Progress towards engineering gene drives for population control

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ABSTRACT

Vector-borne diseases, such as dengue, Zika and malaria, are a major cause of morbidity and mortality worldwide. These diseases have proven difficult to control and currently available management tools are insufficient to eliminate them in many regions. Gene drives have the potential to revolutionize vector-borne disease control. This suite of technologies has advanced rapidly in recent years as a result of the availability of new, more efficient gene editing technologies. Gene drives can favorably bias the inheritance of a linked disease-refractory gene, which could possibly be exploited (i) to generate a vector population incapable of transmitting disease or (ii) to disrupt an essential gene for viability or fertility, which could eventually eliminate a population. Importantly, gene drives vary in characteristics such as their transmission efficiency, confinability and reversibility, and their potential to develop resistance to the drive mechanism. Here, we discuss recent advancements in the gene drive field, and contrast the benefits and limitations of a variety of technologies, as well as approaches to overcome these limitations. We also discuss the current state of each gene drive technology and the technical considerations that need to be addressed on the pathway to field implementation. While there are still many obstacles to overcome, recent progress has brought us closer than ever before to genetic-based vector modification as a tool to support vector-borne disease elimination efforts worldwide.

KEY WORDS: CRISPR, Cas9, Split drive, Medea, ClvR, Homing drives

Introduction

Half of the global population is currently at risk of mosquito-borne diseases such as dengue (Brady et al., 2012; Kraemer et al., 2015) or malaria (World Health Organization, 2019). There have been outstanding reductions in global malaria transmission over the last two decades as a result of integrated mosquito management programs that have focused on reducing human–mosquito contact and suppressing mosquito populations (Bhatt et al., 2015; Cibulskis et al., 2016; World Health Organization, 2019), but other mosquito-borne diseases, such as dengue, are on the rise (Huang et al., 2019; Stanaway et al., 2016). For the majority of arboviruses, there are no available drugs or vaccines, and while there are drugs and vaccines available for malaria, their efficacy at the population level is limited. Mosquito-targeted interventions are the primary methods to prevent disease transmission; however, given the widespread resistance to insecticides, escalating burden of transmission and deaths worldwide related to mosquito-borne disease, new technologies are in high demand. A tool that has sparked significant enthusiasm is gene drive-modified mosquitoes (Champer et al., 2016; Esvelt et al., 2014). Gene drives have promise to be a transformative technology that leads to the elimination of burdensome vector-borne diseases (Feachem et al., 2019; Raban and Akbari, 2017).

Many varieties of engineered gene drives have been developed to date (Fig. 1 and Table 1). Each of these drives are designed to bias the inheritance of a gene or gene modification, known as an effector, so it is inherited more frequently than would be expected by Mendelian segregation. Over time, the gene drive can therefore spread a linked effector into a wild population. In the context of vector–pathogen systems, gene drives are engineered to facilitate biased inheritance of an effector that either confers resistance to a pathogen or suppresses the vector population by generating a non-functional essential sterility or lethality gene. These approaches, known as population replacement and population suppression, respectively, aim to reduce, or possibly eliminate, the targeted mosquito-borne pathogens. While these gene drive systems were initially proposed for population replacement or suppression many years ago (Burt, 2003; Curtis, 1968; Marshall and Akbari, 2016), recent advances in gene editing technologies, principally the clustered regularly interspaced short palindromic repeats (CRISPR) and CRISPR-associated protein 9 (Cas9)-based genome engineering technologies (Jinek et al., 2012), have greatly accelerated the development of proof-of-concept gene drive systems (Esvelt et al., 2014) with demonstrations in mosquitoes (Gantz et al., 2015; Hammond et al., 2016; Kyrou et al., 2018; Li et al., 2019 preprint).

Gene drives may be an essential tool to make genetic mosquito-borne disease control affordable at scale in the field. One of the principal advantages of gene drive systems is that they can be used to overcome fitness costs associated with the effectors and genetic elements required for the stable and heritable transmission of the desirable synthetic genes into a wild population (Burt, 2003; Unckless et al., 2015). In the absence of a gene drive, these fitness costs have caused the rapid elimination of effectors from cage populations (Catteruccia et al., 2003; Lambrechts et al., 2008; Marrelli et al., 2006). Without the biased inheritance directed by the gene drive, it is generally considered infeasible to generate enough individuals at a high enough release frequency to spread effectors to a 100% allele frequency, otherwise known as fixation, in a population (James, 2005).

In this review, we examine the different types of gene drive systems that have been developed to date and the current state of the technology. We focus on two broad categories of drives: ‘localized drives’ and ‘non-localized drives’. Non-localized drives are expected to spread beyond a release site and maintain themselves in the population for many generations. In contrast, localized drives are expected to spread only into local populations and, in some cases, eliminate themselves from the population over time. The ability to limit the spread of localized drives makes these technologies of interest during the trial phase of the technology, when localized control is otherwise desired, or when spread into other populations would be problematic. For instance, if gene drives spread across...
Fig. 1. Schematic illustration of the various non-localized and localized gene drives. (A) Homing endonuclease drives encode an endonuclease that cuts a target sequence in a homologous chromosome. Through a DNA repair mechanism called homology-directed repair (HDR), the drive is then used as a template and is copied into the cut location. (B) Maternal-effect dominant embryonic arrest (Medea) drives encode a maternal toxin (MT) that is deposited into eggs and a zygotic antidote (ZA) to this toxin that is encoded in the early-stage offspring. When mated to a wild-type mosquito, the drive kills all offspring that do not inherit the drive and over time this fitness advantage will drive it into the population. (C) Cleavage drive consists of a germline-specific Cas9 endonuclease and a guide RNAs (gRNAs) targeting a haplosufficient essential gene (toxin) located on another chromosome. The drive also has a cleavage-resistant recoded copy of the essential gene (REG; antidote), which protects offspring that inherit the drive from being killed. (D) X-shredders are engineered to express an endonuclease on the male Y chromosome during spermatogenesis that targets and destroys genes on the X chromosome. This generates male-only progeny, which can be used for population suppression. (E) Translocation drives generate a chromosomal rearrangement or inversion that gives a disadvantage to individuals heterozygous for the drive. This system creates a genetic condition called underdominance, whereby over time the fitness advantage of translocation homozygotes facilitates their spread into a population. (F) Maternal-effect lethal underdominance (UDMEL) drives encode two constructs each possessing a maternally expressed toxin gene (MT1 and MT2), which is active in the embryo, and a zygotic antidote gene (ZA1 and ZA2), which is capable of neutralizing the maternal toxin expressed by the other toxin–antidote construct. This drive is another underdominance system whereby heterozygous females generate mostly inviable offspring, while homozygous females are fully fertile and viable. (G) Split homing or daisy endonuclease drives contain the same components as the standard homing endonuclease drive, but the homing endonuclease and programmable guide components are separated into different lines. Gene conversion only occurs when these lines are crossed to each other and with many releases and over time can be used to drive effectors into a population.
Table 1. Terminology reference for population control technologies

<table>
<thead>
<tr>
<th>Drive class</th>
<th>Type</th>
<th>Goal</th>
<th>Species</th>
<th>Design</th>
<th>Status</th>
<th>Immediate needs for further development</th>
<th>Citations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-Localized Drives</td>
<td>Medea</td>
<td>Population replacement</td>
<td>Low-threshold</td>
<td>D. melanogaster, D. suzuki</td>
<td>Maternal mRNA targeting an essential gene (toxin) linked to an embryonic miRNA-resistant copy of the essential gene</td>
<td>Laboratory proof-of-principle with multigenerational data</td>
<td>Identification of essential genes, maternal and zygotic promoters for each species of interest and improved toxin designs.</td>
</tr>
<tr>
<td>Homing Endonuclease</td>
<td>Population replacement</td>
<td>Low-threshold</td>
<td>D. melanogaster, A. gambiae, A. stephensi</td>
<td>CRISPR-based design with linkage to a beneficial effector (e.g. disrupting malaria transmission)</td>
<td>Laboratory proof-of-principle with some multigenerational data</td>
<td>Development of future effectors to link to the drives, developing next-generation drives with reduced resistance allele generation rates, larger caged trails to determine drive efficacy.</td>
<td>Gantz et al., 2015; Hammond et al., 2016</td>
</tr>
<tr>
<td>Cleaver</td>
<td>Population replacement</td>
<td>Low-threshold</td>
<td>D. melanogaster</td>
<td>Simple CRISPR-based design leveraging cleavage of target gene (toxin) while providing a cleavage-resistant copy of targeted essential gene (antidote). Homing not required for function of this system.</td>
<td>Laboratory proof-of-principle</td>
<td>Develop and demonstrate efficacy in a vector or pest species.</td>
<td>Oberhofer et al., 2019</td>
</tr>
<tr>
<td>X-shredder</td>
<td>Population replacement</td>
<td>Low-threshold</td>
<td>A. gambiae, D. melanogaster</td>
<td>CRISPR-based design, integrated on the Y chromosome, in which redundant X-linked sequences are targeted during spermatogenesis, resulting in removal of X-bearing sperm.</td>
<td>Theoretical</td>
<td>Autosomal linked X-shredders will not spread in a population. Needs to be linked to a Y-chromosome to be effective.</td>
<td>Galizi et al., 2014; Klein et al., 2012; Windbichler et al., 2007; Windbichler et al., 2008; Windbichler et al., 2007; Klein et al., 2012; Windbichler et al., 2008; Galizi et al., 2014; Fasulo et al., 2019 preprint</td>
</tr>
<tr>
<td>Translocations</td>
<td>Population replacement</td>
<td>High threshold</td>
<td>D. melanogaster</td>
<td>Endonuclease-initiated cleavage of transgenes positioned on non-homologous chromosomes, resulting in homology-directed repair and generation of marked translocations</td>
<td>Laboratory proof-of-principle with some multigenerational data</td>
<td>Develop and demonstrate efficacy in a vector or pest species.</td>
<td>Buchman et al., 2018b</td>
</tr>
<tr>
<td>Maternal-effect lethal underdominance</td>
<td>Population replacement</td>
<td>High threshold</td>
<td>D. melanogaster</td>
<td>Underdominant system consisting of inversely positioned maternal miRNAs targeting essential genes (toxins) linked to embryonic miRNA-resistant copies of the essential genes (antidote)</td>
<td>Laboratory proof-of-principle with some multigenerational data</td>
<td>Develop and demonstrate efficacy in a pest species.</td>
<td>Akbari et al., 2013</td>
</tr>
<tr>
<td>Localized Drives</td>
<td>Split homing endonuclease</td>
<td>Population replacement</td>
<td>Transient ‘self-limiting’</td>
<td>D. melanogaster, A. aegypti</td>
<td>CRISPR-based design in which the endonuclease is unlinked from the drive</td>
<td>Laboratory proof-of-principle</td>
<td>Development of next-generation drives with reduced resistance allele generation rates</td>
</tr>
<tr>
<td>Split homing endonuclease</td>
<td>Population replacement</td>
<td>Transient ‘self-limiting’</td>
<td>D. melanogaster, A. aegypti</td>
<td>Local CRISPR-based drive in which the drive components are separated into an interdependent daisy chain</td>
<td>Theoretical</td>
<td>Mathematical models demonstrating efficacy, and demonstration in a model organism</td>
<td>None to date</td>
</tr>
<tr>
<td>Daisy chain</td>
<td>Population replacement/ supression</td>
<td>Transient ‘self-limiting’</td>
<td>D. melanogaster, A. aegypti</td>
<td>Local CRISPR-based drive in which the drive components are separated into an interdependent daisy chain</td>
<td>Theoretical</td>
<td>Develop and demonstrate efficacy in a model organism.</td>
<td>Noble et al., 2019</td>
</tr>
</tbody>
</table>

miRNA, microRNA.
political borders, into ecologically sensitive areas or into areas with conflicting technologies, this may cause significant challenges for regulators and stakeholders. There are a number of varieties of both localized and non-localized drives that have unique attributes that affect their performance and safety and regulation implications.

Overview of the current state of gene drive development
Non-localized drives

Homing endonuclease drives

Homing endonuclease-based gene drives were first described over 15 years ago (Burt, 2003), but in recent years CRISPR-Cas9 genome editing has fostered a resurgence in the interest in these technologies because of their ease of use compared with other genome editing methods (e.g. zinc finger nucleases and transcription activator-like nucleases) and their adaptability to a wide range of organisms (Esvelt et al., 2014). Unsurprisingly, there are now Cas9 endonuclease-based gene drives in yeast (DiCarlo et al., 2015; Roggenkamp et al., 2018), flies (Gantz and Bier, 2015; Kandul et al., 2019 preprint; Oberhofer et al., 2018), mosquito disease vectors (Gantz et al., 2015; Hammond et al., 2016; Kryou et al., 2018; Li et al., 2019 preprint), fungi (Shapiro et al., 2018) and mice (Grunwald et al., 2019). The fundamentals of these gene drive systems have been discussed extensively in other articles (Champer et al., 2016; Esvelt et al., 2014; Gantz and Akbari, 2018; Gantz and Bier, 2016), but in brief, Cas9 is a highly specific endonuclease that can be easily programmed to introduce double-stranded breaks in desired genome regions corresponding to a complementary guide RNA (gRNA) sequence (Fig. 1A). When these breaks are repaired, they can generate insertions or deletions (indels) through a process called non-homologous end joining (NHEJ), or they can be repaired by precise integration of a copy of a corresponding region in the homologous chromosome through a process called homology-directed repair (HDR). This HDR mechanism causes transformation of offspring that would be heterozygous for the drive, by normal Mendelian inheritance, to instead become homozygous for the drive. This process, also known as gene conversion, is key to the efficient, biased inheritance of these drive systems.

The indels caused by NHEJ repair, in contrast, are a primary cause of drive resistance in Cas9-mediated homing endonuclease drives. If the NHEJ-generated indels mutate or omit the genomic sequence targeted by the drive, then the drive may no longer cut at the site. Over time, with accumulation of NHEJ repair events that alter the drive cleavage sites, coupled with any fitness advantage for resistance to the drive, the population will become resistant to the drive. The particular importance of mitigating NHEJ repair in gene drives was recently exemplified in the first example of a CRISPR-Cas9-mediated super-Mendelian inheritance in mice (Grunwald et al., 2019). This was an important initial step towards demonstrating the feasibility of gene drive in mice, but the drive was not functional. Therefore, even though Cas9-based gene drive strategies are seemingly universally applicable, with the accelerated rate of resistance seen against Cas9-mediated homing-based drives in the laboratory, much work is still needed to optimize gene drive components and target selection before this technology can be used to produce reliable drive systems, especially in organisms with limited genetic engineering tools.

One recent advancement in homing endonuclease gene drives is a system that targets highly conserved regions of a sex determination gene, *doublesex (dsx)*, to suppress the principal malaria vector in Africa, *Anopheles gambiae* (Kryou et al., 2018). The *dsx* gene is sex-specifically spliced and encodes sex-specific transcription factors that regulate sexual development in insects (Burtis and Baker, 1989; Salvemini et al., 2011; Shukla and Nagaraju, 2010). In small-population cage studies targeting the female-specific alternatively spliced transcript in *dsx*, Kryou et al. (2018) achieved >95% inheritance in males, which were fully fertile and viable, in addition to demonstrating 99% inheritance in females, which confers a sterile intersex female phenotype. Furthermore, these sterile females were unable to blood feed, which is required for egg production, thereby rendering them incapable of both reproducing and transmitting pathogens.

In the two small-population cage experiments, suppression of the wild-type population was achieved in 12 generations or fewer, but even in as little as 2–5 generations, indels at the target site developed that were resistant to the drive (Kryou et al., 2018). In this case, these resistance mutants were found at a frequency not exceeding 1.16% and did not encode a functional *dsx* transcript required for normal female development, so females were still rendered intersex and sterile. The authors attribute targeting of highly conserved and functionally constrained target sites in *dsx* to the success of this drive despite resistance, and this should be an important consideration for target site selection in future designs, but as this experiment was very small scale (two cages, *N*=300 mosquitoes) and of limited time scale (12 generations), and therefore it is unknown whether resistance mutations that disrupt drive function and restore transcript function will develop in larger populations over a longer periods of time. Moreover, resistance mutation accumulation in other *A. gambiae* gene drive systems were predicted to be incapable of long-term population suppression (Hammond et al., 2016; Marshall et al., 2017).

Medea drives

Maternal-effect dominant embryonic arrest (*Medea*) population replacement drives were first demonstrated over a decade ago (Chen et al., 2007). The design and function of these drives is reviewed extensively elsewhere (Champer et al., 2016; Ward et al., 2011), but briefly, these systems bias inheritance through the tight linkage of a maternally expressed toxin, which to date has been microRNAs (miRNAs) targeting essential genes, and a linked zygotically or maternally expressed toxin, which to date has been microRNAs (miRNAs) targeting essential genes, and a linked zygotically or maternally expressed effector that results in the death of offspring that fail to inherit the *Medea* system (Fig. 1B). Essentially, females with a *Medea* drive deposit a toxin into all their eggs that must be counteracted by an antidote that is expressed early in development, or the offspring dies. In successive generations, this system results in a disadvantage for wild-type alleles and, therefore, if the *Medea* system is linked to an effector to make the population disease resistant, for example in mosquitoes (Buchman et al., 2019a preprint, 2019b; Marshall et al., 2019), or an effector targeting a conditional essential gene, or another conditional lethal effector such as one that confers temperature or small-molecule sensitivity to kill the population, it is predicted to favorably bias inheritance of the
effect and therefore modify the population. The drive could also behave as a non-localized drive when released over a certain threshold, which is dependent on the fitness cost of the gene drive and its associated genetic elements as well as the rate of toxin resistance or natural genetic resistance in the population. The rate of drive resistance accumulation is expected to be lower with these drives compared with homing drives, as offspring that do not receive the full drive are removed from the population, but improvements in drive components are needed to move this technology forward in development. Specifically, there is a need to identify maternal and zygotic/embryonic promoters and appropriate toxin effectors/targets in pest species as the lack of these components has limited the development of these technologies (Akbari et al., 2014; Buchman et al., 2018a), but these needs are not insurmountable. The first Medea drive system in a crop pest species, Drosophila suzukii, was recently demonstrated in the laboratory (Buchman et al., 2018a). The Medea driven was successfully driven to high frequencies into laboratory populations for many generations although natural genetic variation in the laboratory population likely restricted the drive from going to fixation. CRISPR genome engineering technologies such as RNA targeting systems may provide the tools to move these technologies into other organisms (Terns, 2018).

CleaveR drives
A newly developed gene drive system has taken a novel approach to reducing NHEJ-associated drive resistance by eliminating the need for the system to be copied via HDR (Fig. 1C). This CleaveR [Cleave and Rescue (CrvR)] drive has been engineered as proof of concept in D. melanogaster, where it is inserted at a separate location in the genome from the target gene and encodes both a germline-specific Cas9 and gRNA targeting a haplosufficient essential gene (toxin). The drive also encodes a cleavage-resistant, recoded version of the haplosufficient essential gene target (antidote) with limited homology to the wild-type allele to prevent recombination (Oberhofer et al., 2019). The resulting CrvR drive is lethal to offspring that do not inherit the antidote, thereby positively biasing inheritance of the CrvR drive and any linked components. This drive system is notable as its population introgression rate is predicted to be tunable by the introduction frequency. Furthermore, the efficiency of the drive is independent of the repair mechanism and the drive does not require HDR/homing for spread. As previously discussed, the NHEJ repair mechanism results in target disruption and drive resistance, but in this system, NHEJ repair causes lethality in haplosufficient targets and therefore removes these individuals from the population. These CrvR designs may be able to circumvent NHEJ-associated drive resistance, but further multigenerational long-term drive experiments in diverse populations need to be undertaken to determine the true robustness of this system. Additionally, given the requirement for defining key promoters to express the recoded rescue in this drive, the ease of developing this drive in pest species remains to be proven. Finally, it should be noted that while this drive is generally a population replacement type of drive (similar to Medea), it could possibly be used to spread conditional lethal effectors (e.g. temperature sensitivity or sensitivity to a small molecule) and therefore could also be used to replace then suppress populations.

X chromosome shredders
X chromosome shredders (X-shredders) are another potential approach to the development of population suppression drives (Papathanos et al., 2014). These sex-biasing systems can be used in organisms with two different sex chromosomes, known as heterogametic species, where females have two of the same sex chromosomes (XX) and males have two different sex chromosomes (XY). These drives are engineered by expressing an endonuclease from the Y chromosome to target and destroy genes on the X chromosome during spermatogenesis (Fig. 1D). Only the Y-bearing sperm survive to fertilize the egg, so this strategy generates male-only progeny and therefore could be used as a potential drive strategy. In theory, an X-shredder could be designed for any heterogametic species, which would preclude disease vectors such as Aedes aegypti, but be suitable for most species of interest. Although significant attempts to engineer an X-shredder drive in A. gambiae have been undertaken, a complete system has yet to be fully developed (Galizi et al., 2014; Klein et al., 2012; Windbichler et al., 2007, 2008). These efforts have demonstrated that, while targeting the X chromosome during spermatogenesis proved possible, efforts to position the nuclease on the Y chromosome while maintaining functionality is difficult. Future efforts to exploit randomly generated Y chromosome docking strains (Bernardini et al., 2014) or even generate new site-specific docking strains on the Y chromosome using CRISPR (Buchman and Akbari, 2019) may improve functionality of Y chromosome nucleases and thereby expedite the development of the first synthetic X chromosome shredders. Recent work in D. melanogaster unraveled some of the important considerations for designing X-shredder systems, including optimization considerations for Cas9 and gRNA timing and expression, target and target site selection and drive resistance mitigation (Fasulo et al., 2019 preprint). This work also demonstrated the adaptability of X-shredder systems to new organisms as they had only been previously used in one anopheline mosquito vector. As our understanding of X-shredder systems improves, we will certainly see the development of more of these drives.

Localized gene drives
Translocations
CRISPR-Cas9 technologies have also generated useful genome engineering tools that can advance the development of other drive systems, including translocation drives (Gantz and Akbari, 2018). Translocation drives are engineered to generate a reciprocal chromosome rearrangement or inversion that is unlike the chromosomal organization of the wild population (Fig. 1E). When mated to wild-type mosquitoes, translocation heterozygotes produce a large proportion of inviable offspring, as 50% of the offspring do not inherit a balanced set of chromosomes. Homozygotes for the translocation contain a balanced set of chromosomes, so their progeny are viable and consequently have higher fitness than the heterozygotes, a genetic condition known as underdominance. The fitness advantage of homozygotes allows the translocation to spread into the population along with any effector linked to translocation breakpoints (Curtis, 1968). This translocation drive is threshold dependent, however, which means it needs to be released at a high enough frequency to spread into a population. Below that threshold, which is determined by the fitness differentials between heterozygotes and homozygotes and the local population genetics (Marshall and Akbari, 2018), the drive will go extinct in the population.

Translocation drives are confineable to the target species as viable hybridizations between an individual harboring a translocation and another species are exceedingly unlikely. They are also confineable to a local population, as they need to be introduced into a new population at a frequency exceeding its threshold to be maintained in that population, otherwise they are actively driven out. Their introduction is also reversible, as releasing large numbers of wild-type individuals can push the drive below its threshold where over
time it will become extinct in the population. This species specificity and ability to confine and reverse translocation drives make them desirable for field release. Moreover, once they spread into the target population at a high frequency, the drive can halt migration of non-drive alleles as long as these alleles remain below the threshold needed to maintain the drive. Despite these favorable attributes, previous attempts to generate translocations resulted in high fitness costs because of the low precision methods by which these translocations were generated (Asman et al., 1981; Curtis et al., 1972; Lorimer et al., 1972; Robinson, 1976). A recent study in D. melanogaster demonstrated the use of endonuclease engineering technologies to generate site-specific genome translocations that could be reliably integrated into known engineered docking sites in the genome (Buchman et al., 2018a). This is a substantial improvement on the essentially random translocation insertion methods used in the past and thus may be able to mitigate the insertional gene disruption that likely contributed to the fitness issues seen in past engineered translocations. Furthermore, unlike many of the other drives discussed in this review, this drive should be evolutionarily stable, as reversion to the wild-type chromosome arrangement should be exceedingly rare.

**UD^\text{MEL}**

A synthetic maternal-effect lethal underdominance (UD^\text{MEL}) drive is a dual toxin and dual antidote system. This system consists of two transgenic constructs each possessing a maternally expressed toxin gene, which is active in the embryo, and a zygotic antidote gene, which is capable of neutralizing the maternal toxin expressed by the other toxin–antidote construct (Fig. 1F). Similar to the translocation drive, the genetics of the system result in heterozygous females generating mostly inviable offspring, while homozygous females are fully fertile and viable. This disparity in fitness between drive heterozygotes and homozygotes results in bistable dynamics which are characteristic of underdominant systems and can be exploited to drive effectors into populations. To date, UD^\text{MEL} drive has only been built in D. melanogaster as a proof of principle, but as designed, it successfully replaced laboratory populations in a threshold-dependent manner (Akbari et al., 2013). While this system was engineered using synthetic miRNA toxins, recently described CRISPR RNA targeting systems such as C2C2 and Cas13b could help expedite development of this system in other species (Terns, 2018).

**Split homing endonuclease drives**

Split homing endonuclease drives (also referred to as daisy drives when arranged in a chain consisting of several components; Noble et al., 2019) separate the homing endonuclease and gRNA components of a homing endonuclease drive into separate lines, rendering the cleavage and homing inactive until the lines are arranged in a chain consisting of several components; Noble et al., 2019). While initially demonstrated in yeast (DiCarlo et al., 2015), split homing endonuclease drives in D. melanogaster were recently developed that had comparable gene conversion efficiencies to a standard homing endonuclease drive (Champer et al., 2019; Kandul et al., 2019 preprint). The rate of resistance allele formation was higher in the split drive, but this difference is likely due to differences in Cas9 expression between the drives. A split homing endonuclease drive engineered in A. aegypti also demonstrated high transmission rates (up to 94%), but resistance allele formation was also seen in this drive system. Mathematical modeling predicted that this drive could still be maintained at a high population frequency along with its associated cargo for up to 4 years, or 56 generations assuming 14 generations per year, and therefore may be suitable for field trials once effectors are linked (Li et al., 2019 preprint).

While this initial result focuses on split replacement drive, a split homing suppression drive has not been demonstrated yet in the lab but can be designed in multiple ways. Split suppression drives can be developed to target conserved sex-determination genes such as dxx, similar to the drive recently developed for A. gambiae (Kyrou et al., 2018). Alternatively, essential genes could be targeted in this approach (e.g. genes important for female viability, female flight, or fecundity). These results indicate that split drives are likely as adaptable as standard homing endonuclease drives but have the potential to be a safer alternative to these drives. Further research is needed in non-model organisms and in long-term, multi-generational laboratory cage trials to evaluate their efficiency and safety compared with other drives.

**Open questions regarding field performance of drive systems**

To date, gene drives have only been tested under laboratory conditions that are highly uniform and not at all reflective of field conditions. Scaled, confined field trials are desirable to generate the requisite data to better assess the efficacy and safety of these technologies prior to wider scale release. Gene drive technologies have not progressed past the initial proof-of-concept studies in the lab, but there are multiple research groups that plan to move this technology into the field, possibly in the not so distant future. These technologies have the potential to have an enormous impact on human health and livelihood, so their expedited development is essential, but not at the expense of prudent biosafety research and engagement with community members and other relevant stakeholders. From this review, we can see that there are many flavors of gene drives with differing mechanisms and spread dynamics. This suite of drive technologies may prove useful in different contexts. As these technologies advance, there are a number of open questions related to their regulation and implementation. Here, we discuss some of the technical ones.

**How do drives and their components behave over long time scales?**

Studies of gene drive systems thus far tend to be of the order of a few to a dozen generations. There are few if any long-term (>5 years) studies of any gene drives. These studies are essential to evaluate the expected duration of impact and stability of gene drive technologies. Resistant mutants have been identified in homing-based drives over short time scales (1–2 generations) (Champer et al., 2017; Gantz et al., 2015; Hammond et al., 2016; Kandul et al., 2019 preprint; KaramiNejadRunjbar et al., 2018; Kyrou et al., 2018; Li et al., 2019 preprint) and it is expected that these resistant alleles will proliferate over longer time periods and spatial scales. Presuming that resistance allele generation can be significantly reduced, perhaps through some combination of multiplexing of guide RNAs (Marshall et al., 2017) and targeting highly conserved genes (Kyrou et al., 2018), will this be sufficient to interrupt disease transmission in real populations? For homing-based population suppression, even very rare resistance allele generation is expected to interfere with vector population control (Marshall et al., 2017). Larger scale, longer-term studies will be required to determine the resistance allele generation rate and their long-term impact on population suppression strategies. For homing-based population replacement, higher resistance allele generation rates can be tolerated (Noble et al., 2017); but an additional concern is the stability of the drive system and effector gene cargo over time.
(Marshall et al., 2019). If some time elapses between the system reaching fixation in one population and an individual migrating from that population to another, then will the gene drive system and effector still be functional and seed a new wave of spread? What happens if the linked effector forces selection for pathogens that are more virulent (Marshall et al., 2019)? Given these issues and concerns, longer-term studies will be required to determine this as well.

How do drives perform in more genetically diverse populations? Recent laboratory experiments have demonstrated that the genetic strain that a drive is present in can have a significant impact on drive efficacy (Champer et al., 2017; Hammond et al., 2016; Li et al., 2019 preprint). Populations with higher genetic diversity will have a higher probability of containing resistant drive alleles that could block the spread of the drive. Given that the genetic diversity in the wild cannot be entirely captured in a laboratory experiment, it is expected that genetic diversity will have an even more significant impact on the result of gene drive trials and interventions in wild populations. Substantial ecological work and laboratory studies will be required to refine our understanding of the behavior of drives in diverse populations and to ensure that any gene drive released in the wild will be efficacious over the desired treatment area.

The impact of genetic diversity is also a concern for other gene drive systems. A recent study of Medea drive in D. suzuki, a global crop pest, demonstrated that natural genetic variation in D. suzuki populations likely resulted in selection of toxin-resistant variants, which in multi-generational population cage studies resulted in limited drive spread (Buchman et al., 2018a). Similar genetic variation and toxin targeting issues will likely also arise when adapting these technologies to other species, especially as the genetics and population structure of most species are less defined than in D. melanogaster. Potential solutions to these issues include targeting of conserved gene regions, validation of target sites to optimize targeting efficiency and increased multiplexing, but substantial efforts are still needed to identify the minimal requisite components of these Medea systems in other species, including maternal and zygotic/embryonic promoters and appropriate toxin effectors and targets.

Next steps for gene drive technology
For many important vectors, we have developed the genetic tools needed to effectively produce proof-of-principle genetic effectors and gene drive systems for population replacement or suppression. At the current state of the technology, some gene drive technologies seem very adaptable to new species and populations (e.g. homing endonuclease-based drives) while others still need additional genetic innovations before they can be easily transferred to other species of interest (e.g. Medea and other toxin-antidote-based systems).

In many regions of the world, the benefits of current vector-borne disease interventions have plateaued while disease transmission persists. New technologies are needed to control these vectors and eliminate the diseases they transmit. Modeling has demonstrated that gene drive technologies, if they reach certain design requirements, are capable of generating long-term, sustainable vector-borne disease control (Eckhoff et al., 2017); however, drive resistance, genetic diversity in the target populations and other concerns pose challenges to achieving the level of control required to achieve disease elimination. Potential solutions for homing endonuclease-based drives include multiplexing of guide RNAs and targeting highly conserved genes. Work is also required to determine the target profile of each gene drive to meet specific performance and safety needs (Carbalhar-Lejarzá and James, 2017; James et al., 2018) and also to update guidance documents related to research, development and testing of genetically engineered insects and arthropods (Adelman et al., 2017). These profiles and guidance documents may differ between field trials and wide-scale disease control programs and will help to determine the goals that molecular technologies must meet.

Competing interests
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