Gene Drive: Evolved and Synthetic

Austin Burt*† and Andrea Crisanti†‡

†Life Sciences, Imperial College, Silwood Park, Ascot, SL5 7PY, United Kingdom
‡Life Sciences, Imperial College, South Kensington, London, SW7 2AZ, United Kingdom

ABSTRACT: Drive is a process of accelerated inheritance from one generation to the next that allows some genes to spread rapidly through populations even if they do not contribute to—or indeed even if they detract from—organismal survival and reproduction. Genetic elements that can spread by drive include gametic and zygotic killers, meiotic drivers, homing endonuclease genes, B chromosomes, and transposable elements. The fact that gene drive can lead to the spread of fitness-reducing traits (including lethality and sterility) makes it an attractive process to consider exploiting to control disease vectors and other pests. There are a number of efforts to develop synthetic gene drive systems, particularly focused on the mosquito-borne diseases that continue to plague us.

INTRODUCTION TO GENE DRIVE

Most genes are thought to spread and persist in populations because they do something useful for the organisms carrying them, increasing survival and/or reproduction, at least on average. That is, most genes spread in populations by positive Darwinian selection and are maintained in the face of recurrent mutation by purifying selection. Some features of a gene may be selectively neutral—such as which nucleotide is found at a particular silent site—but random drift by itself will not lead to long open reading frames and associated control sequences that produce complex proteins performing a particular function.

There are other genes, a minority, which spread and persist not by natural selection, not by increasing organismal survival or reproduction, but instead by distorting their transmission from one generation to the next. For example, some genes manage to be transmitted to more than half of an individual’s gametes, even when that individual only inherited the gene from one of its two parents. In this case, the frequency of the gene increases due to the process of gene transmission from one generation to the next, and it is this unequal genetic transmission that gives the gene its advantage. Genes or genetic elements showing such transmission ratio distortion, or “drive,” include gametic and zygotic killers, meiotic drivers, homing endonuclease genes, B chromosomes, and transposable elements, each of which has evolved several or many times in different taxa. Moreover, drive has been an important process affecting such genomic features as genome size, base composition, chromosome shape, repeat structure, distribution of recombination hotspots, and centromere structure.

Not only can driving genes spread without doing anything useful for the organisms carrying them, they can even spread if they cause some harm, as long as the effect of the transmission distortion is greater than the effect of the reduced survival and reproduction. For this reason, they are often called selfish genes, or selfish genetic elements. And since they can be harmful to the organism, genes that suppress these genes can themselves spread by natural selection (analogous to the spread of genes suppressing any other parasite), which the selfish gene will then be selected to avoid, potentially leading to an arms race. Occasionally, features of a selfish genetic element may be co-opted to do something useful for the host. Classic examples include mating type switching in yeast, antibody diversification in vertebrates, and telomere maintenance in Drosophila (and in eukaryotes more generally)—the evolution of all of these operations has involved the domestication or co-option of functions of selfish genetic elements.

SYNTHETIC GENE DRIVE SYSTEMS

The fact that gene drive can lead to the spread of fitness-reducing traits makes it potentially useful for controlling disease vectors and other pests. Moreover, the spread can be rapid: in a closed, random mating population, a construct with 100% drive and no fitness effects can increase from 1% to 99% in the population in just nine generations—fast enough to be attractive for public health interventions. Discussions about how to exploit gene drive for pest control date back for decades, long before there was any mechanistic understanding of how they worked, particularly among medical entomologists looking for new ways to control disease vectors. However, classical genetic approaches were not sufficiently flexible to be able to construct a useful gene drive system. Now, with the recent progress in molecular biology, there is renewed interest in trying to make synthetic gene drive elements.
preliminary discussions of potential uses have expanded to agriculture and conservation.\textsuperscript{13} In broad outline, two types of intervention have been considered, either to reduce the size of the target population, or to leave numbers more-or-less intact and genetically modify the population such that it is less harmful (e.g., less able to transmit a pathogen). And three main molecular paradigms are being explored, the use of toxin–antidote systems, chromosomal rearrangements, or sequence-specific nucleases. Toxin-antidote systems and chromosomal rearrangements may be useful for introducing and spreading a new “cargo” gene through a population that makes the population less harmful (e.g., an effector gene that makes mosquitoes unable to transmit a pathogen).\textsuperscript{14,15} Nuclease-based drive systems may also be used to introduce novel genes, or they may be used for population suppression.\textsuperscript{16,17}

**Toxin-Antidote Systems and Chromosomal Rearrangements.** Naturally occurring gene drive systems act as if they produce a toxin and antidote, though often the molecular details are not known. For example, in mice heterozygous for the t-haplotype, and *Drosophila* heterozygous for Segregation Distorter, these elements somehow act during spermatogenesis to sabotage spermatids or sperm carrying the wild-type allele, with the result that each is transmitted to over 90% of the progeny (compared to the Mendelian 50%).\textsuperscript{18,19} In *Tribolium* flour beetles, the *medea* gene acts in heterozygous females to somehow cause progeny that do not inherit the *medea* gene to die.\textsuperscript{20,21} Though the underlying molecular mechanisms are not known, this example stimulated the development of a synthetic gene drive construct in *Drosophila* with the same logic.\textsuperscript{22} The construct combined a microRNA-based repressor of *myd88* (an important protein normally supplied by the mother into the embryo) with a zygotically expressed *myd88* gene that was not affected by the microRNA and supplied the missing protein. As intended, this construct was able to increase in frequency over successive generations in experimental cage populations. Two other *medea* systems, using different components, have also been developed in *Drosophila*,\textsuperscript{23} as have toxin–antidote systems that display maternal-effect lethal underdominance and threshold-dependent invasion into population cages.\textsuperscript{15,24} Recent descriptions of natural toxin–antidote systems in plants, fungi, and nematodes\textsuperscript{25–31} may provide further insights into how these sorts of systems can be engineered.

One way for toxin–antidote systems to work is by generating underdominant fitness interactions, in which the heterozygote is less fit than either of the two homozygotes. Underdominant interactions can also be generated with chromosomal rearrangements such as reciprocal translocations, and if these can be introduced at a sufficient frequency into a population (>50% in the simplest scenario), then they can spread to fixation.\textsuperscript{6} Strains of *Drosophila* with reciprocal translocations have been engineered, and these showed the expected frequency-dependent spread in lab populations.\textsuperscript{32}

**Nuclease-Based Systems: Chromosome Shredding.** In *Aedes* and *Culex* mosquitoes, there is a naturally occurring driving Y chromosome that, in some crosses, is transmitted to more than 90% of a male’s progeny. First described in the 1960s,\textsuperscript{33} there is still no good understanding of how it works at the molecular level, but cytologically it is associated with breaks of the X chromosome at male meiosis, perhaps having something to do with interrupted crossovers.\textsuperscript{34,35} This observation led to the idea that cleavage of the X chromosome during male meiosis might lead to drive of the Y.\textsuperscript{16} In *Anopheles gambiae*, the most important vector of malaria in Africa, the rRNA genes are found in a single cluster of hundreds of copies on the X chromosome, making it an ideal target,\textsuperscript{36} and sure enough, production of a nuclease targeting this sequence during spermatogenesis can produce biased sex ratios, up to 95% males, using both an engineered meganuclease and a CRISPR-based nuclease.\textsuperscript{37,38} A male-biased population sex ratio would be useful because males do not bite people and transmit disease, nor do they contribute as much materially to population productivity, and so total population size is also likely to decline.\textsuperscript{39} The constructs reported to date do not yet constitute a fully functional gene drive system, because the nuclease genes have been inserted on an autosome, and so are themselves still transmitted in a Mendelian manner; the next step is to put them on the Y chromosome, which is challenging because it is highly repetitive and largely suppressed at meiosis, though some progress has been made.\textsuperscript{40,41}

**Nuclease-Based Systems: Homing.** Homing endonuclease genes (HEGs) are a class of natural occurring driving elements for which there is a good understanding about the molecular mechanisms, and these are both simple and general enough to potentially be worth exploiting. HEGs encode a nuclease that recognizes and cuts a sequence that typically occurs just once in the genome. The gene is in the middle of its own recognition sequence, disrupting it and protecting the chromosome it is on from being cut. Therefore, in heterozygotes, only the chromosome not containing the gene is cut; it is then repaired using the HEG-containing homologue as a template, with the result that the HEG is copied across to the chromosome where previously it was absent, converting a heterozygote into a homozygote.\textsuperscript{42,43} This “homing” reaction simply requires a gene encoding a sequence-specific nuclease, with the gene inserted in the middle of its own recognition sequence, and the cell’s DNA repair system takes care of the rest.

HEGs occur naturally in many microbes but have not yet been identified in any insects or vertebrates. Important proof-of-principle experiments demonstrated that the homing reaction can occur in both *Drosophila* and *Anopheles*.\textsuperscript{44–47} These first experiments used meganucleases, and later experiments showed the reaction could also be catalyzed by zinc finger and TALE nucleases,\textsuperscript{48} and, more recently, CRISPR-based nucleases.\textsuperscript{49–51} In principle, the homing reaction can be used for both population-wide knockout of target genes, such as those involved in survival, fertility, or pathogen transmission, or for population-wide knock-in of novel effector genes.\textsuperscript{17,52,53} As with other forms of pest or disease control, due attention must be given to the possibility of resistance evolving.\textsuperscript{54–57}

### PROSPECTS

There are a relatively small number of species for which genetic control methods, including gene drive, may be appropriate. Most prominent are those causing or transmitting diseases. Even now, more than 700 000 people die every year from vector-borne diseases, and there is an additional heavy burden of nonlethal morbidity.\textsuperscript{58} Much of disease control is, ultimately, chemical, with efficacy largely determined by the degree to which production and delivery can be targeted and affordable. Vaccination can be among the most cost-effective of all health interventions because it uses the adaptive immune system to generate and deliver the active agents. The promise of genetic approaches—and gene drive in particular—is again to use
biological processes—mating, meiosis, transcription, translation, etc.—for targeted, cost-effective delivery of appropriate chemicals (e.g., nucleases or antimicrobial peptides) that will substantially reduce disease transmission. Indeed, gene drive may take efficiency a step further, with a single release (perhaps with periodic “booster” releases) giving area-wide, population-level control. Important steps have been made toward realizing this potential, though there remains much more to do.

**Author Information**

**Corresponding Authors**
*E-mail: a.burt@imperial.ac.uk.*
*E-mail: adcrisanti@imperial.ac.uk.*

**Notes**
The authors declare no competing financial interest.

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