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Original article



A gene drive does not spread easily in populations of the honey bee parasite *Varroa destructor*

Nicky R. Faber^{1,2}, Adriaan B. Meiborg¹, Gus R. Mcfarlane³, Gregor Gorjanc¹, and Brock A. Harpur⁴

¹ HighlanderLab, The Roslin Institute and Royal (Dick) School of Veterinary Studies, The University of Edinburgh, Easter Bush Campus, Midlothian EH25 9RG, United Kingdom

² Laboratory of Genetics, Department of Plant Sciences, Wageningen University & Research, Droevendaalsesteeg 1, 6708 PB Wageningen, The Netherlands

³ Burdon Group, The Roslin Institute and Royal (Dick) School of Veterinary Studies, The University of Edinburgh, Easter Bush Campus, Midlothian EH25 9RG, United Kingdom

⁴ Department of Entomology, Purdue University, West Lafayette IN 47907, USA

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Abstract – Varroa mites (*Varroa destructor*) are the most significant threat to beekeeping worldwide. They are directly or indirectly responsible for millions of colony losses each year. Beekeepers are somewhat able to control varroa populations through the use of physical and chemical treatments. However, these methods range in effectiveness, can harm honey bees, can be physically demanding on the beekeeper, and do not always provide complete protection from varroa. More importantly, in some populations varroa mites have developed resistance to available acaricides. Overcoming the varroa mite problem will require novel and targeted treatment options. Here, we explore the potential of gene drive technology to control varroa. We show that spreading a neutral gene drive in varroa is possible but requires specific colony-level management practices to overcome the challenges of both inbreeding and haplodiploidy. Furthermore, continued treatment with acaricides is necessary to give a gene drive time to fix in the varroa population. Unfortunately, a gene drive that impacts female or male fertility does not spread in varroa. Therefore, we suggest that the most promising way forward is to use a gene drive which carries a toxin precursor or removes acaricide resistance alleles.

Varroa destructor / gene drive / genetic population control / modelling

1. INTRODUCTION

When the varroa mite (*Varroa destructor*) jumped from its original host the Eastern honey bee (*Apis cerana*) to the Western honey bee (*Apis mellifera*), it spread rapidly around the globe and caused catastrophic losses of commercial and feral honey bee colonies (Traynor et al., 2020;

Corresponding author: N. R. FABER, nfaber@outlook.com Manuscript Editor: Klaus Hartfelder Buchmann & Nabhan, 1996; Wenner et al., 1996; Kraus & Page, 1995). To this day, varroa mites remain the most highly reported cause of colony loss for commercial beekeepers and hobbyists (Kulhanek et al., 2017; vanEngelsdorp & Meixner, 2010; Molineri et al., 2018; Traynor et al., 2020). There are treatment options available to beekeepers that allow them to control varroa. Unfortunately, currently available treatments do not provide complete protection from varroa and they often harm honey bees or are physically demanding for the beekeeper. For example, acaricides are among the most effective treatments



available and can kill between 49-82% of the varroa within a colony (Pietropaoli & Formato, 2019; Santiago et al., 2000; Roth et al., 2020). Despite their effectiveness, some acaricides also affect honey bees; they reduce honey bee fertility (Rangel & Fisher, 2019), foraging, and immune responses against bacterial infections (Gashout et al., 2020). More concerning still, in some populations varroa mites have developed resistance to acaricides (Sammataro et al., 2005; Elzen et al., 2002; Elzen et al., 2000; Milani, 1999). Beyond chemical treatments, beekeepers can use physical means of varroa control such as drone brood removal, which gives varroa mites limited opportunities to reproduce. However, physical methods can require significant labour and thus may not be feasible on a large scale (Calderone, 2005; Aliano & Ellis, 2005). The unfortunate fact of varroa mite control is that it relies on blunt chemical treatment methods that can harm bees and may not be effective longterm because of evolved resistance. This echoes similar treatment methods available to other pest species around the globe like malarial-vectoring mosquitoes and crop pests like spider mites (Carson, 1962; Prasittisuk & Busvine, 1977; Baker, 1952; Dennehy et al., 1983).

Genetic population controls, like those that can be implemented through the use of a gene drive (Champer et al., 2016), could be a more successful and more sustainable means to control varroa mites and other invertebrate pests than currently available chemical and physical methods (Esvelt et al., 2014). Gene drives are selfish genetic elements that can be engineered to promote the inheritance of desired alleles at rates much greater than conventional Mendelian inheritance (McFarlane et al., 2018). When a gene drive allele is introduced into a population, it spreads through the mating of gene drive carrying individuals with wild-type individuals (Esvelt et al., 2014). A CRISPR-based gene drive element encodes the two components of CRISPR (a Cas nuclease and guide RNA) and can contain a gene of interest one wishes to propagate (Gantz et al., 2015; Buchthal et al., 2019), or it can be targeted to a gene one wants to disrupt (Kyrou et al., 2018; KaramiNejadRanjbar et al., 2018; Lester et al., 2020). In the germline of gene drive carriers, the Cas nuclease and guide RNA are expressed to generate a double-stranded DNA break on the opposing wild-type chromosome at the gene drive locus. This DNA break is repaired through homology-directed repair, using the gene drive harbouring chromosome as the repair template, and thus the gene drive element is copied to the second chromosome (Esvelt et al., 2014). The conversion rates for gene drives in insects can be as high as 100% (Gantz et al., 2015; Hammond et al., 2016; Kandul et al., 2020; Terradas et al., 2021). This process occurs again in the offspring generation and will do so in all subsequent generations, resulting in the gene drive spreading through the target population. A gene drive can be designed to reduce the fitness of individual homozygous carriers with the aim to reduce population size or even achieve extirpation (Champer et al., 2016; Faber et al., 2021).

The introduction of CRISPR-Cas9 gene drives as a management tool for varroa numbers could greatly impact our ability to control them, and technology is progressing to a stage where we could test this strategy. The necessary biochemical and biological research is currently coming together: in vitro-rearing techniques for varroa are being refined (Egekwu et al., 2018; Jack et al., 2020), there is a high-quality reference genome (Techer et al., 2019), and there is a growing list of genes essential to mite survival (Huang et al., 2019). CRISPR-Cas9-mediated mutagenesis has not yet been published for varroa mites but recent work on spider mites demonstrates that this may soon be possible (Dermauw et al., 2020). However, we do not yet know if a gene drive can spread in a varroa population. Prior to any gene drive system being implemented, it is essential to develop a species-specific genetic and demographic model to predict the effectiveness of a drive spreading successfully (James, 2005; Sinkins & Gould, 2006; Prowse et al., 2017; Unckless et al., 2017; Noble et al., 2018; KaramiNejadRanjbar et al., 2018; Lester et al., 2020; Faber et al., 2021). This is especially important in non-model species where mating biology and sex-determination systems can limit the spread of gene drives. In

the case of varroa mites, they can both outbreed and inbreed, and the proportion of each breeding strategy varies throughout the season based on brood cell availability (Bull, 2016; Noble et al., 2018). Inbreeding, along with haplodiploidy (Li et al., 2020) in varroa, reduces the likelihood of a gene drive spreading effectively.

We present a modelling study to investigate the effectiveness of a gene drive given the unique life history of varroa. We estimate the spreading efficiency of a gene drive in a single honey bee colony and identify management techniques beekeepers may have to implement to successfully spread a gene drive in their colonies. We show that spreading a neutral gene drive in varroa is challenging because of the high rate of inbreeding and their exponential growth rate that can quickly overwhelm a honey bee colony. Management strategies like brood breaks and acaricides help to spread gene drive alleles. Unfortunately, we could devise no scenario to spread gene drives that impact fitness traits like male or female fertility. Therefore, we suggest that the most promising way forward is to use a gene drive which carries a toxin precursor or removes acaricide resistance alleles.

2. METHODS

Within R 4.0.5 (Team et al., 2013), we used the package AlphaSimR as a framework for our modelling (Gaynor et al., 2020). AlphaSimR is designed to model the genetics of plant and animal breeding schemes, but lends itself well to general population genetics modelling too. We have created an individual-based, stochastic, day-by-day model of varroa destructor, which consist of three aspects: a static honey bee colony as backbone, a stochastic model of varroa and its life history, and the implementation of a gene drive. Everyday in the model, we track parameters such as the size of the varroa population, the levels of inbreeding, and the allele frequencies at the gene drive locus, among others.

2.1. Honey bee colony simulation

Varroa is a parasite and depends on its honey bee host for reproduction. Therefore, to realistically model a population of varroa, we must also model a honey bee colony. We chose to use a static model for the honey bee colony, as we are primarily interested in the varroa population and not the interaction between parasite and host. We used a honey bee colony model from Calis et al. (1999), who based their model on data from Allen (1965). This model is based on a colony of average size in a Northern European climate and contains the amount of adult honey bees, drone brood, and worker brood over 365 days. At the end of the year, bee and brood numbers are the same as at the start of the year. Therefore, we can model multiple years by replicating this honey bee model several times back to back. We assumed that a honey bee colony would collapse when the varroa population reaches 10,000 individuals, at which point we stopped the model. We also implemented an option to reduce brood amounts through colony management by the beekeeper to manage inbreeding in the varroa population (Büchler et al., 2020). For a variable amount of days, we reduce the brood by a variable percentage of its original amount on those days. In our fixed honey bee colony model, we only change the amount of drone and worker brood and leave the adult bee numbers the same.

2.2. Varroa life history

Our model consists of a number of steps to accurately represent the complex life history of varroa mites:

 Initializing mated females. At the start of the model, we initialize a certain number of mated varroa females. Then, every time when female varroa offspring is created, we assign each varroa a certain number of reproduction cycles it will go through in its life. Current estimates of how many reproduction cycles are completed on average range between 2 and 3 (Martin & Kemp, 1997; Fries & Rosenkranz, 1996). Therefore, we assign each female a number between 1 and 4 randomly, which gives an average of 2.5 reproduction cycles.

2. Brood infestation. The first step in varroa reproduction is the infestation of a honey bee brood cell. For the rate of brood entering, we use a model by Boot et al. (1994), who tested several models to predict this rate. On every day of our model, we calculate the number of infestations (N_i) as:

$$N_i = 1 + e^{-(-2.87 + 0.00385 * \frac{N_b}{N_a} * 10000)^{-1}}, \quad (1)$$

which is dependent on the ratio between available brood (N_h) and the number of adult bees (N_a) (Boot et al., 1994). The biological reasoning behind this model is that varroa are phoretic on adults bees and when those bees get close to available brood cells, the varroa can infest (Boot et al., 1994). When this ratio is low, the probability that an adult bee with a phoretic varroa will pass by an available brood cell is low, and vice versa. Once we have determined the number of varroa that infest brood cells, we assign them to the available drone and worker cells. Varroa prefer drone cells over worker cells, because those are capped for 2 days longer (14 instead of 12 days) (Fries et al., 1994), which enables more varroa offspring to mature. We model a drone cell preference by giving drone cells an eight times higher probability of infestation (Fuchs, 1992). Therefore, by chance any drone or worker cell could be infested by more than one varroa, with the probability of this happening being much higher in drone cells.

3. Generating offspring. Varroa mites first produce a single male offspring, followed by a varying number of female offspring (Traynor et al., 2020). More female offspring are able to mature in drone brood than in worker brood because of the longer capping period of those cells (Rosenkranz et al., 2010). Therefore, we use two separate distributions to determine the number of female offspring per varroa in the two types of brood as described by Ifantidis (1984). These distributions include varroa that produce no offspring as well. The averages of these distributions for female offspring are 1.70 for drone cells and 0.71 for worker cells (Ifantidis, 1984). Excluding the non-productive varroa, the averages of female offspring are 2.77 for drone cells and 1.33 for worker cells (Ifantidis, 1984).

- 4. Mating between offspring. Varroa offspring mate in the brood cell they are born in (Nazzi & Le Conte, 2016). Usually only one varroa infests a cell, which forces offspring to inbreed by full-sibling mating. Occasionally however, especially at the end of the season when varroa numbers are high, multiple varroa infest a single cell, which allows for outbreeding (Beaurepaire et al., 2017). Mated females will generate offspring the rest of their lives with the sperm they save in their spermatheca (Rosenkranz et al., 2010). We model random mating between males and females in a brood cell, where females mate with a single male.
- 5. Emergence from brood. In every brood cell, there is a limit to how many varroa offspring can survive (Martin, 1995). According to data from Martin (1995), the maximum live offspring per cell is 16 in drone cells and 8 in worker cells. Additionally, they show that there is usually one male offspring for every mother mite, so mostly female offspring will not survive in overcrowded brood. This is likely because of competition at the feeding site (Martin, 1995). Therefore, we determine the female offspring survival probability (P_s) per brood cell:

$$P_s = \begin{cases} 0 & f > max - m\\ 1 - \frac{max - m}{f} & f \le max - m \end{cases},$$
(2)

where (*m*) is the number of male offspring, (*f*) the number of female offspring, and (*max*) the maximum number of offspring in that type of brood.

6. Mortality. In our model, we expect 0.5% of varroa to die every day, which is the average between the summer and winter mortality used by Fries et al. (1994). Additionally, we remove varroa who have gone through their final reproduction cycle, after which they are assumed to die (Martin & Kemp, 1997).

2.3. Gene drive implementation

Although AlphaSimR was designed to model large numbers of loci for breeding and quantitative genetics, the framework is perfect for the single locus of a gene drive too. Each individual is modelled with a single gene drive locus on two chromosomes and inheritance is random.

We have implemented a gene drive which homes in the germline and has four potential alleles: wild-type, gene drive, resistance, and non-functional. Like Prowse et al. (2017), we model a probability of cutting (P_C) of 0.95, a probability of non-homologous end joining (P_{NHEJ}), which is variable, a probability that non-functional repair occurs (P_{NFR}) of 0.67, which is the probability of a frame-shift occurring.

3. RESULTS

3.1. Development of a genetic population model of *Varroa destructor*

We first created a realistic, stochastic, population model of *Varroa destructor* that includes genetic inheritance. For an overview and description of the model and life history parameters, see Figure 1 and Methods. Our model has a population trajectory that is similar both in shape and amplitude to previous modelling (Fries et al., 1994; Calis et al., 1999; Martin, 1998) and empirical studies (De Guzman et al., 2007) (Figure 2A). The model begins on day 1 of the calendar year, a period of low or no growth for temperate populations. The population steadily declines due to daily mortality. By the summer, the varroa population grows exponentially. The starting population of varroa greatly influences the speed with which varroa reach threshold levels within a colony. With 100, 10, or 1 initial varroa, it, respectively, takes one, two, or three years longer for the population to reach the threshold of 10,000 individuals where we stop our model. The level of varroa infestation at which beekeepers will typically treat colonies is reached a year earlier. With 1 initial varroa, this single varroa often dies in the winter and therefore, the population grows in only a small number of replicates. Importantly, we observe more variability in models that begin with fewer varroa. This variability is caused by the timing of reproduction of few varroa, where small initial differences will grow bigger with the exponential growth.

We were also able to quantify the seasonal fluctuations in inbreeding in our modelled population (Figure 2B). We estimated the mean homozygosity at 1000 bi-allelic loci (with an initial average allele frequency of 0.5) across a single recombining chromosome. We began each model with a mean homozygosity at the beginning of the year of 0.95 in line with previous estimates for varroa (Beaurepaire et al., 2017). We found that homozygosity remains high throughout most of the beekeeping season but there are pronounced drops in homozygosity during the end of a typical year. This represents a period of time when honey bee colonies are reducing brood production and varroa populations are typically high. This combination increases the amount of mated varroa sharing cells, increases the chance of their offspring outbreeding, and thus reduces homozygosity. Overall, our model is qualitatively similar to expectations for a typical varroa population in a managed honey bee colony living in a temperate climate.

3.2. Inbreeding hinders gene drive spread and a fitness-affecting gene drive cannot spread

We model the release of 1 homozygous gene drive carrying varroa into a population

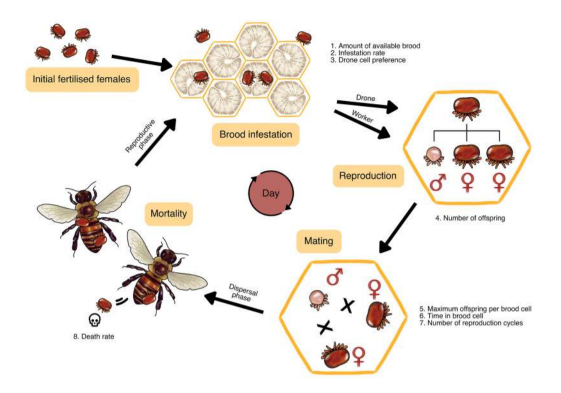


Figure 1. An overview of our varroa demographic model. For full details, see the Methods section. First, we initialize a certain number of fertilized females. Then, we use a backbone model of an average honey bee colony in a temperate climate where a certain amount of new brood cells become available for varroa infestation every day. The varroa infest these cells at a certain rate depending on the number of brood cells and adult bees. Varroa prefer drone cells over worker cells, because those are capped for 2 days longer (14 instead of 12 days), which enables more varroa off-spring to mature. Once in the cell, the fertilized females lay 1 male offspring followed by a varying number of female offspring. Once the females mature, they mate with the male. We assign each female a certain number of reproduction cycles, so one varroa female can infest brood cells multiple times throughout her life. Then, the fully grown bee emerges from the cell with the varroa attached to them, which is the start of the varroa's dispersal phase. At this stage we model a certain mortality rate which accounts for all ways in which a varroa could have died during its life cycle.

of 10 wild-type varroa (gene drive frequency of 0.09), which is relatively high for a nonthreshold dependent gene drive (Prowse et al., 2017; de Jong, 2017). We then track the genotypes and allele frequencies of individual varroa in a single honey bee colony (Figure 2C, D). As can be seen in both plots, the wild-type allele and wild-type genotypes remain the most prevalent even if we allow the model to continue to a population size of 10,000 varroa mites, greatly exceeding population sizes observed in typical colonies (Gatien, 2003). Our model strongly suggests that typical gene drive release frequencies may not be sufficient to spread a gene drive in varroa. This is likely a result of inbreeding, given that gene drive homozygotes are more prevalent than gene drive heterozygotes over the course of the simulation (Figure 2C). As well, gene drive alleles only meaningfully increase in the last days of the model when varroa numbers are high and cell sharing increases. The dynamics described above are consistent even when increasing the initial population size and released gene drive individuals (Figure S1). We found that our model

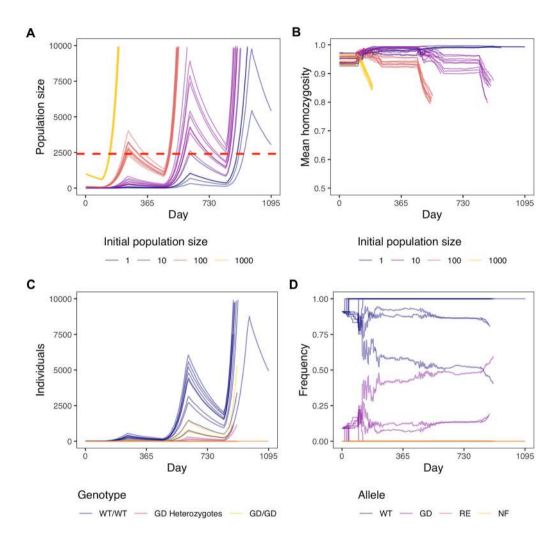


Figure 2. Model of varroa and gene drive spread. For every set of parameters, we run 10 repetitions and stop the model when the varroa population size is over 10,000. **A**) Population size over three years with different initial population sizes. The dashed red line indicates a varroa prevalence of 5% in summer (5 varroa per 100 adult bees), which is used by beekeepers as a "danger threshold" where treatment is necessary for bee colony health. **B**) Mean homozygosity over three years with different initial population sizes. We model a single chromosome with 1000 bi-allelic loci, each with initial average frequency of 0.5. We initiate individuals at 95% homozygosity because varroa have very high inbreeding coefficients of 0.9. **C**) Numbers of individuals with three genotypes over three years: WT = wild-type and GD = gene drive. The initial population size was 10 wild-type varroa with 1 added homozygous gene drive varroa. **D**) Frequencies of gene drive alleles over three years: WT = wild-type, GD = gene drive, RE = resistant, and NF = non-functional. The initial population size was 10 wild-type varroa with 1 added homozygous gene drive varroa, giving an initial gene drive frequency of 0.09.

is not sensitive to parameters influencing the spread of gene drive alleles (Figure S2). In the context of population control, the goal of a gene drive is to reduce population sizes by spreading alleles that reduce fitness. We could not conceive a model that successfully spread a male- or female-specific fitness-reducing drive (Figure S3).

3.3. With high introduction frequencies, a gene drive approaches fixation

When varroa numbers are still low at the start of the year, it is possible to introduce a larger amount of gene drive varroa to immediately obtain a high gene drive allele frequency. More importantly, this higher gene drive allele frequency could ensure that whenever outbreeding occurs, a gene drive varroa is likely involved. Therefore, we modelled a population of 10 wild-type varroa with either 1, 10, or 50 added homozygous gene drive varroa. These amounts, respectively, give initial gene drive frequencies of 0.09, 0.50, and 0.83. We find that the gene drive allele increases most rapidly at an initial release frequency of 0.5, because an outbreeding event is most likely between a gene drive varroa and a wild-type varroa, rather than between two wild-types or between two gene drives (see Figure 3 and S4). Naturally, a high initial gene drive frequency results in the highest gene drive allele frequency in the end. Therefore, a high initial release frequency might be beneficial to spread a gene drive through a varroa population. Unfortunately, we also see that with an initial amount of 50 gene drive varroa, the population reaches 10,000 individuals a year sooner than with 1 or 10 added varroa (see Figure 3).

3.4. Brood breaks increase outbreeding, but do not meaningfully increase the spread of a gene drive

Above, we demonstrate that outbreeding can be impacted by the initial release frequency of gene drive varroa. Ultimately, the amount of cell sharing, and thus outbreeding, depends on three factors: the amount of varroa, the amount of available brood, and the amount of adult honey bees (Boot et al., 1994). Therefore, decreasing the number of available honey bee brood cells can increase outbreeding frequency. Cell availability typically decreases naturally at the end of a beekeeping season when honey bees reduce egg laying. Beekeepers can also artificially change cell availability by preventing or restricting queens from laying eggs, a period called a 'brood break' (Calderone, 2005).

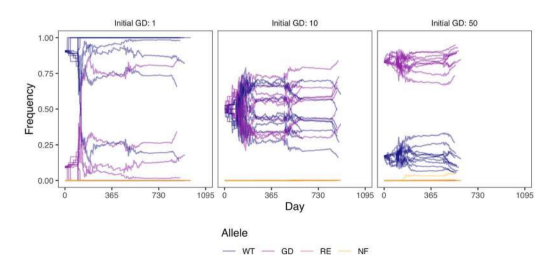


Figure 3. Allele frequencies over three years with different gene drive introductions. The initial population size is 10 wild-type varroa with 1, 10, or 50 added homozygous gene drive varroa, giving respective initial gene drive frequencies of 0.09, 0.50, and 0.83. WT = wild-type, GD = gene drive, RE = resistant, and NF = non-functional. For every set of parameters, we run 10 repetitions and stop the model when the varroa population size is over 10,000.

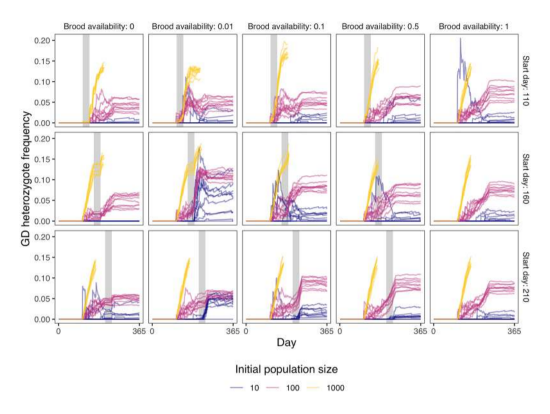


Figure 4. Gene drive (GD) heterozygote frequency over time for different initial population sizes, given different amounts of brood cell availability (as a fraction of the normal amount) and different brood break starting days. The grey bars indicate the brood break. The initial population sizes were 10, 100, or 1000 wild-type varroa with the same number of gene drive varroa on top of that, giving an initial gene drive frequency of 0.5. For every set of parameters, we run 10 repetitions and stop the model when the varroa population size is over 10,000.

We tested two brood break strategies for their effectiveness at increasing outbreeding and the fixation rate of gene drive alleles. For the first strategy we entirely stopped brood production, forcing varroa to stay in the dispersal phase (left-most column in Figure 4). After this brood break, varroa would more likely infest newly available brood with multiple varroa per cell. For the second strategy, we provided a steady but lowered amount of brood throughout the brood break (middle three columns in Figure 4). We also modelled no brood break intervention as a control (right-most column in Figure 4). For each of these strategies, we modelled three different brood break starting days: 110 (early season, when brood production is just starting), 160 (middle season, when brood production is at its maximum), and 210 (late season, just before brood production stops). Both strategies increased the amount of cell sharing (see Figure S6). However, only the strategy where a beekeeper adds in a specific proportion of brood during the break increased the frequency of heterozygous gene drive varroa in a colony relative to the control without brood break (see Figure 4). A brood break with a beekeeper allowing between 0.01 - 0.1 of available cells to be used for brood was the most effective. In practice, this equates to approximately one full frame in a ten-frame Langstoth colony. These results suggest that with some fine-tuning, outbreeding can be increased by the beekeeper and therefore increasing the likelihood of fixing a gene drive.

Gene drive allele frequency should increase after heterozygotes produce offspring, as gene drive homing will occur in these individuals. Thus, during a brood break, we first expect an increase in heterozygotes as outbreeding occurs, followed by an increase in gene drive allele frequency as these heterozygotes reproduce. However, we show in Figure S7 that there is only a modest increase in gene drive allele frequency after the brood break compared to no brood break. This is likely because of the low frequency of heterozygotes, which is lower than 0.2 as can be seen in Figure 4. In this model, we added the same amount of gene drive varroa as there are wild-type varroa, so the allele frequencies are both 0.5. As we showed in Figure 3, this ratio leads to the most rapid increase in gene drive allele frequency. Indeed, in Figure S8 where we model a larger gene drive introduction frequency, the frequency of gene drive heterozygotes is even lower. Despite the high introduction frequency and brood breaks, the gene drive is still not able to fix in the population (see Figure S9). These results show that brood breaks are unlikely to have a large effect on the spread of a gene drive.

3.5. Acaricide treatment may facilitate gene drive fixation

None of the scenarios we ran were able to fix a gene drive before varroa reached threshold levels within a honey bee colony. To that end, we incorporated an acaricide treatment into the model that would be activated anytime a colony reached threshold varroa levels (Figure 5). We found that effective acaricide treatments provide additional time for a gene drive to reach fixation. However, acaricide treatments significantly increase the variability between the model repetitions, which does not disappear when starting the model with a higher number of initial varroa (Figure S10). This means that the observed variability is due to the fact that, by chance, we could be removing more gene drive varroa than wild-types. Therefore, gene drive fixation is not reached very fast and not in all populations.

The best acaricide strategy for gene drive fixation was with 80% acaricide effectivity. With this effectivity varroa populations reach the treatment threshold multiple times within a single year and multiple acaricide treatments are necessary. These repeated relatively ineffective treatments are less prone to variability but probably not desirable in practice. We show that introducing more gene drive carriers after acaricide treatment facilitates faster gene drive fixation and less variability (see Figure S11). At this point gene drive fixation is probably due to population replacement rather than gene drive spread.

4. DISCUSSION

The greatest threat to managed honey bee colonies, globally, is the varroa mite (Kulhanek et al., 2017; vanEngelsdorp & Meixner, 2010; Molineri et al, 2018; Traynor et al., 2020). With the ever-advancing toolkit available to study functional genomics in varroa (Techer et al., 2019; Egekwu et al., 2018; Hasegawa et al., 2021), we suggest that the prospect of genetic control is not far from a reality. We set out to test the feasibility of such a system, in the form of a gene drive, in a modelling study of a population of varroa within a single honey bee colony. We demonstrate that a neutral gene drive could spread in a varroa population in a honey bee colony and open the door to future analysis in exploring how to spread gene drives in nonmodel species with particularly challenging biology.

A gene drive could work in varroa, but it is slow and requires management inputs. Our stochastic model tracked the growth of varroa mite populations each day over several years in a typical temperate honey bee colony. Varroa living in colonies in non-temperate climates will likely need additional modelling given the very different demography that honey bees have in these areas (Medina et al., 2002). We focused on temperate colonies, specifically, because they represent most managed colonies in the United States (Kulhanek et al., 2017) and because temperate climates provide an opportunity for increased outbreeding in varroa. Varroa populations tend to be highest in the fall (DeGrandi-Hoffman & Curry, 2004; Fries et al., 1994; Ifantidis, 1984). During this time, honey bee colonies decrease

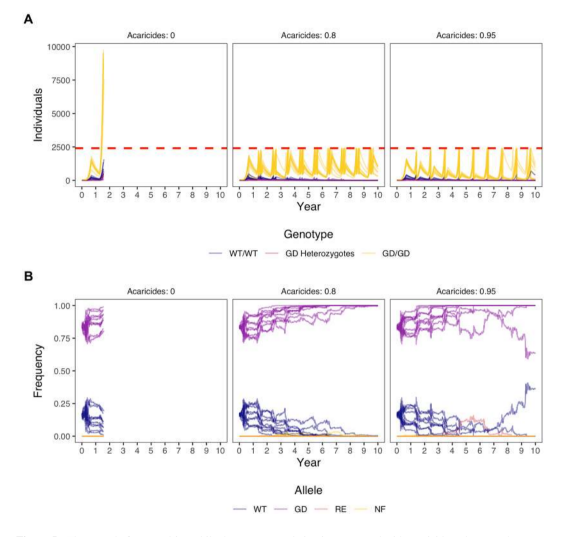


Figure 5. The spread of a gene drive while the varroa population is suppressed with acaricides whenever the varroa prevalence surpasses the danger threshold of 5% in summer (5 varroa per 100 adult bees). The initial population size was 10 wild-type varroa with 50 homozygous gene drive varroa, giving an initial gene drive frequency of 0.83. For every set of parameters, we run 10 repetitions and stop the model when the varroa population size is over 10,000. **A**) Frequencies of gene drive genotypes over time, given different intensities of acaricide treatment when the population surpasses the danger threshold. WT = wild-type, GD = gene drive. **B**) Frequencies of gene drive alleles over time, given different intensities of acaricide treatment when the population surpasses the danger threshold. WT = wild-type, GD = gene drive. **B**) Frequencies of gene drive alleles over time, given different intensities of acaricide treatment when the population surpasses the danger threshold. WT = wild-type, GD = gene drive. **B**) Frequencies of gene drive alleles over time, given different intensities of acaricide treatment when the population surpasses the danger threshold. WT = wild-type, GD = gene drive. **B**) Frequencies of gene drive alleles over time, given different intensities of acaricide treatment when the population surpasses the danger threshold. WT = wild-type, GD = gene drive, RE = resistant, and NF = non-functional.

brood production to prepare for the winter. As we observe and others have empirically demonstrated, varroa mites increase outbreeding rates in the fall because of reduced brood cell availability (Beaurepaire et al., 2017). Outbreeding is critical to the establishment of a varroa gene drive and indeed to any gene drive (Bull, 2016). We could not conceive a model that would successfully spread a lethal gene drive in varroa. The most promising way forward may be to design neutral drives with environmentally induced fitness effects (such as the spreading a toxin precursor), drives which remove acaricide resistance alleles, or drives that target genes involved in varroa-viral interactions. Each of these requires a deeper understanding of varroa functional genomics but may be fruitful for future investigations. Spreading drives that confer varroa with genetic resistance against viruses is a particularly interesting prospect. The threat that varroa mites pose to honey bees is exacerbated by the viruses they introduce into their hosts (Brettell & Martin, 2017; Barroso-Arévalo et al., 2019; Di Prisco et al., 2016).

There are several challenges to establishing a gene drive in varroa that need to be overcome. Natural outbreeding alone was not enough to reliably increase the frequency of gene drive. We attempted to overcome this challenge by incorporating beekeeper management in the form of brood breaks and acaricide treatments. Both influenced the rate of outbreeding and the likelihood of gene drive fixation. Importantly, both of these management practices are used by beekeepers and their incorporation into future gene drive efforts would not be an additional burden. The need for beekeeper management also suggests that a drive has a limited ability to spread beyond the apiary. All gene drive models we attempted faced the additional challenge of concomitantly minimizing population growth. When varroa populations exceed economic thresholds, honey bee colonies produce less honey and have a higher probability of collapsing (Currie & Gatien, 2006; Delaplane & Michael Hood, 1999). Here, we took a very generous threshold of 5 varroa per 100 bees across the year and ran simulations until varroa reached 10,000 mites in a single colony—a level that would almost never be observed in a managed colony. Furthermore, because varroa populations grow exponentially, a honey bee colony can only go without varroa control for a few years at most, depending on the initial infestation level. Controlling varroa growth with acaricides was an effective means to improve the spread of neutral gene drives by providing more time for the gene drives to fix before the honey bee colony reached 10,000 varroa. However, this method in itself is troubling because it does not remove the risk of varroa populations evolving acaricide resistance nor does it remove the risk that some acaricides pose to honey bees. We feel that the addition of management scenarios in our models and others (Lester et al., 2020) is particularly important for the gene drive literature and a feature that could be overlooked. Incorporating the typical management practices into models and understanding how they impact gene drive dynamics may be an important addition to future work.

In summary, our models provide an early look at how gene drives may act in the varroa system. They are by no means comprehensive. Varroa occupy a huge range and experience different colony and apiary environments across it. Location- or management-specific models may reveal that gene drives spread more or less successfully. The genetic background of a honey bee colony and a colony's response to increasing varroa loads were also not modelled. Both could impact the spread of a gene drive. The population dynamics for varroa in varroa-tolerant or resistant colonies is likely different and could impact the spread of a gene drive, perhaps acting like acaricide treatments and providing a longer time for gene drives to spread. Any colony-level responses to increased levels of varroa parasitism could increase or decrease the likelihood of a drive spreading. We also did not explore dynamics outside of a single honey bee colony and did not explore the risks of modified varroa establishing in non-target colonies. Varroa mites are as highly mobile as honey bees and more modelling is necessary to understand the roles of drifting, foraging, robbing, and management in spreading gene drives outside of target colonies (Goodwin et al., 2006; Peck et al., 2016; Peck & Seeley, 2019; Seeley & Smith, 2015). We suggest, given the difficulty we found in spreading drives in a single colony, that the above factors may be unlikely to establish drives in non-target colonies. Even if they could establish outside of target colonies, the spread of gene drive varroa may not be viewed as a major threat, at least in North America. This may not be the case in other parts of its introduced range. In its native range, Varroa destructor can be found in low frequency

in *Apis cerana* colonies where we have little information about its native ecology.

To our knowledge, genetic modification has not been performed in varroa mites and in vitro rearing methods are, so far, unable to maintain a breeding population of varroa (Egekwu et al., 2018). Mutagenesis in chelicerates has recently been accomplished (Dermauw et al., 2020) but transgenesis has yet to be achieved. Gene drives may be many years off for varroa. With more expertise developing in the fields of transgenesis and mutagenesis in arthropods, it is likely that we will see experiments in the varroa system and we hope that our work can help develop ideas about genetic control of this invasive pest species. In the short-term, currently available treatment methods (Currie & Gatien, 2006) and perhaps newer methods (Huang et al., 2019; Leonard et al., 2020) remain the best methods to control varroa.

SUPPLEMENTARY INFORMATION

The online version contains supplementary material available at https://doi.org/10.1007/s13592-021-00891-5.

AUTHOR CONTRIBUTIONS

G.G. and B.A.H. conceived the varroa gene drive project. N.R.F. conducted the modelling with assistance from A.B.M. and G.G. B.A.H. guided the varroa life history aspects and G.R.M. and N.R.F. guided the gene drive aspects of the project. N.R.F. and B.A.H. wrote the manuscript and all authors reviewed it.

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DATA AVAILABILITY

Our generated data and plots can be found on the HighlanderLab GitHub: https://github.com/HighlanderLab/ nfaber_varroa_gd.

CODE AVAILABILITY

Our model code can be found on the HighlanderLab GitHub: https://github.com/HighlanderLab/nfaber_varroa_gd.

DECLARATIONS

Ethical Approval Not applicable.

Consent to Participate Not applicable.

Consent for Publication Not applicable.

Conflict of Interests The authors declare no competing interests.

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