Resistance to natural and synthetic gene drive systems

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Abstract

Scientists are rapidly developing synthetic gene drive elements intended for release into natural populations. These are intended to control or eradicate disease vectors and pests, or to spread useful traits through wild populations for disease control or conservation purposes. However, a crucial problem for gene drives is the evolution of resistance against them, preventing their spread. Understanding the mechanisms by which populations might evolve resistance is essential for engineering effective gene drive systems. This review summarizes our current knowledge of drive resistance in both natural and synthetic gene drives. We explore how insights from naturally
1 | INTRODUCTION

Organisms require networks of cooperating genes. Generally, alleles spread through populations by increasing the reproductive success of the organism as a whole. However, some alleles, defined here as drivers, selfishly bias reproduction to increase their own representation in the next generation, at a cost to the rest of the genome (Burt & Trivers, 2006). For example, "segregation distorters" are a type of driver that subvert the usual rules of Mendelian inheritance in such a way that they are inherited by over 50% of the descendants of heterozygous individuals, and occur naturally in many species including plants, fungi, nematodes, insects and mice (Lindholm et al., 2016). Another example is drive by mitochondria, the key endosymbiont of eukaryotes, which damage male function in many hermaphroditic plants (Burt & Trivers, 2006). This loss of male function diverts resources to seed production, enhancing transmission of the mitochondrial genome, which is typically uniparentally transmitted through ovules but not pollen. Selfish genetic elements likely occur in all species and can have major impacts on the evolution and ecology of their hosts (Burt & Trivers, 2006).

Crucially, the super-Mendelian rate at which gene drivers are transmitted over generations can allow them to spread through populations despite costs. This has inspired researchers to propose using gene drives to solve major biological challenges related to public health, the environment and agriculture (Burt, 2014; Champer, Buchman, & Akbari, 2016; Piaggio et al., 2017; Raban et al., 2020). Two broad types of gene drives have been proposed: population suppression gene drives and population replacement gene drives. Population suppression gene drives can be employed when reduction or elimination of a population is desired. Replacement gene drives offer the potential to alter existing populations for human benefit, for example by spreading alleles or endosymbionts that reduce the ability of mosquitoes to transmit malaria. Strains of the intracellular bacterium Wolbachia reduce the ability of mosquitoes to transmit dengue and other viruses. Wolbachia strains have already been successfully deployed in Australia and elsewhere, spreading through populations by creating mating incompatibilities that disproportionately reduce the fitness of females that do not carry Wolbachia, and reducing the threat of dengue (Nazni et al., 2019; Ryan et al., 2020). New synthetic population suppression and replacement drive systems are being created with increasing regularity, highlighting the enormous promise of CRISPR-Cas9 and other new molecular tools for editing genomes (Champer et al., 2016).

However, gene drives impose costs, certainly on outcompeted alleles, and often on the individual as a whole. Costs at the individual level can arise directly via the mechanism of transmission, for example the costly death of gametes that carry rival alleles, or because the driver carries costs such as associated low fitness alleles or metabolic costs in driving endosymbionts (Burt & Trivers, 2006). The resulting selection can lead to the rapid evolution of resistance traits that prevent the driver from spreading. As a result, many natural drivers have been completely suppressed, only showing drive when crossed into distant relatives that do not carry suppressor alleles (Courret, Chang, Wei, Montchamp-Moreau, & Larracuente, 2019; McDermott & Noor, 2010). This research suggests that we should expect synthetic gene drives, especially those with large fitness effects, to select for resistance, which will potentially undermine their ability to spread, and modify or suppress populations (Barrett et al., 2019; Holman, 2019; Unckless et al., 2017).

For synthetic gene drives to be effectively deployed, we urgently need to understand how quickly resistance will arise. Does resistance usually arise through selection on pre-existing genetic variation, or does it more often involve novel mutations that appear once drive has reached a high frequency? What fraction of natural gene drives reach fixation, go extinct, reach a stable polymorphism or are fully suppressed, and how can we address this question given the difficulties of detection once a gene drive has fixed or been lost? Does resistance to drive typically involve the same fundamental mechanism (e.g. loss of the driver’s target, or "defusing" of the driver by interfering RNAs) across species and types of drivers, or is the resistance mechanism highly idiosyncratic?

In this review, we synthesize what is known about how resistance evolves against both natural and synthetic drives, and point out gaps in our knowledge. We begin by reviewing how resistance has evolved in well-studied natural systems, examining resistance that interferes directly with the molecular mechanisms of drive and then resistance through behaviour and life history. We then turn to the current evidence regarding resistance to synthetic drives. Finally, we discuss the implications for the design of “evolution proof” synthetic gene drives.

2 | RESISTANCE TO GENE DRIVES IN NATURAL SYSTEMS

In any drive system, selection for resistance will act on the target locus itself, genes linked to the target and in some cases on the
The Evolution of **Drive Resistance**

**FIGURE 1** The evolutionary impact of a gene drive, as measured by the magnitude and location of costs imposed (yellow/red gradients). Boxes represent individuals; white rectangles are chromosomes within the organism. Drive creates selection pressure for the three drive resistance mechanisms discussed in this review (blue). The selection pressure for drive resistance is highest at the target locus itself (1a), where rivalling homologous genes suffer both from reduced transmission due to drive (yellow) and (potential) fitness costs to the organism (red). Selection pressure on unlinked loci throughout the genome to disrupt drive will be a function of organismal drive costs (1b). Finally, gene drive may create selection for mechanisms that suppress the drive at the population level (2) [Colour figure can be viewed at wileyonlinelibrary.com]
entire genome. Generally speaking, selection for resistance at the target and linked loci becomes stronger with more biased transmission, whereas the strength of selection for resistance on the rest of the genome increases with higher fitness loss for the organism (Figure 1). These two are often positively related, leading to strong selection for resistance at both the target locus and genome-wide.

We classify resistance as adaptations that reduce the spread of drive elements either by (a) interfering with the molecular mechanism of drive (which we term as “suppression” in this Review) or by (b) altering some aspect of behaviour or life history of carriers which in turn reduces the ability of a driver to spread. We use these categories to structure our review of known drive resistance factors, incorporating natural and synthetic drive systems.

2.1 | Mutations at the target site and suppression of drive machinery

One way to evolve resistance is to modify the target of drive so that it is no longer susceptible. For example, a gene drive that spreads itself by targeting a specific sequence of nucleotides or peptides might impose selection that favours genotypes carrying an altered sequence. Below, we review the evidence for this mode of resistance in nature. For a brief overview of the biological differences between the natural gene drives, see Table 1.

### TABLE 1

| Gene drive system | Mechanism | Key effects |
|-------------------|-----------|-------------|
| Gamete killer     | Drives by killing or damaging gametes that do not carry the driving chromosome. | Reduces sperm number. If on a sex chromosome, can bias population sex ratios. |
| Female meiotic drive | Drive chromosome manipulates meiosis so rival chromosomes are disproportionately discarded in the polar bodies. | Costs relatively unknown, but some well-studied systems associated with low fitness. |
| Transposable elements | Drive sequences copy themselves into other locations in the genome. | Largely deleterious due to gene disruption and DNA breakage. |
| Genetic incompatibility systems | Factors inherited via cytoplasm such as organelles and endosymbionts increase the fitness of females at a cost to males. Mechanisms are extremely diverse. | Effects can include loss of male function, feminization, death of offspring. Can be highly costly. |
| Homing-based systems | Induce targeted double-strand DNA breaks that copy and insert the drive construct during DNA repair. | Effects depend on design. Can include sterility, offspring sex ratio bias, disease resistance. |
| Medea-like systems | Chromosomes bearing a set of lethal loci in which each suppresses the other, killing offspring that do not inherit the system. | Reduced viability if not all loci are inherited. Reduced offspring production. |

2.1.1 | Sex chromosome linked gamete killers

Naturally occurring “gamete killer” meiotic drivers have often been found on sex chromosomes, where they cause distortion in the transmission of the heterogametic sex (Hurst & Pomiankowski, 1991). The evolution of sex chromosome drivers is facilitated by the differentiation between X and Y chromosomes (and Z/W). Driver alleles arising on a well-differentiated sex chromosome therefore have potential targets at many sites that are never linked to the driver: for example, an X-linked driver could promote its own transmission by destroying gametes containing a particular Y-linked locus (Jaenike, 2001). Sex-linked drivers generate especially strong selection for resistance because they alter the population sex ratio. A bias in the population sex ratio creates strong selection favouring individuals/genotypes that produce relatively more of the under-represented sex (Fisher, 1930). This “Fisherian sex ratio selection” confers an additional fitness benefit to alleles that confer resistance to drive in populations showing a biased sex ratio, due to the presence of a sex-linked driver. We therefore expect to see rapid evolution of resistance against sex-linked drivers (Hurst & Pomiankowski, 1991).

We illustrate this using sex chromosome drive systems in *Drosophila simulans*. In the Paris Sex Ratio (SR) system, two X-linked drivers together prevent the disjunction of the Y sister chromatids in the second meiotic division. One of these encodes HP1D2, a
A protein that binds Y chromosome heterochromatin in premeiotic cells, suggesting it targets repeated DNA sequences (Helleu et al., 2016). The Y chromosome of D. simulans exhibits substantial variation in resistance to Paris SR drive, with a wide continuum of phenotypes from high susceptibility (95% female progeny) to complete resistance (50% female progeny; Montchamp-Moreau et al., 2001). These more or less resistant Y chromosomes show extensive structural rearrangements affecting satellite sequences, which strongly suggests that resistance occurs through changes in target repeat sequences (Helleu et al., 2019). In addition, Paris SR is suppressed by yet unidentified autosomal loci (Courret et al., 2018).

The Winters SR is another sex ratio-distorting system in D. simulans, with a drive phenotype different to Paris SR, killing sperm after meiosis (Tao, Araripe, et al., 2007). An X-linked gene, Dox, and likely its progenitor Mdox, are involved in drive (Tao, Araripe, et al., 2007). Winters SR is typically entirely suppressed by high frequencies of the autosomal suppressor locus Nmy. Nmy arose from a retrotransposed inverted repeat of Dox (Tao et al., 2007) and produces an antisense RNA that represses Dox and Mdox through the RNA interference pathway (Lin et al., 2018).

Geographical variation in frequencies of drivers and suppressors has been found in both Paris and Winters systems, including populations that appear completely suppressed (Atlan, Mérçot, Landre, & Montchamp-Moreau, 1997; Bastide, Gérard, Ogereau, Cazemajor, & Montchamp-Moreau, 2013; Kingan, Garrigan, & Hartl, 2010). Under complete suppression, drivers have no transmission advantage but might still impose organismal fitness costs leading to a resultant decline in driver frequency. The Nmy suppressor of Winters SR is dominant and almost at fixation, rendering the driver effectively cryptic, leading to the accumulation of loss-of-function driver mutants (Kingham et al., 2010). Both systems show genetic evidence of positive selection on the drivers, in the Winters systems about 3,000 years ago (Kingham et al., 2010) and in the Paris system within the last century (Bastide et al., 2013; Derome, Baudry, Ogereau, Veuille, & Montchamp-Moreau, 2008). Paris SR, as well as its genetic suppressors, have been recently increasing in frequency in the Middle East (Bastide et al., 2013), and rapid declines in SR frequency have been observed in completely suppressed populations (Atlan et al., 1997; Bastide et al., 2011).

These two SR systems illustrate empirically the dynamic nature of the spread of drivers, followed by the rise of suppressors and then loss of drivers that evolve in a continuous cycle of “red queen” dynamics. However, although many meiotic drive systems we observe in nature have arrived at such a dynamic equilibrium, others have not. There is some evidence that drive can cause extinction, at least in local populations (Pinzone & Dyer, 2013). Other drive systems seem to occur at stable frequencies in different populations, sometimes in geographical clines, for reasons that are not well understood, and there is some evidence that this stability can last for hundreds of generations (Price et al., 2014, 2019).

2.1.2 | Autosomal gamete killers

Autosomal gamete-killing meiotic drivers function by killing gametes that carry alternative alleles (Bravo Núñez, Nuckolls, & Zanders, 2018). Some of the best-studied systems are the spore-killers in various fungal species. First, in Neurospora, an RNA interference-based genome defence mechanism has been shown to be a suppressor of spore-killing alleles (Svedberg et al., 2020). Secondly, there are multiple copies of drivers in the filamentous fungus Podospora anserina, one of which is a known suppressor (Grognet et al., 2014). Likewise, the wtf gene family in Schizosaccharomyces pombe encodes a poison-antidote drive system and has up to 42 copies throughout fission yeast genomes (Eickbush, Young, & Zanders, 2019; López Hernández & Zanders, 2018). Of these copies, some are intact meiotic drivers (alleles that encode both a poison and antidote), some are apparent pseudo-genes, and some are alleles that encode only the antidote (Bravo Núñez et al., 2020). These “antidote-only” wfts act as suppressors of their selfish wtf homologs and are likely to be maintained by selection for resistance to the latter (Bravo Núñez, Lange, & Zanders, 2018). The amplification of the different multi-copy spore-killer systems in fungal genomes is representative of the cycles of amplification of drivers and suppressors often seen in genetic conflict. Patterns of duplication of drivers and suppressors have also been observed on sex chromosomes in Mus musculus house mice (Soh et al., 2014).

Another well-studied system is Segregation Distorter (SD) in Drosophila melanogaster, which contains a driver, enhancers of drive and a target site, found in a region of low recombination (Larracuente & Presgraves, 2012). Males heterozygous for SD and a sensitive wild-type chromosome suffer chromatin condensation defects and dysfunction in wild-type sperm. The target site consists of a large block of tandem repeats. The number of copies of the tandem repeat correlates with sensitivity to drive, and alleles with fewer than 300 repeats are insensitive to drive (Wu et al., 1988). There is substantial variation in target copy number in D. melanogaster populations across the globe. Frequencies of SD are low in natural populations, suggesting a balanced polymorphism, but evidence for genetic sweeps of SD instead suggests rapid turnover of SD chromosomes, either because of competition between SD variants or arms races with suppressors (Brand, Larracuente, & Presgraves, 2015). Unlinked genetic suppressors are known (Hiraizumi & Thomas, 1984), but they have not been studied at the molecular genetic level.

2.1.3 | Female meiotic drive

Female meiotic drive exploits asymmetry in female meiosis to influence which homolog of the chromosome pair is distributed to the egg nucleus as opposed to the excluded polar bodies. Thus, the fitness of the nondriving homolog is reduced, but costs to the organism are small in terms of gamete production. If costs are negligible, then female drivers might readily spread and fix, since only a small region of the genome close to the drive locus would be under selection to
evolve resistance. However, in *Mimulus* monkeyflowers, female drivers impose fitness costs when homozygous (Fishman & Kelly, 2015). In maize (*Zea mays*), the *Kindr* (Ab10) driving knobs system has heterozygous and homozygous fitness costs in seed set and weight (Higgins et al., 2018). Resistant alleles block expression of the *Kindr* complex and are characterized by small interfering RNAs and DNA methylation (Dawe et al., 2018).

### 2.1.4 Transposable elements

Transposable elements (TEs) are DNA sequences that can change their location within a genome, often copying themselves in the process (Feschotte & Pritham, 2007). They have been found in prokaryotes, eukaryotes and even giant viruses (Sun et al., 2015). Transposition is generally deleterious to the individual, resulting in DNA breakage and potentially ectopic recombination, as well as potentially disrupting genes (Feschotte & Pritham, 2007). Mechanisms for suppressing TEs are diverse, and many have ancient origins, such as genome methylation which silences TE expression.

Typically TE invasions follow a cycle, with a novel TE invading a species, or a TE already in the genome escaping suppression (Bousios & Gaut, 2016). The TE rapidly replicates in the genome of the species, imposing costs, which select for suppression. This invasion and suppression can occur extremely quickly. In *Drosophila melanogaster*, a DNA-based TE invaded in the early 1950s and had spread worldwide by the 1980s (Anxolabéhère, Kidwell, & Periquet, 1988). In around the year 2000, this TE jumped to the closely related *D. simulans* and spread even faster worldwide through that species (Hill, Schlötterer, & Betancourt, 2016). RNAi suppression of the TE evolved extremely rapidly in both species, resulting in the TE being largely suppressed in *D. simulans* populations within two decades of invasion. This fast evolution of suppression is facilitated by piRNA clusters in animals that appear to perform a defensive function against TEs (Czech et al., 2018), similar to the CRISPR libraries that provide adaptive immune defence against viruses and plasmid gene drivers in bacteria (Barrangou & Marraffini, 2014). When a TE attacks the organism, sequences from the invading TE are recruited to the piRNA clusters, providing a DNA template that guides RNAi silencing of that TE, preventing it from further replication (Brennecke et al., 2007). The maintenance of these genomic regions as defences against TEs suggests it is possible that other genomic regions may also be maintained over evolutionary time because they defend against TEs or other selfish genetic elements.

### 2.1.5 Genetic incompatibility systems

Cytoplasmic incompatibility can occur between nuclear and mitochondrial DNA (mtDNA), as mtDNA is transmitted almost exclusively from mother to offspring. The most widely recorded example of cytoplasmic incompatibility is cytoplasmic male sterility, in which hermaphroditic plants are rendered male-sterile and are functionally female. Cytoplasmic male sterility is very widely distributed among angiosperm plant species, with populations consisting of both hermaphroditic and female plants (Touzet & Budar, 2004). Nuclear suppressors that restore male fertility (called *R* genes) are commonly found within cytoplasmic male sterility systems. Many *R* genes are members of the pentatricopeptide repeat protein family, involved in processing and editing RNA (Gaborieau, Brown, & Mireau, 2016). They typically act by binding directly to the mitochondrial transcripts, interfering with the production of male sterility proteins (Chen & Liu, 2014). *R* genes show evidence of rapid evolution and diversification (Fujii, Bond, & Small, 2011) suggesting ongoing cycles of conflict with cytoplasmic male sterility genes.

Male-killing caused by some *Wolbachia* bacteria, also inherited via cytoplasm, provides a demonstration of how quickly suppression can spread. Pacific island populations of the butterfly *Hypolimnas bolina* are infected with a *Wolbachia* strain that causes the death of the sons of infected females (Dyson, Kamath, & Hurst, 2002). This benefits infected daughters due to decreased larval competition with siblings, allowing *Wolbachia* to reach extremely high frequencies, resulting in populations with fewer than one male per hundred females (Dyson & Hurst, 2004). A nuclear gene which rescues the male embryos recently appeared and has spread rapidly; in the Samoan *Hypolimnas* population, an equal population sex ratio was restored over the course of 8–10 generations (a single year) after resistance reached the island (Charlat et al., 2007; Hornett et al., 2014).

In another example, feminizing *Wolbachia* in the woodlouse *Armadillidium vulgare* often reach very high frequencies within populations, such that the only males present come from eggs that by chance do not inherit sufficient *Wolbachia* to convert them into females (Leclercq et al., 2016). In these highly female-biased populations, the normal ZW sex determination system is defunct, with *Wolbachia* dose effectively controlling the sex of individuals. This can lead to the loss of the female-determining *W* chromosome; all individuals are ZZ genetic males, but this state is overwritten by the feminization imposed by *Wolbachia*, suggesting that the Z and autosomes have been unable to evolve resistance. In some populations, a novel *W* sex chromosome has appeared; remarkably, this neo-*W* chromosome is a former autosome that now carries a near-complete copy of the *Wolbachia* genome. This neo-*W* is thought to have spread through outcompeting *Wolbachia* feminized ZZ individuals (Cordaux & Gilbert, 2017). Given the likely cost of incorporating a bacterial genome, this illustrates that suppression of gene drives can involve high costs and major genomic rearrangements. Despite these examples of suppression in *Wolbachia*, suppressors against driving organelles and endosymbionts in animals seem to be rare. It is not clear why this is the case, given that cytoplasmic male sterility systems in plants are often suppressed.

### 2.1.6 Systems where suppression has not been found

Although mutations have allowed resistance to evolve in many systems, there are examples of both sex-linked and autosomal
drivers for which little or no suppression has been found. For example, in the well-studied \( t \) haplotype of house mice, distorter loci are bound together in inversions and cause dysregulation of development in sperm carrying the wild-type target allele (Herrmann & Bauer, 2012; Lindholm et al., 2019). Suppression of the \( t \) haplotype has not been found in wild populations (Ardlie & Silver, 1996), although transmission differences have been reported in crosses between laboratory strains (Bennett, Alton, & Artzt, 1983; Gummere, McCormick, & Bennett, 1986). In one closely monitored study population, the \( t \) haplotype declined and went extinct within eight years, which is thought to be due to negative density-dependent effects on fitness (Manser et al., 2011) and positive density-dependent effects on dispersal (Runge & Lindholm, 2018), rather than suppressors of \( t \). The combination of strong distortion and lack of evidence of suppression has led to plans to develop a synthetic sex chromosome driver from the \( t \) haplotype by adding a male sex-determining gene (Sry) to the \( t \), for the purpose of controlling invasive house mouse populations on islands (Backus & Gross, 2016; Campbell et al., 2019).

Similarly, the sex ratio-distorting X chromosome drive system in \textit{Drosophila pseudoobscura} has been studied for almost a century, yet no evidence has been found of target-site variation leading to suppression, or indeed any factors that reduce drive strength (Price et al., 2019). This is puzzling given that \( SR \) reaches 30% frequency in populations in the south-western United States, imposes significant costs on the males that carry it and has apparently existed for hundreds of thousands of years (Kovacevic & Schaeffer, 2000), providing ample time for the evolution of resistance. In the related species \textit{D. subobscura}, only an extremely weak suppressor of drive has been found, again despite a high frequency of drive in natural populations and substantial costs of drive (Verspoor et al., 2018). The same lack of suppressors occurs in \textit{Teleopsis dalmanni} stalk-eyed flies which again have a high frequency \( SR \) drive system which imposes significant viability costs in males and females (Finnegan et al., 2019) and is estimated to be a million years old (Reinhardt et al., 2014). The hybridizing species \textit{D. testacea} and \textit{D. neotestacea} each bear driving X chromosomes, but the former shows strong autosomal suppression (Keais, Lu, & Perlman, 2020), whereas the latter shows no evidence of suppression at all (Pinzone & Dyer, 2013). Surprisingly, in the known \textit{Drosophila} species with \( SR \) gamete-killing drive systems, only about half have evidence of genetic suppression of drive (Courret et al., 2019; Price et al., 2019), even though many of these systems are thought to have existed for tens if not hundreds of thousands of generations (Price et al., 2019).

The existence of multiple well-studied, ancient drive systems showing no evidence of suppression of drive is a mystery. Does this indicate that genetic suppression is unlikely to evolve? One possibility is that the locus that confers susceptibility to drive is small, providing a small mutational target. However, many drivers impose broad costs across the genome (Dyer & Hall, 2019; Finnegan et al., 2019; Hamilton, 1967; Larner, Price, Holman, & Wedell, 2019; Zanders & Unckless, 2019), so loci throughout the genome are predicted to evolve to resist costly gene drives. Here, the lack of resistance mechanisms cannot be due to the small size of the mutational target, suggesting the involvement of other evolutionary constraints. Perhaps effective resistance to the gene drive requires multiple mutations that are not individually beneficial, making resistance evolution less likely. Drive could also target essential sites in the genome that are constrained from evolving, or repetitive DNA that is continually re-created by mutation or transposition, as is thought to be the case for the satellite locus \( Rsp \) that is targeted by the \( SD \) gene drive in \textit{D. melanogaster} (Courret et al., 2019). Another possibility is that some gene drives are involved in ongoing coevolutionary arms races with resistance loci, such that the supposedly unresistable gene drives that we observe are those that have temporarily outpaced their suppressors for a short span of evolutionary time. The \textit{Hypolimnas} example appears to provide an example of this: the costs of \textit{Wolbachia} sex ratio distortion were high and \textit{Wolbachia} was very common, yet for at least a century there was no sign of resistance to the drive. When a resistance allele appeared, it rapidly spread across the species’ range within a few decades (Hornett et al., 2014).

### 2.2 Behavioural and life-history resistance against drive

One explanation for lack of direct suppression of the mechanism of drive is the evolution of indirect resistance involving behavioural or life-history changes. For example, self-medication in which a \textit{Wolbachia}-infected individual might reduce their titre by exposing themselves to heat that impairs \textit{Wolbachia}, or feeding on an antibiotic rich diet (Abbott, 2014; Shikano, 2017; Snook et al., 2000) is a possible but untested idea. There may be many unexplored life-history or behavioural ways to resist drive.

One of the best-known ideas is that noncarriers may avoid drive carriers as mates, preventing offspring from inheriting harmful drivers and improving offspring fitness. Theoretical models support this idea (Lande & Wilkinson, 1999; Manser et al., 2017; Randerson et al., 2000; Reinhold et al., 1999). However, this requires a trait that reliably reveals the presence or absence of drive (Lande & Wilkinson, 1999; Manser et al., 2017). Evidence of mate avoidance of drive carriers is weak or absent from the majority of systems studied. For example, in species where \textit{Wolbachia} induces cytoplasmic incompatibility, uninfected females lose any eggs fertilized by \textit{Wolbachia}-infected males. Despite these costs, there is very little evidence for females avoiding mating with \textit{Wolbachia}-infected males (Champion de Crespigny & Wedell, 2007). Likewise, the only test of populations infected with male-killing \textit{Wolbachia} in \textit{D. inuiliba} also found no evidence that males prefer to mate with uninfected females (Sullivan & Jaenike, 2006). There is also little evidence for female preference against male \textit{Drosophila} carrying \( SR \) drive despite decades of research in several species (Price & Wedell, 2008). In house mice, wild-type females do not avoid mating with \( t \) haplotype-bearing males (Lenington & Coopersmith, 1992; Manser et al., 2015; Sutter & Lindholm, 2016), whereas \( t \)-bearing females have been found to
avoid t-bearing males in some (Lenington & Coopersmith, 1992) but not all studies (Manser et al., 2015; Sutter & Lindholm, 2016).

In these gene drive systems, it is not obvious that any phenotypic characters reliably signal Wolbachia or drive carrier status, which may explain the lack of mate preference. In stalk-eyed flies, female preference is for males with large eyespan, and males carrying SR have a smaller average eyespan (Cotton, Földvári, Cotton, & Pomiankowski, 2014; Johns, Wolfenbarger, & Wilkinson, 2005; Wilkinson et al., 1998), providing a ready-made trait that can distinguish drive from nondrive-carrying males. However, other species of stalk-eyed fly that lack meiotic drive also show female mate preference for exaggerated male eyespan (e.g. Diasemopsis meigenii; Cotton, Rogers, Small, Pomiankowski, & Fowler, 2006), and it has yet to be demonstrated whether mate preference has been strengthened for avoidance of drive carriers. Disentangling general condition-dependent mate preferences from evolved resistance to drive through avoidance of mating with drive carriers can be highly challenging.

In the Winters SR system of D. simulans, the strength of drive declines from 93% to 60% daughters when males are reared at high temperatures, and older males also show a decline in drive (Tao, Masly, et al., 2007). This could promote females evolving a preference for males unlikely to have strong drive due to these nongenetic causes (i.e. high temperature reared or older males), but to date, this has not been examined, although age-based mate choice is common in Drosophila and other organisms (Verspoor et al., 2015). Furthermore, D. simulans females disproportionately discard sperm of Paris SR males after mating (Angelard, Montchamp-Moreau, & Joly, 2008). It is not known whether this post-copulatory selection has evolved due to the benefits of reducing the likelihood of drive bearing offspring, or is a general mechanism selecting against mates that transfer small ejaculates. Perhaps the most convincing evidence for mate choice against drivers comes from feminizing Wolbachia in Armadillidium vulgare. In populations with high Wolbachia frequency, males are rare, and males will benefit from mating with uninfected females who produce more sons. In this case, males have been found to preferentially mate with ZW-uninfected females, rather than genetically male ZZ individuals who have been feminized (Moreau et al., 2001). Whether this has suppressed Wolbachia frequency in populations has not been established. In general, the lack of choice against drive carriers may be due to evolutionary pressure to reduce detectability, with the least detectable gene drive alleles outcompeting ral variants, but this remains to be investigated.

Another route for drive-susceptible females to avoid producing offspring with drive carriers is by increasing the intensity of sperm competition. In several systems of gamete-killing male meiotic drive, drive-carrying males are inferior sperm competitors, because of a reduction in sperm number and quality (Price & Wedell, 2008). For example, in controlled experimental matings, t-carrying males gain only 12% of paternity when a female mates with both a t-carrying and wild-type male (Sutter & Lindholm, 2015). Females could therefore mate with several males indiscriminately and rely on sperm competition to suppress fertilization by drive sperm (Haig & Bergstrom, 1995). An increase in the propensity to mate with multiple males could evolve as a form of resistance to the presence of a driver within the population. Multiple mating potentially evolves more easily than precopulatory mate choice, as no discrimination between driver-carrier and driver-free individuals is required (Haig & Bergstrom, 1995). The evolution of higher remating rates in response to the presence of a sex ratio distorer was seen within 10 generations in a laboratory experiment using D. pseudoobscura (Price et al., 2008). So it is possible that in polyandrous species, sperm competition reduces the success of gamete killers enough that selection for direct genetic suppression is reduced. As yet, there is no concrete evidence for this in nature.

3 | RESISTANCE IN SYNTHETIC SYSTEMS

Synthetic gene drive systems provide some of the most informative studies of the evolution of resistance through sequence changes at target sites. They allow observation of the process of resistance evolution. Sequencing can reveal standing variation and mutations that confer resistance, identify the original resistant individuals and provide empirical evidence of the speed with which resistance arises in the laboratory and, possibly in the future, under field conditions.

3.1 | Mutations at the target site and suppression of drive machinery

3.1.1 | Homing-based drive systems

Many newly engineered systems are based on homing drives that mimic the mode of propagation of homing endonuclease genes (HEGs), a class of naturally occurring selfish genetic elements found in bacteria, fungi and other organisms (Burt & Trivers, 2006). These encode DNA-cleaving enzymes that generate double-strand DNA breaks at target sites with a specific nucleotide sequence as a result of which the sequence is converted into a HEG + allele. Homing potentially converts all target sites in all members of a population. The challenge in generating synthetic drive systems based on HEG proteins (Windbichler et al., 2011) has been alleviated by the production of homing systems based on CRISPR-Cas9 constructs.

Unlike natural homing endonucleases, which tolerate some variation in the fidelity of individual nucleotides in the target sequence, CRISPR-Cas9 cleavage requires a near-perfect match between the ~20 base pair guide RNA sequence and the genomic target site. Hence, single nucleotide differences at target sites can confer resistance. These arise at a high rate, simply through the action of the gene drive. The double-strand breaks induced by Cas9 are typically resolved by homology-directed repair, inserting a copy of the homing agent sequence into the target site. Alternatively, double-strand breaks undergo non-homologous end joining. This process often results in imprecise repair, increasing the rate of mutation at the target site without
insertion of the gene drive. These novel alleles will confer resistance as they have a different sequence, and may preserve gene function. In laboratory experiments with flies and mosquitoes, resistance to CRISPR-Cas9 homing drives emerges rapidly, in particular when the driver targets single sites (Champer et al., 2017; Gantz et al., 2015; Hammond et al., 2017, ; KaramiNejadRanjbar et al., 2018). Functional target gene mutants can be generated at considerable frequency within one generation by in-frame indels (KaramiNejadRanjbar et al., 2018). One approach to delay the evolution of resistance at the target site is to design targets at highly conserved regions in which sequence variation, including in-frame indels, cannot be tolerated because any change is associated with high fitness costs (Kyro et al., 2018). Alternatively, a suite of sites can be targeted by the drive construct.

When the aim is gene replacement rather than population suppression, gene drives are designed to have low fitness costs and avoid disruption of normal host gene function. This should constrain selection for resistance alleles. But the "cargo" of replacement genes is unlikely to be cost-free. Examples of cargoes include genes that encode resistance or susceptibility to disease or toxins, and genes that alter sexual phenotype. All of these will carry costs, and in the long term, they are expected to be lost due to the spread of loss-of-function mutations. When loss of function is caused by deletion, this may even enhance gene drive spread (i.e. of a null allele); replacement gene drives are only useful as long as the cargo remains intact. The assumption is that the replacement gene will spread and persist sufficiently long to provide its public health benefit (Beaghton et al., 2017). Other types of cargo may be more resilient to loss, for example, where the cargo is beneficial to the organism, such as thermal tolerance genes or symbionts (Piaggio et al., 2017). Finally, expression of the endonuclease is unlikely to be without fitness cost and thus subject to mutational decay. This will mostly come to play at the point when the drive construct has already successfully propagated itself in a population. These constraints have hardly been investigated, but seem likely to place limits on the spread and effectiveness of homing gene drives.

3.1.2 | Synthetic sex ratio distorsers

The X chromosome is the target in engineered systems that aim at distorting the sex ratio towards males. One approach, inspired by the mode of action of natural sex distorsors in the mosquitoes *Aedes aegypti* and *Culex pipiens* (Wood & Newton, 1991), operates by targeting the X-linked rDNA cluster with an endonuclease operating in spermatogenesis (Galizi et al., 2016). The lack of target-site resistance, at least when observed at the limited scale of population cage experiments, reflects the use of extremely conserved rDNA target sequences which are present in hundreds of copies on the X chromosome, although even this cannot completely remove the possibility of resistance evolving. Gene drive systems targeting the heterogametic sex chromosome have only been investigated theoretically (Holman, 2019; Prowse et al., 2019) and in preliminary experiments in a house mouse system (Prowse et al., 2019).

3.1.3 | Wolbachia

The cytoplasmic incompatibility wMelPop strain of *Wolbachia* was originally isolated from a laboratory screen of *D. melanogaster*, where it shortens lifespan (Min & Benzer, 1997). It was proposed as a tool for biocontrol of dengue in *Aedes aegypti* mosquitoes as viruses are only transmitted by older mosquitoes. In addition, *Wolbachia* directly inhibits replication of the dengue virus (Walker et al., 2011). wMelPop and other Wolbachia strains have been successfully transinfected into *Aedes aegypti* and their spread is driven by cytoplasmic incompatibility. The spread of wMelPop into natural populations has proven challenging as this strain inflicts a suite of costly effects on its host (Nguyen et al., 2015) but strains with fewer deleterious effects have proven more successful (Hoffmann et al., 2011; Nazni et al., 2019).

There are several avenues to the evolution of resistance to *Wolbachia* infection. Mosquitoes could evolve resistance to cytoplasmic incompatibility or *Wolbachia* itself, *Wolbachia* infections could attenuate over time (there are many examples of weak effects in natural *Wolbachia* infections; Caragata, Tikhe, & Dimopoulos, 2019, or arboviruses could evolve to bypass the inhibitory effect of *Wolbachia* (Bull & Turelli, 2013). Alternatively, as high temperature can eliminate *Wolbachia* infections, it might be possible for mosquitoes to suppress infections by altering their temperature preferences (Ross et al., 2019). However, a trial introduction of *Wolbachia* has seen maintenance of strong cytoplasmic incompatibility and relatively stable frequencies in Australian field populations for seven years since their release, suggesting this may be unlikely, or at least slow to evolve (Ross et al., 2019; Ryan et al., 2020). After nearly a decade of use, there is as yet no evidence of any type of resistance evolving and the ability to block dengue virus has not been lost (Ross, Turelli, et al., 2019; Ryan et al., 2020). A further question is whether *Wolbachia* and dengue will enter a coevolutionary arms race against one another in these populations.

3.1.4 | Medea and underdominance-like systems

Medea-like systems encode a maternal toxin and zygotic antidote, killing offspring that do not inherit the *Medea* gene drive (Beeman, Friessen, & Denell, 1992). Synthetic underdominance systems are conceptually similar, consisting of a set of lethal loci, each associated with a suppressor of the other (Davis, Bax, & Grewe, 2001). Individuals inheriting only one of the loci carry a lethal locus, but not its suppressor, resulting in reduced viability or fertility. Resistance to these systems is likely to occur via changes to the toxin's target. For example, an underdominant maternal-effect lethal introduced
into the soft-fruit pest *D. suzukii* depends on a miRNA toxin and a zygotic antidote to function and will be impaired by variation at the miRNA binding site. Indeed, a recent survey shows natural variation in the miRNA toxin target sites (Buchman, Marshall, Ostrovski, Yang, & Akbari, 2018). Population cage experiments found that the Medea drive was unable to persist in populations, likely due to a combination of significant fitness costs of the driver as well as standing variation in resistance present in the cage populations (Buchman, Marshall, et al., 2018).

In addition to target-site mutation, Medea and similar toxin-antidote systems could also encounter resistance through driver inactivation either through direct suppression or the spread of antidote-only alleles due to mutational inactivation of toxin production. The single study investigating the stability of a *D. melanogaster* underdominance system found no evidence of resistance evolution over > 200 generations (Reed et al., 2018). Finally, there has been recent theoretical proof of principle of other Medea-like systems that rely on either CRISPR-Cas9 transcriptional overactivation of an endogenous target gene as the "toxin" and an insensitive copy of that target as the "antidote" or CRISPR-Cas9 cleavage as the "toxin" and resistant target gene as the "antidote" (Champer, Kim, Champer, Clark, & Messer, 2019). These too will face similar types of resistance (e.g. target-site mutation, driver inactivation). They are not in principle different from other synthetic gene drive systems that utilize CRISPR-Cas9, although their development is still at an early stage and not advanced enough for empirical investigation of resistance evolution.

### 4 | THE STRENGTH OF SELECTION FOR RESISTANCE ACROSS GENE DRIVE SYSTEMS

The strength of selection against a driver can vary dramatically between drive mechanisms and targets. At one extreme, a synthetic driver aimed at killing carriers or preventing reproduction, or distorting sex ratios will create extremely strong genome-wide selection for resistance against drive. In contrast, a biased gene converter that carries no cost to the organism will select for resistance at the target locus and linked sites, but have no effect on the rest of the genome. Drivers may themselves have a range of harmful pleiotropic effects, or be in linkage with deleterious alleles (Burt & Trivers, 2006). Fitness loss is often observed in both males and females, especially when drivers are homozygous (Dyer & Hall, 2019; Finnegan et al., 2019; Hamilton, 1967; Larner et al., 2019; Zanders & Unckless, 2019).

To understand the strength of selection against novel drivers, we need to know their fitness consequences in the field. There is currently a lack of such information for virtually all considered synthetic gene drives. One of the few systems where such information is readily available is for *Wolbachia*-carrying *Aedes* mosquitoes. The fitness costs associated with *Wolbachia* infection have been shown to be exacerbated under field conditions. As an example, the wMelPop *Wolbachia* strain, which invaded mosquito populations in semi-field cage trials, failed in several field trials because infected females had unexpectedly reduced egg viability in the field (Nguyen et al., 2015). This emphasizes the need for field studies of the fitness of drive carriers for the use of gene drive in natural populations.

The spatial structure of target populations is likely to be an important factor in deciding the fate of a gene drive system, as well as the way resistance may arise or spread. For example, Noble et al. (2018) showed that moderate amounts of gene flow between neighbouring populations is sufficient for a HEG-based replacement gene drive to spread between populations, even when resistance systematically arises in each individual population. More generally, we expect not only population genetic structure but also landscape and ecological characteristics to significantly impact the fate of a gene drive. Abiotic barriers (highways, open fields) have been shown to impede the spread of *Wolbachia* infections due to the limited dispersal ability of *Aedes* mosquitoes (Schmidt et al., 2017).

We can also imagine the evolution of tolerance to drive—meaning that the rest of the genome mitigates the deleterious effects of drive without directly interfering with the drive mechanism. For example, in stalk-eyed flies, males with drive invest more in testes to compensate for the loss of half of their sperm caused by the driver (Meade et al., 2020). Such changes do not interfere with drive and may actually enhance its spread. They lessen the deleterious costs of drive to the rest of the genome even though they do not improve fitness for the target chromosome. This reduction in the costs of the gene drive potentially reduces the strength of selection to suppress the driver.

There has been surprisingly little consideration of how all these processes interact when a new driver evolves or enters a population. Does the evolution of an effective defence mechanism against a driver preclude the evolution of other defences? There may be some parallels with the evolution of multiple defences against predators and parasites, which suggests multiple defences commonly evolve (Broom, Higginson, & Ruxton, 2010).

### 5 | STRATEGIES FOR DESIGNING SYNTHETIC DRIVE SYSTEMS TO REDUCE RESISTANCE

Although resistance to synthetic gene drive elements cannot be entirely prevented, it can be anticipated and reduced by careful gene drive design. In particular, the study of both natural and synthetic gene drive systems has taught us some important lessons about how resistance may arise in the face of various drive systems, and what we can do to minimize different types of resistance.

Many gene drives require a target, and evolved changes to this target are an obvious route to resistance. Recent work on CRISPR-Cas9-based homing endonuclease genes (HEGs) and synthetic Medea elements has highlighted the substantial impact that pre-existing and drive-generated target-site changes can have on preventing the spread of synthetic drivers (Champer et al., 2017; Gantz et al., 2015). There are several design strategies that may...
help prevent target-site resistance. Firstly, targets in essential and/or highly conserved sequences/genes may be less tolerant of sequence variability/polyorphism and thus less likely to harbour pre-existing resistance alleles, or to tolerate novel mutational variation (Buchman, Ivy, Marshall, Akbari, & Hay, 2018). For homing-based systems, gene drives could home into genes that are essential, so that incorrect homing events (e.g. nonhomologous end joining) result in lethal products (Bull & Malik, 2017; Esvelt, Smidler, Catteruccia, & Church, 2014; Kyrou et al., 2018). A second strategy would be to target multiple sites. The same principle applies to Medea or other systems with “toxins” that act on specific sequence regions (Champer et al., 2017, 2018; Marshall et al., 2017; Noble et al., 2017). Combining multiple mechanisms, for example a suppressive gene drive that also distorts the sex ratio, could be another way to delay the emergence of resistance (Simoni et al., 2020).

It is also critical to make the driver as stable as possible. For example, reducing the size of a CRISPR-Cas9 HEG transgene increases the likelihood that it will copy itself correctly, and integrating such a drive into endogenous genes may help achieve this goal (Nash et al., 2019; Hoermann et al., 2020). Clearly this may trade off with the benefits of more complicated drivers that reduce resistance evolution by attacking multiple loci. Additionally, repetitive DNA sequence (such as from multiple sgRNA or miRNA backbones) can reduce stability (Bzymek & Lovett, 2001; Marshall et al., 2017; Simoni et al., 2014), and reduction of such repetitiveness can protect against recombination and possible loss of a part of the drive element. It is also important to take into consideration the inherent evolutionary stability of integral gene driver components and mechanisms. For example, using a smaller protein than Cas9 in the drive mechanism could reduce the chance of mutations that inactivate the driver. Additionally, the endogenous homology-directed repair process required for CRISPR-Cas9 HEG function may be error-prone and lead to driver loss of function (Hammond et al., 2016; Oberhofer et al., 2018). Conversely, miRNA or chromosomal rearrangement-based systems may be more evolutionarily stable because they do not rely on large exogenous proteins and error-prone repair pathways to function.

Minimizing any fitness costs of the driver is also likely to reduce selection for resistance. Genomic insertion sites are associated with different costs, so transgenes inserting at a low cost site may create less selection for resistance. It is also advisable to reduce pleiotropic impacts of gene drive, as this can create resistance alleles in some systems. For example, work on CRISPR-Cas9 HEGs suggests that expression of the nuclease in somatic cells can lead to off target-site mutation which reduces the spread of the driver (Beaghton, Hammond, Nolan, Crisanti, & Burt, 2019; Champer et al., 2017; Gantz et al., 2015; Hammond et al., 2017).

It is also important to remember the ecology of the target species, as this may offer novel ideas for making a gene drive system durable, or reveal weaknesses only present in the field. For example, extremely high temperatures in Australia in 2019 may have impaired the transmission of the temperature-sensitive wMel Wolbachia strain used to combat dengue in Queensland mosquitoes. Synthetic drives designed in benign laboratory conditions may struggle in the field during extreme environmental conditions. If a gene drive is unable to penetrate some areas of an environment, due to conditions that prevent drive function or increase its costs, this could provide ideal circumstances for resistance to evolve.

Finally, it is essential to choose the right gene drive for the job. Certain types of drive (e.g. translocations) are much less likely to face resistance, but may spread more slowly than drivers that bias segregation (Buchman, Ivy, et al., 2018; Champer et al., 2016). Additionally, population suppression drives will face considerably stronger evolutionary pressures in terms of resistance than replacement drives (Eckhoff, Wenger, Godfray, & Burt, 2017; KaramiNejadRanbar et al., 2018; Prowse et al., 2017). However, resistance will not always be an impediment to gene drive deployment. For example, if the goal is short-term transformation of a population, then long-term evolution of resistance against the gene drive may not matter (Unkless et al., 2017). Resistance in nontarget populations may make gene drives less likely to spread accidentally (Esvelt et al., 2014). If the target population carries only susceptible alleles, but surrounding populations have a mix of susceptible and resistant alleles, the driver may also be unable to successfully spread to nontarget populations (Sudweeks et al., 2019).

6 | CONCLUSIONS

The evolution of resistance is a key problem in the design and use of gene drives. It is a major challenge faced by natural gene drive systems but remains poorly understood. Resistance based on interference can arise very rapidly, within a single generation, but in some natural systems does not appear to have evolved despite long timeframes.

As illustrated by this review, mechanisms of resistance are very diverse. Although we understand some of the mechanisms that can resist drivers, we rarely have a clear understanding of the forces underlying individual resistance pathways, nor the biological and ecological factors that determine which resistance type or mechanism is more likely to be selected in a given situation. In the context of applied control programmes using specific gene drive approaches, the current lack of knowledge means that robust predictions cannot be made about the types of resistance that are most likely to arise that might impede success.

Nevertheless, combining work from natural and synthetic drive provides an opportunity for a new depth of understanding. The study of synthetic drive systems is limited to short evolutionary timescales and typically small population sizes. In contrast, natural drive systems have evolved over thousands of generations, and several are ancient. Thus, synthetic drives provide a unique opportunity to understand what happens at short time scales when a novel driver arises, with unprecedented understanding of the mechanisms of drive and resistance. Techniques borrowed from synthetic technologies could allow researchers working on natural drive to create constructs based on their driver of interest and
directly test hypotheses related to evolutionary history and fate of natural drive. Conversely, a better understanding of the mechanisms of natural drivers, particularly those that appear hard to suppress, may lead to new generations of synthetic drivers that mimic the successful traits of the natural drivers. Lessons from both synthetic and natural drivers can be combined to produce safer, more effective drive systems.

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CONFLICTS OF INTEREST

The authors state they have no conflicts of interest to declare.

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