Targeting Sex Determination to Suppress Mosquito Populations

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Abstract

Each year, hundreds of millions of people are infected with arboviruses such as dengue, yellow fever, chikungunya, and Zika, which are all primarily spread by the notorious mosquito *Aedes aegypti*. Traditional control measures have proven insufficient, necessitating innovations. In response, here we generate a next generation CRISPR-based precision-guided sterile insect technique (pgSIT) for *Aedes aegypti* that disrupts genes essential for sex determination and fertility, producing predominantly sterile males that can be deployed at any life stage. Using mathematical models and empirical testing, we demonstrate that released pgSIT males can effectively compete with, suppress, and eliminate caged mosquito populations. This versatile species-specific platform has the potential for field deployment to effectively control wild populations of disease vectors.

Introduction

The mosquito *Aedes aegypti* is the principal vector of several deadly viruses, including dengue, yellow fever, chikungunya, and Zika. Unfortunately, its geographic spread has been exacerbated by climate change, global trade, and its preference for anthropogenic habitats, such that 3.9 billion people are presently at risk (Brady et al., 2012). Traditionally, mosquito control has been achieved through insecticides, though developing resistance have rendered them less effective. Moreover, insecticides are harmful to non-target species, such as insect pollinators, and direct exposure can be dangerous for humans and pets (Hemingway et al., 2002; Lu et al., 2020; Sánchez-Bayo, 2021; Trdan, 2016). Alternative integrated approaches for mosquito control include preventative measures designed to reduce mosquito populations through habitat reduction or larval predation, though these can also be impractical or introduce invasive species. Notwithstanding, the continuous expansion of this invasive mosquito species demonstrates that current measures are insufficient.

Alternative genetic biological approaches have been developed to control specific species, including the sterile insect technique (SIT), the Wolbachia-mediated incompatible insect technique (WIIT), \mathcal{Q} -specific release of insects carrying a dominant lethal gene (fsRIDL), inherited female elimination by genetically encoded nucleases to interrupt alleles (Ifegenia), gene drive and precision-guided SIT (pgSIT) (Knipling 1955; Laven 1967; Spinner et al. 2022; Smidler et al. 2023; Li et al. 2017; Kandul et al. 2019; Li et al. 2021). Here, radiation-sterilized males $(\mathcal{J}'s)$, or $\mathcal{J}'s$ harboring *Wolbachia*, are repeatedly mass released to mate with wild females ($\mathcal{Q}'s$) which typically mate only once, thereby reducing mosquito populations over time. The fsRIDL can genetically sex-sort \mathcal{A} 's, and can be released as eggs. If genia disrupts a female-essential gene, femaleless (fle), demonstrating complete genetic sexing via heritable daughter gynecide. Gene drives leverage the CRISPR based precise DNA cutting and repair to spread into populations. Two types of gene drives have been engineered: suppression drives collapses a population, whereas modification drives spread cargo genes that can potentially block the chain of transmission of a targeted disease (Champer et al., 2016; Marshall et al., 2019; Raban and Akbari, 2017; Wang et al., 2021). The pgSIT works by simultaneously disrupting genes essential for Qviability and \mathcal{J} fertility during development to genetically generate sex-sorted and viable sterilized \mathcal{S} 's.

Previously, we showed that disrupting a \bigcirc -specific flight muscle gene (fem-myo) induced a flightless \bigcirc -specific phenotype in *Ae. aegypti* \bigcirc 's (Li et al., 2021). However, this approach would still result in viable flightless transgenic \bigcirc 's being released which may be less desirable than removing females altogether. To further build upon the system, we instead cause \bigcirc lethality, simplifying the process of generating pgSIT \bigcirc 's for release. Our herein-developed next-generation pgSIT approach simultaneously disrupts three genes in *Ae. aegypti: doublessex (dsx)*, *intersex (ix)*, and β -Tubulin 85D (β Tub), to induce \bigcirc lethality, or intersex (\boxdot) transformation, and near-complete \bigcirc/\bigcirc sterility. We demonstrate that pgSIT \bigcirc 's are competitive and multiple releases can suppress cage populations. Notably, mathematical modeling indicates that this technology can control insect populations effectively over a wide range of realistic performance parameters and fitness costs. Overall, this next-generation pgSIT approach provides a viable option for the release of sterile males, bringing the field significantly closer to deployable CRISPR-based vector control strategies. Additionally, we also develop a rapid diagnostic assay to detect pgSIT transgene presence that could be used alongside field tests. Taken together, we have demonstrated that pgSIT could be designed to target essential sex determination and fertility

genes in *Ae. aegypti* to produce sterilized males that can suppress and eliminate caged populations thereby providing an alternate system for controlling this deadly vector.

Results

Disrupting sex-determination and fertility

To induce \mathcal{Q} -specific lethality and/or \mathbb{Z} transformation in *Ae. aegypti*, we targeted either the *dsx* (AAEL009114) or ix (AAEL010217) genes, both of which are associated with sex determination and differentiation in insects (Chase and Baker, 1995; Garrett-Engele et al., 2002; Kyrou et al., 2018; Mysore et al., 2015; Salvemini et al., 2011; Scali et al., 2005; Trujillo-Rodríguez et al., 2021). The dsx gene is located on chromosome 1R and is alternatively spliced to produce two Qspecific $(dsxF_1 \text{ and } dsxF_2)$ and one \bigcirc -specific (dsxM) isoforms (Salvemini et al., 2011). Exon 5b is retained in both \bigcirc -specific $dsxF_1$ and $dsxF_2$ transcripts and is spliced out in the \bigcirc -specific dsxMtranscript. Therefore, to disrupt both \mathcal{Q} -specific dsx isoforms, we selected four gRNAs that target exon 5b (Supplementary Table 1) and assembled the pBac gRNA^{dsx} plasmid (Fig. 1A). The ix gene (AAEL010217) is located on chromosome 3L and is orthologous to the *ix* gene required for ♀-specific sexual differentiation in *Drosophila melanogaster* (*Chase and Baker*, 1995). While the D. melanogaster ix does not undergo sex-specific alternative splicing and is expressed in both sexes, *ix* interacts with *dsxF*, but not *dsxM*, resulting in \mathcal{Q} -specific activity (Garrett-Engele et al., 2002). To disrupt ix, we engineered a gRNA^{ix} cassette harboring two gRNAs targeting ix exon 1 (Fig. 1A, Supplementary Table 1). To determine the effects of targeting these genes, multiple separate gRNA^{dsx} and gRNA^{ix} strains were generated and assessed by crossing with Nup50-Cas9 (hereafter, Cas9) (Li et al., 2021). To explore potential effects of maternal Cas9 carryover, all genetic crosses were conducted reciprocally (Fig. 1B). Remarkably, the disruption of either dsx or *ix* transformed all trans-hemizygous Q's into intersexes (\mathbb{Z} 's), independent of whether Cas9 was inherited from the mother (i.e., \bigcirc Cas9), or father (i.e., \bigcirc Cas9). These gRNA^{dsx}/+; Cas9/+ \square exhibited multiple malformed morphological features, such as mutant maxillary palps, abnormal genitalia, and malformed ovaries (Fig. 1D, Supplementary Fig. 1, Supplementary Table 2). Notably, only ~50% of $gRNA^{dsx}/+$; Cas9/+ \mathbb{Z} 's were able to blood feed, producing very few unhatchable eggs. While a similar fraction of $gRNA^{ix}/+$; Cas9/+ \mathbb{Z} were able to blood feed, the fed Z's laid hatchable eggs (Fig. 1C, Supplementary Table 3). Sanger sequencing confirmed expected mutations at the dsx and ix loci (Supplementary Fig. 2). Importantly, we did not observe changes in the ear anatomy, wing beat frequency (WBF), or phonotactic behaviors in either $gRNA^{dsx}/+$; Cas9/+ \Im 's or $gRNA^{ix}/+$; Cas9/+ \Im 's suggesting that the trans-hemizygous \Im 's have normal courtship behavior (Supplementary Fig. 3, Supplementary Table 3). Taken together, these findings reveal that \mathbb{Z} transformation can be efficiently achieved by disrupting either dsx or ix genes, while the \mathcal{Q} or \mathbb{Z} sterility requires the *dsx* disruption.

Multiplexed target gene disruption

To guide CRISPR-mediated disruption of multiple genes essential for \bigcirc -specific viability and \bigcirc -specific fertility, we engineered the $gRNA^{dsx,ix,\beta Tub}$ construct which combines the effective gRNAs described above (Supplementary Figure 2). More specifically, this construct harbors six gRNAs: three gRNAs targeting the dsx exon 5b; one gRNA targeting the ix exon 1; and two gRNA targeting the β -Tubulin 85D (β Tub, AAEL019894) exon 1 (Li et al., 2021) (Fig. 2A, Supplementary Table 1). The β Tub gene is specifically expressed in mosquito testes (Akbari et al., 2013; Degner et al., 2019; Gamez et al., 2020) and is essential for sperm maturation, elongation, and motility (Chen et al., 2021; Li et al., 2021). Three independent homozygous

 $gRNA^{dsx,ix,\betaTub}$ strains were established and reciprocally crossed to the homozygous *Cas9* strain to measure survival, fitness, and fertility of generated pgSIT (i.e., trans-hemizygous) progeny (Fig. 2B-C, Supplementary Fig. 4). To determine transgene integration sites, we performed genome sequencing of $gRNA^{dsx,ix,\betaTub}\#1$ (hereafter, $gRNA^{dsx,ix,\betaTub}$) and found that this strain harbored a single copy of the transgene inserted on chromosome 3 (Supplementary Fig. 5). Using these sequencing data, we also validated the presence of diverse mutations at the intended target sites (Supplementary Fig. 6). RNA sequencing also revealed the presence of mutations in the target RNA sequences (Supplementary Figs. 7-9, Supplemental Tables 10-11). These findings reveal that single copies of *Cas9* and *gRNA*^{dsx,ix,\betaTub} can induce mutagenesis at six separate DNA targets simultaneously, thereby providing desired pgSIT phenotypic redundancy for increased resilience.

pgSIT \bigcirc 's and \square 's are incapacitated

We next tested the simultaneous disruption of dsx, ix, and βTub , by reciprocally crossing $gRNA^{dsx,ix,\betaTub}$ with *Cas9*, which resulted in the death of most pgSIT \bigcirc 's and \square 's, as no \bigcirc 's and only 5–6% \square 's survived to adulthood (Fig. 2C). Interestingly, around 95% of pgSIT \square 's failed to eclose, independent of whether Cas9 was inherited from the mother or father (termed $pgSIT^{\oplus Cas9}$ or $pgSIT^{\oplus Cas9}$ \square 's, respectively, Supplementary Table 4). The pgSIT \square 's that did eclose had more pronounced abnormal phenotypes than \square 's generated from dsx or ix single-gene disruption (Fig. 1D, Supplementary Fig. 1) and were sterile (Fig. 2C, Supplementary Table 4). Moreover, the vast majority of adult pgSIT \square 's had reduced fitness as they could not blood feed (Fig. 2D, Supplementary Table 5), did not fly as frequently (Fig. 2E, Supplementary Table 6), had smaller abdomens and wings (Supplementary Fig. 1, Supplementary Table 7). Taken together, these results demonstrate that *Cas9* in combination with $gRNA^{dsx,ix,\betaTub}$ induces either the lethality or transformation of pgSIT \bigcirc 's into sterile unfit \square 's.

pgSIT \mathcal{J} 's are mostly sterile with normal longevity and mating ability

We found that ~94% of the eclosed pgSIT mosquitoes were 3° 's due to the majority of the \mathbb{P} 's not surviving to adulthood (Fig. 2C, Supplementary Table 4). Both $pgSIT^{\oplus cas9}$ and $pgSIT^{\oplus cas9} 3^{\circ}$'s were also nearly completely sterile (Fig. 2C, Supplementary Table 4). Surprisingly, despite our assumption that $pgSIT^{\oplus cas9} 3^{\circ}$ and $pgSIT^{\oplus cas9} 3^{\circ}$ would be roughly equivalent, the $pgSIT^{\oplus cas9} 3^{\circ}$'s showed less flight activity than WT (Fig. 2E, Supplementary Table 6), and $pgSIT^{\oplus cas9} 3^{\circ}$'s developmental times were slightly delayed (Supplementary Fig. 11D). Nevertheless, the longevity of pgSIT 3° 's was not significantly different from WT 3° 's (Supplementary Fig. 11, Supplementary Table 7), and both $pgSIT^{\oplus cas9} 3^{\circ}$ and $pgSIT^{\Im cas9} 3^{\circ}$ were competitive and mated with WT 9° 's to effectively suppress the fertility and re-mating ability of WT 9° 's (Fig. 2F, Supplementary Table 8). Taken together, these findings reveal that maternal deposition of Cas9 may induce some fitness costs, however these could be avoided by using a paternal source of Cas9, and despite these observed fitness effects, both $pgSIT^{\oplus cas9} 3^{\circ}$ and $pgSIT^{\Im cas9} 3^{\circ}$ were competitive and both types could likely be capable of inducing population suppression.

pgSIT ♂'s induce population suppression

To further explore whether pgSIT \Diamond 's are competitive enough to mate with WT \heartsuit 's in the presence of WT \Diamond 's to suppress populations, we conducted multigenerational, population cage experiments. At each generation, adult $pgSIT^{\heartsuit cas9}$ or $pgSIT^{\circlearrowright Cas9}$ and WT \Diamond ' mosquitos were released into cages using several introduction frequencies (pgSIT:WT - 1:1, 5:1, 10:1, 20:1, and

40:1; Fig. 3A and Supplementary Table 9). Remarkably, both $pgSIT^{\bigcirc cas9} \circlearrowleft$ or $pgSIT^{\bigcirc Cas9} \circlearrowright$ behaved similarly, and the high-release ratios (20:1, 40:1) eliminated all populations in five generations (Fig. 3B-C). With lower release ratios of 10:1 and 5:1, we also observed significant suppression of the rates of laid and hatched eggs in the cages. No effect was observed in control cages harboring WT mosquitoes with only Cas9 or gRNA mosquitoes (Fig. 3B-C and Supplementary Table 9). To confirm the sterility of pgSIT \circlearrowright 's, hatched larvae were screened for the presence of transgenesis markers each generation. In only 3 out of 30 experimental cages, we scored some larvae harboring transgene markers, though this was very few, indicating that the majority of $pgSIT \checkmark$ were indeed sterile (Supplementary Table 9). These findings confirmed the ability of both generated $pgSIT^{\bigcirc cas9}$ and $pgSIT^{\bigcirc cas9} \circlearrowright$'s to compete with WT \circlearrowright 's for mating with WT \bigcirc 's to gradually suppress and eliminate targeted caged populations.

Theoretical performance of pgSIT in a wild population

To explore the effectiveness of non-optimal pgSIT gene disruption on population suppression outcomes, we simulated releases of pgSIT eggs into a population of 10,000 Ae. aegypti adults (Fig. 4). Weekly releases of up to 500 pgSIT eggs (\bigcirc and \bigcirc) per wild-type adult (\bigcirc and \bigcirc) were simulated over 1-52 weeks. The scale of releases was chosen considering adult release ratios of 10:1 are common for sterile \bigcirc mosquito interventions (Carvalho et al., 2015) and \bigcirc Ae. aegypti produce >30 eggs per day in temperate climates (Mordecai et al., 2019). We assume released eggs have the same survival probability as wild-laid eggs; however if released eggs do have higher mortality, this would be equivalent to considering a smaller release. A preliminary sensitivity analysis was conducted to determine which parameters population suppression outcomes are most sensitive to (Supplementary Fig. 12). The analysis considered probability of elimination (the percentage of 60 stochastic simulations that result in Ae. aegypti elimination) and window of protection (the time duration for which \geq 50% of 60 stochastic simulations result in \geq 90% suppression of the Ae. aegypti population) as outcomes, and explored how these vary with number of releases, size of releases, release interval, gRNA cutting rate, Cas9 maternal deposition rate, pgSIT \mathcal{A} fertility and mating competitiveness, and pgSIT \mathcal{Q} viability. Suppression outcomes were found to be most sensitive to release schedule parameters (number, size and interval of releases), δ fertility and \mathcal{Q} viability. Model-predicted efficacy of pgSIT-induced population suppression was then explored in more detail as a function of these parameters through a series of simulations depicted in Fig. 4. Results from these simulations reveal that a window of protection exceeding two years (a common benchmark for interruption of disease transmission) is generally achieved for pgSIT \mathcal{Q} viability between 0 (complete inviability) and 0.15, and pgSIT \mathcal{A} fertility between 0 (complete sterility) and 0.15, provided the release scheme exceeds ~23 releases of \geq 100 pgSIT eggs per wild adult (Fig. 4A). Achieving a \geq 90% probability of elimination places slightly tighter restrictions on \mathcal{Q} viability and \mathcal{J} fertility - a safe ballpark being \mathcal{Q} viability and \mathcal{J} fertility both in the range 0-0.10, given a release scheme of ~26 releases of 250 pgSIT eggs per wild adult (Fig. 4B). These results suggest a target product profile for pgSIT to be \mathcal{Q} viability and β fertility both in the range 0-0.10. The pgSIT strain described here falls well within these bounds, with \mathcal{Q} viability of 0 and \mathcal{J} fertility of ~0.01.

Mitigation tools for rapid transgene detection

Being able to rapidly identify transgenes in the environment in mixtures of captured mosquitoes will be important for field applications of genetic biocontrol tools. While PCR-based tests do exist, these may be difficult to implement in the field. Therefore, developing an assay with a

simple readout would be desired. To generate a rapid assay for transgene detection that could be used alongside future pgSIT field trials, we adapted the sensitive enzymatic nucleic acid sequence reporter (SENSR) (Brogan et al., 2021a) to rapidly detect pgSIT DNA fragments in *Cas9* and *gRNA*^{dsx,ix,\betaTub} constructs in mosquitoes. In SENSR, the presence of transgenic sequences triggers the activation of the *Ruminococcus flavefaciens* CasRx/gRNA collateral activity that cleaves a quenched probe, resulting in a rapid fluorescent readout (Brogan et al., 2021b; Dalla Benetta et al., 2023) (Brogan et al., 2021a). Multiple ratios of trans-hemizygous *pgSIT* to WT mosquitoes - 0:1; 1:1; 1:5; 1:10; 1:25; 1:50 and a no template control (NTC) were used to challenge SENSR. We found that SENSR effectively detected transgenic sequences from both *Cas9* and *gRNA*^{dsx,ix,βTub} constructs even at the lowest dilution, and no signal was observed in the WT-only and NTC controls (Supplementary Fig. 13A). Notably, in every tested ratio, SENSR rapidly detected the transgenic sequence, reaching the half maximum fluorescence (HMF) in less than 20 min (Supplementary Fig. 13B). Taken together, these results validated SENSR as a rapid and sensitive assay for detection of transgenic sequences in pools of mosquitoes which may be an instrumental tool for future field trials.

Discussion

To develop a robust and stable population control technology for addressing insect disease vectors, we engineered a next-generation pgSIT system that induced near complete \bigcirc -specific lethality and \eth sterility by simultaneously targeting the *dsx*, *ix*, and β *Tub* genes. Our cage experiments indicate that releases of pgSIT \eth 's can induce robust population suppression and elimination, and mathematical models reinforce this finding. To support future field trials of this and other trangene-based technologies, we also developed a rapid and sensitive method, SENSR, for identifying pgSIT transgenes.

Overall, the released pgSIT males were competitive in our cage populations, as repeated releases resulted in population elimination at several different introduction frequencies. That said, we did observe some limitations. Firstly, we determined that pgSIT males were not 100% sterile, with an estimated ~1% still producing some progeny, this may stem from using fewer gRNAs to target βTub , as our previous system that saw complete sterility used four gRNAs while here we used only two (Li et al., 2021). That said, it would not be difficult to build new strains that harbor additional gRNAs targeting βTub or other genes important for male fertility to ensure sterility is achieved. Even though mutations could potentially become fixed within the natural population, the likelihood of off-target effects becoming fixed within the population is exceedingly low. To mitigate potential employed **CHOPCHOP** negative impacts. we V3.0.0 (https://chopchop.cbu.uib.no) for the selection of gRNAs, specifically to minimize the occurrence of genomic off-target cleavage events. Furthermore, our releasing process will be carried out in multiple rounds. Even in the event that an undesired mutant is introduced into the local population, it will either be completely eradicated through subsequent rounds of releases or be naturally eliminated through the process of natural selection over time. Secondly, we also noticed some reduced fitness parameters when Cas9 was inherited maternally, which may result from an overabundance of Cas9 present in the egg (Kandul et al., 2020). That said, these maternal fitness effects can be avoided by ensuring Cas9 is inherited paternally as pgSIT phenotypes can be achieved when Cas9 is inherited from either the father or mother. Moreover, the longevity was not significantly altered, and released pgSIT males still performed well in the population cages, which indicates that these fitness costs and lack of complete male sterility could be tolerated for population suppression. Even in the presence of pgSIT construct imperfections, our mathematical modeling showed that population suppression is very promising, both in terms of the potential to eliminate a mosquito population, or to suppress it to an extent that would largely interrupt disease transmission. Our models determined that having \mathcal{Q} viability and \mathcal{J} fertility both in the range 0-10% provides a reasonable target product profile for pgSIT - criteria that the construct described in this study easily satisfies. Previously published analyses (Li et al., 2021) have also shown fitness costs associated with pgSIT constructs are well-tolerated in the range 0-25%. Overall, our results indicate that perfection is not required for an effective pgSIT system, though improvements to the presented and other systems could allow for reduced introduction ratios.

Before a possible field deployment, there are still several considerations that must be addressed for this pgSIT system (Kandul et al., 2019a). Firstly, our herein-presented system requires a genetic cross to produce the releasable insects, similarly to previous pgSIT systems (Kandul et al., 2022, 2019b; Li et al., 2021) that will require a scalable genetic sexing system. While sophisticated mechanical systems that exploit sex-specific morphological differences have been generated in previous work (Crawford et al., 2020) and are available at www.seneciorobotics.com, these approaches may be cost prohibitive. Therefore, there exists a need to develop inexpensive and scalable genetic sexing systems that could be combined with pgSIT to eliminate its reliance on mechanical adult sexing approaches. Alternatively, if a temperature inducible Cas9 strain was generated in Ae. aegypti then it may be possible to generate a temperature inducible pgSIT (TI-pgSIT) system as was done previously (Kandul et al., 2021) and this would alleviate the need for an integrated genetic sexing approach. Secondly, studies are required to determine the survival and mating competitiveness of released pgSIT males under field conditions, and to optimize their release protocol. Thirdly, although pgSIT is self-limiting and inherently safe, it does require genetic modification, so regulatory authorizations will be necessary prior to implementation. Despite this, we anticipate that obtaining such authorizations will not be insurmountable.

Altogether, we demonstrate that removing females by disrupting sex determinate genes is possible with pgSIT, which can inform the development of future such systems in related species. The self-limiting nature of pgSIT offers a controllable alternative to technologies such as gene drives that will persist and uncontrollably spread in the environment. Moving forward, pgSIT could offer an efficient, scalable, and environmentally friendly next-generation technology for controlling wild mosquito populations, leading to widespread prevention of human disease transmission.

Methods

Mosquito rearing and maintenance

Ae. aegypti mosquitoes used in all experiments were derived from the *Ae. aegypti* Liverpool strain, which was the source strain for the reference genome sequence (Matthews et al., 2018). Mosquito rearing and maintenance were performed following previously established procedures (Li et al., 2021).

Guide RNA design and assessment

To induce \mathcal{Q} -specific lethality, *doublesex* (*dsx*, AAEL009114) and *Intersex* (*ix*, AAEL010217) genes were targeted for CRISPR/Cas9-mediated disruption. For each target gene, the DNA sequences were initially identified using reference genome assembly, and then genomic target sites were validated using PCR amplification and Sanger sequencing (see Supplementary Table 1 for primer sequences). CHOPCHOP V3.0.0 (https://chopchop.cbu.uib.no) was used to select the

gRNA target sites. In total, we selected 4 *dsx* gRNAs and 2 *ix* gRNAs, and assembled the *gRNA*^{*dsx*} and *gRNA*^{*ix*} constructs harboring the corresponding gRNAs. To confirm the activity of the gRNAs *in vivo*, we generated the transgenic strains harboring either the *gRNA*^{*dsx*} or *gRNA*^{*ix*} constructs, and crossed each *gRNA* strain to the *Cas9* strain (Li et al., 2021). We sequenced DNA regions targeted by each selected gRNA in F₁ trans-hemizygous (aka. *gRNA*/+; *Cas9*/+) mosquitoes to assess gRNA/Cas9-induced mutagenesis.

Genetic construct design and plasmid assembly

All plasmids were cloned using Gibson enzymatic assembly (Gibson et al., 2009). DNA fragments were either amplified from available plasmids or synthesized gBlocks[™] (Integrated DNA Technologies) using Q5 Hotstart Start High-Fidelity 2X Master Mix (New England Biolabs, Cat. #M0494S). The generated plasmids were transformed into Zymo JM109 chemically competent E. coli (Zymo Research, Cat # T3005), amplified, isolated (Zymo Research, Zyppy plasmid miniprep kit, Cat. #D4036), and Sanger sequenced. Final selected plasmids were maxiprepped (Zymo Research, ZymoPURE II Plasmid Maxiprep kit, Cat. #D4202) and sequenced thoroughly Oxford Nanopore Sequencing Primordium using at Labs (https://www.primordiumlabs.com). All plasmids and annotated DNA sequence maps are available at www.addgene.com under accession numbers: 200252 (gRNA^{dsx}, 1067B) 200251 $(gRNA^{ix}, 1055J), 200253 (gRNA^{dsx,ix,\betaTub}, 1067L)$ and 164846 (*Nup50-Cas9* or *Cas9*, 874PA).

Engineering transgenic strains

Transgenic strains were generated by microinjecting preblastoderm stage embryos (0.5 - 1 hr old) with a mixture of the *piggyBac* plasmid (100 ng/ul) and a transposase helper plasmid (*pHsp-piggyBac*, 100 ng/ul). Embryonic collection, microinjections, transgenic lines generation and rearing were performed following previously established procedures (Bui et al., 2020; Li et al., 2021, 2020).

Genetic assessment of gRNA strains

To evaluate the activity of $gRNA^{dsx}$ and $gRNA^{ix}$ strains, we reciprocally crossed hemizygous mosquitoes of each generated strains to the homozygous Nup-Cas9 strain (Li et al., 2021), and thoroughly examined F_1 trans-hemizygous \mathcal{Q} progeny (Supplementary Tables 2-3). Each of three established $gRNA^{dsx,ix,\beta Tub}$ strains was subjected to single-pair sibling matings over 5-7 generations to generate the homozygous stocks. Zygosity was confirmed genetically by repeated test crosses to WT. To generate trans-hemizygous progeny with a maternal *Cas9* (*pgSIT*^(Cas9)) and paternal*Cas9*(*pgSIT*^{<math>(Cas9)}), homozygous*gRNA*^{*dsx,ix,βTub*} <math>(cas9) is and (cas9) is were mated to homozygous *Cas9* (cas9) is a constrained of the paternal *Cas9* (*pgSIT*^(Cas9)), homozygous*gRNA*^{*dsx,ix,βTub*} <math>(cas9) is a constrained of the paternal *Cas9* (*pgSIT*^(Cas9)), homozygous*gRNA*^{*dsx,ix,βTub*} <math>(cas9) is a constrained to homozygous *Cas9* (cas9) is a constraine</sup></sup></sup></sup> and δ 's, respectively. Then, the generated trans-hemizygous mosquitoes were mated to WT mosquitoes of the opposite sex, and numbers of laid and hatched eggs were scored to measure the fecundity. Multiple genetic control crosses were also performed for comparisons: WT \Diamond x WT \Diamond ; WT \Diamond x Cas9 \heartsuit ; Cas9 \Diamond x WT \heartsuit ; gRNA \Diamond x Cas9 \heartsuit ; gRNA \heartsuit x Cas9 \Diamond ; gRNA \heartsuit x WT \Diamond ; and gRNA \bigcirc x WT \bigcirc . For the genetic cross, adult mosquitoes were allowed to mate in a cage for 4-5 days, then blood meals were provided, and eggs were collected and hatched. To calculate the normalized percentage of laid eggs per \mathcal{Q} (e.i. fecundity), the number of laid eggs per \mathcal{Q} for each genotype replicate was divided by the maximum number of laid eggs per \mathcal{D} (Supplementary Table 4). The percentage of egg hatching (i.e. fertility) was estimated by dividing the number of laid eggs by the number of hatched eggs. Egg-to-adult survival ratios were the number of laid eggs divided by the total number of eclosed adults. Larvae-to-adult survival rates were calculated by

dividing the number of eclosed adults by the number of larvae. Pupae-adult survival rates were calculated by dividing the number of pupae by the number of eclosed adults. Blood feeding rates were calculated by dividing the number of blood-fed \bigcirc 's or \square 's by the total number of \bigcirc or \square 's. Trans-hemizygous \square 's had many morphological features visibly different from those of WT \bigcirc 's. To assess differences in external and internal anatomical structures, we dissected twenty transhemizygous and control mosquitoes for each genotype and sex (\bigcirc , \bigcirc and \square) in 1% PBS buffer and imaged structures on the Leica M165FC fluorescent stereomicroscope equipped with a Leica DMC4500 color camera. The measurements were performed using Leica Application Suite X (LAS X by Leica Microsystems) on the acquired images (Supplementary Table 2).

Validation of induced mutations

To confirm that the \bigcirc lethality or transformation correlated with specific mutations at the target loci, *dsx* and *ix* gRNA targets were PCR amplified from the genomic DNA extracted from transhemizygous and control mosquitoes. PCR amplicons were purified directly or gel purified using Zymoclean Gel DNA Recovery Kit (Zymo Research Cat. #D4007) and then Sanger sequenced. Presence of induced mutations were inferred as precipitous drops in sequence read quality by analyzing base peak chromatograms (Supplementary Fig. 2). The primers used for PCR and sequencing, including gRNA target sequences, are listed in Supplementary Table 1. In addition, induced mutations and their precise localization to six target sites in *ix*, *dsx*, and β Tub genes were confirmed by sequencing genomes of gRNA^{dsx,ix,\betaTub}#1/Cas9 \Diamond 's and \Box 's using Oxford Nanopore (see below) and aligning them to the AaegL5 genome (GCF_002204515.2). Mutations in the DNA using Nanopore data and mutations in the RNA using RNAseq data were visualized using an integrated genome browser (Supplementary Figs. 7-9).

Ear Johnston's Organ microanatomy and Immunohistochemistry (IHC)

The heads of two-day old mosquitoes were removed, fixed in 4% PFA and 0.25% Triton X-100 PBS, and sectioned using a Leica VT1200S vibratome. The anti-SYNORF1 3C11 primary monoclonal antibody (AB_528479, 1:30, Developmental Studies Hybridoma Bank (DSHB), University of Iowa) and two secondary antibodies, Alexa Fluor 488-conjugated anti-mouse IgG (#A-11029, 1:300, Thermo Fisher) and anti-horseradish peroxidase 555 (anti-HRP, AB_2338959, 1:100, Jackson Immuno Research), were used for IHC. All imaging was conducted using a laser-scanning confocal microscope (FV3000, Olympus). Images were taken with a silicone-oil immersion $60 \times$ Plan-Apochromat objective lens (UPlanSApo, NA = 1.3) as previously described (Su et al., 2018).

Wing Beat Frequency (WBF) measurements

Groups of 30 mosquitoes were transferred to 15cm³ cages containing a microphone array (Knowles NR-23158-000) and provided with a source of 10% glucose water. After two days of LD entrainment, audio recordings of mosquito flight behavior were made using a Picoscope recording device (Picoscope 2408B, Pico Technology) at a sampling rate of 50 kHz. These recordings lasted 30 minutes during periods of peak activity during dusk (ZT12.5-13). Audio recordings were then processed and analyzed using the simbaR package in R (Somers et al., 2022; Su et al., 2020). Four separate cages of mosquitoes were tested for each genotype. Statistical comparisons were made using ANOVA on Ranks tests, followed by pairwise Wilcoxon tests with a Bonferroni correction applied for multiple comparisons.

Phonotaxis assays

Individual \Diamond 's were provided with a 475 Hz pure tone stimulus calibrated to 80 dB SPL (V \approx 5 x 10-4 ms⁻¹) provided for 30 seconds by a speaker. Prior to tone playback, a gentle puff of air was provided to each \Diamond to stimulate flight initiation. Mosquitoes were considered to demonstrate phonotaxis behaviors if they orientated to the speaker and then demonstrated abdominal bending behaviors. A video camera (GoPro) was used to record all responses to stimulation, with responses being scored as either 0 (no approach/no bending) or 1 (approach and bending). Mosquitoes were tested on two consecutive days, with mosquitoes showing behavior considered as a phonotactic response to stimuli on at least one test day being regarded as responders. Three repeats were conducted for each genotype (across 3 generations), with at least 10 \Diamond 's tested per repeat. Statistical comparisons were made using Chi-squared tests.

Transgene copy number and genomic integration site

To infer the transgene copy number(s) and insertion site(s), we performed Oxford Nanopore sequencing of the trans-hemizygous $gRNA^{dsx,ix\beta Tub} \#1/Cas9$ mosquitoes. Genomic DNA was extracted from 10 ♂'s and ♀'s using Blood & Cell Culture DNA Midi Kit (Qiagen, Cat# 13343) and the sequencing library was prepared using the Oxford Nanopore SQK-LSK110 genomic DNA by ligation kit. The library was sequenced for 72 hrs on MinION flowcell (R9.4.1) followed by base calling with Guppy v6.3.2 and produced 16.18 Gb of filtered data with Q score cutoff of 10. To determine transgene copy number(s), reads were mapped to the AaegL5 genome supplemented with transgene sequences $gRNA^{dsx,ix,\betaTub}$ (1067L plasmid) and Cas9 (874PA plasmid) using minimap2 and sequencing depth was calculated using samtools. To identify transgene integration sites, nanopore reads were mapped to the gRNA^{dsx,ix, β Tub} and Cas9 plasmids and the sequences that aligned to the expected regions between the *piggyBac* sites were parsed. These validated reads were then mapped to the AaegL5 genome (GCF 002204515.2) and sites of insertions were identified by examining the alignments in the Interactive Genomics Viewer (IGV) browser. The $gRNA^{dsx,ix,\beta Tub} \# I$ strain harbors a single copy of $gRNA^{dsx,ix,\beta Tub}$ inserted on chromosome 3 (+/+ orientation) between 2993 and 16524 at NC 035109.1:362,233,930 TTAA site (Supplementary Fig. 5). We also confirmed the previously identified insertion site of Cas9 (Li et al.. 2021): chromosome 3 (+/+)orientation) between 2933 and 14573 at NC 035109.1:33,210,107 TTAA.

Transcriptional profiling and expression analysis

To quantify target gene reduction and expression from transgenes as well as to assess global expression patterns, we performed Illumina RNA sequencing. We extracted total RNA using miRNeasy Mini Kit (Qiagen, Cat# 217004) from 20 sexed pupae: WT \bigcirc 's, WT \bigcirc 's, transhemizygous $pgSIT^{\bigcirc Cas9}$ \bigcirc 's, and $pgSIT^{\bigcirc Cas9}$ \bigcirc 's in biological triplicate (12 samples total), following the manufacturer's protocol. DNase treatment was conducted using DNase I, RNasefree (Thermo Fisher Scientific, Cat# EN0521), following total RNA extraction. RNA integrity was assessed using the RNA 6000 Pico Kit for Bioanalyzer (Agilent Technologies #5067-1513), and mRNA was isolated from ~1 µg of total RNA using NEB Next Poly (A) mRNA Magnetic Isolation Module (NEB #E7490). RNA-seq libraries were constructed using the NEBNext Ultra II RNA Library Prep Kit for Illumina (NEB #E7770) following the manufacturer's protocols. Libraries were quantified using a Qubit dsDNA HS Kit (Thermo Fisher Scientific #Q32854), and the size distribution was confirmed using a High Sensitivity DNA Kit for Bioanalyzer (Agilent Technologies #5067- 4626). Libraries were sequenced on an Illumina NextSeq2000 in paired end mode with the read length of 50 nt and sequencing depth of 20 million reads per library. Base calling and FASTQ conversion were performed with DRAGEN 3.8.4. The reads were mapped to

the AaegL5.0 (GCF_002204515.2) genome supplemented with the *Cas9* and *gRNA*^{dsx,ix, β Tub</sub> plasmid sequences using STAR aligner. On average, ~97.8% of the reads were mapped. Gene expression was then quantified using featureCounts against the annotation release 101 GTF downloaded from NCBI (GCF_002204515.2_AaegL5.0_genomic.gtf). TPM values were calculated from counts produced by featureCounts. PCA and hierarchical clustering were performed in R and plotted using the ggplot2 package. Differential expression analyses between pgSIT vs WT samples within each sex were performed with DESeq2 (Supplementary Fig. 13, Supplementary Tables 10-11). Gene Ontology overrepresentation analyses were done with the topGO package.}

Blood feeding assays

Ten mature $WT \ \$'s or pgSIT \mathbb{Z} 's (i.e $gRNA^{dsx+ix+\beta Tub}\#1/Cas9$) generated with Cas9 inherited from either the mother ($pgSIT^{\oplus Cas9}$) or father ($pgSIT^{\oplus Cas9}$) were caged with 100 mature $WT \$'s for two days to insure that every $WT \$ was mated before $WT \$'s were removed from all cages. Then, one anesthetized mouse (similar size and weight) was put into each cage for 20 minutes (from 9:00 am to 9:20 am) daily for five days, and two times were recorded for each mosquito $\$ or \mathbb{Z} . First, the 'time to bite' is the time interval between a mouse placed in a cage and $\$ ' $\$ landing on the mouse. Second, the 'feeding time' is the duration of blood feeding (Supplementary Table 5). When individual $\$ ' $\$'s finished blood feeding they were removed from the cage, while unfed $\$ ' $\$'s were kept in the cage and the blood feeding was repeated the next day. In total, each $\$ A' $\$ had 6000 seconds to initiate bite and feed during 5 days. To assess whether distance affects blood feeding of $pgSIT \$'s, we conducted the experiment in two sizes of cages: small (24.5 X 24.5 X 24.5 cm) and large (60 X 60 X 60 cm). All experiments were repeated 3 times independently.

Flight activity assays

We followed the protocol in our previous study (Li et al., 2021) to assess the flight activity of trans-hemizygous $pgSIT^{\oplus Cas9}$ (i.e., $gRNA^{dsx+ix+\beta Tub}\#1/Cas9$) mosquitoes (Supplementary Table 6).

Mating assays

We followed a previously described protocol (Li et al., 2021), to assess the mating ability of both types of pgSIT males, $pgSIT^{\oplus Cas9}$ and $Cas9 pgSIT^{\oplus Cas9}$, and confirm that prior matings with pgSIT \Diamond 's suppressed WT \bigcirc fertility (Supplementary Table 8).

Fitness parameters and longevity

To assess fitness of pgSIT mosquitoes, we measured developmental times, fertility, and longevity of WT, homozygous $gRNA^{dsx+ix+\beta Tub}\#1$, homozygous *Cas9*, and $pgSIT^{\oplus Cas9}$ and $pgSIT^{\oplus Cas9}$ mosquitoes. The previously published protocol was used to assess fitness parameters and longevity (Li et al., 2021). Briefly, all fitness parameters and longevity were assessed in three groups (3 replicates) of twenty individual mosquitoes (Supplementary Table 7).

Multigenerational population cage trials

To assess the competitiveness of pgSIT \Diamond 's, we performed discrete non-overlapping multigenerational population cage trials as described in our previous study (Li et al., 2021). Briefly, 3/4-day-old mature WT adult \Diamond 's were released along with mature (3/4 days old) pgSIT adult \Diamond 's at release ratios (pgSIT: WT): 1:1 (50:50), 5:1 (250:50), 10:1 (500:50), 20:1 (1000:50),

and 40:1 (2000:50), with three biological replicates for each release ratio (15 cages total). One hour later, 50 mature (3/4 days old) WT adult Q's were released into each cage. All adults were allowed to mate for 2 days. Q's were then blood fed and eggs were collected. Eggs were counted and stored for four days to allow full embryonic development. Then, 100 eggs were randomly selected, hatched, and reared to the pupal stage, and the pupae were separated into a and Qgroups and transferred to separate cages. Three days post eclosion, ratios (pgSIT: WT) of 50 (1:1), 250 (5:1), 500 (10:1), 1000 (20:1), and 2000 (40:1) age-matched pgSIT mature a adults were caged with these mature a's from 100 selected eggs. One hour later, mature Q's from 100 selected eggs were transferred into each cage. All adults were allowed to mate for 2 days. Q's were blood fed, and eggs were collected. Eggs were counted and stored for 4 days to allow full embryonic development. The remaining eggs were hatched to measure hatching rates and to screen for the possible presence of transformation markers. The hatching rate was estimated by dividing the number of hatched eggs by the total number of eggs. This procedure continued for all subsequent generations.

Mathematical modeling

To model the expected performance of pgSIT at suppressing and eliminating local Ae. aegypti populations, we used the MGDrivE simulation framework (Sánchez et al., 2019). This framework models the egg, larval, pupal, and adult mosquito life stages with overlapping generations, larval mortality increasing with larval density, and a mating structure in which \mathcal{Q} 's retain the genetic material of the adult \mathcal{J} with whom they mate for the duration of their adult lifespan. We used the same egg-to-adult survival rate for WT and pgSIT eggs. The inheritance pattern of the pgSIT system was modeled within the inheritance module of MGDrivE. This module includes parameters for \mathcal{Q} pupatory success (as a proxy for viability), and \mathcal{J} fertility and mating competitiveness. We simulated a randomly-mixing Ae. aegypti population of 10,000 adults and implemented the stochastic version of MGDrivE to capture chance effects that are relevant to suppression and elimination scenarios. Density-independent mortality rates for juvenile life stages were calculated for consistency with the population growth rate in the absence of densitydependent mortality, and density-dependent mortality was applied to the larval stage following Deredec et al. (Deredec et al., 2011). Sensitivity analyses were conducted with "window of protection" (mean duration for which the Ae. aegypti population is suppressed by at least 90%), probability of Ae. aegypti elimination, and reduction in cumulative potential for transmission as the outcomes, and model parameters sampled describing the construct (gRNA cutting rate and maternal deposition frequency), release scheme (number, size and frequency of releases), Qviability, and \mathcal{J} fertility and mating competitiveness. Model parameters were sampled according to a latin-hypercube scheme with 2^{16} parameter sets and 20 repetitions per parameter set. Regression models were fitted to the sensitivity analysis data using multi-layered perceptrons as emulators of the main simulation in predicting the window of protection outcome. Subsequent simulations focused on the most sensitive parameters from the sensitivity analysis (those describing the release scheme, \mathcal{Q} viability, and \mathcal{J} fertility) and were explored via factorial simulation design. Complete model and intervention parameters are listed in Supplementary Table 12-13.

Transgene detection method in mosquitoes

SENSR assays were performed as described previously (Brogan et al., 2021c) using a two-step nucleic acid detection protocol. Target sequences were first amplified in a 30 min isothermal

preamplification reaction using recombinase polymerase amplification (RPA). RPA primers were designed to amplify 30 bp gRNA spacer complement regions flanked by 30 bp priming regions from the transgenic elements (Cas9 coding sequence or gRNA scaffold for Cas9 or gRNA^{dsx,ix,\betaTub} construct, respectively) while simultaneously incorporating a T7 promoter sequence and "GGG" into the 5' end of the dsDNA gene fragment to increase transcription efficiency (Brieba et al., 2002). The RPA primer sequences and synthetic target sequences can be found in Supplementary Table 14. The RPA reaction was performed at 42°C for 30 min by using TwistAmp Basic (TwistDx #TABAS03KIT). The final conditions for the optimized (28.5% sample input) RPA protocol in a 10-µL reaction are as follows: 5.9 µL rehydration buffer (TwistDx), 0.35 µL primer mix (10 µM each), 0.5 µL MgOAc (280 mM), and 50 ng of genomic DNA. The RPA reaction was then transferred to a second reaction, termed the Cas Cleavage Reaction (CCR), which contained a T7 polymerase and the CasRx ribonucleoprotein. In the second reaction, in vitro transcription was coupled with the cleavage assay and a fluorescence readout using 6carboxyfluorescein (6-FAM). A 6-nt poly-U probe (FRU) conjugated to a 5' 6-FAM and a 3' IABlkFQ (Iowa Black Fluorescence Quencher) was designed and custom-ordered from IDT(Brogan et al., 2021d). Detection experiments were performed in 20-µL reactions by adding the following reagents to the 10 µL RPA preamplification reaction: 2.82 µL water, 0.4 µL HEPES pH 7.2 (1 M), 0.18 µL MgCl2 (1 M), 1 µL rNTPs (25 mM each), 2 µL CasRx (55.4 ng/µL), 1 µL RNase inhibitor (40 U/µL), 0.6 µL T7 Polymerase (50 U/µL), 1 µL gRNA (10 ng/µL), and 1 µL FRU probe (2 µM). Experiments were immediately run on a LightCycler 96 (Roche #05815916001) at 37°C for 60 min with the following acquisition protocol: 5 sec acquisition followed by 5 sec incubation for the first 15 min, followed by 5 sec acquisition and 55 sec incubation for up to 45 min. Fluorescence readouts were analyzed by calculating the backgroundsubtracted fluorescence of each timepoint and by subtracting the initial fluorescence value from the final value. Statistical significance was calculated using a one-way ANOVA followed by specified multiple comparison tests (n = 4). The detection speed of each gRNA was calculated from the Half-Maximum Fluorescence (HMF) (Brogan et al., 2021d). Raw SENSR assays data can be found in Supplementary Table 15.

We generated 12 gRNAs, six target regions within *Cas9* or *gRNA* transgenes, to assess the detection capabilities in a SENSR reaction for each selected gRNA using *Cas9* and *gRNA*^{dsx,ix,\betaTub} plasmids for detection of *Cas9* and *gRNA*^{dsx,ix,\betaTub} transgenic sequences, respectively. After identifying the best gRNA for detection of *Cas9* and *gRNA* sequences, we run SENSR of DNA preps made from *Ae. aegypti pgSIT*^{QCas9} and WT mosquitoes harboring both *Cas9* and *gRNA*^{dsx,ix,\betaTub} constructs at different ratios of *pgSIT*^{QCas9} to WT mosquitoes: 1:0; 1:1; 1:5; 1:10; 1:25; 1:50; 0:1; and No Template Control (NTC). We challenged the best gRNA (A4 and B3) in a SENSR assay and found that both gRNA detected transgenic DNA even at the lowest dilution and no signal was observed in both controls, only WT mosquitoes and NTC.

Statistical analysis

Statistical analysis was performed in JMP8.0.2 by SAS Institute Inc and Prism9 for macOS by GraphPad Software, LLC. We used at least three biological replicates to generate statistical means for comparisons. P values were calculated using a two-sided Student's t test with equal variance. The departure significance for survival curves was assessed with the Log-rank (Mantel-Cox) texts. Statistical comparisons of WBF for each sex were made using ANOVA on Ranks tests, followed by pairwise Wilcoxon tests with a Bonferroni correction applied for multiple

comparisons. Chi-squared test was used to assess significance of departure in phonotaxis assay. All plots were constructed using Prism 9.1 for macOS by GraphPad Software and then organized for the final presentation using Adobe Illustrator.

Ethical conduct of research

All animals were handled in accordance with the Guide for the Care and Use of Laboratory Animals as recommended by the National Institutes of Health and approved by the UCSD Institutional Animal Care and Use Committee (IACUC, Animal Use Protocol #S17187) and UCSD Biological Use Authorization (BUA #R2401).

Reporting summary

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

Data availability

Complete sequence maps assembled in the study are deposited at www.addgene.org under ID# 200252 ($gRNA^{dsx}$, 1067B), 200251 ($gRNA^{ix}$, 1055J) and 200253 ($gRNA^{dsx,ix,\betaTub}$, 1067L)). Illumina and Nanopore sequencing data have been deposited to the NCBI-SRA under BioProject ID PRJNA942966, SAMN33705824-SAMN33705835; PRJNA942966, SAMN33705934. Data used to generate figures are provided in the Supplementary Materials/Source Data files (Supplementary Tables 1-11). *Aedes* transgenic lines are available upon request. Source data is provided with this paper.

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Author Contributions

O.S.A, M.L., N.P.K. conceptualized and designed experiments; T.Y., L.A., S.G., E.D.B., D.J.B., performed molecular analyses; M.L., J.R.E., T.W., H.L., M.B. performed genetic experiments; Y.Z, N.D., C.M. performed behavioral experiments; Y.M.L., M.P.S., performed IHC and phonotaxis assays; H.M.S.C, J.M.M performed mathematical modeling; I.A. performed bioinformatics: O.S.A, N.P.K, M.L., R.S. analyzed and compiled the data. All authors contributed to writing and approved the final manuscript.

Competing interests

O.S.A. is a founder of Agragene, Inc. with equity interest. O.S.A., M.L., and N.P.K are founders of Synvect with equity interest. The terms of this arrangement have been reviewed and approved by the University of California, San Diego in accordance with its conflict of interest policies. All remaining authors declare no competing interests.

Additional information

The online version contains supplementary material available at XX.

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Figure 1. CRISPR/Cas9-mediated disruption of *dsx* **or** *ix* **affects female mosquito morphology and fecundity.** A Schematic map of gRNA genetic constructs. The $gRNA^{dxx}$ and $gRNA^{ix}$ constructs harbor a 3xP3-tdTomato marker and multiple gRNAs guiding Cas9 to the female-specific *doublesex* (*dsx*) gene or the female-active *intersex* (*ix*) gene, respectively. **B** A schematic of the genetic cross between the homozygous *Cas9* and hemizygous $gRNA^{dxx}/+$ or $gRNA^{ix}/+$ mosquitoes to generate trans-hemizygous F_1 females (\bigcirc 's). Reciprocal genetic crosses were established and two types of trans-hemizygous $F_1 \bigcirc$'s were generated: $gRNA/+; \bigcirc Cas9/+ \bigcirc$'s inherited a maternal Cas9; and $gRNA/+; \bigcirc Cas9/+ \bigcirc$'s inherited a paternal Cas9. Then, both trans-hemizygous \bigcirc 's were crossed to wildtype (WT) males (\bigcirc 's), and their fecundity was assessed. **C** A comparison of the fecundity and male ratio of trans-hemizygous, hemizygous *Cas9* or $gRNA \bigcirc$'s to those of WT \bigcirc 's. The bar plot shows means and one standard deviation (±SD).

The data presents one transgenic strain of each construct, $gRNA^{dsx}\#1$ and $gRNA^{ix}\#1$, as all strains induced similar results (all data can be found in Supplementary Table 3). **D** All $gRNA^{dsx}\#1/+$; Cas9/+ and $gRNA^{ix}\#1/+$; Cas9/+ transhemizygous \bigcirc mosquitoes exhibited male-specific features (red arrows in **D**), had reduced fecundity, and were transformed into intersexes (\square 's). Statistical significance of mean differences was estimated using a two-sided Student's *t* test with equal variance (ns: $p \ge 0.05$, *p < 0.05, *p < 0.01, and ****p < 0.001). Source data is provided as a Source Data File.



Figure 2. The pgSIT cross results in nearly complete female lethality and male sterility. A Schematic maps of targeted genes (box) and $gRNA^{dsx,ix,\betaTub}$ construct. Red arrows show relative locations of gRNA target sequences (box). gRNA^{dsx,ix,βTub} harbors a 3xP3-tdTomato marker and six gRNAs to guide the simultaneous CRISPR/Cas9mediated disruption of dsx, ix, and βTub genes. Violet exon boxes 5a and 5b represent \mathcal{Q} -specific exons. **B** A schematic of the reciprocal genetic cross between the homozygous Cas9, marked with Opie2-CFP, and homozygous $gRNA^{dsx,ix,\betaTub}$ to generate the trans-hemizygous F₁ (aka. pgSIT) progeny. Relative positions of dsx, ix, and βTub target genes (bar color corresponds to gRNA color), and transgene insertions in the Cas9 (Nup50-Cas9 strain¹) and $gRNA^{dx,ix,\betaTub}$ #1 strains are indicated in the three pairs of Ae. aegypti chromosomes. To assess the fecundity of generated pgSIT mosquitoes, both trans-hemizygous \mathcal{Q} 's and \mathcal{A} 's were crossed to the WT \mathcal{A} 's and \mathcal{Q} 's, respectively. C Comparison of the survival, sex ratio, and fertility of trans-hemizygous, hemizygous Cas9 or gRNA^{dsx,ix, βTub #1, and} WT mosquitoes. The bar plot shows means \pm SD over triple biological replicates (n = 3, all data in Supplementary Table 4). **D** Blood feeding assays using both types of trans-hemizygous intersexes (\mathbb{Z} 's): $pgSIT^{\bigcirc Cas9}$ and $pgSIT^{\bigcirc Cas9}$ \mathbb{Z} 's. To assess blood feeding efficiency, individual mated \mathbb{Q} 's or \mathbb{Z} 's were allowed to blood feed on an anesthetized mouse inside a smaller $(24.5 \times 24.5 \times 24.5 \text{ cm})$ or larger cage $(60 \times 60 \times 60 \text{ cm})$, and we recorded the time: (i) to initiate blood feeding (i.e., time to bite); and (ii) of blood feeding (i.e., feeding time). The plot shows duration means \pm SD over 30 \bigcirc 's or \square 's (n = 30) for each genetic background (Supplemental Table 5). E Flight activity of individual mosquitoes was assessed for 24 hours using vertical Drosophila activity monitoring (DAM, Supplemental Table 6). F Mating assays for fertility of offspring produced via crosses between trans-hemizygous 3° 's that inherited a maternal *Cas9* (*pgSIT*^(2Cas9)) or paternal *Cas9* (*pgSIT*^{$(3Cas9)}) and WT <math>^{(2)}$'s (Supplemental Table 8). The plot shows fertility means</sup></sup> \pm SD over three biologically independent groups of 50 WT \bigcirc 's (n = 3) for each experimental condition. Statistical significance of mean differences was estimated using a two-sided Student's t test with equal variance. (ns: $p \ge 0.05^{ns}$ *p < 0.05, **p < 0.01, and ***p < 0.001). Source data is provided as a Source Data File.



Figure 3. Population suppression in multigenerational population cage experiments. A Multiple pgSIT:WT release ratios, such as 1:1, 5:1, 10:1, 20:1, and 40:1, were tested in triplicate. A schematic diagram depicts the cage experiments for the 5:1 ratio. To start the 1st generation (G1) of multigenerational population cages, 250 mature pgSIT adult 3° 's were released with 50 similarly aged wildtype (WT) adult 3° 's into a cage, and in one hour 50 virgin 2° 's were added. The mosquitoes were allowed to mate for 2 days before all 3° 's were removed, and 2° 's were blood fed and eggs were collected. All eggs were counted, and 100 eggs were selected randomly to seed the next generation (G2). The sex-sorted adult 3° 's that emerged from 100 eggs (N₃) were released with five times more (5N₃) pgSIT adult 3° 's into a cage, and N₂ 2° 's were added later. The remaining eggs were hatched to measure hatching rates and score transgene markers. This procedure was repeated for five generations for each cage population (Supplemental Table 9). Two types of pgSIT 3° 's were assessed: those that inherited a maternal *Cas9* (*pgSIT*^{2° Cas9}), and those that inherited a paternal *Cas9* (*pgSIT*^{3° Cas9}). **B-C** Multigeneration population cage data for each release ratio plotting the normalized percentage of laid eggs (**B**) and hatched eggs (**C**). Points show means \pm SD for each triplicate sample. Source data is provided as a Source Data File.



Figure 4. Model-predicted efficacy of pgSIT egg releases on *Ae. aegypti* **population suppression as a function of release scheme, male sterility and female viability.** Weekly releases were simulated in a randomly-mixing population of 10,000 adult mosquitoes using the MGDrivE simulation framework² and parameters described in Table S10. Population suppression outcomes were identified as being most sensitive to model parameters describing the release scheme, male fertility and female viability (Supplementary Fig. 12). A These parameters were varied in factorial experiments assessing suppression outcomes including probability of elimination (the percentage of 60 stochastic simulations that result in *Ae. aegypti* elimination), and window of protection (the time duration for

which $\geq 50\%$ of 60 stochastic simulations result in $\geq 90\%$ suppression of the *Ae. aegypti* population). Female viability was varied between 0 (complete inviability) and 0.15, male fertility was varied between 0 (complete sterility) and 0.15, release size was varied between 0 and 500 eggs released per wild adult, and the number of weekly releases was varied between 0 and 52. Regions of parameter space for which the probability of elimination exceeds 90% are depicted in purple, and in which the window of protection exceeds two years in light blue. Time-series of population dynamics for select parameter sets are depicted in a-h. Here, the total female population is denoted in red, and the Cas9-carrying female population is denoted in blue. The light blue shaded region represents the window of protection lasts for several years for male fertility and female viability in the range 0-0.15 for simulated release schemes. **B** Regions of parameter space for which the probability of elimination exceeds 90% are depicted as a function of male fertility (x-axis), female viability (y-axis), and the minimum number of weekly releases required to achieve this (shadings, see key). Release size is set to 250 eggs per wild adult. The shaded square depicts the region of parameter space in which male fertility is between 0-10% and female viability is between 0-10%. A $\geq 90\%$ elimination probability is achieved with ~20-32 weekly releases for pgSIT systems having these parameters.



Supplementary Figure 1. Changes in morphological structures induced by disruption of targeted genes. External and internal structures were measured in mosquitoes of each genotype and sex. The trans-hemizygous mosquitoes harboring the maternal Cas9 ($Cas9^{\circ}$) were used for all measurements. The plot shows means and \pm SD over fifteen mosquitoes (n = 15) for each morphological structure. All data can be found in Supplementary Table 2.

Statistical significance of mean differences was estimated using a two-sided Student's *t* test with equal variance. ($p \ge 0.05^{15}$, $p < 0.05^*$, $p < 0.01^{**}$, and $p < 0.001^{***}$). Source data is provided as a Source Data File.



Supplementary Figure 2. Induced mutations at the targeted sequences in *dsx* and *ix*. The gene sequences targeted by gRNAs were PCR amplified from pgSIT mosquitoes and Sanger sequenced. Good quality of Sanger read chromatogram indicates the presence of a single consistent temple sequence (no cleave) at a targeted region, for example $gRNA^{dsx}#3$ (A) or $gRNA^{is}#2$ (B). An active CRISPR/gRNA-mediated cleavage at a particular DNA target results in re-ligation of cut fragments leading to the origin of diverse mutation-bearing alleles. These induced mutations are localized around the gRNA cleavage site and result in the drop of Sanger read quality (cleavage) as indicated for $gRNA^{dsx}#1$, $gRNA^{dsx}#2$, $gRNA^{dsx}#4$, (A) and $gRNA^{is}#1$ (B).



Supplementary Figure 3. Ae. aegypti dsx or ix disruption does not result in significant alterations in ear anatomy, Wing Beat Frequency (WBF), and male phonotactic behaviors. Immunohistochemistry and microanatomic comparison of mosquito Johnston's Orgran (cup-like structure) with chordotonal neurons responsible for auditory transduction in females (\bigcirc 's, **A**) or males (\bigcirc 's, **B**) among different genotypes. Female JO's contained far fewer neurons than males across all genotypes, and distributions of synaptic punctate assessed by Anti-Synapsin (3C11, in green) were restricted to the somata, as opposed to the male-type distribution observable in the gap between the cilia and somata tracked by anti-Horseradish Peroxidase (HRP, in red). WBFs are different between Ae. aegypti \mathcal{Q} 's and \mathcal{A} 's, and are shown on separate plots, (C) and (D) respectively. Each point represents the estimated frequency of a single fly-by (aka, the frequency of sound produced by one event of a mosquito flying past the microphone). WBF box plots indicate the median (middle bar), and 25% and 75% quartiles (lower/upper boundaries of box). The average \bigcirc WBF is 468 Hz. No significant differences were observed between \bigcirc groups, or between $\sqrt[3]{}$ groups, regardless of the genotypes tested (ANOVA on ranks, P > 0.05). The male phonotaxis assay was conducted, in which individual flying 3's showed attractive or no response (scored as 1 or 0, respectively) toward the female-specific WBF produced for 30 secs by a speaker (E). Three batches (aka. 3 replicate groups) of at least 10 3's were examined for each genotype. "'s across all genotypes exhibited robust responses to sound. The point plot shows the mean (middle bar) and one Standard Deviation (\pm SD) for the proportion of \vec{c} 's responding to the auditory stimulus. No significant differences were observed across groups (*Chi*-squared test, P > 0.05).



Supplementary Figure 4. Genetic Characterization of independent transgenic strains for CRISPR/Cas9mediated disruption of *dsx, ix,* and βTub . (A) Schematic maps of targeted genes (box) and $gRNA^{dsx,ix,\betaTub}$ construct. Relative locations of gRNA target sequences are depicted with red arrows (box). $gRNA^{dsx,ix,\betaTub}$ harbors a 3xP3*tdTomato* marker and six gRNAs to guide the simultaneous CRISPR/Cas9-mediated disruption of *dsx, ix,* and βTub genes. (B) A schematic of the reciprocal genetic cross between the homozygous *Cas9*, marked with *Opie2-CFP*, and homozygous $gRNA^{dsx,ix,\betaTub}\#1$ to generate the trans-hemizygous F_1 (aka. pgSIT) progeny. Relative positions of *dsx, ix,* and βTub genes (bar's color corresponds to gRNA's color), and transgene insertions in the *Cas9* (*Nup50-Cas9* strain) and $gRNA^{dsx,ix,\betaTub}\#1$ strains are indicated in the three pairs of *Ae. aegypti* chromosomes. (C) Three independent $gRNA^{dsx,ix,\betaTub}$ strains were generated and assessed by crossing to the *Cas9* strain and comparing the pgSIT phenotypes induced in each trans-hemizygous progenies. The survival, sex ratio, and fertility of trans-hemizygous and hemizygous $gRNA^{dsx,ix,\betaTub}$ mosquitoes were scored for each of three $gRNA^{dsx,ix,\betaTub}$ strains and compared to the corresponding values found for $gRNA^{dsx,ix,\betaTub}\#1$ strain. The bar plot shows means and \pm SD over triple biological replicates (n = 3, all data can be found in Supplementary Table 4). The bars show means and \pm SD over three biologically independent groups of 50 WT Q's (n = 3) for each experimental condition. Statistical significance of mean differences was estimated using a two-sided Student's *t* test with equal variance. ($p \ge 0.05^{ms}$, $p < 0.05^{*}$, $p < 0.01^{**}$, and $p < 0.001^{***}$). Source data is provided as a Source Data File.



Supplementary Figure 5. Determination of transgene copy number for the $gRNA^{dsx,ix,\betaTub}$ #1 strain using Oxford Nanopore genome sequencing. tandard box plot depicting the coverage distributions of the three chromosomes and the *Cas9* and $gRNA^{dsx,ix,\betaTub}$ transgenes in trans-hemizygous $gRNA^{dsx,ix,\betaTub}$ #1/*Cas9* (aka. pgSIT) mosquitoes. The center line is median, first and third quartiles are the bounds of the box, upper and lower whiskers extend from the box to the largest and lowest observed value, but no further than 1.5* Interquartile Range (IQR) from the box. Sequencing depths for chromosomes 1, 2, 3 and for Cas9 and $gRNA^{dsx,ix,\betaTub}$ transgenes were 9.71, 9.47, 9.17, 12.23, 9.50, respectively. Normalized sequencing depths were 1.03, 1.00, 0.97, 1.30, 1.01 consistent with the transgenes being present at single copy. The Nanopore sequencing data have been deposited to the NCBI sequence read archive (SRA) under BioProject ID PRJNA942966 and BioSample ID SAMN33705934.



Supplementary Figure 6. Oxford Nanopore sequencing results validating disruptions in the target sites. Integrated genome browser snapshots are zoomed in at genome regions containing gRNA target sites at dsx (A), ix (B), and βTub (C) genes in pgSIT β 's and φ 's. The direction of DNA strand and scale is indicated above generated reads. Sampled deletion mutations localize to gRNA target sites (yellow overlay) (A).



Supplementary Figure 7. Integrative genome browser snapshot of *dsx* validating gRNA target disruption in both the DNA and RNA. Oxford Nanopore reads are aligned on the top and RNAseq reads for the various genotypes indicated are aligned. On the bottom in blue are the gRNA target sites.



Supplementary Figure 8. Integrative genome browser snapshot of *ix* validating gRNA target disruption in both the DNA and RNA. Oxford Nanopore reads are aligned on the top and RNAseq reads for the various genotypes indicated are aligned. On the bottom in blue is the gRNA target site.



Supplementary Figure 9. Integrative genome browser snapshot of βTub validating gRNA target disruption in both the DNA and RNA. Oxford Nanopore reads are aligned on the top and RNAseq reads for the various genotypes indicated are aligned. On the bottom in blue are the gRNA target sites.



Supplementary Figure 10. Transcription profiling and expression analysis of *Ae. aegypti* liverpool and pgSIT mosquito samples. (A) PCA analysis and (B) hierarchical clustering of twelve samples used for RNA sequencing. MA-plots showing the differential expression patterns between: (C) pgSIT 3's vs WT 3's, and (D) pgSIT 2's vs WT 4's. Blue bots indicate significantly differentially expressed genes (FDR < 0.05), while non-significantly differentially expressed genes are depicted by the larger blue or gray dots.



Supplementary Figure 11. Longevity, fecundity, and developmental times of the generated pgSIT mosquitoes. (A) Survival plots of 20 adult \bigcirc 's or \square 's either cohabitated with \bigcirc 's or not, and (B) 20 adult \bigcirc 's either cohabitated with \bigcirc 's or not over three independent experiments (n=3). Survival means ± standard errors (SE) over days following adult eclosion are plotted. Vertical lines and values present median survivals for each tested group. Mosquito survival curves for each tested group were compared to the curve for WT's mosquitoes of the corresponding sex. The departure significance was assessed with the Log-rank (Mantel-Cox) test and is indicated above median values. (C) Female fecundity plots, measured as egg laying and hatching rates, of 60 adult \bigcirc 's or \square 's of each tested group (n = 60). (D) Plots of larva-to-pupa and pupa-to-adult developmental times were measured in 60 \bigcirc 's and \bigcirc 's or \square 's for each tested group (n = 60). Both point plots in panels C and D show mean and ± SD (Supplemental Table 7). Statistical significance of mean differences was estimated using a two-sided Student's *t* test with equal variance.($p \ge 0.05^{ns}$, $p < 0.05^*$, $p < 0.01^{**}$, and $p < 0.001^{***}$). Source data is provided as a Source Data File.



Supplementary Figure 12. Sensitivity of pgSIT population suppression outcomes to model parameters. A Partial dependency (thin traces) and individual conditional expectation plots³ (magenta dashed lines) are depicted for regression models fitted to three model outcomes: i) probability of elimination (POE), ii) window of protection (WOP) (measured in years), and iii) reduction in cumulative potential for transmission (RPT). These three outcomes were evaluated for several parameters - number of releases, size of releases (as a proportion of the number of wild adults in the environment), release interval (in days), gRNA cutting rate, Cas9 maternal deposition rate, male fertility, male mating competitiveness (relative to wild-type males), and female viability. Population suppression outcomes are most sensitive to release schedule parameters (number, size and interval of releases), male fertility and female viability. **B** Statistical sensitivity analysis metrics on the simulated data are compared to importance metrics for surrogate models. For the first-order statistical sensitivity analysis, we used Delta⁴, Fourier Amplitude Sensitivity Test (FAST)⁵, and High-Dimensional Model Representation (HDMR)⁶; whilst for the emulators we used Permutation feature importance⁷, and SHapley Additive exPlanations (SHAP)⁸. The normalized importance barcharts are in good agreement with the variable-to-outcome rankings. (Supplementary Tables 10-11). Source data are provided as a Source Data File.



Supplementary Figure 13. Sensitive and rapid detection of transgenic DNA fragments with the Sensitive Enzymatic Nucleic acid Sequence Reporter (SENSR) assay. (A) Intensity of signal from SENSR assay for both *Cas9* and *gRNA*^{dsx,ix, β Tub} constructs along a concentration gradient, as ratio of $pgSIT^{QCas9}$ to WT mosquitoes, and No Template Control (NTP). The fluorescence signal represents the background-subtracted signal (n = 4). (B) Speed of cleavage along a ratio of $pgSIT^{QCas9}$ to WT mosquitoes, as target abundance. The speed of collateral activity is represented using an HMF analysis (n = 4). (Supplementary Tables 12-13). Source data is provided as a Source Data File.

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