seeded at 12.5% allelic frequency (Supplementary Fig. 4). SDGD005958 failed to persist in the populations and disappeared after two generations. SDGD⁰¹¹³⁷⁷ was stable for eight generations, owing to a better balance of drive and fitness costs. This, in turn, generated low-level population suppression by maintaining a sex ratio of approximately 65% males (Supplementary Fig. 4).

Optimization of temporal and spatial characteristics and level of expression of Cas9 and I-PpoI. Our initial findings revealed that SDGD constructs targeting female fertility genes could bias both their own inheritance and the sex ratio of progeny. However, fitness costs, most likely associated with non-optimal spatial and temporal activity of both the Cas9 and I-PpoI genes, impaired SDGD

spread into mosquito populations. To minimize the ectopic activity of Cas9, we replaced the vasa promoter with the regulatory regions of the zero population growth (zpg) gene (AGAP006241). The zpg promoter has previously been applied to regulate Cas9 expression in gene-drive constructs, increasing the fertility of heterozygous individuals as compared to those harboring constructs using vasa^{17,19}. Previous studies have also shown that the expression levels of I-PpoI during spermatogenesis are crucial in determining whether the outcome is sex bias or sterility; high levels of activity correlate with male sterility^{13,21}. The destabilized version of I-PpoI (W124L¹³) used in this study was previously found to confer the highest levels of fertility while maintaining strong male bias from at least three independent genomic loci13. However, this I-PpoI variant impaired male fertility when expressed under the transgenic beta2-tubulin promoter inserted into the AGAP011377, AGAP007280 and AGAP005958 loci. To reduce the transcriptional activity of the beta2-tubulin promoter, we generated three variants by inserting a G+C-rich sequence of 100 bp in proximity to conserved sequences at position -244, -271 or -355 with respect to the ATG start codon (Supplementary Fig. 5). Each variant was tested for expression using a dual-fluorescence reporter system in vivo (Supplementary Fig. 5). For subsequent experiments, we selected beta2-tubulin promoter variant 244 (beta2244), which showed transcriptional activity that was about 8.1% that of the wild-type promoter sequence (Supplementary Fig. 6). The initial SDGD plasmid was then modified to replace the *vasa* promoter with the *zpg* regulatory sequences (as described in ref. 17), while the beta2-tubulin promoter was replaced with beta2244.

An SDGD targeting the dsx gene. To maximize the performance of the SDGD, we developed the construct SDGD^{dsx}, containing the zpg-Cas9 transcription unit, beta2244-I-PpoI and a guide RNA (gRNA) designed to target the intron 4-exon 5 boundary of the dsx gene (AGAP004050), because we previously reported that this site minimizes the development of resistance to a gene drive¹⁷. In addition, females that are homozygous for dsxF exhibit an 'inter-sex' phenotype and are viable but unable to bite¹⁷; therefore, this affects the vector competence of the population earlier than an SDGD targeting a standard female fertility locus, in which homozygous females are sterile but can bite and transmit. Unlike SDGD⁰⁰⁷²⁸⁰, SDGD⁰¹¹³⁷⁷ and SDGD⁰⁰⁵⁹⁵⁸, SDGD^{dsx} had no measurable impact on the fertility of heterozygotes: the larval output of SDGD^{dsx} males was comparable to that of controls (126.7 \pm 50.7 (s.d.) and 140.8 \pm 40.8 (s.d.), respectively; P = 0.39; Fig. 2a and Supplementary Table 1). The fertility of SDGD^{dsx} heterozygous females, measured as viable



Fig. 2 | Fertility and sex and inheritance bias of an SDGD targeting the female isoform of the sex-determination *dsx* **gene (AGAP004050). a**, Counts of eggs and hatched larvae determined in individual crosses ($n \ge 33$) of SDGD^{dsx}-heterozygous females and males to wild-type (wt) mosquitoes. While male fertility was comparable to that of wild type (male fertility 0.86; no significant difference, NS), females showed a 37% reduction in larval output as compared to wild type (female fertility of 0.627; *P=0.0124, Kruskal-Wallis test). Values on the right indicate the mean count \pm s.e.m., with larval hatching rate in parentheses. **b**, Scatterplots showing the fraction of SDGD^{dsx} transgene inheritance (*y* axis) against sex bias (*x* axis) in the progeny of individual SDGD^{dsx} trans-heterozygous males (left; n=63) and females (right; n=39) crossed to wild-type individuals. Individual blue and pink dots represent the progeny derived from a single female, and the red dot indicates the average of the population. Error bars correspond to s.d. Both male and female SDGD^{dsx}-heterozygous mosquitoes showed a strong bias in sex ratio toward males (0.93 \pm 0.09). Dotted lines indicate the fraction of males (*x* axis) and the fraction of SDGD (*y* axis) as expected by Mendelian inheritance.

offspring, was reduced as compared to controls (98.8 \pm 63(s.d.) and 140.8 ± 40.8 (s.d.), respectively; P = 0.012), although it was still sufficient to produce a large number of fertile individuals (Fig. 2a). High levels of maternal nuclease deposition can affect the fertility of the female progeny^{17,18,20}; however, we did not observe a significant difference in fertility when comparing females inheriting the transgene from a transgenic female parent to those inheriting the trangene from a male parent (Supplementary Fig. 7). As expected, we observed a marked male bias $(93.1\% \pm 0.08\% (s.d.))$ in the offspring of SDGD^{dsx} heterozygous males (Fig. 2b). The sex-distortion phenotype was stably transmitted from male mosquitoes to their transgenic male offspring, and no differences were observed as compared to males that inherited the construct from a female or a male (Supplementary Table 3). Strong super-Mendelian inheritance of the construct of 96.0% \pm 0.08% (s.d.) and 99.9% \pm 0.01% (s.d.) was observed from both males and females, respectively, based on the frequency of red fluorescent protein (RFP)⁺ progeny from heterozygous parents (Fig. 2b), making SDGD^{dsx} suitable for population suppression experiments.

SDGD^{dsx} **invades caged mosquito populations.** We used fertility, inheritance bias, sex-distortion data and mutant phenotype information to develop both deterministic and stochastic discrete-generation models (Methods and Supplementary Tables 2, 4 and 5) to predict the spread of SDGD^{dsx} into mosquito populations, simulating the release of 10% and 50% SDGD^{dsx}-heterozygous mosquitoes into

caged populations of 600 individuals. The stochastic model predicted that the transgene would quickly invade the population, reaching 100% allelic frequency and leading to collapse of the population in 93% and 98% of 10,000 simulations after 30 generations from a 10% and 50% SDGD release, respectively (Fig. 3). The deterministic model, however, showed differences in outcome depending on the values for the fertility of heterozygous females and males, ranging from population elimination to suppression and to the disappearance of SDGD^{dsx} if male fertility was below 0.5 compared to wild type (Supplementary Figs. 8 and 9). To test the model prediction, we released SDGD^{dsx} heterozygotes at either 2.5% or 25% allelic frequency into two populations of 600 caged mosquitoes, each in two replicates. At each generation, larvae were screened for the presence of the fluorescence marker linked to the transgene, and subsequently the fraction of males and females in the population was assessed. We observed a rapid spread of SDGD^{dsx} in all four populations, with the transgene reaching 100% allelic frequency between 4 and 12 generations. The spread of SDGD^{dsx} induced a strong bias of the population sex ratio toward males, accompanied by a progressive reduction of egg output, which led to population elimination at generations 5 and 6 for the replica cages that started with 25% SDGDdsx allelic frequency and at generations 9 and 13 for the replica cages that started with 2.5% SDGD^{dsx} release (Fig. 3).

Fitness of female progeny in SDGD^{dsx} males. SDGD-heterozygous males generated <6% female progeny, and female offspring

NATURE BIOTECHNOLOGY

ARTICLES



Fig. 3 | Kinetics of SDGD^{dsx} **spread in target mosquito populations.** The spread of SDGD^{dsx} was investigated in two different experiments starting with an allelic frequency of 2.5% (10% male release) and 25% (50% male and female release), in replica (cage A and cage B). The 10% release cages were set up with a starting population of 300 wild-type females, 270 wild-type males and 30 SDGD^{dsx}-heterozygous males. The 50% release cages were started with 150 wild-type females, 150 sDGD^{dsx}-heterozygous males and 150 SDGD^{dsx}-heterozygous females (allelic frequency of 25%). Each consecutive generation was established by selecting 600 larvae. The frequency of the transgene (fraction of RFP⁺ individuals), the sex ratio (female/ male) and the relative egg output (fraction of eggs produced relative to the first generation) were recorded at each generation. **a**, The bar plots represent the fraction of males and females (blue and pink shading, respectively) for each population, and the striped pattern shows the fraction of transgenic individuals. Black lines indicate the total fraction of individuals containing SDGD^{dsx} (as a fraction of RFP⁺) individuals. **b**, The frequency of the transgene, the sex ratio and the relative egg output superimposed on both a deterministic model (black dashed lines) and 20 representative stochastic simulations (gray solid lines) of the dynamics of invasion of SDGD^{dsx} based on release scenarios of 25% and 2.5% SDGD^{dsx} allelic frequency. In 93% and 98% of the stochastic simulations (of 10,000 runs), the release of SDGD-heterozygous individuals at a starting frequency of 2.5% and 25%, respectively, is predicted to collapse the population within 30 generations. Dotted lines indicate the expected Mendelian distribution of sex. Fitness and life history parameter estimates are provided in Supplementary Table 2.

inherited an X chromosome from male gametes exposed to the I-PpoI nuclease during spermatogenesis. We investigated whether the inheritance of a potentially damaged X chromosome affected female fertility and the SDGD homology-directed repair rate. We crossed females that carried one 'I-PpoI-exposed' X chromosome from the father to wild-type males and compared their fertility parameters to those of daughters of SDGD^{dsx} females that carried two unaffected copies of the X chromosome. We observed that females inheriting one I-PpoI-exposed X chromosome did not significantly differ in fertility (measured as the number of hatched larvae) nor in drive inheritance, suggesting that, if there is a contribution to fitness of a damaged X chromosome in females, this was not detectable in our assay. To further investigate the potential

impact of I-PpoI-exposed X chromosomes, we modeled additional fitness reductions in individuals with a damaged X chromosome using deterministic discrete-generation cage simulations of a theoretical scenario of SDGD^{dsx} release (10% males and 50% males/50% females) into a caged population (Supplementary Fig. 10). The model predicted little or no effect during the initial spread of the transgene, but a reduction in the suppression load that correlated with the cost of the damaged X chromosome was detected (Supplementary Fig. 10).

Dynamics of sex-distorter drive. Driving a sex distorter into a female fertility locus could impose a sufficiently high load on the population to the point that the population is suppressed and

eliminated. However, the dynamics of an SDGD are complex and depend not only on the fertility of SDGD^{dsx}-heterozygous individuals (Supplementary Figs. 8 and 9) but also on the rate of male bias (Supplementary Fig. 11), and in certain scenarios these dynamics are not intuitive. For example, when female (W/D, where W is the wild-type allele and D represents the SDGD allele) fertility is reduced, such as below 0.5, the load on the population increases with increasing sex distortion, whereas for higher female fertility, such as above 0.5, the load is greater when there is no sex distortion (equivalent to a gene drive without a sex distorter, m=0.5; Supplementary Fig. 11). The sex distorter allows the SDGD construct to spread at low (or even zero) female fertility, imposing a substantial load (Supplementary Fig. 12). This is because the male bias mitigates the effect of low female fitness. Overall, increasing sex distortion makes the construct less sensitive to variation in female heterozygous fertility (Supplementary Fig. 12). At the limit of complete male bias (male progeny=100%), the load is independent of female fertility because no SDGD females are created and only SDGD-heterozygous males can pass on the construct. Based on our experimental parameter estimates for SDGD^{dsx}, the SDGD allele is predicted to be present in an intermediate equilibrium with wild-type and nonfunctional resistance alleles at a sufficient frequency to induce a dramatic population reduction and possibly prevent reinvasion events (Supplementary Fig. 13).

Discussion

Our results show that SDGD^{dsx} functions as a sex-distorter autosomal gene drive. In four cage experiments, SDGD^{dsx} progressively biased the sex ratio toward males, with eventual population collapse. Notably, we did not observe the development of functional mutations at the target dsx site that blocked the spread of the distorter. This observation further supports the notion that the dsx sequence at the intron 4-exon 5 boundary is highly functionally constrained and validates its use as a target sequence for gene-drive solutions in anopheline mosquitoes. It should also be noted that a sex distorter that simultaneously destroys the female isoform of the dsx gene while reducing the female population also decreases the opportunity of resistant mutations to arise (because they are not selected in males). In addition, targeting a sequence present in hundreds of copies on the X chromosome reduces the likelihood that nuclease-induced resistance will evolve to block the sex-distorter component.

Our SDGD solution also combines a number of features in terms of efficacy, robustness and predicted time to impact (on disease transmission), which differ from those for previously described gene drives or autosomal sex-distorter systems, making it particularly attractive for field implementation (Table 1). In two replicate caged experiments, SDGD^{dsx} consistently induced population collapse starting from an allelic frequency of 2.5%. For field experiments, this translates into mosquito numbers to be released that are within the range of production capability; recent studies modeling the impact of hypothetical X-shredder Y-drive mosquitoes on a national scale predict that the release of as few as ten males in 1% of human settlements will achieve over 90% population suppression after 4 years²².

SDGD^{dsx} is predicted to show a higher level of robustness than a gene drive alone, even if one of the critical components breaks down or mutates, due to the synergy of the components. Loss or inactivation of the I-PpoI sequence will result in the generation of functional dsx gene drive that will also contribute to population suppression (Supplementary Fig. 12), and loss of function of either of the two drive components (Cas9 or gRNA) will produce nonfunctional dsx alleles (R) that, in heterozygous individuals, will still contribute to the production of male-biased progeny owing to the presence of functional I-PpoI. Mutations and recombination events of the constructs involving both the drive and distorter will generate R nonfunctional *dsx* mutations. These R mutations are constantly generated at the target locus by the action of the nuclease¹⁷, but they are not selected because they do not restore function of the *dsx* gene and homozygous R females are sterile; therefore, they are continuously lost as they arise.

Modeling based on our experimental data shows that SDGD^{dsx} offers some important advantages in short-term drive dynamics and long-term outcomes. Importantly, the number of transmission-competent (that is, biting) females is reduced more quickly by SDGD^{dsx} than by a standard gene drive targeting the same locus (Supplementary Fig. 1), which could lead to a strong effect on disease transmission (time to impact after release). In comparing a distorting and a non-distorting gene drive, the equilibrium load imposed by SDGD^{dsx} is less sensitive to female fitness costs, which is particularly relevant given the uncertain extrapolation of fitness effect measurements from the lab to the field.

Previous modeling of gene drive without the sex distorter showed that under certain conditions (for example, leaky expression of the drive construct) there can be an accumulation of nonfunctional cleavage-resistant sequences, which prevents the transgene from going to fixation^{17,23}. Deterministic modeling of SDGD^{dsx} indicates that there is also the potential for the transgene to go to an intermediate equilibrium frequency and population suppression, rather than complete fixation and elimination (Supplementary Figs. 8, 9 and 12). The lower the SDGD fertility in heterozygous individuals, the more likely an intermediate equilibrium is reached. For the observed fertility values of SDGDdsx-heterozygous females, stochastic models predicted population elimination for finite cage populations in 93-98% of the simulations, with kinetics of spread in line with observed data. Under field or semi-field conditions, the fertility estimates of heterozygous individuals could differ and tilt the balance one way or the other toward population reduction rather than population elimination. Achieving a strong population reduction may be regarded as less effective than elimination in a field scenario; however, it could help achieve long-term stable vector control via a higher tolerance to repopulation through migration as compared to a system that quickly eliminates an entire target population.

Males carrying a non-driving I-PpoI construct designed to cause dominant male sterility²¹ were recently released in a field location of Burkina Faso²⁴ as part of a phased, step-by-step assessment of novel genetic approaches to malaria control, following independent guidance and recommendations^{25,26}. This opened the way to the use of an I-PpoI-based distorter for the implementation of genetic vector control measures.

We believe that SDGD^{dsx} outperforms other anopheline gene drives, combining efficacy, resistance management and robustness, and is well suited as an anti-malaria intervention.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41587-020-0508-1.

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References

- 1. Helleu, Q., Gérard, P. R. & Montchamp-Moreau, C. Sex chromosome drive. Cold Spring Harb. Perspect. Biol. 7, a017616 (2015).
- Taylor, J. E. & Jaenike, J. Sperm competition and the dynamics of X chromosome drive: stability and extinction. *Genetics* 160, 1721–1731 (2002).
- Burt, A. & Trivers, R. Genes in Conflict: the Biology of Selfish Genetic Elements (Harvard University Press, 2009).
- 4. Hamilton, W. D. Extraordinary sex ratios. Science 156, 477-488 (1967).

NATURE BIOTECHNOLOGY

E874-E880 (2011).

- 5. Deredec, A., Godfray, H. C. J. & Burt, A. Requirements for effective malaria control with homing endonuclease genes. Proc. Natl Acad. Sci. USA 108,
- 6. Lyttle, T. W. Experimental population genetics of meiotic drive systems. I. Pseudo-Y chromosomal drive as a means of eliminating cage populations of Drosophila melanogaster. Genetics 86, 413-445 (1977).
- Price, T. A. R., Hodgson, D. J., Lewis, Z., Hurst, G. D. D. & Wedell, N. Selfish genetic elements promote polyandry in a fly. Science 322, 1241-1243 (2008).
- 8 Pinzone, C. A. & Dyer, K. A. Association of polyandry and sex-ratio drive prevalence in natural populations of Drosophila neotestacea. Proc. Biol. Sci. 280, 20131397 (2013).
- Hastings, I. M. Selfish DNA as a method of pest control. Philos. Trans. R. Soc. 9 Lond. B Biol. Sci. 344, 313-324 (1994).
- 10. Sinkins, S. P. & Gould, F. Gene drive systems for insect disease vectors. Nat. Rev. Genet. 7, 427-435 (2006).
- 11. Hickey, W. A. & Craig, G. B. Genetic distortion of sex ratio in a mosquito, Aedes aegypti. Genetics 53, 1177-1196 (1966).
- 12. Sweeny, T. L. & Barr, A. R. Sex ratio distortion caused by meiotic drive in a mosquito, Culex pipiens L. Genetics 88, 427-446 (1978).
- 13. Galizi, R. et al. A synthetic sex ratio distortion system for the control of the human malaria mosquito. Nat. Commun. 5, 3977 (2014).
- 14. Galizi, R. et al. A CRISPR-Cas9 sex-ratio distortion system for genetic control. Sci. Rep. 6, 31139 (2016).
- 15. Turner, J. M. A. Meiotic sex chromosome inactivation. Development 134, 1823-1831 (2007).
- 16. Taxiarchi, C. et al. High-resolution transcriptional profiling of Anopheles gambiae spermatogenesis reveals mechanisms of sex chromosome regulation. Sci. Rep. 9, 1-12 (2019).
- 17. Kyrou, K. et al. A CRISPR-Cas9 gene drive targeting doublesex causes complete population suppression in caged Anopheles gambiae mosquitoes. Nat. Biotechnol. 36, 1062-1066 (2018).
- 18. Hammond, A. et al. A CRISPR-Cas9 gene drive system targeting female reproduction in the malaria mosquito vector Anopheles gambiae. Nat. Biotechnol. 34, 78-83 (2016).
- 19. Hammond, A. M. et al. Improved CRISPR-based suppression gene drives mitigate resistance and impose a large reproductive load on laboratory-contained mosquito populations. Preprint at bioRxiv https://doi. org/10.1101/360339 (2018).

- 20. Gantz, V. M. et al. Highly efficient Cas9-mediated gene drive for population modification of the malaria vector mosquito Anopheles stephensi. Proc. Natl Acad. Sci. USA 112, E6736-E6743 (2015).
- 21. Windbichler, N., Papathanos, P. A. & Crisanti, A. Targeting the X chromosome during spermatogenesis induces Y chromosome transmission ratio distortion and early dominant embryo lethality in Anopheles gambiae. PLoS Genet. 4, e1000291 (2008).
- 22. North, A. R., Burt, A. & Godfray, H. C. J. Modelling the potential of genetic control of malaria mosquitoes at national scale. BMC Biol. 17, 26 (2019).
- 23. Beaghton, A. K., Hammond, A., Nolan, T., Crisanti, A. & Burt, A. Gene drive for population genetic control: non-functional resistance and parental effects. Proc. Biol. Sci. 286, 20191586 (2019).
- 24. Scudellari, M. Self-destructing mosquitoes and sterilized rodents: the promise of gene drives. Nature 571, 160-162 (2019).
- 25. James, S. et al. Pathway to deployment of gene drive mosquitoes as a potential biocontrol tool for elimination of malaria in sub-Saharan Africa: recommendations of a scientific working group. Am. J. Trop. Med. Hyg. 98, 1-49 (2018).
- 26. Committee on Gene Drive Research in Non-Human Organisms: Recommendations for Responsible Conduct, Board on Life Sciences, Division on Earth and Life Studies, & National Academies of Sciences, Engineering, and Medicine. Gene Drives on the Horizon: Advancing Science, Navigating Uncertainty, and Aligning Research with Public Values (National Academies Press, 2016).

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Methods

Ethics statement. Generation of SDGD constructs. To create SDGD vectors p172 (vas2; GenBank accession MT270142) and p182 (zpg), the β 2-eGFP(F2A)I-PpoI transcription unit from pBac[3xP3-DsRed] β 2-eGFP:II-PpoI-124L¹³ was excised by AscI digestion and cloned into AscI-digested p165 (vas2-CRISPR^h (ref. ¹⁸): GenBank accession KU189142) and p174 (zpg-CRISPR^h (ref. ¹⁹); GenBank accession MH541847), respectively. SDGD vectors were further modified by BsaI-mediated Golden Gate assembly to contain gRNA spacers targeting AGAP011377 (GCAGACGTAGAAATTTTC), AGAP007280 (GGAAGAAAGT GAGGAGGA), AGAP005958 (GAGATACTGGAGCCGCGAGC)¹⁸ and AGAP004050 (GTTTAACACAGGTCAAGCG]¹⁷. To include the beta2²⁴⁴ promoter modification, plasmid p182 was further modified to generate p182–244 (GenBank accession MT270141) according to the beta2²⁴⁴ variant described below. Additional sequences of all vectors are available as Supplementary Information.

Microinjection of embryos and selection of transformed mosquitoes. All mosquitoes were reared under standard conditions of 80% relative humidity and 28 °C. The mosquitoes were blood fed on anesthetized mice or by Hemotek, and freshly laid embryos were aligned and used for microinjections as described previously²⁷. To generate SDGD mosquitoes, we injected respective docking line^{17,18} embryos with solution containing p174 or p182–244 and a plasmid-based source of PhiC31 integrase²⁸ (at 200 ng μ l⁻¹ and 400 ng μ l⁻¹, respectively). All surviving G₀ larvae were crossed to wild-type mosquitoes, and G₁ positive transformants were identified using a fluorescence microscope (Nikon, Eclipse TE200) as RFP⁺ larvae for the recombination-mediated cassette exchange (RMCE) events.

Containment of gene-drive mosquitoes. All mosquitoes were housed at Imperial College London in an insectary that is compliant with Arthropod Containment Guidelines Level 2 (ref. ²⁹). All genetically modified (GM) work was performed under institutionally approved biosafety and GM protocols. In particular, GM mosquitoes containing constructs with the potential to show gene drive were housed in dedicated cubicles, separated from the external environment by at least six doors requiring two levels of security card access. Moreover, because the insectary is located in a city with a northern temperate climate, *A. gambiae* mosquitoes are also ecologically contained. The physical and ecological containment of the insectary are compliant with guidelines set out in a recent commentary calling for safeguards in the study of synthetic gene-drive technologies³⁰.

Mutagenesis of the *beta2-tubulin* **promoter.** Bioinformatic analysis of the regulatory region of the *beta2-tubulin* gene (*AGAP008622*) was performed using the Promoter2.0 Prediction Server³¹ and the Neural Network Promoter Prediction tool³² to identify the conserved region. A synthetic 100-bp DNA sequence with a G+C content of 65% (sequence reported in Supplementary Fig. 3) was designed using Geneious R11 (https://www.geneious.com/) and cloned into the *beta2-tubulin* promoter at position -244, -271 or -355 with respect to the ATG start codon using site-specific mutagenesis of plasmid pBac[3xP3-DsRed] β 2-eGFP::I-PpoI-124L¹³ by nested PCR with primer pairs B2-355_r and B2-355_f, B2-271_r and B2-271_f, and B2-244_r and B2-244_f followed by Spac-fwd and Spac-rev, for the *beta2³⁵⁵*, *beta2²⁷¹* and *beta2²⁴⁴* variants, respectively. A second unmodified copy of the *beta2-tubulin* promoter was cloned to express the mCherry gene.

Name	Sequence
B2-355_r	GGCCAACTCGGGTCCGAGTCGTCTTCTTGGATGGGATGATG
B2-355_f	CGCCAGCACTCTCAGACTCAATACGAATTTATTTGTGGCATCG
B2-271_r	GGCCAACTCGGGTCCGAGTCATATGACTACTATGATCAT CTTTTGC
B2-271_f	CGCCAGCACTCTCAGACTCAGAG CCG TAC GTG CCG G
B2-244_r	GGCCAACTCGGGTCCGAGTCCACGAAATGATCCGGCAC
B2-244_f	CGCCAGCACTCTCAGACTCACAGAACCTTCAGAGACGTTG
Spac-fwd	GTGAGAAGTGCGCGTCTCGTTCCCGCAGCTCGCCAGCACTC TCAGACTCA
Spac-rev	CATCCGCCCTAACTCCGCCCGTGGGTCGTTGGCCAACTCGGG TCCGAGTC

Dual-fluorescence assay experiment. Three- to five-day-old adult male heterozygous mosquitoes were collected in Falcon tubes and anesthetized on ice 5 min before dissection. Testes were micro-dissected using an Olympus SZX7 optical microscope, and pictures of gonads were taken using the EVOS imaging system (Thermo-Fisher) with magnification of ×20 and the following exposure settings: bright field: gain 50%; GFP channel: gain 30%, 120 ms; RFP channel: gain 80%, 120 ms. Unmodified pictures were then analyzed using ImageJ software³³. Testis areas were selected using the freeform selection tool, and integrated density and mean gray values were measured for the GFP and RFP channels independently using the same selection area. A reading for the background (same selection area)

was then subtracted from the integrated density value for each testis to remove background noise. The value for fluorescence intensity was measured as the ratio between the GFP reading and the mCherry reading and normalized to the value of the unmodified *beta2-tubulin* control.

Phenotypic assays. Phenotypic assays designed to examine SDGD inheritance and relative fecundity in mosquitoes were carried out essentially as described before^{17,18}. Briefly, the offspring of heterozygous individuals crossed to wild-type counterparts were screened by RFP expression. Non-fluorescent progeny were kept as controls. Groups of 50 male and 50 female blood-fed mosquitoes were mated to an equal number of wild-type mosquitoes for 5 d, and a minimum of 40 females were allowed to lay individually. The entire egg and larval progeny was counted for each lay. Females that failed to give progeny and had no evidence of sperm in their spermathecae were excluded from the analysis. To determine the inheritance and sex-ratio bias of SDGD, the entire larval progeny was screened for the presence of DsRed, which is linked to the SDGD allele, and all the progeny was sexed at the pupal stage to determine the sex ratio. Statistical differences between genotypes were assessed using the Kruskal–Wallis test.

Cage trial assays. To perform cage trials of SDGD⁰¹¹³⁷⁷ and SDGD⁰⁰⁵⁹⁸, we introduced 100 heterozygous transgenic males into a population of 100 wild-type males and 200 wild-type females (transgenic allelic frequency of 12.5%) in triplicate. As a control, 100 heterozygous transgenic males from the autosomal self-limiting sex-distorter ^{gfn}124L-2 line¹³ were released at the same frequency in a separate population, in triplicate. In addition, a population of 200 wild-type males and 200 wild-type females served as a negative control.

For the starting generation only, age-matched male and female pupae were allowed to emerge in separate cages and were mixed only when all the pupae had emerged. Mosquitoes were left to mate for 5 d before they were blood fed on anesthetized mice. After 2 d, the mosquitoes were set to lay in a 300-ml egg bowl filled with water and lined with filter paper. The eggs produced from the cage were photographed and counted using JMicroVision v1.27. Before counting, the eggs were dispersed using gentle water spraying in the egg bowl to homogenize the population, and 450 eggs were randomly selected to seed the next generation. Larvae emerging from the 450 eggs were counted and screened for the presence of the RFP marker to score the transgenic rate of the progeny. All the pupae were sexed to determine the sex ratio of the population.

To perform cage trials of SDGD^{dsx}, we set up two different experiments, in replicate. The 10% release cages were set up with a starting population of 300 wild-type females, 270 wild-type males and 30 SDGD^{dsx}-heterozygous males (starting allelic frequency of 2.5%). The 50% release cages were started with 150 wild-type males, 150 SDGD^{dsx}-heterozygous males and 150 SDGD^{dsx}-heterozygous males (allelic frequency of 2.5%). For the starting generation only, age-matched male and female pupae were allowed to emerge in separate cages and were mixed only when all the pupae had emerged. Mosquitoes were left to mate for 5 d before blood feeding. After 2 d, the mosquitoes were set to lay in a 300-ml egg bowl filled with water and lined with filter paper. All larvae were allowed to hatch, and each consecutive generation was established by randomly selecting 600 larvae, split into 3 trays of 200 larvae each. All 600 larvae were secreed for the presence of the RFP marker, and the pupae from one tray were sexed to determine the sex ratio. On day 8, mosquitoes were offered a second blood meal, and all the eggs produced were photographed and counted using Egg counter software³⁴.

Statistical analysis. Statistical analysis was performed as indicated using GraphPad Prism version 7.0.

Population genetics model. Discrete time. To model the results of the cage experiments, we use discrete-generation recursion equations for the genotype frequencies, with males and females treated separately, similarly to Kyrou et al.¹⁷. We extend the previous study17 to model the SDGD by including a sex bias and possible X-chromosome damage in the progeny of SDGD males, although here we do not include parental effects on fitness (as these effects were not strongly observed). We consider three alleles at the female fertility target site, W (wild type), D (driving sex distorter) and R (nonfunctional nuclease resistant). We also differentiate between the two possible types of X chromosome: x (wild type) and X, which denotes an X chromosome that has passed through an SDGD male and survived X-shredding but may be damaged, resulting in an additional fitness cost to the individual carrying it. $F_{ij,pq}(t)$ and $M_{ij,qY}(t)$ denote the genotype frequency of females and males, respecitvely, in the total population, where the first set of indices denotes alleles at the target site {WW, WD, WR, DD, DR, RR} and the second set denotes the sex chromosomes, $pq = \{xx, xX, XX\}$ for females and $q = \{x, X\}$ for males. There were 18 female genotypes and 12 male genotypes; six types of eggs: $E_{W,x}$, $E_{D,x}$, $E_{R,x}$, $E_{W,X}$, $E_{D,X}$ and $E_{R,X}$, where the first index refers to the target site allele and the second index to the sex chromosome; and eight types of sperm: S_{W,X}, S_{R,X} (no S_{D,x}, because we assume that SDGD males only contribute X chromosomes), S_{W,X}, S_{D,X}, S_{R,X}, S_{W,Y}, S_{D,Y} and S_{R,Y}.

Homing. Adults of genotype W/D at the target site produced gametes at meiosis in the ratio W:D:R as follows:

$$(1 - d_f)(1 - u_f) : d_f : (1 - d_f)u_f$$
 in females

$$(1 - d_m)(1 - u_m) : d_m : (1 - d_m)u_m$$
 in males

Here d_f and d_m are the rates of transmission of the driver allele in the two sexes and u_f and u_m are the fractions of non-drive gametes at the target site that are repaired by meiotic end-joining and are nonfunctional and resistant to the drive (R). In all other genotypes, inheritance at the target site is Mendelian.

Sex distortion. The SDGD X-shredder only affects the sex ratio of the progeny if it is in males. It destroys the X chromosome while males are making their sperm, resulting in mostly Y-bearing sperm. From male SDGD heterozygotes, progeny will therefore consist of m_1 ($1/2 < m_1 \le 1$) males and $(1 - m_1)$ females; from male SDGD homozygotes (D/D), the progeny will be m_2 ($1/2 < m_2 \le 1$) males and $(1 - m_2)$ females. For simplicity, when comparing to the experiment, we assume $m_1 = m_2 = m$. We assume no mutations cause loss of function of the sex distorter from the construct or resistance to X-shredding.

All X chromosomes contributed by SDGD males that survived X-shredding are assumed to be 'damaged' X chromosomes (versus wild-type X chromosomes), reflected in the reduced reproductive fitness of the individual carrying it (see 'Fitness'). We assume a damaged X chromosome was susceptible to further shredding if it was inherited by an SDGD male and, for simplicity, that the fitness cost of carrying a damaged X chromosome is the same no matter how many times the chromosome passes through an SDGD male and survives X-shredding.

Fitness. We let $w_{ij,pq}, w_{ij,qY} \le 1$ represent the reproductive fitnesses of female and male genotypes relative to a fitness of 1 for wild-type homozygotes, where $\{ij\}$ denotes alleles at the target site of the construct {WW, WD, WR, DD, DR, RR} and the second set of indices denotes $pq = \{xx, XX, XX\}$ for females and $q = \{x, X\}$ for males. While all fitness parameters are retained in the recursion equations for generality, for comparison with the experiment, we assume that the target gene is needed for female fertility; thus, females with D/D, D/R and R/R at the target site are sterile. There is no reduction in fitness in W/R females from carrying only one copy of the target gene (W/R), but W/D females have reduced fitness due to the presence of the SDGD construct, as observed experimentally (Supplementary Table 2). We assume no costs to males with no copies of the SDGD (W/D, D/R) have a fitness reduction consistent with experimental observation (Supplementary Table 2).

If the individual also carries a damaged X chromosome, we assume that this imposes an additional cost that affects reproductive success. To calculate the overall fitness of the genotype, the fitness value associated with carrying the damaged X chromosome is multiplied by the fitness value associated with D (or R) alleles at the target site (Supplementary Tables 4 and 5). Reduced fitness in males with a copy of the damaged X chromosome is $(1 - s_{X,m})$ and in females with two copies of the damaged X chromosome is $(1 - s_{X,m})$, with $s_{X,P}, s_{X,m} = 0$ if there is no cost and 1 if the damaged X chromosome causes sterility. For females with one damaged X chromosome and one wild-type x chromosome, the reduction is $(1 - h_{X,x})$, where $h_{X,i}$ is the dominance coefficient (0 for fully recessive; 1 for fully dominant). For baseline parameters, we assume these costs are equal to zero.

Recursion equations. We first consider the gamete contributions from each genotype. The proportions $E_{k,l}(t)$ with allele $k = \{W, D, R\}$ at the target site and sex chromosome $l = \{x, X\}$ in eggs produced by females participating in reproduction are given in terms of the female genotype frequencies $F_{ij,pq}(t)$:

$$E_{k,l}(t) = \frac{\sum_{i=1}^{3} \sum_{j=i}^{3} \sum_{pq=xx;xX;XX} \left(c_{ij,pq}^{k,l} w_{ij,pq} F_{ij,pq}(t) \right)}{\sum_{i=1}^{3} \sum_{j=i}^{3} \sum_{pq=xx;xX;XX} \left(w_{ij,pq} F_{ij,pq}(t) \right)}$$
(1)

where *i* and *j* are each summed such that {1, 2, 3} corresponds to {W, D, R}. The coefficients $c_{ij,pq}^{k,l}$ in equation (1) correspond to the proportion of the gametes from female individuals of genotypes (*ij*, *pq*) that carry alleles (*k*, *l*), as shown in Supplementary Table 4 (rows correspond to genotypes; columns correspond to alleles).

The proportions $S_{k,l}(t)$ with allele $k = \{W, D, R\}$ at the target site and sex chromosome $l = \{x, X, Y\}$ in sperm are given in terms of the male genotype frequencies $M_{ij,qY}(t)$:

$$S_{k,l}(t) = \frac{\sum_{i=1}^{3} \sum_{j=i}^{3} \sum_{q=x,X} \left(c_{ij,qY}^{k,l} w_{ij,qY} M_{ij,qY}(t) \right)}{\sum_{i=1}^{3} \sum_{j=i}^{3} \sum_{q=x,X} \left(w_{ij,qY} M_{ij,qY}(t) \right)}$$
(2)

where, again, *i* and *j* are each summed such that {1, 2, 3} corresponds to {W, D, R}. The coefficients $S_{ij,qY}^{k,l}$ in equation (2) correspond to the proportion of the gametes from male individuals of type (*ij*, *qY*) that carry alleles (*k*, *l*), as shown by the rows and columns, respectively, in Supplementary Table 5. Note that $S_{Dx}(t) = 0$ because SDGD males only contribute damaged X chromosomes, so no entry for this is included in Supplementary Table 5.

We define the proportion of females in the population as

$$F(t) = \sum_{i=1}^{3} \sum_{j=i}^{3} \sum_{pq=xx;xX;XX} F_{ij,pq}(t)$$

and the average female reproductive fitness as

$$\bar{w}_f(t) = \sum_{i=1}^3 \sum_{j=i}^3 \sum_{pq=xx;xX;XX} \left(\frac{w_{ij,pq} F_{ij,pq}(t)}{F(t)} \right)$$

Analogously, for the male proportion, we define the proportion of males in the population as

$$M(t) = \sum_{i=1}^{3} \sum_{j=i}^{3} \sum_{q=x,X} M_{ij,qY}(t)$$

and the average male fitness as

$$ar{w}_m(t) = \sum_{i=1}^3 \sum_{j=i}^3 \sum_{q=x,X}^3 \left(rac{w_{ij,qY} M_{ij,qY}(t)}{M(t)}
ight)$$

Note that in equations (1) and (2), the normalization factor in the denominator is therefore $\bar{w}_f(t)F(t)$ and $\bar{w}_m(t)M(t)$, respectively.

The load on the population incorporates reductions in female and male fertility and decreased frequency of females due to the SDGD spreading in the population, and at time (t) is defined as

$$L(t) = 1 - 2F(t)\bar{w}_{\rm f}(t)\bar{w}_{\rm m}(t)$$

This equals zero when only wild-type individuals are present and one if the SDGD has been established and the average female fitness, or fraction of females present, is equal to zero. We note that increases in load predicted by the cage model do not predict absolute changes in population density in the field but can be an indication of comparative potential reductions³⁵.

To model cage experiments, we start with an equal number of males and females. For 50% release, the initial frequency for wild-type females and males is $F_{WW,xx} = M_{WW,xY} = 1/4$ and for heterozygote drive females and males is $F_{WD,xx} = M_{WD,xY} = 1/4$. For 10% release of males only, $M_{WW,xY} = 9/20$ and $M_{WD,xY} = 1/20$ and all females are wild type, $F_{WW,xx} = 1/2$. Assuming random mating, we obtain the following recursion equations for the female genotype frequencies in generation (t + 1):

$$F_{ij,pq}(t+1) = \left(1 - \frac{\delta_{ij}}{2}\right) \left(1 - \frac{\delta_{pl}}{2}\right) \\ \left(E_{i,p}(t)S_{j,q}(t) + E_{j,p}(t)S_{i,q}(t) + E_{i,q}(t)S_{j,p}(t) + E_{j,q}(t)S_{i,p}(t)\right)$$

where $pq = \{xx, xX, XX\}$, and δij is the Kronecker delta. The factors $\left(1 - \frac{\delta_{ij}}{2}\right), \left(1 - \frac{\delta_{ij}}{2}\right)$ account for the factor of 1/2 for homozygosity at the target site (for $ij = \{W,W, D/D, R/R\}$) and at the sex chromosomes (for $pq = \{xx, XX\}$). We obtain the following recursion equations for the male genotype frequencies:

$$M_{ij,qY}(t+1) = \left(1 - \frac{\delta_{ij}}{2}\right) \left(E_{i,q}(t)S_{j,Y}(t) + E_{j,q}(t)S_{i,Y}(t)\right)$$

where $q = \{x, X\}$ and $\left(1 - \frac{\delta_{ij}}{2}\right)$ accounts for the factor of 1/2 for homozygosity at the target site (for $ij = \{W/W, D/D, R/R\}$).

Stochastic version. In the stochastic version of the model described above, random values for probabilistic events are taken from the appropriate multinomial distributions, with probabilities estimated from the experiment where applicable (Supplementary Table 2). To model the cage experiments, 150 female and 150 male wild-type adults (or 300 females and 270 males for 10% release of males only) along with 150 female and 150 male heterozygotes (or no females and 30 males for 10% release) are initially present. Females may fail to mate or mate once in their life with a male of a given genotype, according to its frequency in the male population, chosen randomly and with replacement such that males may mate multiple times. The number of eggs from each mated female is multiplied by the egg production of the male relative to the wild-type male, to account for experimental observations of reduced egg production from SDGD fathers. The eggs may hatch or not, with a probability that depends on the product of larval hatching values from the mother and father, relative to wild type. To start the next generation, 600 larvae are randomly selected, unless fewer than 600 larvae have hatched, in which case the smaller amount initiates the next generation, following the experiment. The probability of subsequent survival to adulthood is assumed to be equal across genotypes. Assuming very large population sizes allows results for the genotype frequencies that are indistinguishable from the deterministic model. For the deterministic egg count, we use the large population limit of the stochastic model.

Population dynamics model (continuous time). To model changing population sizes in the field (for Supplementary Fig. 1), we use a continuous-time population dynamics

model with one life stage and logistic density dependence in the recruitment rate based on models developed previously^{36,37}. Here n(t) represents the abundance of adult individuals, f(t) and m(t) represent the total abundances of adult females and males, and $f_{ij,pq}(t)$ and $m_{ij,qY}(t)$ are the genotype abundances where, as above, the first set of indices denotes alleles at the target site and the second set denotes the sex chromosomes. Populations are normalized to the prerelease wild-type population size such that n(t=0)=1, and time is continuous and measured in generations. The dynamics of the total population size are given by the following differential equation:

$$\frac{dn(t)}{dt} = 2\left(\frac{R_{\rm m}}{1 + 2(R_{\rm m} - 1)\bar{w}_{\rm m}(t)\bar{w}_{\rm f}(t)f(t)}\right)\bar{w}_{\rm m}(t)\bar{w}_{\rm f}(t)f(t) - n(t)$$

The total recruitment rate of adults incorporates a density-dependent factor (the term in parentheses) based on Deredec et al.⁵ and depends on the total number of females, f(t), multiplied by the average female fitness, $\bar{w}_{\rm f}(t) = \sum_{i=1}^3 \sum_{j=i}^3 \sum_{pq=xx;XX;XX} (w_{ij,pq}f_{ij,pq}(t)/f(t))$. Because SDGD males may have reduced fertility, the recruitment rate is also dependent on the average male fitness, $\bar{w}_{\rm m}(t) = \sum_{i=1}^3 \sum_{j=i}^3 \sum_{q=x,X} (w_{ij,qY}m_{ij,qY}(t)/m(t))$ (we assume that the number of males is not limiting and that all males participate in mating). $R_{\rm m}$ is the intrinsic growth rate of the population per generation at low density.

The equations for the individual genotype populations for females and males are

$$\begin{array}{l} \frac{\eta_{ipq}(t)}{dt} = 2 \begin{pmatrix} \frac{R_m \bar{w}_m(t) \bar{w}_i(t) f(t)}{1+2(R_m-1) \bar{w}_m(t) \bar{w}_i(t) f(t)} \end{pmatrix} \begin{pmatrix} 1 - \frac{\delta_{ij}}{2} \end{pmatrix} \begin{pmatrix} 1 - \frac{\delta_{jq}}{2} \end{pmatrix} \\ \begin{pmatrix} e_{i,p}(t) s_{j,q}(t) + e_{j,p}(t) s_{i,q}(t) + e_{i,q}(t) s_{j,p}(t) + e_{j,q}(t) s_{i,p}(t) \end{pmatrix} - f_{ij,pq}(t) \end{array}$$

$$\frac{dm_{ij,qY}(t)}{dt} = 2 \left(\frac{R_{m} \bar{w}_{m}(t) \bar{w}_{i}(t) f(t)}{1 + 2(R_{m} - 1) \bar{w}_{m}(t) \bar{w}_{i}(t) f(t)} \right) \left(1 - \frac{\delta_{ij}}{2} \right) \\ \left(e_{i,q}(t) s_{j,Y}(t) + e_{j,q}(t) s_{i,Y}(t) \right) - m_{ij,qY}(t)$$

Egg and sperm proportions $e_{k,l}(t)$ and $s_{k,l}(t)$ are as defined in equations (1) and (2) in the discrete-generation model above, with $f_{ij,pq}(t)$ and $m_{ij,pq}(t)$ instead of frequencies $F_{ij,pq}(t)$ and $M_{ij,pq}(t)$.

All calculations were carried out using Wolfram Mathematica³⁸.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

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The full sequence of vectors is provided through the NCBI database. The GenBank accession codes for vectors p172 and p182–244 are MT270142 and MT270141, respectively. Sanger sequencing of vector p182–244 is available as Supplementary Information. Additional vector sequences are provided as Supplementary Note.

References

 Fuchs, S., Nolan, T. & Crisanti, A. Mosquito transgenic technologies to reduce *Plasmodium* transmission. *Methods Mol. Biol.* 923, 601–622 (2013).

- Volohonsky, G. et al. Tools for Anopheles gambiae transgenesis. G3 5, 1151–1163 (2015).
- Scott, T. W. Containment of arthropod disease vectors. *ILAR J.* 46, 53–61 (2005).
- Akbari, O. S. et al. Safeguarding gene drive experiments in the laboratory. Science 349, 927–929 (2015).
- Knudsen, S. Promoter2.0: for the recognition of PolII promoter sequences. Bioinformatics 15, 356–361 (1999).
- Reese, M. G., Harris, N. L. & Eeckman, F. H. Large scale sequencing specific neural networks for promoter and splice site recognition. In *Proceedings of the Pacific Symposium on Biocomputing* (eds. Hunter, L. & Klein, T.) 737–738 (World Scientific, 1996).
- Schneider, C. A., Rasband, W. S. & Eliceiri, K. W. NIH Image to ImageJ: 25 years of image analysis. *Nat. Methods* 9, 671–675 (2012).
- Mollahosseini, A. et al. A user-friendly software to easily count Anopheles egg batches. Parasit. Vectors 5, 122 (2012).
- Deredec, A., Burt, A. & Godfray, H. C. J. The population genetics of using homing endonuclease genes in vector and pest management. *Genetics* 179, 2013–2026 (2008).
- Beaghton, A., Beaghton, P. J. & Burt, A. Vector control with driving Y chromosomes: modeling the evolution of resistance. *Malar. J.* 16, 286 (2017).
- Beaghton, A. et al. Requirements for driving antipathogen effector genes into populations of disease vectors by homing. *Genetics* 205, 1587–1596 (2017).
- 38. Wolfram Research, Inc. Mathematica (Wolfram Research, Inc., 2019).

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Author contributions

A.S., A.M.H., T.N. and A.C. designed the research, and A.S., A.M.H., R.G., K.K. and C.T. performed the research. A.K.B. and A.B. performed modeling; D.M., M.G. and G.M. performed insect husbandry; A.S., A.M.H. and A.C. analyzed data; and A.S. and A.C. wrote the paper, with input from all authors.

Competing interests

The authors declare no competing interests.

Additional information

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For	all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.
n/a	Confirmed
	\bigtriangledown The exact sample size (<i>n</i>) for each experimental group/condition, given as a discrete number and unit of measurement
	A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
	The statistical test(s) used AND whether they are one- or two-sided Only common tests should be described solely by name; describe more complex techniques in the Methods section.
\ge	A description of all covariates tested
	A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
	A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
	For null hypothesis testing, the test statistic (e.g. <i>F</i> , <i>t</i> , <i>r</i>) with confidence intervals, effect sizes, degrees of freedom and <i>P</i> value noted <i>Give P values as exact values whenever suitable</i> .
\ge	For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
\boxtimes	For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
\boxtimes	Estimates of effect sizes (e.g. Cohen's <i>d</i> , Pearson's <i>r</i>), indicating how they were calculated
	Our web collection on statistics for biologists contains articles on many of the points above

Software and code

Policy information about availability of computer code		
Data collection	No customized software was used	
Data analysis	No customized software was used	

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All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

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All data generated or analysed during this study are already included in this published article (and its supplementary information files).

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

🔀 Life sciences

Behavioural & social sciences

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Life sciences study design

Sample size	Sample size was chosen consistent with the previous literature reporting similar assays . Sample size was maximised within the feasibility of performing biological assays with live insects. Starting frequencies in population experiments were chosen in order to minimise the effect of stochastic noise.
Data exclusions	No data exclusion
Replication	Biological replicates were performed as indicated int eh main text
Randomization	Randomization was not appropriate for this experimental approach
Blinding	Blinding was not performed in this study. Many genotypes had distinguishing physical characteristics

All studies must disclose on these points even when the disclosure is negative.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems Methods Involved in the study Involved in the study n/a n/a \boxtimes Antibodies \mathbf{X} ChIP-seq Eukaryotic cell lines \boxtimes \boxtimes Flow cytometry \boxtimes Palaeontology \boxtimes MRI-based neuroimaging Animals and other organisms \ge Human research participants \boxtimes Clinical data

Animals and other organisms

Policy information about <u>stud</u>	ies involving animals; ARRIVE guidelines recommended for reporting animal research	
Laboratory animals	For the purpose of giving mosquitoes a bloodmeal anesthetized mice (CD1 strain) were used. All animal work was conducted according to UK Home Office Regulations and approved under Home Office License PPL 70/8914.	
Wild animals	This study did not involve wild animals	
Field-collected samples	This study did not involve field-collected samples.	
Ethics oversight	For the purpose of giving mosquitoes a bloodmeal anesthetized mice (CD1 strain) were used. All animal work was conducted according to UK Home Office Regulations and approved under Home Office License PPL 70/8914.	

Note that full information on the approval of the study protocol must also be provided in the manuscript.