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Safeguarding gene drive experiments in the laboratory

Multiple stringent confinement strategies should be used whenever possible

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Gene drive systems promote the spread of genetic elements through populations by assuring they are inherited more often than Mendelian segregation would predict (see the figure). Natural examples of gene drive from *Drosophila* include sex-ratio meiotic drive, segregation distortion, and replicative transposition. Synthetic drive systems based on selective embryonic lethality or homing endonucleases have been described

previously in *Drosophila melanogaster* (1–3), but they are difficult to build or are limited to transgenic populations. In contrast, RNA-guided gene drives based on the CRISPR/Cas9 nuclease can, in principle, be constructed by any laboratory capable of making transgenic organisms (4). They have tremendous potential to address global problems in health, agriculture, and conservation, but their capacity to alter wild populations outside the laboratory demands caution (4–7). Just as researchers working with self-propagating pathogens must ensure that these agents do not escape to the outside world, scientists working in the laboratory with gene drive constructs are responsible for keeping them confined (4, 6, 7).

Two of us recently used a CRISPR/Cas9-based gene drive system to generate a *Drosophila* strain homozygous for a loss-of-function mutation [the mutagenic chain reaction (6)] (see the figure). Even though *D. melanogaster* ordinarily poses no threat to human health or agriculture, the accidental release of flies carrying gene drive constructs from the laboratory could have unpredictable ecological consequences. This study there-

fore used institutionally approved stringent barrier methods. Only one experimenter handled the flies, inside an Arthropod Containment Level 2 insectary suitable for work with mosquitoes carrying human pathogens. Because barrier protocols can be vulnerable to human error (8), these authors suggested (6) that additional molecular confinement methods described (4) and used by others of us in budding yeast (9) could further reduce risks. That these studies documented highly efficient RNA-guided gene drive in flies and yeast underscores the potential of the technology and the risk resulting from an accidental release.

As concerned scientists working in related areas, we engaged in collective discussions to identify and publicize interim safety recommendations for laboratory

research involving potential gene drive systems while formal national guidelines are developed. Although we cannot claim to represent all researchers, we share a commitment to the safe and responsible development of gene drive technology. Although we differ in our assessments of the types of precaution needed, we recognize that any single confinement strategy could fail. We therefore unanimously recommend that future studies use a combination of stringent confinement strategies (see the table) whenever possible and always use safeguards adequate for preventing the unintentional release of synthetic gene drive systems into natural populations.

RECOMMENDATIONS. RNA-guided gene drive systems are created by delivering into the germline a DNA cassette encoding Cas9 and a single synthetic guide RNA (sgRNA) that is flanked by sequences matching those on either side of the sgRNA target site (4). Cas9 nuclease-stimulated copying of the cassette into the target allele leads to continued Cas9+sgRNA expression and subsequent copying of the cassette into the other allele (6, 9). The recurrent conversion of heterozygotes into homozygotes permits spread through populations (see the figure).

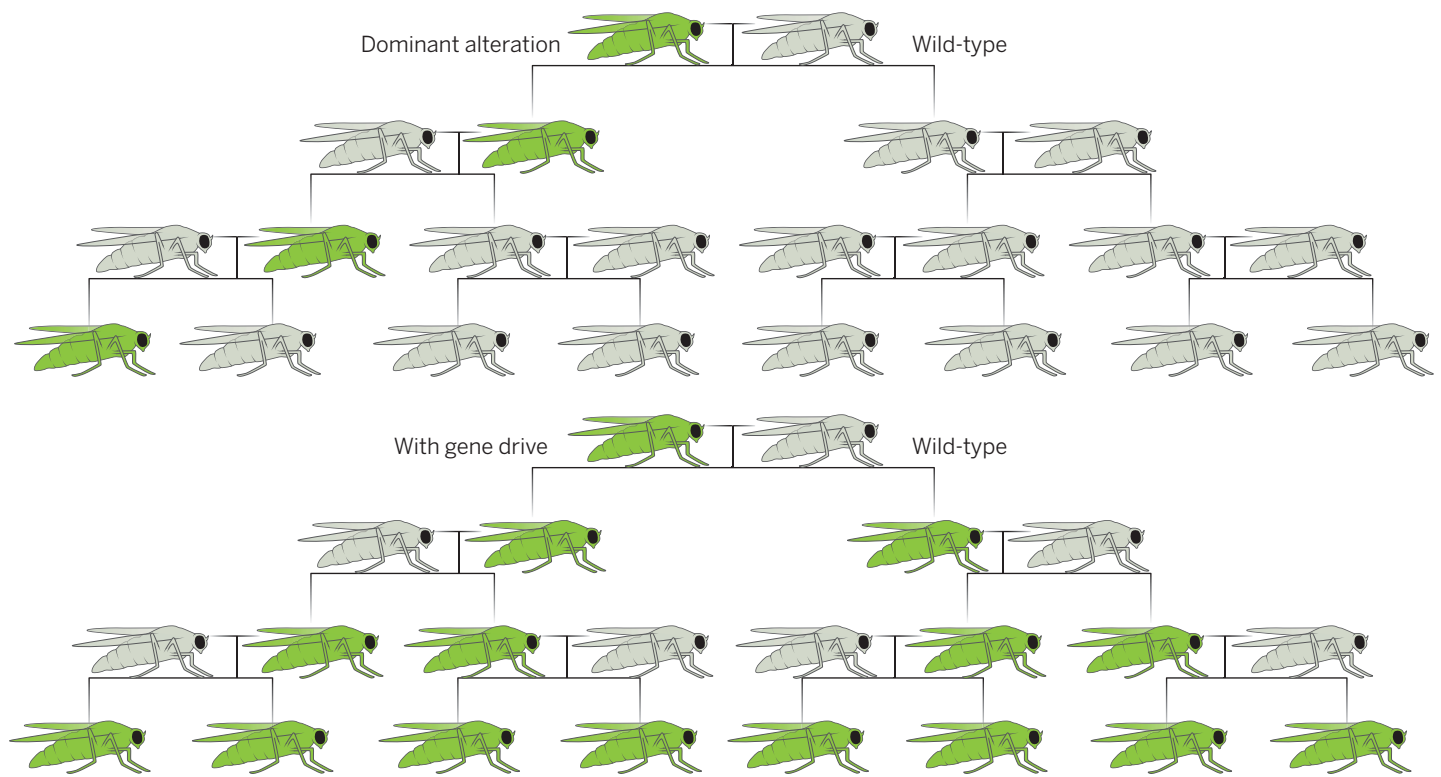
The vast majority of recent genome engineering approaches developed in model organisms neither involve nor risk the creation of gene drive systems. For example, *Drosophila* mutants can be readily generated by injecting sgRNAs or sgRNA-encoding plasmids into transgenic embryos expressing Cas9 (10–13) or by crossing

Potentially stringent confinement strategies for gene drive research

Multiple stringent confinement strategies should be used whenever possible.

TYPE	STRINGENT CONFINEMENT STRATEGY	EXAMPLES
Molecular	Separate components required for genetic drive Target synthetic sequences absent from wild organisms	sgRNA and Cas9 in separate loci (8) Drive targets a sequence unique to laboratory organisms (3,4,8)
Ecological	Perform experiments outside the habitable range of the organism Perform experiments in areas without potential wild mates	<i>Anopheles</i> mosquitoes in Boston <i>Anopheles</i> mosquitoes in Los Angeles
Reproductive	Use a laboratory strain that cannot reproduce with wild organisms	<i>Drosophila</i> with compound autosomes*
Barrier	Physical barriers between organisms and the environment • Remove barriers only when organisms are inactive • Impose environmental constraints • Take precautions to minimize breaches due to human error	Triply nested containers, >3 doors (6) Anesthetize before opening (6) Low-temperature room, air-blast fans Keep careful records of organisms, one investigator performs all experiments (6)

*An example of reproductive confinement would be *Drosophila* laboratory strains with a compound autosome, where both copies of a large autosome are conjoined at a single centromere. These strains are fertile when crossed inter se but are sterile when outcrossed to any normal or wild-type strain because all progeny are monosomic or trisomic and die early in development.



The spread of RNA-guided gene drive systems. Unlike the population dynamics of normal genomic alterations, gene drive systems can spread changes through wild populations by converting heterozygotes into homozygotes in each generation.

sgRNA-expressing strains to Cas9-expressing strains (12–14). These approaches do not risk creating a gene drive system because cassettes encoding Cas9 and sgRNA are not inserted into the cut site or located adjacent to one another in the genome and can thus be safely used by researchers without additional precautions. Given the availability of efficient alternatives and the potential risks, we recommend that gene drive approaches to genome engineering be strictly reserved for cases that require their use.

The safest approach for using gene drives creates biallelic mutations with an sgRNA-only cassette that can spread only when combined with an unlinked Cas9 transgene (4). In such a “split gene drive system,” homozygous individuals lacking the Cas9 gene can be easily isolated in subsequent generations. The efficiency of gene drive exhibited by a split system in yeast is equivalent to that of a construct encoding both Cas9 and sgRNA (9). Split drive systems present a much lower

risk if organisms are accidentally released because the population frequency of the Cas9 gene will be determined by normal, nondrive dynamics, consequently limiting the spread of the sgRNA cassette.

Nevertheless, any mutational event that moves the Cas9 gene into or directly adjacent to the sgRNA cassette could create an autonomous Cas9+sgRNA drive system by allowing the Cas9 gene to be copied into the target locus along with the sgRNA cassette upon repair of Cas9-induced DNA cleavage. Although the probability of such an event is extremely low, we recommend that at least one additional form of stringent confinement be used (see the table) and that the strains be continually monitored.

Other forms of stringent confinement include performing experiments in an area lacking wild populations (4) and, when the goal is to study gene drive systems in the laboratory, exclusively targeting synthetic sequences not found in natural populations

(3, 4, 9). Because these strategies suffer from independent vulnerabilities, the safety improvements afforded by combining them will be multiplicative. Thus, the great majority of gene drive experiments can be performed with minimal risk of altering wild populations. Accordingly, we strongly recommend that

1) All work involving potential gene drive systems should be preceded by a thorough assessment by the relevant biosafety authorities of the risk of unwanted release from the laboratory. We encourage these authorities to seek guidance from external experts and make their evaluation available to others.

2) All laboratory gene drive experiments should employ at least two stringent confinement strategies (see the table) whenever possible to minimize the risk of altering wild populations. Using one form of confinement may be justified only if relevant biosafety authorities determine that

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it will reduce the probability of release to a level that is acceptably low. This probability must be defined on a case-by-case basis. The analyses necessary to confidently predict the efficacy of confinement strategies for gene drive systems are in a nascent form. Therefore, any proposal to use one rather than multiple forms of confinement requires even greater scrutiny and extensive deliberation between regulatory authorities and scientists.

3) Organisms carrying gene drive constructs that could spread if the reproductively capable life stages were to escape in transit should not be distributed to other institutions until formal biosafety guidelines are established. Whenever possible, laboratories should instead send DNA constructs or information sufficient to reconstruct the gene drive. Protocols for distributing materials should be established in discussion with the wider research community and other relevant stakeholders.

Broadly inclusive and ongoing discussions among diverse groups concerning safeguards, transparency, proper use, and public involvement should inform expert bodies as they develop formal research guidelines for gene drive research in the laboratory and potential transitions to open field trials. We applaud the U.S. National Academy of Sciences for committing to provide recommendations for responsible gene drive research (15). By recommending strong safeguards and encouraging discussion of this technology, we hope to build a foundation of public trust for potential future applications in public health, sustainable agriculture, and ecological conservation. ■

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MICROBIOME

Microbiota ROR γ t Regulates intestinal suppressor T cells

Gut microbes influence the balance of regulatory T cell subtypes to control inflammation

By Ahmed N. Hegazy^{1,2} and Fiona Powrie^{1,2}

The immune system in the intestine is highly adapted to resist invading pathogens while residing peacefully with the abundant and diverse commensal bacteria that colonize the gastrointestinal tract. In turn, bacterial signals shape immunity in the intestine, promoting intestinal homeostasis in part by inducing and expanding specialized regulatory T (T_{reg}) cells that prevent aberrant inflammatory responses to self and environmental stimuli (1). On pages 989 and 993 of this issue, Ohnmacht *et al.* (2) and Sefik *et al.* (3), respectively, report the development and function of a subpopulation of T_{reg} cells found primarily in the large intestine, and characterized by expression of the nuclear hormone receptor retinoic acid receptor-related orphan receptor γ (ROR γ t). This is surprising because ROR γ t classically promotes the differentiation of T helper 17 (T_H17) cells, a population associated with tissue inflammation in many inflammatory diseases (4). Both studies show that microbiota-derived signals induce the expression of ROR γ t in T_{reg} cells that control intestinal inflammation (see the figure). These findings highlight the diversity of colonic T_{reg} cells, their complex transcriptional programs, and their important role in the intestine.

T_{reg} cells express the forkhead transcription factor Foxp3, which promotes their differentiation, maintenance, and function (5). Alongside anti-inflammatory functions, they control nonimmunological processes including tissue repair and metabolism in the parenchyma (6). T_{reg} cells also adapt to environmental cues by expressing canonical effector T cell-associated transcription factors to control pathogenic immune responses (7).

Both Ohnmacht *et al.* and Sefik *et al.* found that in mice, a large fraction of intestinal T_{reg} cells express ROR γ t. These cells were distinct from colonic T_{reg} cells that express the transcription factor GATA3 and are poised to respond to the cytokine interleukin (IL)-33 after tissue damage (8, 9). However, ROR γ t-expressing T_{reg} cells had an activated phenotype similar to that of GATA3-expressing T_{reg} cells, and bore markers related to T_{reg} cells

residing in lymphoid and non-lymphoid tissues (6). Strikingly, the microbiota was an absolute requirement for the induction and maintenance of ROR γ t-expressing T_{reg} cells in these animals. This T_{reg} cell population was markedly reduced in germ-free mice, and colonization with a diverse microbiota or consortia of symbionts was sufficient for the induction of ROR γ t-expressing T_{reg} cells. Sefik *et al.* went further and recolonized germ-free mice with 22 different bacterial species, and showed that a number of them (not belonging to any specific phylum or genus) elicited ROR γ t-expressing T_{reg} cells at comparable frequencies to a diverse microbiota. Short-chain fatty acids, which are common bacterial metabolites, can selectively expand intestinal T_{reg} cells (10). Ohnmacht *et al.* could increase ROR γ t-expressing T_{reg} cells by feeding mice a diet rich in the short-chain fatty acid butyrate.

“These studies...are an important stepping stone to deciphering the complex dynamics of different tissue-resident T_{reg} cell subsets...”

Which signals promote ROR γ t expression in T_{reg} cells? The T_H17-favoring cytokines IL-6 and IL-23 were required for accumulation of ROR γ t-expressing T_{reg} cells, which raises the question of what tips the balance toward these T cells rather than T_H17 cells. The vitamin A metabolite retinoic acid promotes T_{reg} cell generation in vivo and ROR γ t-expressing T_{reg} cells in vitro (11, 12). Consistent with this, Ohnmacht *et al.* show that vitamin A metabolism influences the differentiation equilibrium by favoring the development of ROR γ t-expressing T_{reg} cells in vivo. Although both T_{reg} cells and T_H17 cells express ROR γ t, analysis of all the transcripts expressed by each population revealed marked differences, suggesting that the transcriptional footprint of ROR γ t is context-dependent in different T cells.

What is the function of ROR γ t-expressing