

Novel Synthetic *Medea* Selfish Genetic Elements Drive Population Replacement in *Drosophila*; a Theoretical Exploration of *Medea*-Dependent Population Suppression

Omar S. Akbari,^{†,‡} Chun-Hong Chen,^{§,‡} John M. Marshall,^{‡,‡} Haixia Huang,[†] Igor Antoshechkin,[†] and Bruce A. Hay^{*,†}

[†]Division of Biology, MC 156-29, California Institute of Technology, Pasadena, California 91125, United States

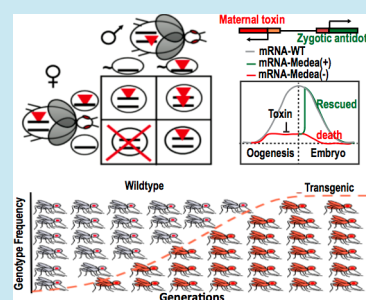
[‡]MRC Center for Outbreak Analysis & Modeling, Department of Infectious Disease Epidemiology, Imperial College London, London W2 1PG, U.K.

[§]Institute of Molecular and Genomic Medicine, National Health Research Institutes, 35 Kayen Road Zhunan Miaoli, Taiwan

Supporting Information

ABSTRACT: Insects act as vectors for diseases of plants, animals, and humans. Replacement of wild insect populations with genetically modified individuals unable to transmit disease provides a potentially self-perpetuating method of disease prevention. Population replacement requires a gene drive mechanism in order to spread linked genes mediating disease refractoriness through wild populations. We previously reported the creation of synthetic *Medea* selfish genetic elements able to drive population replacement in *Drosophila*. These elements use microRNA-mediated silencing of *myd88*, a maternally expressed gene required for embryonic dorso-ventral pattern formation, coupled with early zygotic expression of a rescuing transgene, to bring about gene drive. *Medea* elements that work through additional mechanisms are needed in order to be able to carry out cycles of population replacement and/or remove existing transgenes from the population, using second-generation elements that spread while driving first-generation elements out of the population. Here we report the synthesis and population genetic behavior of two new synthetic *Medea* elements that drive population replacement through manipulation of signaling pathways involved in cellular blastoderm formation or Notch signaling, demonstrating that in *Drosophila* *Medea* elements can be generated through manipulation of diverse signaling pathways. We also describe the mRNA and small RNA changes in ovaries and early embryos associated from *Medea*-bearing females. Finally, we use modeling to illustrate how *Medea* elements carrying genes that result in diapause-dependent female lethality could be used to bring about population suppression.

KEYWORDS: selfish genetic element, synthetic biology, maternal effect, gene drive, mosquito, dengue, malaria



Insects act as vectors for a number of important diseases of humans, animals, and plants. Examples include malaria and West Nile (humans and birds); dengue, yellow fever, lymphatic filariasis, chikungunya, and Chagas disease (humans); Rift valley fever and trypanosomiasis (humans and livestock); and plant diseases such as Huanglongbing (citrus), Almond leaf scorch, Pierce's disease (grapes), and zebra chip disease (potato).¹ Replacement of the wild population with insects engineered to be unable to transmit disease provides a method of disease prevention that complements traditional vector control methods. Population replacement is attractive as a disease prevention mechanism because it is species-specific and does not involve the use of chemicals, involve gross modification of the environment, or require direct contact with the disease hosts (humans, animals or plants). Perhaps most importantly, it is in principle self-sustaining, providing protection over large areas even if other vector control strategies suffer gaps in coverage.

Transgenes that mediate disease refractoriness are unlikely to confer an overall fitness benefit on insects that carry them.^{2–4}

Therefore, an essential component of any population replacement strategy is a gene drive mechanism that will ensure the spread of linked transgenes to genotype or allele fixation in a modest number of generations following release. A number of gene drive mechanisms have been proposed,^{5,6} but only one, *Medea*, has been engineered and shown to drive replacement in a wildtype population, in *Drosophila*.⁷ A *Medea* selfish genetic element consists of two chromosomally located, tightly linked transgenes: one that encodes a toxin inherited by all progeny of *Medea*-bearing mothers and a second that encodes an antidote active in the zygote. *Medea* spreads by causing the death of non-*Medea*-bearing progeny of *Medea*-bearing mothers (Figure 1A), thereby causing a relative increase in the population frequency of the *Medea*-bearing chromosome (*Medea*-bearing chromosomes and their non-*Medea*-bearing

Special Issue: Design and Operation of Mammalian Cells

Received: August 30, 2012

Published: November 30, 2012

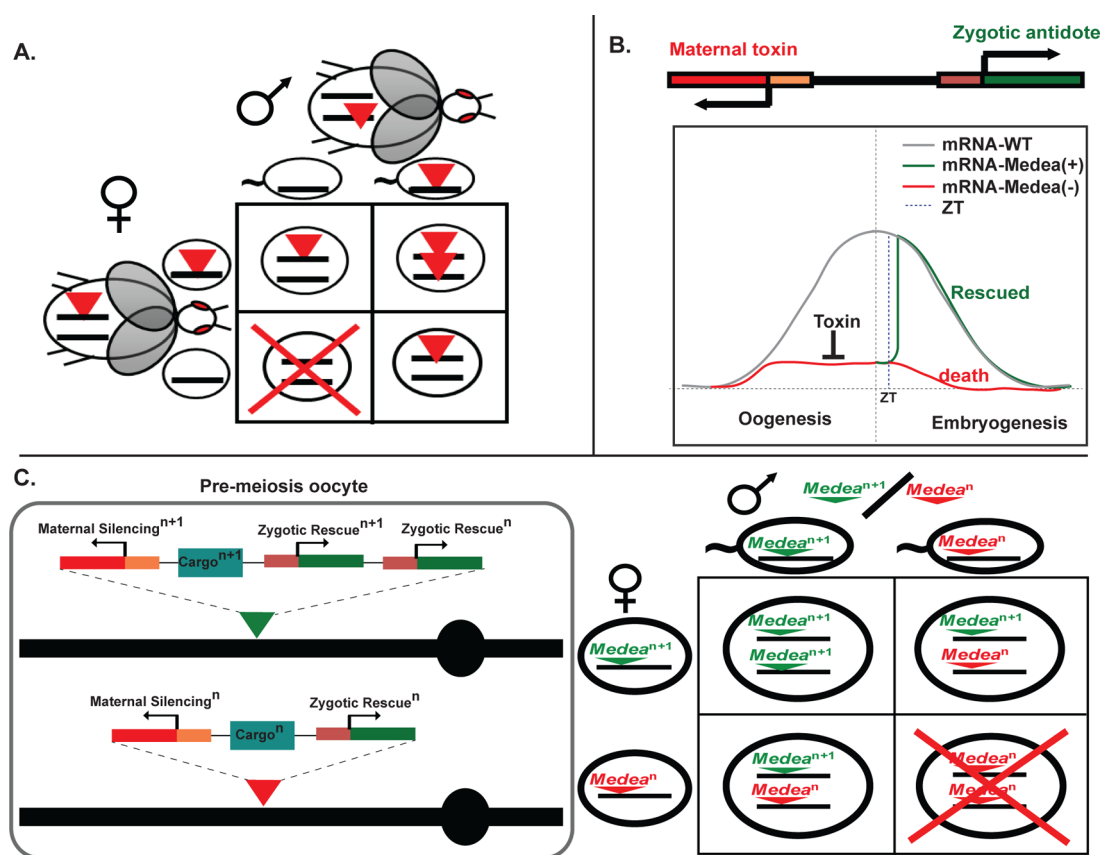


Figure 1. *Medea* genetics, molecular basis of synthetic elements, and cycles of population replacement. Mothers carrying *Medea* cause the death of all progeny that fail to inherit *Medea* (A). Synthetic *Medea* elements consist of two genes. Maternally expressed miRNAs (the toxin) silence (red line during oogenesis) the expression of a maternally expressed transcript (gray line) that normally provides a product essential for early embryonic development. Rescue of *Medea*-dependent maternal-effect lethality occurs when progeny inheriting *Medea* express a version of the silenced maternal mRNA sufficient to rescue normal development (green line). Progeny that fail to inherit *Medea* die because the endogenous levels of the maternally deposited mRNA (red line during embryogenesis) are insufficient for normal development (B). Second-generation *Medea* elements ($n + 1$) carrying a new cargo, a new toxin, a new antidote, and a copy of the antidote from the previous generation (n) drive into a population at the expense of a first generation element (n), when both elements are located at the same position in the genome so as to force them to compete for germline transmission (C).

counterparts are hereafter referred to as *Medea* and non-*Medea* alleles, respectively).^{8–12} In the absence of family level selection, *Medea*-bearing individuals and alleles experience no direct benefit from this killing, but non-*Medea* alleles experience *Medea*-dependent death (a fitness loss) in each generation that is dependent on the *Medea* allele frequency. Whenever the average cost associated with the *Medea* allele is less than that associated with the non-*Medea* allele, a *Medea* element located on an autosome spreads, with the population ultimately consisting of *Medea*-bearing heterozygotes and homozygotes. When *Medea* is located on the X chromosome in a X/Y male heterogametic species, *Medea* is predicted to spread to allele fixation, with wildtype alleles being completely eliminated.¹²

We previously generated synthetic *Medea* elements through maternal expression of microRNAs designed to silence the expression of a maternally expressed transcript, *myd88*, whose product is required in the embryo for dorso-ventral pattern formation. Maternal expression of these small RNAs created a pretoxic state, a loss of maternal *myd88*, which if left unopposed led to the death of all embryos. Zygotic rescue in this case was mediated by expression of a tightly linked transgene, encoding a microRNA-insensitive version of the *myd88* transcript lacking target sites present in the endogenous transcript, driven by a transient early zygote-specific promoter (Figure 1B).⁷

There are several reasons why it is important to be able to create *Medea* elements that work through additional mechanisms. First, while the Toll pathway and Myd88 are likely to be required for dorso-ventral pattern formation in many if not all insects, key components may be supplied zygotically rather than maternally in some species (for an example, see ref 13), which would prevent maternal silencing from causing embryonic lethality, a prerequisite for *Medea*-dependent drive. In addition, the genes that make up synthetic *Medea* elements and their transgene cargo will inevitably be subject to mutation. First, the toxin encoding miRNAs, or the promoter driving their expression, can mutate to inactivity. This leads to the creation of antidote-only alleles, which are resistant to *Medea*-dependent killing but do not drive. If these chromosomes have a higher fitness than those carrying intact *Medea* elements, the former will spread at the expense of the latter, potentially resulting in the appearance of wildtype individuals capable of disease transmission.¹¹ Second, genes mediating disease refractoriness can also mutate to inactivity. If this loss of function results in a *Medea*-bearing chromosome with increased fitness as compared to chromosomes carrying an intact *Medea* element, the former will spread at the expense of the latter, also resulting in the appearance of individuals capable of spreading disease. Finally, principles of risk management argue that it is always desirable

to be able to greatly decrease the frequency of, or eliminate, a particular transgenic modification from a wild population, if desired.

One strategy that addresses the above issues, albeit not always perfectly, involves creating second-generation *Medea* elements that carry a new toxin, a new antidote, and if needed a new set of transgenes designed to bring about disease refractoriness. If second generation elements are located at a different position in the genome from first generation elements, the former will spread, creating a population with both elements. If instead a second generation element is located at the same position in the genome as the first generation element – and it also carries a copy of the first generation antidote – this element should spread into the wild population at the expense of the first generation element in the same way first generation element spread at the expense of wildtype⁷ (Figure 1C). Thus, for an autosomal element carrying a fitness cost, the frequency of a first generation element will be greatly decreased but will not be eliminated (it will persist in heterozygous combination with the second generation element at a frequency that depends on the relative fitness difference between first and second generation elements).¹² In contrast, an X-linked second-generation element is predicted to completely eliminate a first generation element from the population. These manipulations do not restore the pretransgenic state but do allow one to carry out multiple cycles of population replacement, with distinct cargoes, or an element with no cargo (when the goal is primarily to decrease the frequency of, or remove, a particular gene for disease refractoriness from the population) being driven into the population during each cycle. A critical prerequisite for each of these strategies is the availability of multiple, independent toxin–antidote combinations able to show *Medea*-based gene drive.

■ RESULTS AND DISCUSSION

Engineering Novel Synthetic *Medea* Elements. In order to generate a synthetic *Medea* element through zygotic rescue of a maternal loss-of-function phenotype, one needs genes whose expression occurs maternally, but whose products are not required until after the initiation of zygotic transcription. Two genes with these characteristics are *discontinuous actin hexagons* (*dah*, CG6157) and *O-fucosyltransferase 1* (*o-fut1*, also known as neurotic, CG12366). *Dah* is a membrane-associated protein required in *Drosophila* for proper formation of the metaphase furrow during syncytial blastoderm stages and cellularization during blastoderm formation.¹⁴ Maternal loss of *dah* results in 100% embryo lethality, and embryonic death is not rescued when embryos inherit a wildtype copy of the gene from the father, indicating that *dah* mRNA and/or protein must normally be provided maternally. *Dah* is a member of the dystrotelin branch of the family of proteins that also includes dystrophins and dystrobrevins.¹⁵ Clear *Dah* homologues are present in all sequenced *Drosophila* species and in mosquitoes but are not obvious in other sequenced insect species such as beetles (*Tribolium*), wasps (*Nasonia*), or honeybees (*Apis mellifera*). More distantly related dystrotelins are found throughout the animal kingdom, though functions for many remain to be determined. *O-fut1* is required for Notch signaling in *Drosophila*^{16,17} with maternal loss resulting in a neurogenic phenotype, in which excess neurons are produced at the expense of embryonic epidermis.¹⁶ As with *dah*, loss of maternally provided *O-fut1* cannot be rescued through inheritance of a wildtype copy of the gene from the father.

O-fut1 fucosylates Notch, and this modification is required for many Notch-dependent signaling events. *O-fut1* also plays a non-enzymatic role as an ER chaperone that promotes Notch folding^{18,19} and as an extracellular component essential for Notch endocytic trafficking. *O-fut1* homologues are conserved throughout the animal kingdom, as is its role in Notch-dependent signaling.²⁰

We engineered synthetic *Medea* elements based around *dah* and *o-fut1* using the same architecture used to generate *Medea*^{myd88}.⁷ In order to silence the expression of *dah* or *o-fut1*, we generated for each gene a transcription unit (the toxin) encoding an 2-mer of two synthetic miRNAs, designed to base pair with 100% complementarity to the relevant target transcript. Synthetic miRNAs were generated using the mir6.1 backbone, as described previously.⁷ The expression of each miRNA multimer was driven by a modified version of the maternal-specific bicoid (*bcd*) promoter. Adjacent to each of these transcription units we placed a second transgene (the antidote) encoding a version of either *dah* or *o-fut1* recoded so as to be resistant to silencing by the maternally deposited synthetic miRNAs, with expression being driven by the transient, early promoter from the *bottleneck* (*bnk*) gene.²¹ Elements carrying both toxin and antidote genes are known as *Medea*^{dah} and *Medea*^{o-fut1}. Flies carrying these constructs are characterized below.

We also generated *Medea* elements based around maternal silencing and zygotic rescue of a number of other genes known to have maternal-effect lethal loss-of-function phenotypes (unpublished observations). For seven of these, *pelle*,²² *tube*,²³ *almondex*,²⁴ *bloated tubules*,²⁵ *concertina*,²⁶ *anillin*,²⁷ and *torso*,²⁸ we did not observe maternal effect killing, presumably due to inefficient gene silencing, possibly coupled with low levels of the maternal product being sufficient to provide sufficient activity for normal development. These elements were not characterized further. For two other genes, *groucho* (*gro*), a transcriptional corepressor required for multiple aspects of early embryonic development, including neurogenesis, sex determination, and segmentation,²⁹ and *trunk* (*trk*), a ligand for the receptor Torso, required for specification of the embryonic termini,³⁰ we were able to create *Medea* elements that showed strong maternal-effect lethality, with 100% of embryos derived from matings between heterozygous mothers and wildtype fathers dying (construct details provided in the methods). *Medea*^{gro} also showed zygotic rescue behavior. However, rescue was imperfect and often resulted in male-biased sex ratios. These elements did not show drive when introduced into a non-transgenic population at an allele frequency of 25%, as described previously.⁷ In the case of *Medea*^{trk} elements, no zygotic rescue was obtained.

The above results indicate that it can be difficult to silence maternal gene expression and that even when good maternal silencing is achieved, it may also prove difficult to replace the needed product through zygotic expression in a way that restores normal development. In the case of *gro*-based *Medea* elements, it is likely that specific zygotic levels of *gro* are important, with *bnk*-drive expression of the rescuing transgene resulting in inappropriate levels that disrupt multiple signaling pathways.²⁹ In the case of *trk*-based *Medea* elements the reasons for lack of rescue are unclear. Injection of *trk* mRNA into embryos from homozygous mutant *trk* mothers has been shown to rescue embryonic pattern formation, albeit at a low frequency.³¹ It is possible that, as with *gro*, timing and levels of protein expression are critical. While we cannot draw strong

Table 1. *Medea*^{dah} and *Medea*^{o-fut1} Show *Medea*-like Selfish Genetic Behavior^a

parental genotype		inherited by		adult M progeny %		embryo survival %	
		oocyte	embryo	predicted	observed (<i>dah</i> , <i>o-fut1</i>)	predicted	observed (<i>dah</i> , <i>o-fut1</i>) ± sd
male	female	maternal toxin	zygotic antidote %				
M/+	+/+	0	50(0); 50(1)	50	50 (<i>n</i> > 12,000), 50 (<i>n</i> > 11,000)	100	98.05 ± 2.3, 96.28 ± 2.8
M/M	M/M	2	100(2)	100		100	98.97 ± 6.0, 92.74 ± 6.0
+/+	M/+	1	50(0); 50(1)	50	100 (<i>n</i> > 10,000), 100 (<i>n</i> > 12,000)	50	49.56 ± 0.5, 47.43 ± 2.0
M/M	M/+	1	50(1); 50(2)	100		100	98.05 ± 2.3, 97.35 ± 2.0
M/+	M/+	1	25(0); 50(1); 25(2)	75		75	73.62 ± 1.5, 73.9 ± 1.7
M/+	M/M	2	50(1); 50(2)	100		100	97.7 ± 1.0, 98.41 ± 1.5
+/+	M/M	2	100(1)	100		100	99.47 ± 1.5, 94.51 ± 2.0
M/M	+/+	0	100(1)	100		100	95.22 ± 1.1, 96.99 ± 2.5
+/+	+/+	0	100(0)	0		100	93.80 ± 3.19

^aProgeny of crosses between parents of several different genotypes are shown. M refers to the *Medea*^{dah}- or *Medea*^{o-fut1}-bearing chromosome; + refers to the non-*Medea*-bearing homolog. The maternal copy number (0 to 2) of *bcd*-driven miRNAs (maternal toxin) targeting the endogenous *dah* or *o-fut1*, and the zygote copy number (0 to 2), and percentage of embryos inheriting *bnk*-driven *dah* or *o-fut1* (zygotic antidote) are indicated, as are the adult progeny genotypes predicted for Mendelian inheritance of *Medea*^{dah} or *Medea*^{o-fut1}. The percent embryo survival was normalized to that of wildtype (*w*¹¹⁸), which was 93.80 ± 3.19. blank, not measured.

conclusions from this limited data set, our results may suggest that genes whose products provide a necessary condition for a process, but whose expression is not sufficient to drive the process (adaptors such as *myd88* and *dah*), are better candidates than those whose expression drives a process in a dose-dependent manner (signal transduction effectors such as *gro* and *trk*).

***Medea*^{dah} and *Medea*^{o-fut1} Show Maternal-Effect Lethal and Zygotic Rescue Behavior.** Multiple transgenic lines were generated for each of the above constructs. One line each was selected for further characterization. Matings between heterozygous *Medea*^{dah}/+, or *Medea*^{o-fut1}/+ males (where + indicates a wildtype chromosome) and +/+ females resulted in high levels of embryo viability, similar to those for the *w*¹¹⁸ strain used for transformation (Table 1). In addition, ~50% of adult progeny carried *Medea*, as expected for Mendelian segregation without dominant fitness costs. Matings among homozygotes for *Medea*^{dah} or *Medea*^{o-fut1} also resulted in high levels of egg viability. In contrast, when heterozygous *Medea*^{dah}/+ or *Medea*^{o-fut1}/+ females were mated with homozygous +/+ males, ~50% of embryos died and all adult progeny were *Medea*-bearing. These observations, together with the results of several other crosses (Table 1), indicate that a single copy of each *Medea* toxin is sufficient to induce 100% maternal-effect lethality and a single copy of each rescue transgene is sufficient to rescue normal development of embryos derived from mothers expressing one or two copies of the toxin.

***Medea*^{dah} and *Medea*^{o-fut1} Drive Population Replacement.** The above observations suggested that *Medea*^{dah} and *Medea*^{o-fut1} should drive population replacement. To test this prediction we mated equal numbers (1:1 ratio) of heterozygous *Medea*^{dah}/+ or *Medea*^{o-fut1}/+ males to WT (+/+) females in separate bottles in triplicate, giving rise to progeny populations with *Medea*^{dah} or *Medea*^{o-fut1} present at an allele frequency of ~25%. Populations were followed for a total of 20 generations, counting the number of *Medea*-bearing adults each generation, as described previously.⁷ In all cages non-*Medea*-bearing individuals disappeared from the population between generations 12 and 16 (Figure 2).

The observed changes in *Medea* frequency trailed the predicted rate of increase for a *Medea* element with no fitness cost, with the delay being more extreme for *Medea*^{dah} than for *Medea*^{o-fut1}. The magnitude of the fitness costs were estimated

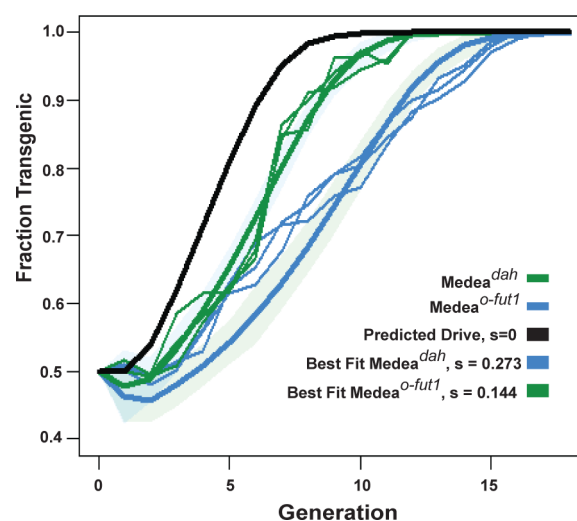


Figure 2. *Medea*^{dah} and *Medea*^{o-fut1} drive population replacement. Fraction of the adult population that is transgenic is plotted versus number of generations for both *Medea*^{dah} (thin blue lines) and *Medea*^{o-fut1} (thin green lines). Expected transgenic frequencies for a *Medea* with no fitness cost (*s* = 0) are indicated by the black line. Predicted behavior of *Medea* elements with fitness costs corresponding to best-fit estimates *Medea*^{dah} (bold blue line) and *Medea*^{o-fut1} (bold green line) were derived from actual *Medea*^{dah} and *Medea*^{o-fut1} behavior. 95% confidence intervals are indicated by light green and blue shading. *s* = additive fitness cost in *Medea* homozygotes.

by calculating the expected dynamics for a variety of fitness costs (both additive and dominant), calculating the likelihood of the observed data for each fitness cost, and selecting the fitness cost having the greatest likelihood (for more details, see the materials and methods). Confidence intervals for the fitness costs were then obtained using a Markov Chain Monte Carlo sampling procedure.³² We found that additive fitness costs provided a much better fit to the data for *Medea*^{dah}, while for *Medea*^{o-fut1}, both additive and dominant fitness costs performed equally well. Assuming additive fitness costs, we estimate a cost in homozygotes of 27.3% for *Medea*^{dah} (95% confidence interval: 26.8–27.8%) and of 17.4% for *Medea*^{o-fut1} (95% confidence interval: 16.5–18.2%). The basis for these costs is unknown. It could reflect insertion site-dependent effects on the expression of neighboring genes. It could also reflect the

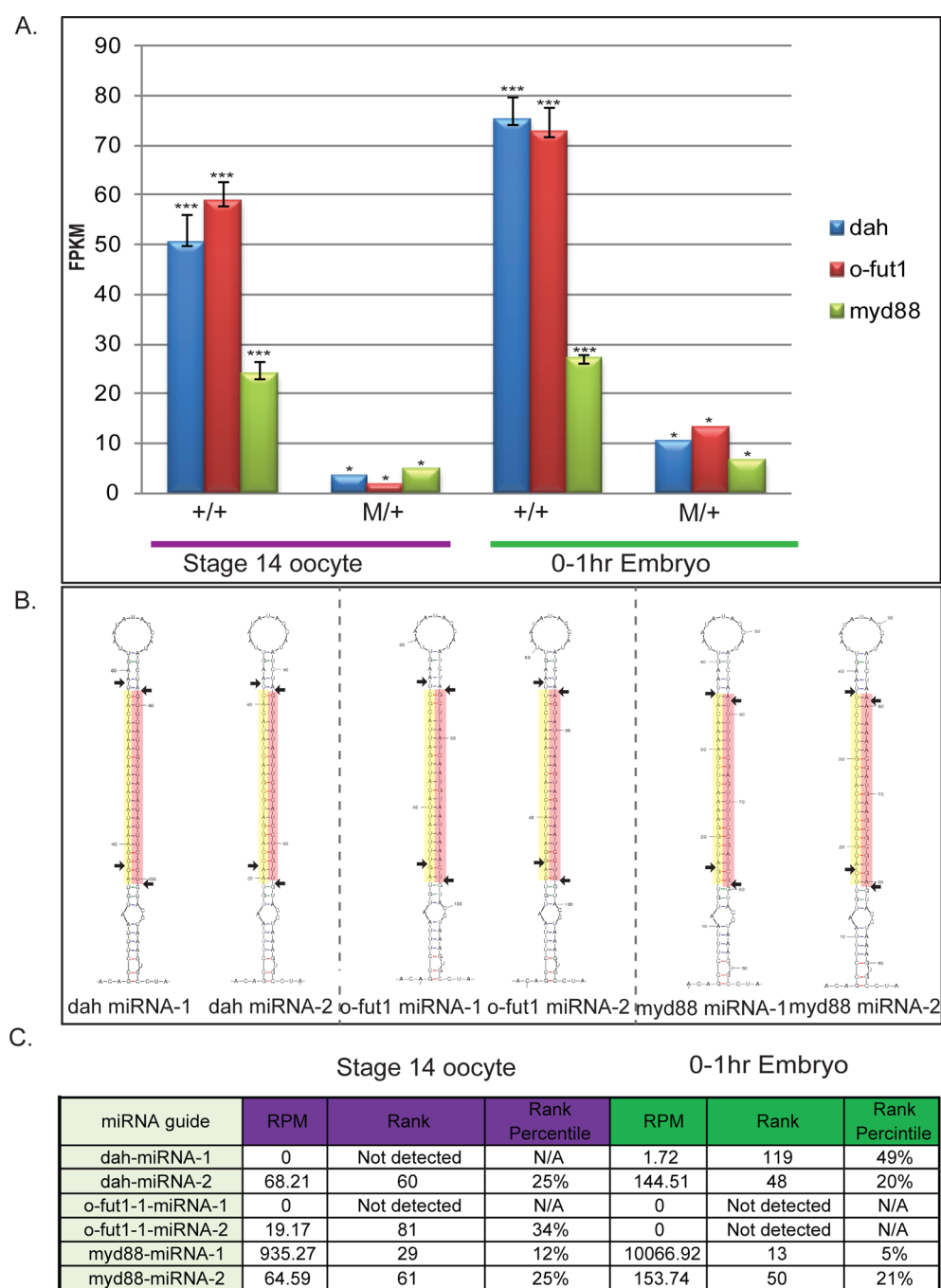


Figure 3. Molecular characterization of stage 14 oocytes and 0–1 h embryos from mothers of specific genotypes. Levels of *myd88*, *o-fut-1*, and *dah* mRNA determined using RNA-seq (expressed as fragments per kilobase per million reads, FPKM) in wildtype stage 14 oocytes and 0–1 h embryos from wildtype mothers and wildtype fathers (+/+), and from oocytes and embryos derived from heterozygous *Medea*^{myd88} (green bars), *Medea*^{dah} (blue bars), or *Medea*^{o-fut1} (red bars) mothers crossed to (+/+) males. Asterisks indicate the number of replicates (A). The structures of the miRNAs used to silence *dah*, *o-fut1*, and *myd88* are indicated. The orange shaded region indicates the predicted miRNA guide strand; the yellow shaded region represents the miRNA* strand. Horizontal arrows indicate sites of Drosha and Dicer cleavage (B). Small RNA reads for synthetic miRNAs were expressed as reads per million reads (RPM), which normalizes read count to the size of each library. To provide a sense of scale, the expression levels of synthetic miRNAs were also compared to those of endogenous miRNAs expressed in the ovary, both in terms of absolute rank and in terms of rank percentile.

effects of manipulating the developmental expression of the targeted genes (imperfect rescue, for example) or other off-target effects on unknown genes. The results of crosses tabulated in Table 1 do not identify any obvious fitness costs associated with survival of embryos or adults carrying *Medea*^{dah} or *Medea*^{o-fut1}. However, many aspects of life history such as response to stress, longevity, and fertility/fecundity remain to

be explored. Characterization of identical *Medea* elements inserted at new locations should provide insight into whether the costs observed are site-specific or intrinsic to these elements.

These results, in conjunction with our earlier work, demonstrate that *Medea* selfish genetic behavior in *Drosophila* can be engineered through the manipulation of diverse

signaling pathways. Each of the pathways targeted, embryonic dorso-ventral pattern formation,³³ blastoderm formation,³⁴ and Notch signaling,³⁵ requires the products of multiple genes in order to bring about normal early embryonic development. Many of these are also provided maternally. In consequence, it may, with the caveats noted above, be possible to generate *Medea* elements through maternal silencing and zygotic rescue of those whose activity is not required in some other context during oogenesis. Elements generated this way, even those that target different components of the same pathway, will function as independent, non-cross-rescuing *Medea* elements.

Our positive results notwithstanding, it remains to be demonstrated that synthetic *Medea* elements can be generated using the above strategy in pest insects. In unpublished work we have tried and thus far been unable to bring about maternal effect killing in the mosquito *Aedes aegypti* in response to maternal expression of miRNAs designed to silence *Aedes myd88*, *dah*, or *o-fut1*. While transcript levels of the targeted genes are decreased in response to miRNA expression, it is possible that remaining levels of maternally deposited product are sufficient to bring about normal development. Extensive transcriptional profiling of *Aedes* oogenesis and early embryonic development (unpublished observations) also leaves open the possibility that early zygotic transcription of *myd88*, *dah*, and *o-fut1* may provide some rescuing activity, limiting our ability to kill through maternal silencing alone. These difficulties highlight the fact that the approach to *Medea* development described herein requires some detailed knowledge of oogenesis and embryonic development with respect to when specific transcripts/proteins are required and how they are provided. This information exists for the genes we targeted in *Drosophila* and provided the basis for our use of them, but it does not exist for other insects. An alternative approach to *Medea* development involves the use of proteins or RNAs that provide a toxic dominant gain of function activity that acts in the zygote. If these are generally toxic to insect embryos (though not oocytes), this strategy would appear to require less knowledge of the insect under study (primarily information relating to promoters and UTRs that provide control over RNA stability and protein translation during oogenesis), which may make the system more portable across species. In summary, while our results demonstrate that multiple synthetic *Medea* elements can be generated through maternal silencing and zygotic rescue, it remains unclear if this design strategy provides a straightforward path to creation of similar elements in other insects, for which more limited information is available.

Molecular Characterization of *Medea*-Dependent Gene Silencing. An important unanswered question with respect to the synthetic *Medea* elements generated here and previously⁷ is the extent to which the targeted maternal transcripts are being silenced and the levels of synthetic miRNAs associated with these levels of silencing. MicroRNA-dependent silencing can occur through translational inhibition and/or transcript degradation.³⁶ To provide a minimum estimate of the extent to which maternal *myd88*, *dah*, and *o-fut1* were being silenced, we used next generation sequencing technology (RNA-seq) to measure transcript levels in both the mature oocytes (stage 14, s14) of *Medea*/+ mothers and early embryos (0–1 h) derived from crosses between these mothers and +/+ males. In s14 oocytes transcript levels were decreased by 5.0-fold (*myd88*), 14.27-fold (*dah*), or 30-fold (*o-fut1*) with respect to the levels of the endogenous transcripts. Similar, though somewhat lower levels of silencing were also observed

in the early embryo (0–1 h): 4.1-fold for (*myd88*), 7.12-fold for (*dah*), and 5.53-fold for (*o-fut1*) (Figure 3A). The somewhat lower level of knockdown in the 0–1 h early embryo versus the s14 oocyte may reflect poly A addition to previously non-polyadenylated transcripts.³⁷ It may also reflect, in part, the initiation of zygotic transcription and expression of the rescue transgenes.

To begin to identify the characteristics of synthetic miRNAs able to bring about effective silencing of the maternal transcripts discussed above, we sequenced libraries of small RNAs isolated from s14 oocytes and 0–1 h embryos derived from crosses between mothers heterozygous for *Medea*^{myd88}, *Medea*^{dah}, or *Medea*^{o-fut1} and +/+ fathers (Supplementary Table 1). We detected the expression of 154 (64.7%) of the 238 total *Drosophila melanogaster* annotated miRNAs in miRbase (v19), with the top 15 expressed miRNAs constituting the bulk, between 89.14% and 97.81%, of the total reads mapping to miRNAs. modENCODE does not have an equivalent data set, the only related one being small RNAs from whole ovaries, which includes all stages of ovarian development.³⁸ Therefore, to gain a sense of reproducibility from sample to sample we calculated correlation coefficients, comparing levels of miRNAs in wildtype s14 oocytes or 0–1 h embryos with comparable stages derived from *Medea*^{myd88}, *Medea*^{dah} or *Medea*^{o-fut1} mothers (Supplement Tables 2 and 3). Correlation coefficients were high for DAH (s14 = 0.84 and 0–1 h = 0.96) and *myd88* (s14 = 0.91 and 0–1 h = 0.97), but were significantly lower for *o-fut1* (s14 = 0.66 and 0–1 h = 0.92). Consistent with this, the rank abundance of the top 20 miRNAs in DAH and *Myd88* samples were similar (though not identical) to those from wildtype, while those from *o-fut1* samples were much less similar. The basis for the low correlation coefficients observed with *o-fut1* samples requires further exploration. It could be that expression of synthetic miRNAs targeting *o-fut1*, and/or position effects related to the site of *Medea* insertion in the genome, resulted in significant changes in the patterns of small RNA gene expression. Alternatively, differences in sample preparation may be responsible.

We calculated levels of synthetic miRNAs in terms of reads per million total reads (RPM) and noted their rank percentile abundance with respect to endogenous miRNAs expressed in the ovary or embryo to provide a sense of scale (Figure 3C). For all synthetic miRNAs we only detected mature miRNAs from the guide strand (Figure 3B). For *Medea*^{o-fut1}, both synthetic miRNAs were undetectable in early embryos, and only one, *o-fut1*-miRNA-2, was observed at low levels in the s14 oocyte (rank percentile of 34). For *Medea*^{dah}, *dah*-miRNA-1 was not detected in the ovary and was found only at very low levels in the embryo. In contrast, *dah*-miRNA-2 was found at somewhat higher levels in the oocyte and embryo (rank percentile of 25% and 20%, respectively). Finally, for *Medea*^{myd88} we observed that one miRNA, *myd88*-miRNA-1, was present at relatively high levels in the ovary and embryo (rank percentile of 12% and 5%), while *myd88*-miRNA-2 was expressed at much lower levels.

Given the high correlation coefficients observed in *myd88* and *dah* samples, the observed levels of synthetic miRNAs may be an accurate reflection, at least to some extent, of their endogenous levels. This cannot be said for the *o-fut1* samples. There are several possible explanations for why the levels of specific miRNAs in the *myd88* and *dah* samples might be low and/or discordant with those of other miRNAs expressed from the same transcript. First, miRNAs loaded into Ago1-containing

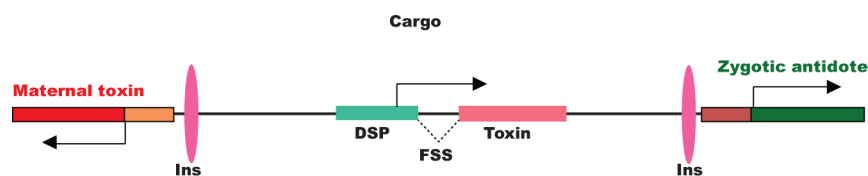


Figure 4. Hypothetical *Medea* elements carrying a cargo transgene cassette able to induce cue-dependent female killing. The *Medea* illustrated carries a cargo gene consisting of a diapause-specific promoter (DSP) driving a toxin that is synthesized as an intact protein only in response to female-specific splicing (B). Chromatin insulators (Ins), female specific splicing events (FSS), and 3' untranslated regions (UTR) are indicated.

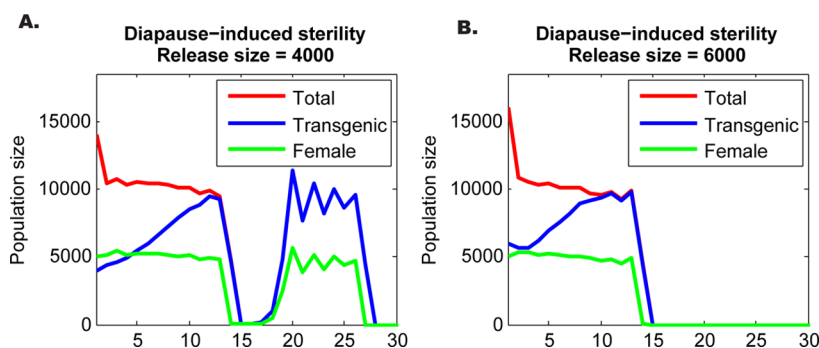


Figure 5. Modeling the ability of *Medea* elements carrying genes that cause cue-dependent female lethality to bring about population eradication. 4,000 males homozygous for a *Medea* carrying a transgene cassette that induces diapause-dependent female killing are released into a total wild population of 10,000. The fate of the total population (red line), transgenics (blue line), and females (green line) are followed for 30 generations. Diapause occurs in generation 13 and generation 26 (A). 6,000 transgenic males are released into the wild population as above (B).

RISC complexes and bound to targets with perfect complementarity are degraded through a tailing and trimming process, while those bound to Ago2-containing RISC undergo 2'-O-methylation at their 3' ends, which prevents degradation through this pathway^{39,40}. All of our miRNAs are designed to have perfect complementarity with their targets, and while we designed them using rules thought to promote loading into Ago2,^{41–43} their actual loading preferences have not been determined. Thus, it is possible that Ago-loading preferences are influencing their fate. That said, if the low levels of specific synthetic miRNAs we observe in the ovary and early embryo result from their degradation subsequent to binding to fully complementary mRNA target transcripts while in association with Ago1, the fact that we still observe good maternal transcript silencing would suggest that these miRNAs have significant ability to silence prior to degradation. In previous work in S2 cells, Ago1-bound miRNAs with perfectly complementary to their targets were unable to bring about detectable silencing.⁴⁴ It will be interesting to determine if Ago1-bound miRNAs behave differently in the ovary.

Preferential processing of specific miRNAs from the polycistronic transcript may also contribute the low and/or discordant levels observed for specific miRNAs in a polycistron. For example, once processing of one of the miRNAs occurs, if processing of the second miRNA is delayed, exonuclease activity may result in degradation of the cleaved precursor before processing of the second miRNA occurs.⁴⁵ Finally, active turnover may promote the degradation of specific mature miRNAs.⁴⁶ Clearly more work is needed in order to understand which miRNAs are bringing about silencing and which Ago-RISC complex they are associated with. Generation of *Medea* elements that drive population replacement and that express single miRNAs, in conjunction with immunoprecipitation of Ago1- and Ago2-bound miRNAs from *Medea*-bearing females, should provide additional insights into the characteristics of highly active maternal miRNAs.

Using *Medea* To Bring about Environmental Cue-Dependent Population Suppression.

Here we consider a second possible use of *Medea*: to bring about population suppression, and perhaps eradication, following population replacement with a transgene cassette that results in female-specific lethality in response to a specific environmental cue such as diapause, which allows many insects to survive long periods of adverse conditions such as cold or drought.^{47–49} An attractive feature of an environmental cue-dependent population reduction strategy is that, in contrast to the use of insecticides, the strategy is species-specific. A cassette to be driven into the population for such a strategy might consist of a gene expressed under the control of a diapause-induced promoter, encoding a transcript that undergoes female-specific splicing to produce a toxin in females but not males (Figure 4). We use discrete-generation difference equations to theoretically explore this idea. We consider the *Medea* element as a single allele, which we denote as “M” and refer to the corresponding position on the wildtype chromosome as “m.” A one-locus model can then be used for the discrete-generation dynamics. We assume random mating, 100% toxin efficiency, and equal fitness costs in males and females having the element in the absence of the external cue. We also consider density dependence. We model the effects of an external cue by modulating the fitness costs based on the presence or absence of the cue. In the absence of the cue, we consider additive fitness costs of 17.4% in homozygous males and females (i.e., $s_m = s_f = 0.174$ and $h_f = h_m = 0.5$), corresponding to fitted estimates for *Medea*^{o-fut1}, which suffered smaller fitness costs than *Medea*^{dah}. These fitness costs are unlikely to reflect fitness in the wild. They should perhaps be thought of as providing a minimum estimate and are incorporated for illustrative purposes only. In the presence of the cue, we consider the case of dominant, female-specific lethality, so that male fitness costs are unchanged and female fitness costs are 100% in both heterozygotes and homozygotes (i.e., $s_f = 1$ and $h_f = 1$). To

focus on mosquitoes as a specific example, if the female-specific toxin is under the control of a diapause-specific promoter, then there should be little or no fitness cost associated with the presence of this transgene during the ~12 generations of the rainy season, while female-specific lethality should result during the ~1 generation of the dry season in which diapause occurs. We consider a release at the beginning of the rainy season, neglecting seasonal changes in the environmental carrying capacity. Details of the model are found in the Methods. Here we focus on several important conclusions.

We consider a release of homozygous transgenic males at the beginning of the rainy season (generation 1). The rainy season lasts 12 generations, during which time the transgene spreads into the population. Then, as the dry season begins (generation 13), transgenic females suffer a lethal fitness cost such that the female population is greatly reduced by generation 14 (the beginning of the next rainy season), with a consequent drop in total population size by generation 15. This scenario is shown in Figure 5A,B. In Figure 5A 4,000 transgenic males are released into a wildtype population of 10,000. By generation 13, the transgene has almost reached transgene fixation (i.e., almost all individuals have either one or two copies of the *Medea* element), but because transgene fixation is not complete, a few wildtype females survive diapause and the population rebounds during the next rainy season. However, by the time of the next diapauses (generation 26), the population is entirely transgenic and so, this time, the female population collapses at generation 27 and there is a total population crash by generation 28. Figure 5B illustrates a scenario in which 6,000 transgenic males are released. In this case the population is entirely transgenic by the time of the first diapause (generation 13). As a result a permanent population crash occurs by generation 15.

To summarize, the key result from our modeling is the finding that that an environmental cue can be used to bring about a population crash, provided that *Medea* is given time to spread to transgene fixation before the cue appears. Appearance of the cue prior to transgene fixation may lead to transient population suppression, but the continued presence of wildtype females allows the population to ultimately rebound. However, because *Medea*-bearing males are not killed by the cue, population replacement continues, and cue-dependent suppression can be initiated the next season. While the numbers of *Medea*-bearing males that need to be released in order for population replacement to happen within one season is high (~1:1 with respect to wildtype males), these numbers are still significantly lower than those associated with sterile male release programs designed to bring about population suppression and do not constitute a fundamental limitation of the technology.⁵⁰

Can the components required to bring about *Medea*-dependent population suppression be created? In the above model the encoded toxin would only be active in females as a result of a female-specific splicing event. Genes with these splicing characteristics have been created to bring about female-specific killing for other vector-control strategies^{51–53} suggesting such a toxin can be generated. However, diapause-specific genes and promoters have not yet been identified. In addition, because the cue-dependent transgene causes lethality, mutants that lack this activity, or the *Medea* toxin, which is required for drive, will be strongly selected for. Multiple copies of the cue-dependent lethality cassette, and the gene encoding the *Medea* toxin (the maternal promoter and its associated poly miRNA encoding transcript, each copy of which may include multiple

miRNAs, targeting different regions of the target transcript), can delay the effects of mutation and selection. However, they cannot prevent them. Therefore, use of such a technology to bring about a population crash would likely be most effective in contexts such as islands in which population size is limited and migration of wildtypes into the population is negligible. More detailed ecological models will be required to identify specific species and environments in which such a strategy for population suppression might be successful.

METHODS

Construction of miRNAs. The *Drosophila* miRNA mir6.1 stem-loop was modified to target *dah*, *o-fut1*, *gro*, or *trk*. To generate a mir6.1 stem-loop backbone that generates a mature miRNA complementary to one or the other of these target sites we annealed pairs of primers. For example, to make a miRNA that targets *dah* site 1, primers *dah-1-f* (5'-CTT AAT CAC AGC CTT TAA TGT AGG GAA ATA TAT AAC AAT ACA CTA AGT TAA TAT ACC ATA TCT-3') and *DAH-1-r* (5'-ATG TTA GGC ACT TTA GGT ACA GGG AAA TAT ATA ACA ATA AAC TAG ATA TGG TAT ATT AAC TTA G-3') were annealed; to target *dah* site 2, primers *dah-2-f* (5'-TTA AAC TTA ATC ACA GCC TTT AAT GTA ACC AGG ATG CGA ACT ATA CAC TAA GTT AAT ATA CCA TAT CTA G-3') and *dah-2-r* (5'-AAT GAT GTT AGG CAC TTT AGG TAC AAC CAG GAT GCG AAC TAT AAA CTA GAT ATG GTA TAT TAA CTT AG-3') were annealed; to target *o-fut1* target site 1, primers *o-fut1-1-f* (5'-AAA CTT AAT CAC AGC CTT TAA TGT AGT TTT ATT ACA TTG ATT ACG CTA AGT TAA TAT ACC ATA TCT AG-3') and *o-fut1-1-r* (5'-AAT GAT GTT AGG CAC TTT AGG TAC AGT TTT ATT ACA TTG ATT AAG CTA GAT ATG GTA TAT TAA CTT AGC G-3') were annealed; to target *o-fut1* target site 2, primers *o-fut1-2-f* (5'-ATC ACA GCC TTT AAT GTC AGG ATT ATC TAC TTA AAT CCT TAA GTT AAT ATA CCA TAT CTA AGT-3') and *o-fut1-2-r* (5'-ATG ATG TTA GGC ACT TTA GGT ACC AGG ATT ATC TAC TTA AAT ACT TAG ATA TGG TAT ATT AAC TTA AGG A-3') were annealed; to target *gro* target site 1, primers *gro-1-f* (5'-CTT AAT CAC AGC CTT TAA TGT TAC CAT AAA ACG CTG GCA ACC ATA AGT TAA TAT ACC ATA TCT ATG TT-3') and *gro-1-r* (5'-TGA TGT TAG GCA CTT TAG GTA CTA CCA TAA AAC GCT GGC AAA CAT AGA TAT GGT ATA TTA ACT TAT GG-3') were annealed; to target *gro* target site 2, primers *gro-2-f* (5'-TTA ATC ACA GCC TTT AAT GTG GAT CAA CTG CCA TCG AAA CAT TAA GTT AAT ATA CCA TAT CTA ATT T-3') and *gro-2-r* (5'-ATG TTA GGC ACT TTA GGT ACG GAT CAA CTG CCA TCG AAA AAT TAG ATA TGG TAT ATT AAC TTA ATG T-3') were annealed. To build up the miRNAs specific to *trk* we designed a polycistronic strand of 6 unique miRNAs. To target *trk* target site 1, primers *trk-1-f* (5'-CTT AAA CTT AAT CAC AGC CTT TAA TGT GCC AGC CAT GTT TCT GCG TCT ATA AGT TAA TAT ACC ATA TCT ATA TAC-3') and *trk-1-r* (5'-TAA TGA TGT TAG GCA CTT TAG GTA CGC CAG CCA TGT TTC TGC GTA TAT AGA TAT GGT ATA TTA ACT TAT AGA-3') were annealed; to target *trk* target site 2, primers *trk-2-f* (5'-CTT AAA CTT AAT CAC AGC CTT TAA TGT AGA AGA GCT ATC ATC TGG GCT ATA AGT TAA TAT ACC ATA TCT ATA TC-3') and *trk-2-r* (5'-TAA TGA TGT TAG GCA CTT TAG GTA CAG AAG AGC TAT CAT CTG GGA TAT AGA TAT GGT ATA TTA ACT TAT AGC CC-3') were annealed; to target *trk* target site 3,

primers *trk-3-f* (5'-CTT AAA CTT AAT CAC AGC CTT TAA TGT ACT GGA TTG ACT TAG GCT TCA ATA AGT TAA TAT ACC ATA TCT ATT TA-3') and *trk-3-r* (5'-TAA TGA TGT TAG GCA CTT TAG GTA CAC TGG ATT GAC TTA GGC TTA AAT AGA TAT GGT ATA TTA ACT TAT TGA-3') were annealed; to target *trk* target site 4, primers *trk-4-f* (5'-CTT AAA CTT AAT CAC AGC CTT TAA TGT ATG CTA TAC TAG TAG CGC CCT TTA AGT TAA TAT ACC ATA TCT AAA TG-3') and *trk-4-r* (5'-TAA TGA TGT TAG GCA CTT TAG GTA CAT GCT ATA CTA GTA GCG CCA TTT AGA TAT GGT ATA TTA ACT TAA AGG G-3') were annealed; to target *trk* target site 5, primers *trk-5-f* (5'-CTT AAA CTT AAT CAC AGC CTT TAA TGT TAT ACT AGT AGC GCC ATT TCA TTA AGT TAA TAT ACC ATA TCT AAT TA-3') and *trk-5-r* (5'-TAA TGA TGT TAG GCA CTT TAG GTA CTA TAC TAG TAG CGC CAT TTA ATT AGA TAT GGT ATA TTA ACT TAA TGA-3') were annealed; to target *trk* target site 6, primers *trk-6-f* (5'-CTT AAA CTT AAT CAC AGC CTT TAA TGT ATA CAC GTT TAA AAT GTC ACC ATA AGT TAA TAT ACC ATA TCT ATG TT-3') and *trk-6-r* (5'-TAA TGA TGT TAG GCA CTT TAG GTA CAT ACA CGT TTA AAA TGT CAA CAT AGA TAT GGT ATA TTA ACT TAT GGT-3') were annealed. These primary products were all then amplified using primers *mir6.1 5' NotI/FseI/BglII* (5'-TCG GGC GGC CGC ATT TGG CCG GCC AAA GAT CTT TTA AAG TCC ACA ACT CAT CAA GGA AAA TGA AAG TCA AAG TTG GCA GCT TAC TTA AAC TTA ATC ACA GCC TTT AAT GT-3') and *mir6.1 3' EcoRI/AscI/BamHI* (5'-TGA AGA ATT CAT TGG CGC GCC TTT GGA TCC AAA ACG GCA TGG TTA TTC GTG TGC CAA AAA AAA AAA TTA AAT AAT GAT GTT AGG CAC TTT AGG TAC-3'). These primers add *mir6.1* flanking sequences that are thought to promote miRNA processing, as well as several restriction sites. PCR products were purified with Qiagen (Valencia, CA) PCR purification columns, and then digested with several restriction enzymes. For *dah-1*, *o-fut1-1*, *gro-1*, and *trk-1* these were *EcoRI* and *BamHI*, for *dah-2*, *o-fut1-2*, *gro-2*, and *trk-2* these were *BglII* and *NotI*. Digested products were then ligated into *sry-bcd-GMR*⁷ cut with *EcoRI* and *NotI*, generating *pBcd-mir6.1-ofut1*, *pBcd-mir6.1-dah*, *pBcd-mir6.1-gro*, and *pBcd-mir6.1-trk2*. The *pBcd-mir6.1-trk2* was then digested with *AscI/BamHI*, and the *trk-3* miRNA was digested with *BglII/AscI* and cloned. This digest/cloning was repeated to sequentially clone in *trk4*, *trk5*, and *trk6* producing a plasmid with 6 miRNAs targeting *trk*: *pBcd-mir6.1-trk6*.

Construction of the Antidotes and Final *Medea* Constructs. Plasmids *pBcd-mir6.1-ofut1*, *pBcd-mir6.1-dah*, *pBcd-mir6.1-gro*, and *pBcd-mir6.1-trk6* were cut with *XhoI*. The gypsy insulator was amplified from genomic DNA using primers *gyp-f* (5'-AAG AGA TGT AGA GAT GGC ACA ATT GGT CGA CCT CGA GTC ACG TAA TAA GTG TGC GTT GAA TTT ATT-3') and *gyp-r* (5'-ATG AGG CGT CCA GGA TCC CAT GGG GTT CAT CTA ATG TTT AAA CAA TTG ATC GGC TAA ATG GTA TGG CAA-3'). The *bnk* promoter was amplified similarly using primers *bnk-f* (5'-TTT TCT TGC CAT ACC ATT TAG CCG ATC AAT TGT TTA AAC ATT AGA TGA ACC CCA TGG GAT CCT GG-3') and *bnk-r* (5'-AGA AGT AAG GTT CCT TCA CAA AGA TCC TGG CCG GCC TCG CCG AAT TCG TTG ACG GTT GAA GTA C-3'). The SV40 UTR was amplified using primers *SV40-f* (5'-CGT ACT TCA ACC GTC AAC GAA TTC GGC GAG GCC GGC CAG GAT CTT TGT GAA GGA ACC TTA CTT C-3') and *SV40-r* (5'-ATA ATT TGC GAG TAC GCA AAG CTT

GGC TGC AGG TCG ACG GAT CCA GAC ATG ATA AGA TAC ATT GAT G-3'). All three fragments were ligated together using one-step recombination technology,³⁴ producing a unique *fse-1* cloning site between the *bnk* promoter and SV40 3' UTR, resulting in plasmids *pBcd-mir6.1-ofut1-gyp-Bnk-fes1-SV40*, *pBcd-mir6.1-gro-gyp-Bnk-fes1-SV40*, *pBcd-mir6.1-dah-gyp-Bnk-fes1-SV40*, and *pBcd-mir6.1-trk-gyp-Bnk-fes1-SV40*.

The *O-fut1* coding region was amplified from a cDNA library using primers *o-fut1-anti-f* (5'-CAA CAG CAC ATT CGT ACT TCA ACC GTC AAC GAA TTC GGC ATG CAG TGG CTC AAA ATG AAG C-3') and *o-fut1-anti-r* (5'-ATG TCA CAC CAC AGA AGT AAG GTT CCT TCA CAA AGA TCC TTT ACA GCT CCT CGT GCA CGT TTG T-3'). One-step recombination technology was used to introduce these into *fse-1* digested *pBcd-mir6.1-ofut1-gyp-Bnk-fes1-SV40*, creating *Medea^{o-fut1}*. Note that because this *o-fut1* transcript lacks a 3' UTR present in the endogenous *o-fut1* transcript, it is not silenced by *mir6.1-ofut1* or *mir6.1-ofut1-2*, which target the *o-fut1* 3' UTR. The *dah* coding region was amplified from a cDNA library using primers *dah-anti-f-1* (5'-TCA ACA GCA CAT TCG TAC TTC AAC CGT CAA CGA ATT CGG CAT GCT GAG ATC GTC GGT GCC CGT-3') and *dah-anti-r-1* (5'-GTT GCC CTG TCC AAC TTG TAA TTG GCG TCT TGA TTG AAA TGG CCT AGT TTC TCG CAG GC-3'), and *dah-anti-f-2* (5'-GCC TGC GAG AAA CTA GGC CAT TTC AAT CAA GAC GCC AAT TAC AAG TTG GAC AGG GCA AC-3') and *dah-anti-r-2* (5'-ATG TCA CAC CAC AGA AGT AAG GTT CCT TCA CAA AGA TCC TGC TCA CGT GCT GAT GCG CCG CT-3'). One-step recombination was used to introduce these products into *fse-1* digested *pBcd-mir6.1-dah-gyp-Bnk-fes1-SV40*, creating *Medea^{dah}*. Note that because this *dah* transcript lacks a 5' UTR present in the endogenous *dah* transcript, it is not silenced by *mir6.1-dah-1*, which targets the *dah* 5' UTR. However, *mir6.1-dah-2* targets exon 2 of the CDS in *dah*, and therefore in order to make the antidote insensitive to this miRNA we recoded the nucleotide sequence in *dah* in such that it codes for the same amino acid sequence using different codons. The *gro* coding region was amplified from cDNA library using primers *gro-anti-f* (5'-CAT TCG TAC TTC AAC CGT CAA CGA ATT CGG CTT AAT TAA ATG TAT CCC TCA CCG GTG CGC-3') and *gro-anti-r* (5'-CAC ACC ACA GAA GTA AGG TTC CTT CAC AAA GAT CCT GCG GCC GCT TAA TAA ATA ACT TCG-3'). One-step recombination was used to introduce these products into *fse-1* digested *pBcd-mir6.1-gro-gyp-Bnk-fes1-SV40*, creating *Medea^{gro}*. Note that because this *gro* transcript lacks a 5' UTR present in the endogenous *gro* transcript, it is not silenced by *mir6.1-gro-1* or *mir6.1-gro-2*, which are engineered to target *gro* 5' UTR. The *trk* coding region was amplified in 3 fragments to recode target sites from cDNA library using primers for 268bp fragment 1 *trk-anti-1-f* (5'-GTC AAC GAA TTC GGC TTA ATT AAT GTT CTT AAG AAT TCT GTG TCC AAC TAT GAA ATC CCA AAG-3') and *trk-anti-1-R* (5'-GGG CAG CTC GTA CGA CGA TCG CTT ATA GCC TAA GTG GTA TGA TTT TTC CTC GGG CTC GCC C-3') and 296 bp fragment 2 *trk-anti-2-f* (5'-CGA GCC CGA GGA AAA ATC ATA CCA CTT AGG CTA TAA GCG ATC GTC GTA CGA GCT GCC CTT C-3') and *trk-anti-2-R* (5'-ATA GCG TGG AAA GTA GTT GAG TCC GAG ATC GAT CCA ATT GAT CTT CGA CGA GCA CTC CCA-3') and 310 bp fragment 3 *trk-anti-3-f* (5'-TCG TCG AAG ATC AAT TGG ATC GAT CTC GGA CTC AAC TAC TTT CCA CGC

TAT ATC CGT TCG-3') and *trk*-anti-3-R (5'-CAT TCT AGT TGT GGT TTG TCC AAA CTC ATC AAT GTC TAG TAT AGC ATA ACA CAT TCA CAG-3'). One-step recombination was used to introduce these products into *fse-1* digested pBcd-mir6.1-*trk*-gyp-Bnk-*fes1*-SV40, creating *Medea^{trk}*. Note that because this *trk* transcript lacks a 5' and 3' UTR present in the endogenous *trk* transcript, it is not silenced by mir6.1-*trk*-1, which targets the *dah* 5' UTR, or mir6.1-*trk*-4, mir6.1-*trk*-5, or mir6.1-*trk*-6, which target the 3' UTR. However, mir6.1-*trk*-2 and mir6.1-*trk*-3 target exon 1 of the CDS; in order to make the antidote insensitive to these miRNAs, we recoded the nucleotide sequence in *trk* such that it codes for the same amino acid sequence using different codons.

Transgenesis and Population Cage Experiments.

Germline transformants were generated in a *w¹¹¹⁸* background using standard techniques, by Rainbow Transgenic Flies, Inc. (www.rainbowgene.com, Newbury Park, CA). All fly experiments were carried out at 25 °C, ambient humidity in 250 mL bottles containing Lewis medium supplemented with live dry yeast. Fly rearing was carried out in a light-tight chamber placed in an incubator or in a darkened incubator. For all drive experiments three populations of 50 wildtype *w¹¹¹⁸* (+/+) males and 50 males heterozygous for either *dah* (*w¹¹¹⁸/Y*; *Medea^{dah}/+*) and *o-fut1* (*w¹¹¹⁸/Y*; *Medea^{o-fut1}/+*) were each independently crossed with 100 wildtype *w¹¹¹⁸* females in separate bottles in triplicate. For all sets of experiments flies were allowed to lay eggs for 4 days, after which adults were removed. Progeny were allowed to develop, eclose, and mate for another 10 days. All adult progeny were collected at this single time point, and their genotypes were determined using eye color as a marker (no *Medea*, *w¹¹¹⁸* = white eyed; *Medea*/+ = yellow/orange eyed; *Medea/Medea* = darker red eyed). Following counting, progeny were transferred to fresh bottles and allowed to lay eggs for 4 days, and the cycle was repeated.

Embryo and Adult Viability Determination. Adult viability for the crosses presented in Figure 2 was determined as follows. Fifty adult males of the indicated genotype were allowed to mate with 50 virgin females in bottles supplemented with dry yeast for 3 days. Ten bottles were established for each cross. Adults were then removed. Adult progeny from each bottle were collected, genotyped, and counted (either directly or by weighing and comparing with a standard) for 10 days following eclosion of the first progeny. For embryo viability counts, 2–4-day-old adult virgin females were allowed to mate with males of the relevant genotypes for 2–3 days in egg collection chambers supplemented with wet yeast paste. On the following day, a 3 h egg collection was carried out, after first having cleared old eggs from the females through a precollection period on a separate plate for 3 h. Embryos were isolated into groups of 100 and kept on an agar surface at 25 °C for 48–72 h. The percent survival was then determined by counting the number of unhatched embryos. Four groups of 100 embryos per cross were scored in each experiment, and each experiment was carried out three times. The results presented are averages from these three experiments. Embryo survival was normalized with respect to the percent survival observed in parallel experiments carried out with the *w¹¹¹⁸* strain used for transgenesis.

Total RNA Isolation. Total RNA was extracted from WT (+/+) and heterozygous *Medea^{M-DAH/+}* or *Medea^{M-O-fut1/+}* females stage 14 oocytes and 0–1 h staged embryos. Samples were flash frozen at specific time points, and total RNA was extracted using the Ambion mirVana mRNA isolation kit

(Ambion/Applied Biosystems, Austin, TX). All sample collections were staged in the incubator at a relative humidity 60%, 25 °C with a 12 h/12 h light cycle until the desired time point was reached and then immediately flash frozen. Following the extraction, RNA was treated with Ambion Turbo DNase (Ambion/Applied Biosystems, Austin, TX). The quality of RNA was assessed using a Bioanalyzer 2100 (Agilent Technologies, Santa Clara, CA) and a NanoDrop 1000 UV-vis spectrophotometer (NanoDrop Technologies/Thermo Scientific, Wilmington, DE). RNA was then prepared for sequencing using an Illumina mRNA-Seq Sample Preparation Kit (Illumina San Diego, CA).

Small RNA Extraction, Cloning, and Sequencing.

Twenty micrograms of total RNA from each time point was size fractionated in 15% TBE-urea polyacrylamide gels. For both the s14-oocytes and 0–1 h embryo time points, an 18–32 nt band was excised and sequenced. Ethanol-precipitated RNA was ligated to an HPLC-purified 3' linker using T4 RNA ligase (Ambion/Applied Biosystems, Austin, TX) and then purified in a 15% TBE-urea polyacrylamide gel. Ligation products were recovered by high-salt elution following electrophoresis. A HPLC-purified 5' RNA linker was ligated to the RNA using T4-RNA ligase, and the RNA was purified in a 15% TBE-urea polyacrylamide gel. Ligation products were recovered by high-salt elution following electrophoresis. Reverse transcription using SSIII (Invitrogen, Carlsbad, CA) was performed, and cDNA was amplified using Phusion polymerase (Finnzymes Oy, Espoo, Finland). Amplified cDNA libraries were purified using a 2% agarose gel prior to sequencing using the Illumina Genome Analyzer II system. Linker and primer sequences are indicated below: 3' Ligation linker: Modban (AMP-5'p=5'pCTG TAG GCA CCA TCA ATdideoxyC-3'); 5' Ligation Solexa Linker (5'-rArCrAr CrUrC rUrUrU rCrCrC rUrArC rArCrG rArCrG rCrUrC rUrUrC rCrGrA rUrC-3'); 3' RT primer: BanOne (5'-ATT GAT GGT GCC TAC AG-3'); 5' PCR primer Sol_5_SBS3 (5'-AAT GAT ACG GCG ACC ACC GAA CAC TCT TTC CCT ACA CGA CG-3'); 3' PCR primer Sol_3_Modban (5'-CAA GCA GAA GAC GGC ATA CGA TTG ATG GTG CCT ACA G-3').

Poly(A+) Read Alignment and Quantification. The poly(A+) reads were processed and aligned to a reference index we generated for the *Drosophila melanogaster* genome (obtained from www.flybase.org), using TopHat v1.4.1.⁵⁵ Reads were aligned using both default parameters. TopHat incorporates the Bowtie v0.12.7 algorithm to perform the alignment.⁵⁶ The aligned read files were processed by Cufflinks v0.9.3⁵⁷ Cufflinks uses the normalized RNA-Seq fragment counts to measure the relative abundances of transcripts. The unit of measurement is fragments per kilobase of exon per million fragments mapped (FPKM).

Small RNA Read Alignment and Quantification. The 5' and 3' adapter sequences for the small RNA reads were removed using custom Perl scripts requiring a minimal match to the adapter sequence of 6 bp and a minimal size of 18 bp and maximum size of 32 bp (sequences for the adapters are noted above). The trimmed sequences were aligned to the *Drosophila melanogaster* reference genome using bowtie v0.12.7, allowing no mismatches and unlimited alignments/read. We determined the small RNA abundance using custom in house Perl scripts in which we quantified the read count per million of mapped reads (RPM) for each genomic position. Each endogenous miRNA was determined by merging the genomic coordinates from miRbase v19 with our genomic coordinates using custom

Perl scripts. Levels of all endogenous miRNAs were quantified in terms of RPM and ranked according to expression level. Levels of synthetic miRNAs were quantified in terms of RPM and in terms of rank and rank percentile with respect to the subset of endogenous miRNAs.

Fitness Cost Calculation. We modeled *Medea* dynamics assuming random mating, discrete generations, and 100% toxin efficiency. The proportions of the k th generation that are WT, heterozygous and homozygous for *Medea* are denoted by u_k , v_k , and w_k , respectively. Considering all possible mating pairs and taking into account that WT offspring of heterozygous mothers are unviable, the genotypes of embryos in the next generation are described by the ratio $\hat{u}_{k+1}:\hat{v}_{k+1}:\hat{w}_{k+1}$, where

$$\hat{u}_{k+1} = u_k^2 + 0.5u_kv_k \quad (1)$$

$$\hat{v}_{k+1} = 2u_kv_k + 0.5v_k^2 + u_kv_k + w_kv_k \quad (2)$$

$$\hat{w}_{k+1} = w_k^2 + w_kv_k + 0.25v_k^2 \quad (3)$$

Normalizing these ratios and taking into account fitness costs, the genotype frequencies in the next generation are given by

$$u_{k+1} = \hat{u}_{k+1}/W_{k+1} \quad (4)$$

$$v_{k+1} = \hat{v}_{k+1}(1 - hs)/W_{k+1} \quad (5)$$

$$w_{k+1} = \hat{w}_{k+1}(1 - s)/W_{k+1} \quad (6)$$

Here, s and hs represent the fitness costs associated with being homozygous or heterozygous for the element. For additive fitness costs, $h = 0.5$, and for dominant fitness costs, $h = 1$. W_{k+1} is a normalizing term given by

$$w_{k+1} = \hat{u}_{k+1} + \hat{v}_{k+1}(1 - hs) + \hat{w}_{k+1}(1 - s) \quad (7)$$

The likelihood of the data was calculated by assuming a binomial distribution of WT and red-eyed individuals and by using the model predictions to generate expected proportions for each fitness cost, i.e., by calculating the log-likelihood,

$$\log L(s) = \sum_{i=1}^3 \sum_{k=1}^{18} \log \left(\frac{R_{i,k} + WT_{i,k}}{R_{i,k}} \right) + R_{i,k} \log(v_k(s) + w_k(s)) + WT_{i,k} \log(u_k(s)) \quad (8)$$

Here, $R_{i,k}$ and $WT_{i,k}$ are the number of red-eyed and WT individuals at generation k in experiment i , and the expected genotype frequencies are fitness cost-dependent. The best estimate of the fitness cost is that having the highest log-likelihood. 95% confidence intervals were estimated using a Markov Chain Monte Carlo sampling procedure.

Modeling a *Medea*-Dependent Population Crash. *Population Frequency Model.* Releases of *Medea* individuals are likely to be all-male, since male mosquitoes do not bite and male agricultural pests do not lay eggs. Additionally, the external cues envisaged may induce a gender-specific fitness cost. For this reason, we consider a model allowing for different genotype frequencies among males and females. We denote the proportion of the k th generation that are females having the genotypes mm, Mm, and MM by $u_{f,k}$, $v_{f,k}$, and $w_{f,k}$, respectively. The corresponding proportions for males are $u_{m,k}$, $v_{m,k}$, and $w_{m,k}$. By considering all possible mating pairs, the genotype frequencies in the next generation are given by

$$u_{f,k+1} = (v_{m,k}u_{f,k} + 2u_{m,k}u_{f,k})/\sigma_k \quad (9)$$

$$u_{m,k+1} = (v_{m,k}u_{f,k} + 2u_{m,k}u_{f,k})/\sigma_k \quad (10)$$

$$v_{f,k+1} = (2u_{m,k}w_{f,k} + v_{m,k}w_{f,k} + u_{m,k}v_{f,k} + v_{m,k}v_{f,k} + 2w_{m,k}u_{f,k} + v_{m,k}u_{f,k} + w_{m,k}v_{f,k})(1 - h_f s_f)/\sigma_k \quad (11)$$

$$v_{m,k+1} = (2u_{m,k}w_{f,k} + v_{m,k}w_{f,k} + u_{m,k}v_{f,k} + v_{m,k}v_{f,k} + 2w_{m,k}u_{f,k} + v_{m,k}u_{f,k} + w_{m,k}v_{f,k})(1 - h_m s_m)/\sigma_k \quad (12)$$

$$w_{f,k+1} = (2w_{m,k}w_{f,k} + v_{m,k}w_{f,k} + w_{m,k}v_{f,k} + 0.5v_{m,k}v_{f,k})(1 - s_f)/\sigma_k \quad (13)$$

$$w_{m,k+1} = (2w_{m,k}w_{f,k} + v_{m,k}w_{f,k} + w_{m,k}v_{f,k} + 0.5v_{m,k}v_{f,k})(1 - s_m)/\sigma_k \quad (14)$$

Here, s_f and $h_f s_f$ represent the fitness costs associated with being homozygous or heterozygous for the *Medea* element in females, s_m and $h_m s_m$ represent the corresponding fitness costs in males, and σ_k represents the proportion of embryos that survive to maturity. This is given by

$$\sigma_k = 2v_{m,k}u_{f,k} + 4u_{m,k}u_{f,k} + (2u_{m,k}w_{f,k} + v_{m,k}w_{f,k} + u_{m,k}v_{f,k} + v_{m,k}v_{f,k} + 2w_{m,k}u_{f,k} + v_{m,k}u_{f,k} + w_{m,k}v_{f,k})(2 - h_f s_f - h_m s_m) + (2w_{m,k}w_{f,k} + v_{m,k}w_{f,k} + w_{m,k}v_{f,k} + 0.5v_{m,k}v_{f,k})(2 - s_f - s_m) \quad (15)$$

Using these difference equations, the equilibria, thresholds, and time-series dynamics of the *Medea* element can be calculated. From previous analyses^{7,12} we know that a *Medea* element released into a population above a certain fitness-dependent threshold will spread to transgene fixation, meaning that all individuals will eventually be either heterozygous or homozygous for the construct. The threshold is particularly difficult to derive but is 0 for a fitness cost of 0 and increases monotonically with fitness cost magnitude.

Discrete Population Model. A stochastic model can be used to theoretically explore the idea of using *Medea* to induce a population crash because, in this way, discrete populations and the concept of 0 population size can be considered. Density dependence is also an important consideration because, at low population sizes, larval competition is reduced and a single female can produce more offspring that survive to adulthood. We adapt a general density-dependent model⁵⁸ of the form

$$N_{k+1} = R_0 N_k e^{-\alpha N_k} \quad (16)$$

Here, N_k represents the absolute population size at generation k , R_0 represents the average number of offspring produced per insect that survive to adulthood in a density-independent population (or the average number of female offspring that each female gives birth to), and α represents the strength of density dependence. This last term is related to the carrying capacity, K , of the habitat by

$$\alpha = \ln(R_0)/K \quad (17)$$

For agricultural pests under field conditions, R_0 typically ranges between 3 and 11 but could be much higher in some populations.^{50,59} We assume a default value of $R_0 = 7$ and a carrying capacity of 10,000.

We adapt this model to account for the fact that the number of offspring in the next generation is determined by the number of adult females, an important consideration as we consider male-only releases of *Medea* homozygotes, and the external cue we envisage causes female-specific lethality. We also adapt the model to account for the fact that the maternal toxin is early acting, leading to offspring lethality prior to larval competition, and hence unviable offspring do not contribute to density dependence. On the other hand, we assume that fitness costs associated with the construct are late-acting, manifesting themselves at the adult stage and hence not reducing the strength of density dependence. Similar assumptions are made in models of other genetic population suppression systems.⁶⁰

Taking these considerations into account, the total expected population size at generation $k + 1$ is given by

$$\bar{N}_{k+1} = 2R_0 N_{f,k} \sigma_k e^{-2\alpha N_{f,k} \sigma_{l,k}} \quad (18)$$

Here, $N_{f,k}$ represents the female population size at generation k , and $2R_0 N_{f,k}$ represents the number of male and female offspring that females give birth to, neglecting offspring lethality and density dependence. The population size that is subject to density dependence is given by $2R_0 N_{f,k} \sigma_{l,k}$, where $\sigma_{l,k}$ represents the proportion of offspring that survive to the larval stage. Finally, σ_k represents the proportion of offspring that survive to mate and is given in eq 15. It is worth noting that, for an entirely wildtype population, we recover the general density-dependent model in eq 16 since $2N_{f,k} = N_k$ and $\sigma_k = \sigma_{l,k} = 1$.

With this density-dependent framework in place, we integrate the population genetics of the *Medea* element into the model. At generation k , we denote the number of females having genotypes mm, Mm, and MM at generation k by $U_{f,k}$, $V_{f,k}$, and $W_{f,k}$ respectively. The corresponding numbers for males are $U_{m,k}$, $V_{m,k}$, and $W_{m,k}$. This framework allows us to express the female population size at generation k by

$$N_{f,k} = U_{f,k} + V_{f,k} + W_{f,k} \quad (19)$$

and the total population size at generation k by

$$N_k = U_{f,k} + V_{f,k} + W_{f,k} + U_{m,k} + V_{m,k} + W_{m,k} \quad (20)$$

The genotype frequencies at generation k are therefore

$$\begin{aligned} & (u_{f,k}, u_{m,k}, v_{f,k}, v_{m,k}, w_{f,k}, w_{m,k}) \\ &= (U_{f,k}, U_{m,k}, V_{f,k}, V_{m,k}, W_{f,k}, W_{m,k}) / N_k \end{aligned} \quad (21)$$

Equations 1–7 then describe the expected genotype frequencies in the next generation, and the proportion of offspring that survive to the larval stage is essentially eq 15 without fitness costs, i.e.,

$$\begin{aligned} \sigma_{l,k} &= 4u_{m,k}u_{f,k} + 2u_{m,k}v_{f,k} + 4u_{m,k}w_{f,k} + 4v_{m,k}u_{f,k} \\ &+ 3v_{m,k}v_{f,k} + 4v_{m,k}w_{f,k} + 4w_{m,k}u_{f,k} + 4w_{m,k}v_{f,k} \\ &+ 4w_{m,k}w_{f,k} \end{aligned} \quad (22)$$

Having calculated the expected total population size at generation $k + 1$, using eq 18, the expected numbers of individuals having each genotype at generation $k + 1$ is given by

$$(\bar{U}_{f,k+1}, \bar{V}_{f,k+1}, \bar{W}_{f,k+1}) = \bar{N}_{k+1} (u_{f,k+1}, v_{f,k+1}, w_{f,k+1}) \quad (23)$$

$$(\bar{U}_{m,k+1}, \bar{V}_{m,k+1}, \bar{W}_{m,k+1}) = \bar{N}_{k+1} (u_{m,k+1}, v_{m,k+1}, w_{m,k+1}) \quad (24)$$

Stochastic effects are modeled by sampling the number of individuals having each genotype from a Poisson distribution with a mean equal to the expected number of individuals having this genotype. This has the effect of discretizing the population, which is appropriate for modeling a population crash. Illustrating this calculation for Mm females, an absolute population size of $V_{f,k+1}$ can be calculated from an expected population size of $\bar{V}_{f,k+1}$ by sampling from the Poisson distribution:

$$\Pr(V_{f,k+1} | \bar{V}_{f,k+1}) = e^{-\bar{V}_{f,k+1}} \frac{(\bar{V}_{f,k+1})^{V_{f,k+1}}}{V_{f,k+1}!} \quad (25)$$

This procedure can be repeated for all six genotypes, so that the total population size at generation $k + 1$ becomes

$$\begin{aligned} N_{k+1} &= U_{m,k+1} + U_{f,k+1} + V_{m,k+1} + V_{f,k+1} + W_{m,k+1} \\ &+ W_{f,k+1} \end{aligned} \quad (26)$$

Considering a release of $X_{m,0}$ males homozygous for the *Medea* element at generation 0, the initial condition is given by

$$(U_{f,0} + V_{f,0} + W_{f,0}) = (0.5K, 0, 0) \quad (27)$$

$$(U_{m,0} + V_{m,0} + W_{m,0}) = (0.5K, 0, X_{m,0}) \quad (28)$$

Subsequent releases of $X_{m,k}$ male homozygotes at generation k can be modeled by making the following substitution:

$$(U_{m,k}, V_{m,k}, W_{m,k}) \leftarrow (U_{m,k}, V_{m,k}, W_{m,k} + X_{m,k}) \quad (29)$$

We consider a single, all-male release representing 50% of the population at the time of release, i.e., $X_{m,0} = K$.

■ ASSOCIATED CONTENT

📄 Supporting Information

Supplementary tables containing microRNA expression profiles for s14 ovary and 0–1 h embryo, top 20 miRNAs in s14 ovary, and top 20 miRNAs in 0–1 h embryo. This material is available free of charge via the Internet at <http://pubs.acs.org>.

■ AUTHOR INFORMATION

Corresponding Author

*E-mail: haybruce@caltech.edu.

Author Contributions

#These authors contributed equally to this work.

Notes

The authors declare no competing financial interest.

■ ACKNOWLEDGMENTS

We would like to thank Lorain Schaeffer and Vijaya Kumar for help with library preparations and sequencing. This work was supported by grants to B.A.H. from the NIH (DP10D003878) and to J.M.M. from the Medical Research Council, U.K., and in part by a grant from the Foundation for the NIH through the Grand Challenges in Global Health initiative (Anthony A. James, PI, UC Irvine).

■ REFERENCES

- (1) Nicholson, G. M. (2007) Fighting the global pest problem: preface to the special Toxinon issue on insecticidal toxins and their potential for insect pest control. *Toxinon* 49, 413–422.
- (2) Schmid-Hempel, P. (2005) Evolutionary ecology of insect immune defenses. *Annu. Rev. Entomol.* 50, 529–551.
- (3) Tripet, F., Aboagye-Antwi, F., and Hurd, H. (2008) Ecological immunology of mosquito-malaria interactions. *Trends Parasitol.* 24, 219–227.
- (4) Lambrechts, L., Koella, J. C., and Boete, C. (2008) Can transgenic mosquitoes afford the fitness cost? *Trends Parasitol.* 24, 4–7.
- (5) Braig, H. R., and Yan, G. (2001) The spread of genetic constructs in natural insect populations, in *Genetically Engineered Organisms: Assessing Environmental and Human Health Effects* (Letourneau, D. K., and Burrows, B. E., Eds.), pp 251–314, CRC Press, Boca Raton, FL.
- (6) Sinkins, S. P., and Gould, F. (2006) Gene drive systems for insect disease vectors. *Nat. Rev. Genet.* 7, 427–435.
- (7) Chen, C. H., Huang, H., Ward, C. M., Su, J. T., Schaeffer, L. V., Guo, M., and Hay, B. A. (2007) A synthetic maternal-effect selfish genetic element drives population replacement in *Drosophila*. *Science* 316, 597–600.
- (8) Beeman, R. W., Friesen, K. S., and Denell, R. E. (1992) Maternal-effect selfish genes in flour beetles. *Science* 256, 89–92.
- (9) Hastings, I. M. (1994) Selfish DNA as a method of pest control. *Philos. Trans. R. Soc. London, Ser. B* 344, 313–324.
- (10) Wade, M. J., and Beeman, R. W. (1994) The population dynamics of maternal-effect selfish genes. *Genetics* 138, 1309–1314.
- (11) Smith, N. G. (1998) The dynamics of maternal-effect selfish genetic elements. *J. Theor. Biol.* 191, 173–180.
- (12) Ward, C. M., Su, J. T., Huang, Y., Lloyd, A. L., Gould, F., and Hay, B. A. (2011) *Medea* selfish genetic elements as tools for altering traits of wild populations: a theoretical analysis. *Evolution* 65, 1149–1162.
- (13) Maxton-Kuchenmeister, J., Handel, K., Schmidt-Ott, U., Roth, S., and Jackle/Jackle, H. (1999) Toll homologue expression in the beetle *Tribolium* suggests a different mode of dorsoventral patterning than in *Drosophila* embryos. *Mech. Dev.* 83, 107–114.
- (14) Zhang, C. X., Lee, M. P., Chen, A. D., Brown, S. D., and Hsieh, T. (1996) Isolation and characterization of a *Drosophila* gene essential for early embryonic development and formation of cortical cleavage furrows. *J. Cell Biol.* 134, 923–934.
- (15) Jin, H., Tan, S., Hermanowski, J., Bohm, S., Pacheco, S., McCauley, J. M., Greener, M. J., Hinitis, Y., Hughes, S. M., Sharpe, P. T., and Roberts, R. G. (2007) The dystrotelin, dystrophin and dystrobrevin superfamily: new paralogues and old isoforms. *BMC Genomics* 8, 19.
- (16) Sasamura, T., Sasaki, N., Miyashita, F., Nakao, S., Ishikawa, H. O., Ito, M., Kitagawa, M., Harigaya, K., Spana, E., Bilder, D., Perrimon, N., and Matsuno, K. (2003) *neurotic*, a novel maternal neurogenic gene, encodes an O-fucosyltransferase that is essential for Notch-Delta interactions. *Development* 130, 4785–4795.
- (17) Okajima, T., and Irvine, K. D. (2002) Regulation of notch signaling by o-linked fucose. *Cell* 111, 893–904.
- (18) Okajima, T., Xu, A., Lei, L., and Irvine, K. D. (2005) Chaperone activity of protein O-fucosyltransferase 1 promotes notch receptor folding. *Science* 307, 1599–1603.
- (19) Okajima, T., Reddy, B., Matsuda, T., and Irvine, K. D. (2008) Contributions of chaperone and glycosyltransferase activities of O-fucosyltransferase 1 to Notch signaling. *BMC Biol.* 6, 1.
- (20) Shi, S., and Stanley, P. (2003) Protein O-fucosyltransferase 1 is an essential component of Notch signaling pathways. *Proc. Natl. Acad. Sci. U.S.A.* 100, 5234–5239.
- (21) Schejter, E. D., and Wieschaus, E. (1993) bottleneck acts as a regulator of the microfilament network governing cellularization of the *Drosophila* embryo. *Cell* 75, 373–385.
- (22) Shelton, C. A., and Wasserman, S. A. (1993) *pelle* encodes a protein kinase required to establish dorsoventral polarity in the *Drosophila* embryo. *Cell* 72, 515–525.
- (23) Letsou, A., Alexander, S., Orth, K., and Wasserman, S. A. (1991) Genetic and molecular characterization of *tube*, a *Drosophila* gene maternally required for embryonic dorsoventral polarity. *Proc. Natl. Acad. Sci. U.S.A.* 88, 810–814.
- (24) Michellod, M. A., Forquignon, F., Santamaria, P., and Randsholt, N. B. (2003) Differential requirements for the neurogenic gene *almondex* during *Drosophila melanogaster* development. *Genesis* 37, 113–122.
- (25) Johnson, K., Knust, E., and Skaer, H. (1999) *bloated tubules* (blot) encodes a *Drosophila* member of the neurotransmitter transporter family required for organisation of the apical cytocortex. *Dev. Biol.* 212, 440–454.
- (26) Parks, S., and Wieschaus, E. (1991) The *Drosophila* gastrulation gene *concertina* encodes a G alpha-like protein. *Cell* 64, 447–458.
- (27) Field, C. M., Coughlin, M., Doberstein, S., Marty, T., and Sullivan, W. (2005) Characterization of anillin mutants reveals essential roles in septin localization and plasma membrane integrity. *Development* 132, 2849–2860.
- (28) Sprenger, F., Stevens, L. M., and Nusslein-Volhard, C. (1989) The *Drosophila* gene *torso* encodes a putative receptor tyrosine kinase. *Nature* 338, 478–483.
- (29) Paroush, Z., Finley, R. L., Jr., Kidd, T., Wainwright, S. M., Ingham, P. W., Brent, R., and Ish-Horowicz, D. (1994) *Groucho* is required for *Drosophila* neurogenesis, segmentation, and sex determination and interacts directly with hairy-related bHLH proteins. *Cell* 79, 805–815.
- (30) Casanova, J., Furriols, M., McCormick, C. A., and Struhl, G. (1995) Similarities between trunk and spatzle, putative extracellular ligands specifying body pattern in *Drosophila*. *Genes Dev.* 9, 2539–2544.
- (31) Casali, A., and Casanova, J. (2001) The spatial control of Torso RTK activation: a C-terminal fragment of the Trunk protein acts as a signal for Torso receptor in the *Drosophila* embryo. *Development* 128, 1709–1715.
- (32) Ross, S. (2009) *A First Course in Probability*, 8 ed., Prentice Hall, New York.
- (33) Moussian, B., and Roth, S. (2005) Dorsoventral axis formation in the *Drosophila* embryo—shaping and transducing a morphogen gradient. *Curr. Biol.* 15, R887–899.
- (34) Mazumdar, A., and Mazumdar, M. (2002) How one becomes many: blastoderm cellularization in *Drosophila melanogaster*. *Bioessays* 24, 1012–1022.
- (35) Guruharsha, K. G., Kankel, M. W., and Artavanis-Tsakonas, S. (2012) The Notch signalling system: recent insights into the complexity of a conserved pathway. *Nat. Rev. Genet.* 13, 654–666.
- (36) Huntzinger, E., and Izaurralde, E. (2011) Gene silencing by microRNAs: contributions of translational repression and mRNA decay. *Nat. Rev. Genet.* 12, 99–110.
- (37) Benoit, P., Papin, C., Kwak, J. E., Wickens, M., and Simonelig, M. (2008) PAP- and GLD-2-type poly(A) polymerases are required sequentially in cytoplasmic polyadenylation and oogenesis in *Drosophila*. *Development* 135, 1969–1979.
- (38) Berezikov, E., Robine, N., Samsonova, A., Westholm, J. O., Naqvi, A., Hung, J. H., Okamura, K., Dai, Q., Bortolamiol-Becet, D., Martin, R., Zhao, Y., Zamore, P. D., Hannon, G. J., Marra, M. A., Weng, Z., Perrimon, N., and Lai, E. C. (2011) Deep annotation of *Drosophila melanogaster* microRNAs yields insights into their processing, modification, and emergence. *Genome Res.* 21, 203–215.
- (39) Ameres, S. L., Horwich, M. D., Hung, J. H., Xu, J., Ghildiyal, M., Weng, Z., and Zamore, P. D. (2010) Target RNA-directed trimming and tailing of small silencing RNAs. *Science* 328, 1534–1539.
- (40) Ameres, S. L., Hung, J. H., Xu, J., Weng, Z., and Zamore, P. D. (2011) Target RNA-directed tailing and trimming purifies the sorting of endo-siRNAs between the two *Drosophila* Argonaute proteins. *RNA* 17, 54–63.
- (41) Reynolds, A., Leake, D., Boese, Q., Scaringe, S., Marshall, W. S., and Khvorova, A. (2004) Rational siRNA design for RNA interference. *Nat. Biotechnol.* 22, 326–330.

- (42) Okamura, K., Liu, N., and Lai, E. C. (2009) Distinct mechanisms for microRNA strand selection by *Drosophila* Argonautes. *Mol. Cell* 36, 431–444.
- (43) Czech, B., and Hannon, G. J. (2011) Small RNA sorting: matchmaking for Argonautes. *Nat. Rev. Genet.* 12, 19–31.
- (44) Forstemann, K., Horwich, M. D., Wee, L., Tomari, Y., and Zamore, P. D. (2007) *Drosophila* microRNAs are sorted into functionally distinct argonaute complexes after production by Dicer-1. *Cell* 130, 287–297.
- (45) Ballarino, M., Pagano, F., Girardi, E., Morlando, M., Cacchiarelli, D., Marchioni, M., Proudfoot, N. J., and Bozzoni, I. (2009) Coupled RNA processing and transcription of intergenic primary microRNAs. *Mol. Cell. Biol.* 29, 5632–5638.
- (46) Chatterjee, S., and Grosshans, H. (2009) Active turnover modulates mature microRNA activity in *Caenorhabditis elegans*. *Nature* 461, 546–549.
- (47) Clark, M. S., and Worland, M. R. (2008) How insects survive the cold: molecular mechanisms—a review. *J. Comp. Physiol., B* 178, 917–933.
- (48) Hahn, D. A., and Denlinger, D. L. (2011) Energetics of insect diapause. *Annu. Rev. Entomol.* 56, 103–121.
- (49) Clements, A. N. (1992) *The Biology of Mosquitoes. Vol 1, Development, Nutrition and Reproduction*, Chapman & Hall, New York.
- (50) Dyck, V. A., Hendrichs, J., and Robinson, A. S. (Eds.) (2005) *Sterile Insect Technique: Principles and Practice in Area-Wide Integrated Pest Management*, Springer, Dordrecht, The Netherlands.
- (51) Fu, G., Condon, K. C., Epton, M. J., Gong, P., Jin, L., Condon, G. C., Morrison, N. I., Dafa'alla, T. H., and Alphey, L. (2007) Female-specific insect lethality engineered using alternative splicing. *Nat. Biotechnol.* 25, 353–357.
- (52) Fu, G., Lees, R. S., Nimmo, D., Aw, D., Jin, L., Gray, P., Berendonk, T. U., White-Cooper, H., Scaife, S., Kim Phuc, H., Marinotti, O., Jasinskiene, N., James, A. A., and Alphey, L. (2010) Female-specific flightless phenotype for mosquito control. *Proc. Natl. Acad. Sci. U.S.A.* 107, 4550–4554.
- (53) Schetelig, M. F., and Handler, A. M. (2012) A transgenic embryonic sexing system for *Anastrepha suspensa* (Diptera: Tephritidae). *Insect Biochem. Mol. Biol.* 42, 790–795.
- (54) Gibson, D. G., Young, L., Chuang, R. Y., Venter, J. C., Hutchison, C. A., 3rd, and Smith, H. O. (2009) Enzymatic assembly of DNA molecules up to several hundred kilobases. *Nat. Methods* 6, 343–345.
- (55) Trapnell, C., Pachter, L., and Salzberg, S. L. (2009) TopHat: discovering splice junctions with RNA-Seq. *Bioinformatics* 25, 1105–1111.
- (56) Langmead, B., Trapnell, C., Pop, M., and Salzberg, S. L. (2009) Ultrafast and memory-efficient alignment of short DNA sequences to the human genome. *Genome Biology* 10, R25.
- (57) Trapnell, C., Williams, B. A., Pertea, G., Mortazavi, A., Kwan, G., van Baren, M. J., Salzberg, S. L., Wold, B. J., and Pachter, L. (2010) Transcript assembly and quantification by RNA-Seq reveals unannotated transcripts and isoform switching during cell differentiation. *Nat. Biotechnol.* 28, 511–515.
- (58) Bellows, T. S. (1981) The descriptive properties of some models for density dependence. *J. Anim. Ecol.* 50, 139–156.
- (59) Dye, C. (1984) Models for the population dynamics of the yellow fever mosquito, *Aedes aegypti*. *J. Anim. Ecol.* 53, 247–268.
- (60) Alphey, N., Bonsall, M. B., and Alphey, L. (2011) Modeling resistance to genetic control of insects. *J. Theor. Biol.* 270, 42–55.