# Modeling the impact of rainfall and temperature on sterile insect control strategies in a Tropical environment

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#### Abstract

The sterile insect technique (SIT) is a biological control technique that can be used either to eliminate or decay a wild mosquito population under a given threshold to reduce the nuisance or the epidemiological risk. In this work, we propose a model using a differential system that takes into account the variations of rainfall and temperature over time and study their impacts on sterile males releases strategies. Our model is as simple as possible to avoid complexity while being able to capture the temporal variations of an Aedes albopictus mosquito population in a domain treated by SIT, located in Réunion island. The main objective is to determine what period of the year is the most suitable to start a SIT control to minimize the duration of massive releases and the amount of sterile males to release, either to reduce the mosquito nuisance, or to reduce the epidemiological risk. Since sterilization is not 100% efficient, we also study the impact of different levels of residual fertility within the released sterile male population. Our study shows that rainfall plays a major role in the dynamics of the mosquito and the SIT control, that the best period to start a massive SIT treatment lasts from July to December, that residual fertility has to be as small as possible, and that increasing the size of the releases is not always necessarily interesting. We also highlight the importance of combining SIT with mechanical control, i.e. the removal of breeding sites, in particular when the initial mosquito population is large.

**Keywords:** Vector control; *Aedes spp*; Sterile Insect Technique; Temperature; Rainfall; Residual fertility; Nuisance reduction; Epidemiological risk; Mathematical modeling; Numerical simulation.

### 1 Introduction

Being vectors of many diseases, like malaria, dengue, Lymphatic filariasis, zika, chikungunya, yellow fever, and japanese encephalitis, mosquitoes are one of the biggest killers in the world, and particularly in tropical and subtropical areas. Indeed, Female mosquitoes need blood meals to provide the nutriments for egg development. That is why they bite either during the night, like anopheles spp, at sunset or sunrise, like aedes albopictus, or along the day, like aedes aegypti, inside or outside houses. Since they can survive several weeks, they will bite several times depending on their gonotrophic cycle [14]. While opportunistic feeders, Anopheles and Aedes mosquitoes prefer biting mammals and, preferably, humans [25], with sometimes multiple blood feeding [8].

To eradicate mosquito-borne diseases, the initial option was the massive use of chemicals in the 1950's and 1960's, mainly against anopheles. Despite some successes, like in Réunion island where malaria has been eradicated since the fifties, thanks to the use of DDT [13], we know now that this was a huge mistake: even if the use of adulticides was successful against mosquitoes, the damages on the biodiversity were important. In addition, resistance to some adulticides raised, such that, in several places around the World, there is no option left to fight the mosquitoes, like in the French West Indies [17]. Fortunately, other eco-friendly control methods, more respectful of the biodiversity, have been developed. Among them, the Sterile Insect Technique (SIT) is the most promising one. This is an old control technique, proposed in the 30s and 40s by three key researchers in the USSR, Tanzania and the USA and, first, applied in the field in the 50's [11]. SIT has been used more or less successfully on the field against various kinds of Pests or Vectors (see [5, 11] for various examples). The classical SIT consists of mass releases of males sterilized by ionizing radiation. The released sterile males transfer

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their sterile sperms to wild females, which will have no viable offspring, resulting in a progressive decay of the targeted population. For mosquitoes, other sterilization techniques have been developed using either genetics (release of insects carrying a dominant lethal technique, in short, RIDL technique [24]) or bacteria (SIT-IIT: cytoplasmic incompatibility thanks to Wolbachia bacteria) [23]. Many SIT projects against mosquitoes are now ongoing around the World. It is important to keep in mind that controlling a vector-population could have an impact on the ecosystem equilibrium. In fact, it depends if the targeted vector is an invasive species, i.e. established recently, of if it is established for a long time. If the establishment is recent (a couple of years), then with elimination, we can expect to recover the initial state of the ecosystem, before the invasion starts. In contrary when the vector is established, then elimination is not necessarily the goal. Reduction is preferable in order to reduce the epidemiological risk. We will explore these two objectives in this work.

Several mosquito models have been developed, from very simple models to more sophisticated ones, according to the number of biological states that are taken into account. These models consider the mosquito population either at the individual level (see for instance [2]) or at the population level (see for instance [16]), taking eventually into account the spatial component (see for instance [3] for a brief review). Each of these approaches has advantages and disadvantages. However, it is always important to keep in mind that biological parameters are difficult to obtain in the field, such that for too complex models, it might be impossible to set the (right) values of the parameters. In addition, most of the biological parameters are linked to environmental parameters, like rainfall, humidity and temperature. The main goal of the present article will be to take into account these parameters in the mathematical model to find the best control.

Mathematically, it can make sense to consider constant or periodic parameters to derive the main dynamics of the mosquito population. However, for practical reasons, i.e. anticipating when the mosquito population is growing or decaying, it is better to have a mosquito model that takes into account the most important parameters, like rainfall, humidity and temperature, to adapt the control, i.e. for instance the size of the releases or/and the period of the releases.

Since our work takes place within an ongoing SIT feasibility program (TIS 2B) against *Aedes albopictus* in La Réunion, a tropical French island located in the Indian Ocean, we will focus on this species in the rest of the paper. In La Réunion, *Aedes albopictus* has become the main vector of dengue and chikungunya [27]. However, we believe that our approach is sufficiently generic to be applied to other mosquito species.

While links between Aedes albopictus parameters and temperature have been studied in laboratory [9], for a fixed (laboratory) humidity, it is more challenging to take into account the effect of rainfall (and humidity).

Some weather-dependent mosquito models have been developed, mainly with Temperature-dependent parameters (see for instance [7,10] and references therein) and very few with temperature and rainfall-dependent parameters (see [26] and references therein). However, in general, these last models are quite complex: they relied on statistical approaches and on the user's subjective choices, such that the calibration (of many parameters), with respect to the environmental parameters, is not generic and might not be able to provide a unique set of valuable values. We firmly believe that simple (but not too simple) models can rapidly provide useful and reliable information to help field experts to manage vector control campaigns.

That is why we consider a minimalistic model (minimal in terms of stages and thus parameters), based on [4], to build an approximation of the mosquito population over many years. Then, we show that the density variation (in time) of the mosquito population can help to improve vector-control strategies combining SIT with mechanical control, i.e. the removal of breeding sites to reduce the larvae carrying capacity.

It is well known that sterilization does not necessarily induce that the sperm of the sterilized males is 100% sterile (see for instance [20, 34]): it depends on the radiation source, the dose-rate, and the container. That is why quality control after sterilization is set up to control that the sterile males remain competitive and efficient. Efficiency is thus related to the effective sterility or the residual fertility (RF, shortly), i.e. the percentage of sperms that is still fertile despite the irradiation, that is equivalent to say, at the population level, that a small proportion,  $\varepsilon$ , of the sterile males is fertile. In our model, we will take into account the fact that residual fertility can occur within the sterile male population to study its impact on SIT control.

In general, in most of the SIT models (except in [5] and the present paper), residual fertility is never taken into account, while it can have a strong impact on SIT efficiency. Indeed, in [5], using a 2-dimensional mosquito model with constant parameters, the authors showed that the residual fertility,  $\varepsilon$ , is strongly linked to the basic offspring number,  $\mathcal{N}$ , of the wild population: when

$$\varepsilon < \frac{1}{N},$$
 (1)

then SIT is efficient and the mosquito population can be lowered under any given threshold, provided that enough sterile males are released. However, when  $\varepsilon$  is below but close to  $\frac{1}{\mathcal{N}}$ , the amount of sterile males increases almost exponentially [5]. On the contrary, when  $\varepsilon > \frac{1}{\mathcal{N}}$ , then, whatever the size of the sterile males releases, the wild population cannot be lowered under a certain threshold value, that can be roughly estimated [5]. In fact, we suspect that a too-large residual fertility could partly explain failures in some SIT programs. For instance,

in [34], a SIT program conducted on Mauritius island, which is close to Réunion island, the residual fertility was experimentally estimated at around 3.05% (2.29% - 3.92%), which could explain why mitigated results were obtained, before a cyclone occurred and broke the experiment. We believe that 3% of RF is too large such that the reduction of the wild population with SIT is limited. Indeed, in [5], the authors showed that if RF is too large, then, whatever the size of the sterile males releases, the population cannot be lowered under a given threshold.

The paper is organized as follows: In section 2, we build a temperature and rainfall-dependent entomological model and an impulsive periodic SIT model. Then, in section 3, we provide several numerical simulations to discuss the impact of the temperature, the rainfall, the residual fertility and the mechanical control on SIT starting period and duration to reduce either the nuisance or the epidemiological. Finally, in section 4, we end the paper with some conclusions and perspectives.

### 2 Rainfall, humidity, and temperature dependent SIT model

The first aim of the present work is to develop a temperature-rainfall-dependent entomological model, to take into account real field data, including mean daily rainfall and mean daily temperature.

Following [4], we will consider the following model

$$\begin{cases}
\frac{dA}{dt} = \phi(Temp)F - (\gamma(Temp) + \mu_{1,A}(Temp) + \mu_{A,2}(Temp, Rain)A) A, \\
\frac{dM}{dt} = (1 - r(Temp))\gamma(Temp)A - \mu_{M}(Temp)M, \\
\frac{dF}{dt} = r(Temp)\gamma(Temp)A - \mu_{F}(Temp)F,
\end{cases}$$
(2)

where A, M, and F represent, respectively the aquatic (larvae, pupae) stage, the adult (male and mated and fertile female) stages. Here, we implicitly assume that all emerging females will mate and become fertile. Note carefully that compartment A does not include the eggs since we consider a density mortality rate  $\mu_{A,2}$  that concerned only the larval and pupal stages. Thus, the parameter  $\phi$  represents the daily amount of deposited eggs that hatch to become larvae. All parameters of model (2) are described in Table 2. This model has been studied

Table 1: Description of the entomological parameters

Symbol	Description	Unit
$\phi$	Number of hatched eggs at each deposit per capita	$\mathrm{Day}^{-1}$
$\gamma$	Maturation rate from larvae to adult	$\mathrm{Day}^{-1}$
$\mu_{A,1}$	Density independent mortality rate of the aquatic stage	$\mathrm{Day}^{-1}$
$\mu_{A,2}$	Density dependent mortality rate of the aquatic stage	Day <sup>-1</sup> Individuals <sup>-1</sup>
r	Sex ratio	-
$1/\nu_m$	Average extrinsic incubation period (EIP)	Day
$1/\mu_F$	Average lifespan of fertilized and eggs-laying females	Day
$1/\mu_M$	Average lifespan of males	Day

in [4] when the parameters are supposed to be constant. In [4]the authors developed a new strategy to maintain the wild population under a certain threshold, using a permanent and sustainable low level of SIT control, thanks to a massive-small releases strategy. We set  $\mathcal{N}_{\max} = \max_{t \in [0, +\infty)} \mathcal{N}(t)$ , where  $\mathcal{N}$  represents the basic offspring number, defined as follows

$$\mathcal{N} = \frac{r\phi\gamma}{(\gamma + \mu_{A,1})\,\mu_F}.$$

In Réunion island, a network of Weather stations (from Météo France but also from CIRAD) allows us to estimate some weather parameters all around the island, and in particular where real SIT field experiments started in July 2021, in the site of Duparc, a 20-hectare urban area located within the commune of Sainte Marie in the northern district of La Réunion [15]. These releases consist of manually weekly releases of 150 000 to 250 000 sterile males (once a week) produced and irradiated in La Réunion. The efficacy of sterile males releases is assessed over time by monitoring the induced sterility in eggs using ovitraps and the subsequent population suppression using adult traps, and, from time to time, Mark-Release-Recapture experiments. This experiment lasted until September 2022.

From the Weather station located in La Mare, close to the site of Duparc (1km), we are able to obtain the following daily data: the rainfall, the average daily temperature and also the humidity. Thus, following [29], we first define the breeding site carrying capacity to define the density death-rate  $\mu_{A,2}$ . Obviously, the persistence of breeding sites is a key factor for the mosquito population's survival. Indeed, rainfall creates breeding sites, while evapotranspiration tends to shrink them. Following [29], we define the variable H(t) as the amount of water available at day t and defined as follows

$$H(t+1) = \begin{cases} 0 & \text{if } H(t) + \Delta(t) \leq 0, \\ H_{max} & \text{if } H(t) + \Delta(t) \geq H_{max}, \\ H(t) + \Delta(t) & \text{otherwise,} \end{cases}$$

with

$$H_{\max} := \max_{t \in [t_0, t_0 + T]} \{Rain(t)\}$$

and

$$\Delta(t) := Rain(t) - Evap(t),$$

where Rain(t) is the daily rainfall and Evap(t) the daily evaporation,  $t_0$  is the starting date of the simulations, and T the total duration. Following [29], the evaporation function is defined as follows

$$Evap(t) = k \times (25 + Temp(t)^{2}) \times (100 - Hum(t)),$$

where Temp(t) is the average temperature and Hum(t) the humidity. Finally, the carrying capacity is defined

$$K(t) = K_{max} \frac{H(t)}{H_{max}} + K_0,$$

where  $K_0 > 0$  can be seen as the fixed artificial carrying capacity, i.e. rainfall-independent, human-made, by watering, for instance, plants around houses (flower pots, plates, and vases), and  $K_{\text{max}}$ , the natural maximal carrying capacity.

The initialization in time of the variables (in particular of H) is explained in the simulations section, page 6. We notice that, in general, in the literature, the question of initialization is not always taken into account, while it can have a strong influence over several months on the population dynamic. We illustrate this fact in Figure 2, page 7.

In order to estimate  $\mu_{2,A}$ , we consider the positive equilibrium related to the carrying capacity, like in [10, 30, 31]. Thus, for a fixed value of K, the aquatic stage at equilibrium is given by

$$A_K^* = \left(1 - \frac{1}{\mathcal{N}}\right) K. \tag{3}$$

In our model, the aquatic stage at equilibrium is defined as follows

$$A^* = \frac{\gamma + \mu_{A,1}}{\mu_{A,2}} \left( \mathcal{N} - 1 \right), \tag{4}$$

such that considering the equality between both equilibria given in (3) and (4), we derive the following relationship between  $\mu_{A,2}$  and K, that is

$$\mu_{A,2} = \frac{\mathcal{N}(\gamma + \mu_{A,1})}{K} = \frac{r\gamma\phi}{\mu_F K}.$$

Thus, taking into account the dependency of the parameters to Temperature and Rainfall, we deduce that

$$\mu_{A,2}(Temp, Rain) = \frac{r(Temp)\gamma(Temp)\phi(Temp)}{\mu_F(Temp)K(Temp, Rain)}.$$
 (5)

As it is well known, eggs are deposited above the waterline. They hatch once they are flooded by rainfall. However, the hatching rate can be seasonal. From A. albopictus eggs, picked up at Duparc, the mean hatching proportion is around 90% (G. Legoff, personal communication, TIS2B project).

For the other parameters, we will consider the data recalled in Table 9, page 18, as used in [31] to obtain the parameters estimates given in Table 10, page 19. See [31] for some explanations on the formulas.

Then, we can use simple interpolation polynomials (with cubic spline, like in [32]) to estimate these parameters for any given temperature Temp, within the range  $[15^{\circ}, 35^{\circ}]$ .

We will assume constant or periodic releases of sterile males, at rate  $u_S(t)$  (either constant or variable), and  $\mu_S$ , the sterile male mortality rate, which is supposed to be similar to the wild males mortality rate [33]. The dynamic of the sterile males is modeled by

$$\frac{dM_S}{dt} = u_S(t) - \mu_S(Temp)M_S. \tag{6}$$

Thus, based on model (2), we consider the following temperature and rainfall dependent SIT-model

$$\begin{cases}
\frac{dA}{dt} = \phi(Temp)F - (\gamma(T) + \mu_{1,A}(T) + \mu_{A,2}(T,R)A) A, \\
\frac{dM}{dt} = (1 - r(T))\gamma(T)A - \mu_{M}(T)M, \\
\frac{dF}{dt} = r(T)\gamma(T)\frac{M + \varepsilon\beta M_{S}}{M + \beta M_{S}}A - \mu_{F}(T)F, \\
\frac{dM_{S}}{dt} = u_{S}(t) - \mu_{S}(Temp)M_{S}.
\end{cases} (7)$$

The parameter  $\beta$  is the competition parameter that is deduced from the Fried index [12]. Following [20], we will consider  $\beta=1$ , which means that a sterile male is as competitive as a wild male. The Fried index is estimated through a protocol described in [12] (see also [28]). Last, as explained in the introduction, all irradiated males are not 100% sterile: a small proportion of their sperm can stay fertile. At the population scale, this is modeled by the parameter  $\varepsilon$ , which represents the proportion of sterile males that remain fertile and thus can fertilize emerging females. As already explained in [39][Remark 4], A. albopictus female's got 3 spermathecae that allow to stock sperms such that, in general, one mating is sufficient to fulfill, at least 2, spermathecae and to have enough sperms to fertilize eggs along its lifespan. Thus the nonlinear term related to SIT treatment occurs when females emerge from compartment A and not in the birth rate. Considering SIT impact on the birth-rate, i.e.  $\frac{M+\varepsilon\beta M}{M+\beta M}\phi$ , is very convenient from the mathematical point of view but it is not biologically realistic. We prefer

to stick to the biological reality, at least for A. albopictus. Thus, altogether, in model (7), the term  $\frac{M + \varepsilon \beta M}{M + \beta M} \phi$  represents the probability for an emerging female to be fertilized and thus to enter compartment F.

#### 2.1 Impulsive SIT massive-small releases strategy

In the rest of this paper, we will consider the so-called massive-small releases strategy developed in [4]. Indeed, once a few sterile males are released, this induces a strong Allee effect, such that the elimination equilibrium,  $\mathbf{0}$ , and a positive equilibrium E are both asymptotically stable. In [4], we showed that it is possible to estimate the minimal amount of sterile males to release to stay in the basin of attraction of  $\mathbf{0}$  when the wild population is sufficiently small. This means that massive releases have first to be used over a sufficient time period in order to enter in a particular subset of the basin of attraction of  $\mathbf{0}$ , that is estimated thanks to a given (small) release rate value. Once inside the basin of attraction of  $\mathbf{0}$ , it is possible to switch from massive releases to small releases. This strategy is particularly useful since SIT cannot be stopped in order to prevent the re-establishment of the targeted vector, either by introduction or by migration.

According to this massive-small releases strategy, we want to find the best period to start SIT control (without or with Mechanical Control) to minimize the number of massive periodic impulsive releases, and thus the amount of sterile males to release.

Thanks to [4], and assuming that  $N\varepsilon \leq 1$ , system (7) has the following long term behavior

- There exists a release rate threshold,  $u^*$ , such that when  $u_S(t) > u^*$  then (A, M, F) converges to 0.
- When  $0 < u_S(t) < u^*$ , then there exist two positive equilibria,  $\mathbf{E_1}$  and  $\mathbf{E_2}$  such that  $\mathbf{E_1} < \mathbf{E_2}$ . Moreover  $\mathbf{E_1}$  is unstable while  $\mathbf{E_2}$  is stable and  $[\mathbf{0}, \mathbf{E_1})$  lies in the basin of attraction of  $\mathbf{0}$ .

We will consider two different levels of massive releases (6000 or 12000 sterile males per ha), such that  $u > u^*$  for all  $t \ge 0$ , to decrease the wild population below a threshold set,  $[\mathbf{0}, \mathbf{E_1})$ , defined by (inexpensive) small releases (say 100 sterile males per ha), in a minimum time. We want to know how the duration of massive releases is influenced by temperate and rainfall conditions during the year, the residual fertility and the level of mechanical control. In other words, we search a control with the form

$$u(t) = \tau \Lambda_{\text{massive}} \sum_{i=1}^{N} \delta_{t_0 + (i-1)\tau}(t), \tag{8}$$

where  $\delta(t)$  is the Dirac function,  $t_0$  is the starting time of the massive releases,  $\tau$  the periodicity of the releases (here,  $\tau=7$ ), and N the number of weakly massive releases. The massive release  $\Lambda_{\rm massive}$  and the small releases  $\Lambda_{\rm small}$  are fixed. For a given  $t_0$  there exists a  $t_1(t_0)=t_0+N_1(t_0)\tau$  (according to [4]) such that, after this time, the wild population (A,M,F) remains in the box  $[\mathbf{0},\mathbf{E_{1,min}}(\tau\Lambda_{\rm small}))$  where  $\mathbf{E_{1,min}}(\tau\Lambda_{\rm small})$  is defined as follow: For a given  $\Lambda_{\rm small}$ , for each time t, we compute the equilibrium  $\mathbf{E_1}(\mathbf{t})$  of the system associated to the parameters at time t, and we define  $\mathbf{E_{1,min}}(\tau\Lambda_{\rm small})=\min_{\mathbf{t}}\mathbf{E_1}(\mathbf{t})$ , where the minimum is taken between the beginning and the end of the time interval considered in the simulation.

The main goal is to find the (best) starting time of the massive releases,  $t_0$ , in order to minimize the duration of the releases and thus, the number of massive releases.

System (6)-(8) can be rewritten as an impulsive differential system with fixed moments of impulse effect, that

is

$$\begin{cases}
\frac{dM_S}{dt} = -\mu_S(T)M_S, & t_0 + i\tau < t \le t_0 + (i+1)\tau \\
M(t^+) = M(t) + \tau \Lambda_{massive}, & t = t_0 + i\tau,
\end{cases}$$
(9)

for i=0,...,N-1. Since, the right-hand side of (7)-(9)<sub>1</sub> is locally Lipschitz-continuous on  $\mathbb{R}^4$ , we can use a classical existence Theorem (for instance Theorem 1.1 in [36], or Theorem 2.1 in [35]), to deduce that there exists  $T_e>0$  and a unique solution of system (7)-(9), defined from  $(t_0,T_e)\longrightarrow\mathbb{R}^4$ .

#### 3 Numerical simulations and discussion

System (7)-(9) is solved thanks to odeint of the python library scipy.integrate. The codes and the data corresponding to the temperature and the rainfall in Réunion island are available on

https://github.com/michelduprez/Impact-of

-Rainfall-and-Temperature-on-IT-control-strategies.git

We consider temperature, rainfall and humidity data based on (noised) data recorded in weather stations close to the site of Duparc (a 20 ha place), a neighborhood of Sainte Marie, located in the North of Réunion island from the 1st of January 2009 to the 14th of July 2021.

In La Réunion, since the sterile males are produced on-site, everything from the eggs to the sterilization is controlled (qualitatively), such that since the production started, the residual fertility is, in general, less than 1%, with an average value of around 0.6%, a very good results compared to other SIT projects, like [34]. That is why, in the forthcoming simulations, we will consider three cases of RF, namely 0%, 0.6% and 1.2%.

In Fig. 1, we consider the following values:  $K_{\text{max}} = 20 \times 10000$ ,  $K_0 = 20 \times 100$  to derive the carrying capacity, K, and thus  $\mu_{A,2}$ .

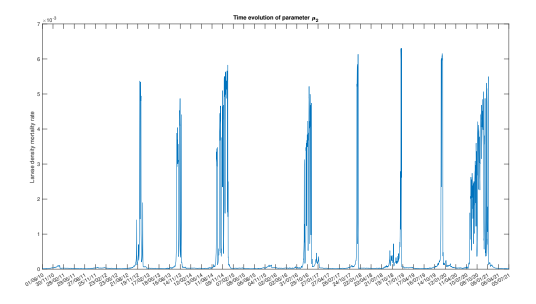


Figure 1: Time evolution of  $\mu_{A,2}$  from the 1st of September 2010 until the 14th of July 2021

It is interesting to see the behavior of  $\mu_{A,2}$  in the second-half of 2020, which was a particularly dry period compared to previous years in La Réunion. Note also that when mechanical control is considered, it will impact  $\mu_{A,2}$ : for instance, thanks to Formula (5), 40% of Mechanical control increases  $\mu_{A,2}$  by 66.7%.

The choice of the initial condition for H and the initial conditions for the mosquitoes will impact respectively the initialization of the carrying capacity and the dynamic of the system for at most 12 months (see Fig. 2, page 7, where we have considered extreme initial values for H(0) and for the population). That is why, in order to have a reliable estimate of the wild mosquito population at the beginning of the control, i.e. the 1st of Sept. 2010, we start the simulations at the beginning of January 2009.

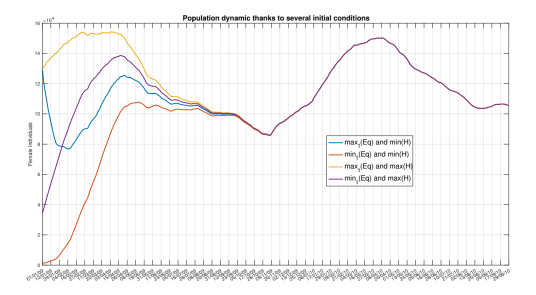


Figure 2: Temperature and rainfall dependent model - Simulations of the mosquito dynamics with several initial conditions, thanks to the initial rainfall data and the initial size of the mosquito population

Using the parameters values and the estimates of  $\mu_{A,2}$  given in Fig. 1, page 6, we derive the dynamic of the mosquito population in Duparc without release: see Fig. 3(c), page 22. In Fig. 3(a)-(b), page 22, we also show the time evolution if the mean daily Temperature and the mean daily rainfall (mm). As expected, periods, where the rainfall is low, imply a rapid decay of the population size, leading to an "almost" oscillatory behavior. This result is confirmed by Mark-Release-Recapture experiments derived in Duparc [15], where the ratio between the mosquito density/ha within the dry period and the mosquito density/ha within the wet period is a factor 10, which highlights the importance to consider a temporal dynamic in the parameters.

#### 3.1 Global sensitivity analysis

Before starting the simulations, we provide a global sensitivity analysis (GSA) using Latin hypercube sampling (LHS) and Partial rank correlation coefficient(PRCC), within the range of values for the temperature and rainfall given in Fig. 3(a)-(b), page 22. The main objective of this GSA is to identify key parameters that most drive the variables and the dynamics of our system. Briefly, the objective of the LHS-PRCC sensitivity analysis is to identify key parameters whose uncertainties contribute to prediction imprecision and to rank these parameters by their importance in contributing to this imprecision. We refer the interested reader to [1, 18] for further information about LHS-PRCC analysis. The forthcoming LHS-PRCC figures are made using R [21] and R-studio [22].

In Fig. 4, page 23, the main sensitive parameters for all variables of model (2) are  $\mu_{A,2}$ ,  $\Phi$  and  $\nu_A$ , as expected. An important outcome is that since  $\mu_{A,2}$  is strongly related to the temperature and rainfall, mechanical control, i.e. the removal of breeding sites, is more than useful to increase  $\mu_{A,2}$ , and, thus, to impact (negatively) all compartments. This fact is also confirmed in the next LHS-PRCC analysis of the SIT model with continuous nand periodic releases: see Fig. 5, page 24, and 6, page 25. Also, as expected  $\Lambda_S$  and  $\beta$  are sensitive parameters in the SIT models. Contrary to the initial entomological model, the parameter  $\mu_F$  has become more sensitive in the SIT models. All these information show that a combined effort mixing SIT and mechanical control will have a strong impact on the wild population control. Last, if an additional control tool against females is able to increase  $\mu_F$ , using, for instance, killing-traps, the overall control will be even more efficient. Last but not least, for  $\tau$  varying from 1 (daily releases) to 14, its impact on the periodic SIT-model is negligible: what matter are the size of the releases and the competition parameters.

#### 3.2 Reducing mosquito nuisance

As studied in [4], we consider the massive-small strategy, which consists first of periodic massive releases, such that  $\tau \Lambda_{massive} = 20 \times 6000 \text{ or } 20 \times 12000 \text{ individuals, until the wild population has become lower than } \mathbf{E}_{1,\min}(\boldsymbol{\Lambda}_{small})$ ,

where here  $\tau \Lambda_{small} = 20 \times 100$  individuals, such that

$$\mathbf{E_{1,min}}(\boldsymbol{\Lambda_{small}}) \approx \left\{ \begin{array}{ll} (37.0, 29.8, 0.511) & \text{if } \varepsilon = 0.0, \\ (17.97, 14.47, 0.24) & \text{if } \varepsilon = 0.006, \\ (1.38, 0.350, 0.0059) & \text{if } \varepsilon = 0.012. \end{array} \right.$$

We notice that, when the residual fertility increases, then the size of the box  $[0, \mathbf{E_{1,min}}(\Lambda_{small})]$  becomes smaller: SIT introduces a strong Allee effect while the residual fertility weakens it. Thus, in the forthcoming simulations, it will not be surprising to find large duration time values when the residual fertility is large. Once  $[0, \mathbf{E_{1,min}}(\Lambda_{small})]$  is reached, the periodic releases continue at rate  $\Lambda_{small}$ . Using the numerical simulations, we are able to estimate the date where the system can switch from massive releases to small releases and thus evaluate the duration of the (very) massive releases.

According to the time variation of the parameters, the basic offspring number  $\mathcal{N}$  will vary between 29.7 and 85.7 with a mean of 48.41 over the considered period. Thus even if Formula (1) is not verified at each time, it is satisfied in mean, i.e.

$$\varepsilon < \frac{1}{T} \int_0^T \frac{1}{\mathcal{N}(t)} dt = 0.0222. \tag{10}$$

An important issue in SIT control is to estimate the duration of the massive releases. Thanks to contrasted environmental conditions, within the period [2010 - 2020], we show that the minimal time to decay the wild population under a certain threshold can greatly vary: see Figs. 8-9, from page 27 to page 28, for two weakly release rates (6000 Ind/ha and 12000 Ind/ha), for different levels of Mechanical Control (0%, 20% and 40%) and three levels of residual fertility (0%, 0.6% and 1.2%).

In general, in the literature, many SIT simulations are done with constant parameters, except, for instance, in [10,37]. If we consider the mean value for each parameter over the whole period, we derive  $\mathcal{N} \approx 49.3$  which is a quite large basic offspring number, but not surprising within a tropical context.

Before starting the simulations, we would like to emphasize that our meteorological data last from 01/01/2009 and 14/07/2021. Our simulations intend to derive an estimate of SIT treatment duration within this interval, when the SIT control starts 01/09/2010. Thus, the reader does not have to be surprised to see that our results curves do not end at the same time: this is simply because the simulations were not able to enter  $[0, \mathbf{E_{1.min}}(\Lambda_{small})]$  before the 14th of July 2021. Thus the simulation stops.

In the forthcoming simulations of model (7)-(9), we will consider four cases to illustrate the importance of considering all (or not) environmental parameters:

- 1. Model 1: model (7)-(9) with temperature and rainfall dependent parameters.
- 2. Model 2: model (7)-(9) with temperature-dependent parameters only.
- 3. Model 3: model (7)-(9) with a constant average temperature and rainfall-dependent parameters.
- 4. Model 4: model (7)-(9) with average parameters values estimated from September 2010 to mid-July 2021.

We first provide Fig. 7, page 26, where simulations of the four models are given for different values of RF, 0%, 0.6% and 1.2%, and without mechanical control. It is interesting to notice that model 1 and model 3 behave similarly more or less: this is due to the fact that rainfall is taken into account. For RF equal to 0% and 0.6%, model 3 is close to the variations of model 1. Of course, model 4 provides a constant value. Model 2, where the parameters are only temperature dependent, provide the worst estimates, except in the years where rainfall was abundant, i.e. 2010 and 2011. However, along the decade, except when RF is equal to 1.2%, model 2 is the worst one. We discuss and compare these four cases:

1. We now comment Fig. 8, page 27, and Fig. 9, page 28.

When mechanical control is considered, there is a gain in the duration of massive releases. However, the impact of mechanical control is more important in periods where the mosquito population is large, whatever the residual fertility: compare the years 2010-2011 with the years 2016, 2019, and 2020, in Fig. 8, page 27. As also showed in Fig. 8, page 27, the residual fertility is a very important parameter: if RF increases from 0% to 0.6%, this leads to an increase of the duration (of the massive releases) by 30%. Then, increasing RF from 0% to 1.2% increases the duration (of massive releases) by 250% (and even 350% when a minimal time is reached).

Thus, for a given number of sterile males (here 6000 Ind/ha) to release, the impact of RF is important, such that it seems natural to increase the size of the releases. This is done in Fig. 9, page 28 with  $\Lambda_M=12000$  Ind/ha. Surprisingly, the benefit in duration is quite low. It is only when RF=1.2% (see Fig. 9(c), page 28), that the gain (in time) is significant, but only in periods when the mosquito population is large. Thus, increasing the release rate reduces a bit the duration of the massive releases, but the gain is very little compared to the production effort and thus to the cost increase. In fact, above a certain amount, it seems that a saturation effect occurs, such that very massive releases over a long time are not necessarily (always) an appropriate strategy.

The death rate of mosquitoes is mainly linked to the temperature, we can intuitively think that the best period to act is during the Austral winter, i.e. from the end of June to September. According to Figs. 8 and 9, page 27 and 28, the best intervention period, i.e. the SIT starting-time, lasts, in general, from July to December. Indeed, in this figures, we observe that the lowest number of weeks needed are in this period. Thus, the window to start the SIT control is larger than expected, and is not necessarily reduced to the Austral winter period (from June to September) but can also include the Spring period (from October to December).

2. In Table ??, page ??, we provide the results obtained with the average values for the biological parameters, computed thanks to their mean values over the period [2010, 2020]: we obtain a mean estimate of the duration of the massive releases according to the levels of mechanical control, the levels of residual fertility and for the two weekly release rates. We remark that it requires only **one** computation and thus can help to derive a first (rough) approximation in terms of releases strategy as well as the minimal amount of sterile males to produce over the massive releases period.

While for small residual fertility, i.e. 0% and 0.6%, we obtain comparable average duration estimate (compare Table ??), for a large residual fertility, the result is clearly different, and certainly far from the reality.

Increasing the sterile males release rate, i.e. switching from  $6000 \ Ind/ha$  to  $12000 \ Ind/ha$ , is not necessarily useful: it is very costly in terms of sterile males with a little gain with respect to the duration of the massive releases, especially when mechanical control is strong, i.e. at 40%.

Of course, as expected, mechanical control has a positive effect with, in mean, a gain of time between 5 and 10 weeks, depending on the level of residual fertility. Again, this is to balance between the cost of mechanical control and the effective gain in time, and thus the gain in terms of sterile males.

This "mean constant" simulation provides reasonable values, at least for residual fertility less than or equal to 0.6%, that can help to evaluate the maximal duration of the massive releases with and without mechanical control. However, this approach does not provide any information on the best period to start the SIT treatment to reduce the duration of the massive releases and, thus, optimize the sterile males production.

3. In the third case, we consider, like in many other so-called "realistic" models, temperature-dependent parameters. We use the approach developed in [10], where the carrying capacity, K, depends on the temperature only. It was defined as follows: the capacity is at its maximum, i.e.  $K = K_{max}$ , at  $27^{\circ}C$ , which is the mean temperature at the end of the rainy season in Réunion Island. It is assumed that  $15^{\circ}C$  corresponds to austral winter, when the precipitation is low (dry season), and the capacity is at the lowest,  $K = K_{\min}$ . We assumed that  $K_{\min} = 0.1 \times K_{\max}$ . Therefore, we assume that K increases when the temperature is goes from  $15^{\circ}C$  to  $27^{\circ}C$ . Then, when the temperature is above  $27^{\circ}C$ , it is assumed that K decreases, either due to evaporation and to the fact that, in the period where high temperature occurs, heavy rains occur and can be detrimental to breeding sites, such that  $K = 0.75 \times K_{\max}$  at  $35^{\circ}C$ . Thus, a continuous relation between K and the temperature is obtained using linear interpolation.

This leads to Fig. 10, page 29 and Fig. 11, page 30. Compared to the simulations obtained for the temperature and rainfall dependent model, the dynamics is rather different with a smoothing effect such that the amplitudes between the dry and the rainy seasons are strongly reduced: compare with Fig 8, page 27. This does not allow us to determine the best period to start the releases.

In Table ??, page ??, we derive the average values for the duration and the total amount of sterile males to release. We notice that the temperature-dependent model provides almost similar results whatever the level of residual fertility: compare with Table ?? to Table ??. The over-estimate of the duration might come from the way we estimate the carrying-capacity according to temperature only.

Like in the previous cases, increasing the sterile males release rate does not provide, on average, a great benefit: compare the values in Table ??, page ??. However, on the contrary, mechanical control might have an positive impact.

The main disappointment with this temperature-dependent only model comes from the fact that the interannual periodic behavior, for which the best period to start SIT would (roughly) be between May and September. This is exactly the same conclusion reached by entomologists.

4. In the fourth and last case, we consider the average values for the parameters related to temperature, but rainfall is taken into account following the description given in the previous section. The simulations are provided in Figs 12, page 31, and 13, page 32.

We notice that, like in Figs. (8) and (9), we recover larger amplitude in the oscillations between rainy and dry periods. This allows to capture years where rainfall was more abundant than usual (like in 2010 and 2011, for instance), and, also to capture, like for the temperature and rainfall-dependent model, the best periods to start SIT.

Mechanical control already has an impact on massive duration treatment

However, while for small residual fertility, less than or equal to 0.6%, the rainfall-model seems to provide an equivalent result to the full model, for large residual fertility, namely 1.2%, the results seem to be worst, and, even worse than those obtained for the average parameters model (case 2).

Table 2: Simulations with constant (average) temperature and rainfall-dependent parameters: massive releases duration and total mean amount of sterile males to release over the 20 hectares: (a) 0% of Residual Fertility; (b) 0.6% of Residual Fertility; (b) 1.2% of Residual Fertility

	6000 Ind/ha		12000 Ind/ha	
$\varepsilon = 0$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	52	6 240 000	45	10 800 000
20% of MC	48	5760000	42	10080000
40% of MC	44	5280000	40	9600000

(b)

	6000 Ind/ha		12000 Ind/ha	
$\varepsilon = 0$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	60	7200000	52	12480000
20% of MC	55	6 600 000	49	11760000
40% of MC	51	6120000	46	11040000

(c)

6000  Ind/ha		12000 Ind/ha	
Mean number of	Total mean amount of	Mean number of	Total mean amount of
releases	sterile males released	releases	sterile males released
75	9 000 000	65	15 600 000
69	8280000	61	14640000
63	7560000	57	13680000
	Mean number of releases  75 69	Mean number of releasesTotal mean amount of sterile males released759000000698280000	Mean number of releasesTotal mean amount of sterile males releasedMean number of releases7590000006569828000061

To summarize: from the four cases, we derive contrasted results. On average, and for small residual fertility, we obtain almost similar results: see Tables ??, Table ??, Table 2, and Table ??. In particular, the average (constant) model is not so bad. So, the use of constant parameters is pertinent in this case. However, since we are not yet capable of predicting weather data accurately, it is important to have an initial estimate for the duration of the massive treatment and then refine this estimate while the process is in progress.

Based on the average values given in the previous Tables, we can estimate, roughly at least, the duration and the amount production of sterile males needed for the massive releases to reduce the nuisance. In general, the amount of sterile males to produce is considerable. However, all along the treatment, the size of the massive releases can be adapted to the wild male population, using Mark-Release-Recapture experiments, as well as the periodicity of the releases, like in [5,38].

Last, if the residual fertility is too large, say to 2%, then it is impossible to reduce (in a reasonable amount of time) the wild population under the given threshold, such that the wild population can be controlled with small releases of  $\tau \Lambda_M = 20 \times 100$  individuals. The recommendation is thus to improve the sterilization process to have a residual fertility as small as possible, at least, less than 0.6%, for instance.

Our simulations took place in the context of nuisance reduction, i.e. to reduce the wild mosquito population in order to reduce the number of bites. It is not always necessary to reach this objective. In particular, in a tropical context, where people are used to mosquitoes. There, the most important goal is to reduce the epidemiological risk.

#### 3.3 Reducing the epidemiological risk

In the previous simulations, we derive numerics to lower the mosquito population under a given release threshold for sterile males, for instance, 100 Ind/ha/week. As explained, in a tropical context, another option is to reduce the population in order to reduce the epidemiological risk, to prevent the risk of an epidemic. Since dengue is often circulating in La Réunion, we can couple our entomological model with a dengue model, like the one

Table 3: *DENV* epidemiological constant parameter values [39]

Symbol	$\mu_h$	$\eta_h$	$N_h$
Value	$\frac{1}{365 \times 78}$	1/7	2000
Unit	$day^{-1}$	$day^{-1}$	Ind

developed in [39]. According to the epidemiological model developed in [39], and recalled in Appendix B, we derive the following formula for the SIT basic reproduction number

$$\mathcal{R}_{0,SIT}^{2} = \frac{\nu_{m}}{\nu_{m} + \mu_{F}} \frac{B\beta_{mh}}{\mu_{F}} \frac{B\beta_{hm}}{\eta_{h} + \mu_{h}} \frac{F_{S}^{*}}{N_{h}}, \tag{11}$$

where  $F_S^*$  is the amount of susceptible adult females at the Disease Free Equilibrium (DFE) equilibrium, estimated according to the value taken by  $\tau\Lambda_{Massive}$ . In [39], the authors showed that  $F_S$  is either strictly positive or equal to zero, depending on the amount of sterile males released. The positive parameters  $\mu_h$  and  $1/\nu_h$  represent respectively the average human mortality rate and the average viremic period. Since no disease-induced mortality is considered, the total human population is supposed to be constant and equal to  $N_h$ . The average rate of mosquito bites per individual is denoted B>0, and  $\beta_{mh}>0$  ( $\beta_{hm}>0$ ) is the probability of dengue transmission from an infected female mosquito (human) to a susceptible human (mosquito) during such an event. The positive parameter  $\nu_m>0$  is the extrinsic incubation rate (EIR). However, it is well known that the previous epidemiological parameters, B,  $\nu_m$ ,  $\beta_{hm}$ , and  $\beta_{mh}$ , are Temperature-dependent. Using results from [40], we consider the following transmission probability for an infected Aedes albopictus to transmit DENV-2, using a Lactin-1 function

$$\beta_{mh}(T) = \exp(\alpha \times T) - \exp\left(\alpha \times T_{max} - \frac{(T_{max} - T)}{\delta_T}\right),$$

with  $\alpha = 0.20404$ ,  $T_{max} = 37.354$ , and  $\delta_T = 4.89694$ . The probability of transmission from humans to mosquitoes is negligible for "low" temperatures, increases linearly to one at a maximum temperature and remains at one for higher temperatures. Thus, following [41], we consider

$$\beta_{hm}(T) = \frac{T^7}{T^7 + \beta_h^7},$$

with  $\beta_h = 18.9871$ . Last but not least, the EIP (Extrinsic incubation Period) decreases according to the temperature. Using again [40], we derive the following interpolation:

$$\nu_m(T) = aT^2 + bT + c,$$

with a = -0.001, b = 0.0670, and c = -0.866. All other (epidemiological) parameters are supposed to be constant

Finally, following [19] [Table A in the supplementary material], we consider the following Brière function for the biting rate B:

$$B(T) = c \times T \times (T - T_{\min}) \sqrt{T_{\max} - T},$$

where c = 1.93e - 04,  $T_{\min} = 10.25$ , and  $T_{\max} = 38.32$  are the parameters related to Aedes albopictus mosquito. The threshold  $\mathcal{R}_{0,SIT_c}^2$  is related to the long-time behavior of the system. For practical purpose, we will consider  $\mathcal{R}_{eff}$ , the effective reproduction number, that is defined as follow

$$\mathcal{R}_{eff}(t) = \frac{\nu_m(t)}{\nu_m(t) + \mu_F(t)} \frac{B(t)^2 \beta_{mh}(t) \beta_{hm}(t)}{\mu_F(t) (\eta_h + \mu_h)} \frac{F_S(t)}{N_h}.$$
 (12)

In fact, assuming that  $t_{DENV}$  is the time where a DENV virus starts circulating, we will estimate  $\mathcal{R}_{eff}$  at time  $t_{DENV}$ . Clearly, if  $\mathcal{R}_{eff}(t_{DENV}) < 1$  and  $\mathcal{R}_{0,SIT_c}^2 < 1$ , then no epidemics will occur. In contrary, even if  $\mathcal{R}_{0,SIT_c}^2 < 1$  but  $\mathcal{R}_{eff}(t_{DENV}) > 1$  then an outbreak may occur.

In the forthcoming simulations, we will estimate the time needed to lower  $\mathcal{R}_{eff}$  below 0.5 for different sizes of massive releases, i.e. 6000 or 12000 sterile males per ha, and for different residual fertility, with and without mechanical control. Thus, it suffices to find  $t^*$  such that the wild female population verifies

$$F(t^*) < \frac{\nu_m(t^*) + \mu_F(t^*)}{\nu_m(t^*)} \frac{\mu_F(t^*) (\eta_h + \mu_h)}{B(t^*)^2 \beta_{mh}(t^*) \beta_{hm}(t^*)} \frac{N_h}{2}, \tag{13}$$

for a given  $N_h$ . For the numerical simulations, we will consider the parameters values given in Table 3, page 12.

In Fig. 14, page 33, and Fig. 15, page 34, for a given SIT starting date, we compute the number of releases necessary to reach  $R_{eff}(t^*) < 0.5$  for the full model. Of course, the duration of the SIT control. When  $N_h = 2000$ , it is very interesting to see that the results differ from the previous objective of reducing the nuisance. Indeed, depending on the year and the period within the year, the number of releases varies between 2 and 34 (22), when 6000 (12000) sterile males are released every week, whatever the residual fertility. This shows that the objective of reducing the epidemiological risk is easier to reach, even when residual fertility occurs than reducing the nuisance. In addition, releasing more sterile males is only beneficial in years where the wild population is very large (2010, 2011, for instance), but globally there is no important gain. Last but not least, residual fertility is not really an issue here, and, again, mechanical control is only useful when the wild population is large. Otherwise, when the population is small (from September to December), SIT alone could work.

Simulations with the temperature-only and the rainfall-only dependent models are also given in Fig. 16, page 35 and Fig. 17, page 36, for a weakly release rate of 6 000 Indiv/ha. We recover the same results as with the full model, i.e. a very low impact of the residual fertility; in some periods, reaching  $R_{eff}(t^*) < 0.5$  can be fast

Like in the nuisance reduction section, it is to compare the results obtained with the full model with results obtained with the temperature-dependent or rainfall-dependent parameters SIT model. As expected from the previous computations the amplitudes of oscillations with this temperature-dependent model are small compared to the rainfall-temperature model, while they are almost similar with the rainfall-dependent model. Thus, the number of releases varies between 20 (14) and 34 (23) at most, when 0% (40%) mechanical control occur: see Fig. 16<sub>1</sub>. Also, the simulations in Fig. 16, page 35, confirm that the residual fertility has less negative effect to reach  $\mathcal{R}_{eff} < 0.5$  than to reach the objective of nuisance reduction. Still, with 1.2% residual fertility, the amount of releases varies between 21 (14) and 36 (24), almost the same values as those obtained with 0% residual fertility.

In fact, the results provided by the temperature-only model are quite acceptable for periods where the mosquito population is large but seems not so accurate to derive the appropriate amount of releases when the population is (strongly) regulated by rainfall, i.e. in periods where there is a rain deficit.

We also derive Table 4, page 13, for the average values model in order to compare with the estimates obtained for the temperature-dependent model (Table 5, page 14), the rainfall-dependent model (Table 6, page 15), and the temperature and rainfall dependent model (Table 7, page 16). The conclusions are almost the same: the impact of residual fertility is (very) low, as well as the impact of mechanical control and very massive releases (12 000 MS/ha). However, of course, this approach can not provide the best period to start in order to minimize the amount of the release.

Of course, once  $\mathcal{R}(t^*)$  < 0.5 is reached, it will be necessary to continue to release enough sterile males to maintain  $\mathcal{R}(t^*)$  below 0.5, as long as needed. This can be evaluated through numerical simulations.

Remark 1. In fact for this particular objective of reducing the epidemiological risk, and because in La Réunion we have some seasonality, the meaningful strategy, as already proposed in [10, Fig.4], would be to consider massive releases only over the wet period to maintain the mosquito population at a level corresponding to the population size during the dry period, the Austral winter, where, in general, the epidemiological risk is low because the vector population is low, except, of course, when rainfalls occur during this period, like in 2010 and 2011. Thus, we would have a "massive and stop" SIT strategy, contrary to the nuisance reduction where a "massive and small" releases strategy seems more appropriate. According to the temperature and rainfall dependent model, the duration of the massive releases could occur from October-November to May-June, that is over 8 months, with, eventually, a reduction of the size of the massive releases, using, for instance, a closed-loop control strategy

### 4 Conclusion

We build a minimalistic rainfall-temperature entomological model to derive the dynamics of *Aedes albopictus* population in the place of Duparc (Sainte-Marie), La Réunion island. Since we are in a sub-tropical environment, our simulation shows that rainfall definitively plays a great role in the dynamics of the mosquito population, with rapid decay or growth. In fact, it seems that they are periods where the mosquito dynamics are mainly driven by temperature, while in other periods the dynamic is more driven by rainfall. Other approaches, based either on constant, temperature-dependent or rainfall-dependent parameter values, provide similar results on average. However, they are not satisfactory enough to be an "accurate" alternative.

Quality control within SIT is an important issue: if it fails, then release cannot occur. Within the quality control, we focus on residual fertility. We show that it may have an important impact on SIT duration and, eventually on its efficacy. In nuisance reduction, the lower the residual fertility, the lower the number of massive releases needed to switch from a massive releases strategy to a small releases strategy. However our results show that the massive releases duration for "high" residual fertility can greatly vary depending if the model takes into account the temperature, the rainfall or both. Also, residual fertility seems to be less detrimental when it comes to consider a SIT strategy to reduce the epidemiological risk.

Table 4: Reduction of the epidemiological risk. Simulations with the mean values parameters: massive releases duration and total mean amount of sterile males to release over the 20 hectares: (a) 0% of Residual Fertility; (b) 0.6% of Residual Fertility; (b) 1.2% of Residual Fertility

	6000  Ind/ha		12000 Ind/ha	
$\varepsilon = 0$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	23	2 760 000	18	4320000
20% of MC	19	2280000	16	3840000
40% of MC	16	1920000	14	3360000

6000  Ind/ha		12000 Ind/ha	
Mean number of	Total mean amount of	Mean number of	Total mean amount of
releases	sterile males released	releases	sterile males released
23	2760000	18	4320000
20	2240000	16	3840000
17	2040000	14	3360000
	Mean number of releases  23	Mean number of releases  23 20 20 Total mean amount of sterile males released 2760 000 2 240 000	Mean number of releasesTotal mean amount of sterile males releasedMean number of releases232 760 00018202 240 00016

(c)

	6000 Ind/ha		12000 Ind/ha	
$\varepsilon = 0.012$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	24	2880000	19	4560000
20% of MC	20	2400000	17	4080000
40% of MC	17	2040000	15	3600000

When SIT is considered, coupled or not with Mechanical control, we recommend starting the releases of sterile males within a period that last from July to December, when the mosquito population is, in general, at its lowest. However, the duration of the SIT treatment or the number of periodic releases, constant in size, of sterile males may vary thanks to the environmental parameters.

Clearly, combining Mechanical control with SIT is strongly recommended, in particular when the SIT treatment starts within a humid year: see for instance the years 2010 and 2011 where the dry period was more rainy than usual.

Last, the massive-small releases strategy is only useful to reduce substantially the nuisance due to mosquitoes, i.e. to reach nearly elimination and to maintain the population under a certain level, related, here, with the number of sterile male individuals considered in the small releases. If the objective of SIT control is only to reduce the epidemiological risk, i.e.  $\mathcal{R}_{eff} < 0.5$ , then the duration of the SIT treatment, with massive releases, will be short compared to elimination. In addition, having non-zero residual fertility, as long as  $\varepsilon < 1/\mathcal{N}$ , seems to be less problematic to reach  $\mathcal{R}_{eff} < 0.5$ , than to reach elimination. From a practical point of view, this can be very convenient. However, once  $\mathcal{R}_{eff} < 0.5$  or elimination is reached, it will be necessary to continue to release a sufficient amount of sterile males in order to keep the wild population under a certain threshold. At this stage, a closed-loop control (taking into account feedback from the system, like the size of the mosquito population through Mark Release and Recapture experiments) can be used from time to time, in order to reduce the overall cost, as described in [5,38].

Our model, while minimalistic from the variables and parameters point of view, captures relatively well the dynamics of the *Aedes albopictus* population along the year, and this without too many details. However, as with all models, improvements could be made. For instance, by taking into account the migration of males and females from neighboring places. Indeed, we have considered our area closed, in the sense that no external mosquitoes can invade the treated area. Migration is another big concern in SIT treatment. We recently showed that if migration is little, then SIT treatment can work [6]. Otherwise, it is mandatory to isolate the targeted domain. This is not so easy with mosquitoes because, so far, no (killing) attractant is efficient.

Last, we assume that no sterile females are released. In fact, we know that this is not the case: there is always a (small) percentage of sterile females that are released. If, in the past, only less than 5% of sterile females was acceptable [42], this is not the case now. IAEA recommends not to release more than 1% of sterile females.

Table 5: Reduction of the epidemiological risk. Simulations with temperature-dependent parameters: massive releases duration and total mean amount of sterile males to release over the 20 hectares: (a) 0% of Residual Fertility; (b) 0.6% of Residual Fertility; (b) 1.2% of Residual Fertility

	6000 Ind/ha		12000 Ind/ha	
$\varepsilon = 0$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	29	3 480 000	21	5040000
20% of MC	24	2880000	18	4320000
40% of MC	20	2400000	16	3840000

	6000 Ind/ha		12000 Ind/ha	
$\varepsilon = 0.006$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	29	3 480 000	21	5040000
20% of MC	24	2880000	19	4560000
40% of MC	20	2400000	16	3840000

(c)

	6000 Ind/ha		12000 Ind/ha	
$\varepsilon = 0.012$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	30	3 600 000	22	5280000
20% of MC	25	3000000	19	4560000
40% of MC	20	2400000	17	4080000

This is part of the control quality process. Thus, following [39], accidental releases of females could be taken into account, in order to derive how they could impact the SIT releases strategy along a year, from the nuisance reduction or epidemiological point of view.

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Table 6: Reduction of the epidemiological risk. Simulations with rainfall-dependent parameters: massive releases duration and total mean amount of sterile males to release over the 20 hectares: (a) 0% of Residual Fertility; (b) 0.6% of Residual Fertility; (b) 1.2% of Residual Fertility

	6000 Ind/ha		12000 Ind/ha	
$\varepsilon = 0$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	21	2520000	17	4 080 000
20% of MC	18	2160000	15	3600000
40% of MC	15	1800000	13	3120000

	6000 Ind/ha		12000 Ind/ha	
$\varepsilon = 0.006$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	22	3 480 000	17	4080000
20% of MC	19	2880000	15	3600000
40% of MC	16	2400000	13	3120000

(c)

	6000 Ind/ha		12000 Ind/ha	
$\varepsilon = 0.012$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	22	3 600 000	17	4080000
20% of MC	19	3000000	16	3840000
40% of MC	16	2400000	14	3360000

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Table 7: Reduction of the epidemiological risk. Simulations with temperature and rainfall dependent parameters: massive releases duration and total mean amount of sterile males to release over the 20 hectares: (a) 0% of Residual Fertility; (b) 0.6% of Residual Fertility; (b) 1.2% of Residual Fertility

( )	v			
	6000 Ind/ha		12000 Ind/ha	
$\varepsilon = 0$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	20	2 400 000	16	3 840 000
20% of MC	17	2040000	14	3360000
40% of MC	14	1680000	12	2880000

	6000 Ind/ha		12000 Ind/ha	
$\varepsilon = 0.006$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	20	2 400 000	16	3 840 000
20% of MC	17	2040000	14	3360000
40% of MC	15	1800000	13	3120000

(c)

	6000  Ind/ha		12000 Ind/ha	
$\varepsilon = 0.012$	Mean number of	Total mean amount of	Mean number of	Total mean amount of
	releases	sterile males released	releases	sterile males released
0% of MC	21	2520000	16	3 840 000
20% of MC	18	2160000	14	3360000
40% of MC	15	1800000	13	3120000

Table 8: Description of the epidemiological and entomological parameters

Symbol	Description	Unit
$1/\mu_h$	Average lifespan of human	Day
$1/\eta_h$	Average viremic period	Day
В	Daily average mosquito bites	$\mathrm{Day}^{-1}$
$\beta_{mh}$	Transmission probability from infected mosquito	=
$\beta_{hm}$	Transmission probability from infected human	=
φ	Number of eggs at each deposit per capita	$\mathrm{Day}^{-1}$
$\gamma$	Maturation rate from larvae to adult	$\mathrm{Day}^{-1}$
$\mu_{A,1}$	Density independent mortality rate of the aquatic stage	$\mathrm{Day}^{-1}$
$\mu_{A,2}$	Density dependent mortality rate of the aquatic stage	Day <sup>-1</sup> Individuals <sup>-1</sup>
r	Sex ratio	-
$1/\nu_m$	Average extrinsic incubation period (EIP)	Day
$1/\mu_F$	Average lifespan of fertilized and eggs-laying females	Day
$1/\mu_M$	Average lifespan of males	Day
$\Lambda_M$	Sterile male release rate	Individuals $\times$ Day <sup>-1</sup>

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Table 9: Entomological parameters of Aedes albopictus at different temperatures (from [9])

Symbol	Name	$T=15^{o}$	$T=20^{\circ}$	$T=25^{o}$	$T=30^{\circ}$	$T=35^{o}$
$r_{viable}$	Proportion of viable eggs (E-L1)	8.2	66.9	49.2	51.4	10
$N_{eggs}$	Number of eggs deposited	0	50.8	65.3	74.2	48.7
$ au_{gono}$	Duration of the gonotrophic cycle	NA	8.1	3.1	3.9	1.3
$ au_A$	Time from hatching to emergence	35	14.4	10.4	8.8	12.3
$s_A$	Survivorship from larva first instar to adult	50	77.5	76.3	67.5	2.5
$ au_M$	Adult male half-life	15.45	10.25	9.6	8.55	7.4
$ au_F$	Adult female half-life	19.65	15.15	15.3	16.9	10

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### 5 Appendix: Parameters values

See Table 9, page 18, and Table 10, page 19.

## 6 Appendix A: Equilibria

• Without release

In the case of constant coefficients, the equilibria of System (2) are 0 and

$$\begin{cases}
A^* = \frac{\gamma + \mu_{A,1}}{\mu_{A,2}} (\mathcal{N} - 1), \\
M^* = \frac{(1 - r)\gamma A^*}{\mu_M} = Q(\mathcal{N} - 1), \\
F^* = \frac{r\gamma A^*}{\mu_F},
\end{cases} (14)$$

Table 10: Parameter values for system (2) deduced from Table 9, page 18

			mean	mean	mean	mean	mean
Symbol	Name	Formula	value at				
			$15^{o}$	$20^{o}$	$25^{o}$	$30^{o}$	$35^o$
φ	Effective fecundity	$\frac{r_{viable}N_{eggs}}{ au_{gono}}$	0	4.1957	10.3637	9.7792	3.7462
$\mu_{A,1}$	Aquatic death rate	$-rac{log(s_A)}{ au_A}$	0.0198	0.0177	0.0260	0.0447	0.2999
$ u_A$	Aquatic to adult	$rac{1}{ au_A}$	0.0286	0.0694	0.0962	0.1136	0.0813
	transition rate						
$\mu_M$	Adult male death rate	$rac{\log(2)}{ au_M}$	0.0449	0.0676	0.0722	0.0811	0.0937
$\mu_F$	Adult female death rate	$rac{log(2)}{ au_F}$	0.0353	0.0458	0.0453	0.0413	0.0693

The formula given in Table 10 are similar to those given in [31]. Let us explain how we deduce  $\mu_M$  from  $\tau_M$ . Since  $\tau_M$  is the adult-male half-life, it means that half of the population has disappeared in  $\tau_M$  days. Then, assuming that the dynamic of adult males follows an exponential law, i.e.  $x(t) = x(0)e^{-\mu_M t}$ , it is straightforward to deduce that  $\mu_M$  is solution of  $\frac{1}{2} = e^{-\mu_M \times \tau_M}$ , that is  $\mu_M = \frac{\log 2}{\tau_M}$ . The other parameters values follow with the same reasoning.

where 
$$Q = \frac{(1-r)\gamma\left(\gamma + \mu_{1,A}\right)}{\mu_{2,A}\mu_{M}}$$
.

#### With release

In the case of constant coefficients, some calculations are needed to find the release threshold,  $M_{T_1,\varepsilon}$  and also to derive the equilibria. We have to solve

$$\begin{cases} \phi F = (\gamma + \mu_{1,A} + \mu_{2,A}A) A, \\ (1 - r)\gamma A = \mu_M M, \\ \frac{M + \varepsilon \beta M_T}{M + \beta M_T} r \gamma A = \mu_F F. \end{cases}$$

Using the first and the third equalities leads to

$$\frac{M + \varepsilon \beta M_T}{M + \beta M_T} r \gamma = \frac{\mu_F}{\phi} \left( \gamma + \mu_{1,A} + \mu_{2,A} A \right),$$

and using the fact that

$$M = \frac{(1-r)\gamma}{\mu_M} A,$$

we have

$$\frac{(1-r)\gamma A + \varepsilon \mu_M \beta M_T}{(1-r)\gamma A + \mu_M \beta M_T} r \gamma = \frac{\mu_F}{\phi} \left( \gamma + \mu_{1,A} + \mu_{2,A} A \right),$$

that is

$$r\gamma\phi\left((1-r)\gamma A + \varepsilon\mu_M\beta M_T\right) = \mu_F\left((1-r)\gamma A + \mu_M\beta M_T\right)\left(\gamma + \mu_{1,A} + \mu_{2,A}A\right),$$

leading to second order polynomial

$$\mu_F(1-r)\gamma\mu_{2,A}A^2 + [\mu_F(1-r)\gamma(\gamma + \mu_{1,A}) + \mu_F\mu_M\beta M_T\mu_{2,A} - r\gamma\phi(1-r)\gamma]A + \mu_F\mu_M\beta M_T(\gamma + \mu_{1,A}) - r\gamma\phi\varepsilon\mu_M\beta M_T = 0$$

$$(1-r)\gamma\mu_{2,A}A^{2} + \left[ (1-r)\gamma(\gamma + \mu_{1,A}) + \mu_{M}\beta M_{T}\mu_{2,A} - \frac{r\gamma\phi}{\mu_{F}}(1-r)\gamma \right]A$$
$$+ \mu_{M}\beta M_{T}(\gamma + \mu_{1,A}) - \frac{r\gamma\phi}{\mu_{F}}\varepsilon\mu_{M}\beta M_{T} = 0$$

$$(1 - r)\gamma\mu_{2,A}A^{2} + [(1 - r)\gamma(\gamma + \mu_{1,A})(1 - \mathcal{N}) + \mu_{M}\beta M_{T}\mu_{2,A}]A + \mu_{M}\beta M_{T}(\gamma + \mu_{1,A})(1 - \varepsilon\mathcal{N}) = 0,$$

or equivalently

$$\frac{(1-r)\gamma}{\mu_M}A^2 - \left[Q\left(\mathcal{N}-1\right) - \beta M_T\right]A + \beta M_T \frac{(\gamma+\mu_{1,A})}{\mu_{2,A}}\left(1-\varepsilon\mathcal{N}\right) = 0,$$

that is, using (14),

$$\frac{(1-r)\gamma}{\mu_M} A^2 - [M^* - \beta M_T] A + \beta M_T \frac{(\gamma + \mu_{1,A})}{\mu_{2,A}} (1 - \varepsilon \mathcal{N}) = 0.$$
 (15)

We compute the discriminant of the last equation

$$\Delta(\varepsilon) = \left[M^* - \beta M_T\right]^2 - 4Q\beta M_T \left(1 - \varepsilon \mathcal{N}\right).$$

We will distinguish three cases:

- Assume  $\varepsilon > 1/\mathcal{N}$ , then  $\Delta(\varepsilon) > 0$ , such that there always exists one positive equilibrium

$$A_{\varepsilon}^* = \mu_M \frac{M^* - \beta M_T + \sqrt{\Delta(\varepsilon)}}{2(1-r)\gamma}.$$

After straightforward computations, we can show that even with very massive releases, the aquatic equilibrium is bounded from below by

$$2\frac{\varepsilon \mathcal{N}-1}{\mathcal{N}-1}A^*$$

- Assume  $\varepsilon = 1/\mathcal{N}$ . Then  $\Delta(\varepsilon) = 0$  iff  $\beta M_T = M^*$ . In fact if  $\beta M_T > M^*$ , then  $A_{\varepsilon}^* = 0$ .
- Assume  $\varepsilon < 1/\mathcal{N}$ . Setting  $y = \beta M_T$ , we derive

$$\begin{array}{lll} \Delta(\varepsilon) & = & \left[Q\left(\mathcal{N}-1\right)-y\right]^2 - 4Q\left(1-\varepsilon\mathcal{N}\right)y \\ & = & y^2 - 2Q\left(\left(\mathcal{N}-1\right) + 2\left(1-\varepsilon\mathcal{N}\right)\right)y + \left(Q\left(\mathcal{N}-1\right)\right)^2 \end{array}$$

The discriminant of the equation  $\Delta(\varepsilon) = 0$  following the variable y is given by

$$\begin{array}{lcl} \delta(\varepsilon) & = & (2Q)^2 \left( (\mathcal{N}-1) + 2 \left( 1 - \varepsilon \mathcal{N} \right) \right)^2 - \left( 2Q \left( \mathcal{N}-1 \right) \right)^2 \\ & = & 16 \left( 1 - \varepsilon \mathcal{N} \right) \left( 1 - \varepsilon \right) Q^2 \mathcal{N}. \end{array}$$

Since  $\varepsilon < 1/\mathcal{N}$ , then  $\delta(\varepsilon) > 0$ , and we obtain two roots

$$\beta M_{T_1,\varepsilon} = Q \left( \mathcal{N} + 1 - 2\varepsilon \mathcal{N} - 2\sqrt{(1 - \varepsilon \mathcal{N})(1 - \varepsilon) \mathcal{N}} \right)$$

and

$$\beta M_{T_2,\varepsilon} = Q \left( \mathcal{N} + 1 - 2\varepsilon \mathcal{N} + 2\sqrt{(1 - \varepsilon \mathcal{N})(1 - \varepsilon) \mathcal{N}} \right).$$

When  $\varepsilon = 0$ , we recover the result obtained in [4]

$$\beta M_{T_1,0} = Q \left( \sqrt{\mathcal{N}} - 1 \right)^2.$$

Assume  $0 < \varepsilon < \frac{1}{N}$ . Then, when  $0 < M_T < M_{T_1,\varepsilon}$ , we have  $\Delta(\varepsilon) > 0$ , and thus two equilibria

$$A_{1,\varepsilon} = \mu_M \frac{Q\left(\mathcal{N} - 1\right) - \beta M_T - \sqrt{\left[Q\left(\mathcal{N} - 1\right) - \beta M_T\right]^2 - 4Q\beta M_T\left(1 - \varepsilon \mathcal{N}\right)}}{2(1 - r)\gamma},$$

and

$$A_{2,\varepsilon} = \mu_M \frac{Q\left(\mathcal{N} - 1\right) - \beta M_T + \sqrt{\left[Q\left(\mathcal{N} - 1\right) - \beta M_T\right]^2 - 4Q\beta M_T\left(1 - \varepsilon \mathcal{N}\right)}}{2(1 - r)\gamma}.$$

Since

$$A_{1,\varepsilon} + A_{2,\varepsilon} = \frac{\mu_M}{2(1-r)\gamma} \left( Q\left(\mathcal{N} - 1\right) - \beta M_T \right) > \frac{\mu_M}{2(1-r)\gamma} \left( Q\left(\mathcal{N} - 1\right) - \beta M_{T_1,\varepsilon} \right) = \frac{\mu_M}{2(1-r)\gamma} \left( Q\left(2\varepsilon\mathcal{N} + 2\left(\sqrt{(1-\varepsilon\mathcal{N})(1-\varepsilon)\mathcal{N}} - 1\right)\right) \right).$$

We remark that

$$(1 - \varepsilon \mathcal{N}) (1 - \varepsilon) \mathcal{N} - 1 = \mathcal{N} - 1 - (\mathcal{N} + 1) \varepsilon + \mathcal{N} \varepsilon^{2}.$$

The discriminant of the last polynomial in  $\varepsilon$  is given by

$$\Delta_{\varepsilon} = (\mathcal{N} + 1)^2 - 4\mathcal{N}(\mathcal{N} - 1) = -3\mathcal{N}^2 + 6\mathcal{N} + 1$$

In general,  $\mathcal{N}$  is large, i.e.  $\mathcal{N} >> 1 + \frac{2}{\sqrt{3}}$ , such that  $\Delta_{\varepsilon} < 0$ , such that the  $\varepsilon$ -polynomial is always positive. Then  $A_{1,\varepsilon} + A_{2,\varepsilon} > 0$ . Moreover, since the last term in (15), we have also  $A_{1,\varepsilon}A_{2,\varepsilon} > 0$ . Thus the two roots  $A_{1,\varepsilon}$  and  $A_{2,\varepsilon}$  are positive. Therefore the system has two equilibria  $E_{1,2} = (A_{1,2}, M_{1,2}, F_{1,2})$  such that  $\mathbf{0} << E_1 << E_2$ .

In practice, we are, in general, in the case  $\varepsilon < 1/\mathcal{N}$ , see (10) and  $\mathcal{N} > 1 + \frac{2}{\sqrt{3}}$ .

**Appendix B: The epidemiological Model** We briefly recall the SIR-SEI model of dengue transmission, studied in [39], without taking into account the accidental releases of sterile females. The evolution of the human population is given by the following SIR model, with Susceptible, Infected, and Recovered compartments:

$$\begin{cases}
\dot{S}_{h} = \mu_{h} N_{h} - B \beta_{mh}(T) F_{I} \frac{S_{h}}{N_{h}} - \mu_{h} S_{h}, \\
\dot{I}_{h} = B \beta_{mh}(T) F_{I} \frac{S_{h}}{N_{h}} - (\eta_{h} + \mu_{h}) I_{h}, \\
\dot{R}_{h} = \eta_{h} I_{h} - \mu_{h} R_{h},
\end{cases} (16)$$

Extending the model of evolution of the mosquito population described in the previous sections, we use here a SEI model for the wild female mosquitoes, with Susceptible, Exposed and Infected compartments, adapted from (7):

$$\begin{cases}
\frac{dA}{dt} = \phi(T)(F_S + F_E + F_I) - (\gamma(T) + \mu_{1,A}(T) + \mu_{A,2}(T,R)A) A, \\
\frac{dM}{dt} = (1 - r(T))\gamma(T)A - \mu_M(T)M, \\
\frac{dF_S}{dt} = r(T)\gamma(T)\frac{M + \varepsilon\beta M_S}{M + \beta M_S}A - B\beta_{mh}(T)F_S\frac{I_h}{N_h} - \mu_F(T)F, \\
\dot{F}_E = B\beta_{mh}(T)F_S\frac{I_h}{N_h} - (\nu_m(T) + \mu_F(T))F_E \\
\dot{F}_I = \nu_m(T)F_E - \mu_F(T)F_I.
\end{cases}$$
(17)

where  $M_S$  is driven by (6).

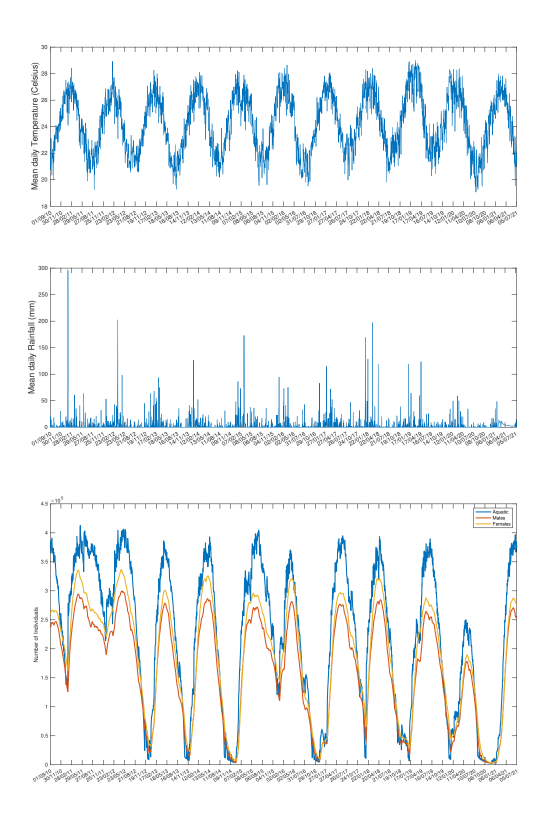
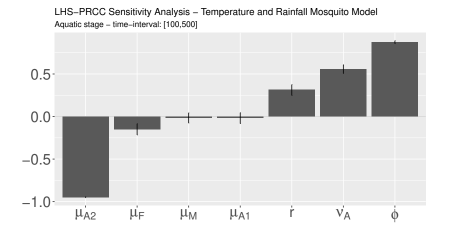
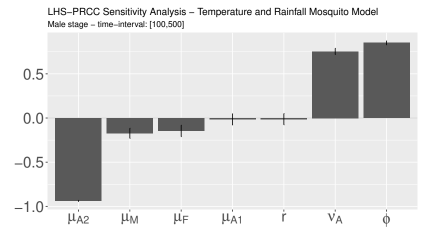


Figure 3: Temperature and rainfall dependent model: (a) Mean Daily Temperature (Celsius) - (b) Mean daily rainfall (mm) - (c) Mosquito population dynamics from the 1st of September 2010 until the 14th of July 2021





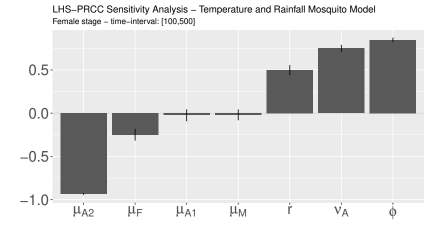
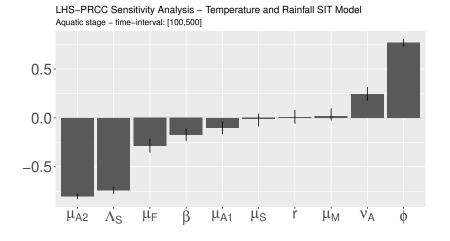
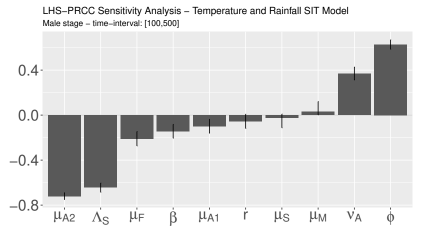


Figure 4: LHS-PRCC analysis of model (2)





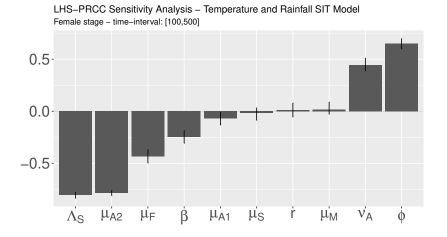


Figure 5: LHS-PRCC analysis of the SIT-model (7)

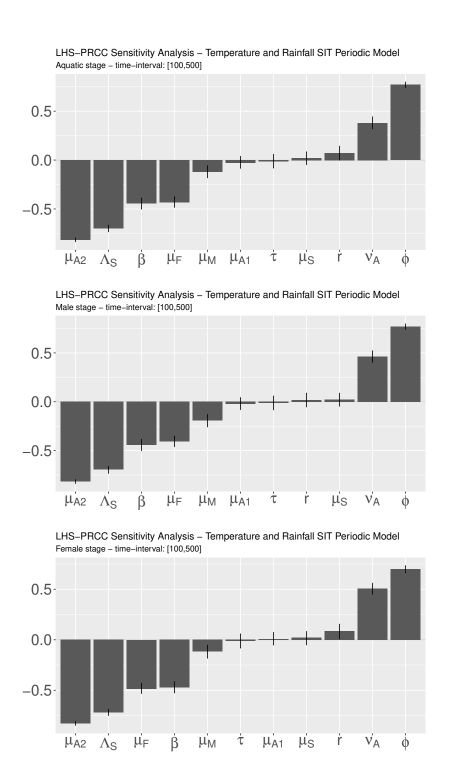


Figure 6: LHS-PRCC analysis of the SIT-model (7) with periodic releases

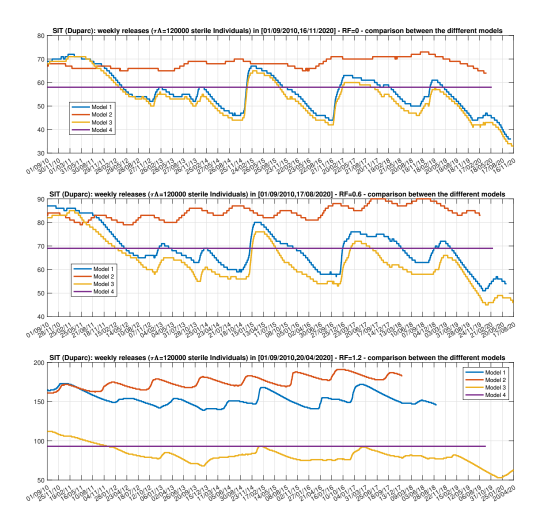


Figure 7: Number of releases needed for mosquito reduction - Comparison between the four models: (a) with 0% residual fertility; (b) with 0.6% residual fertility; with 1.2% residual fertility

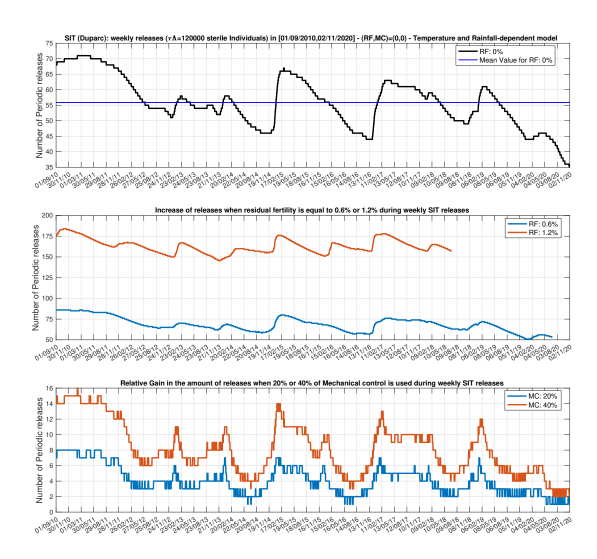


Figure 8: Temperature and rainfall dependent model - Weekly SIT control with 6000 sterile Ind/ha: (a) Duration and mean duration for 0% residual fertility; (b) Additional duration when 0.6% or 1.2% residual fertility occurs ithout mechanical control; (c): SIT in combination with 20% and 40% of mechanical control and 0% of residual fertility: gain in the releases amount

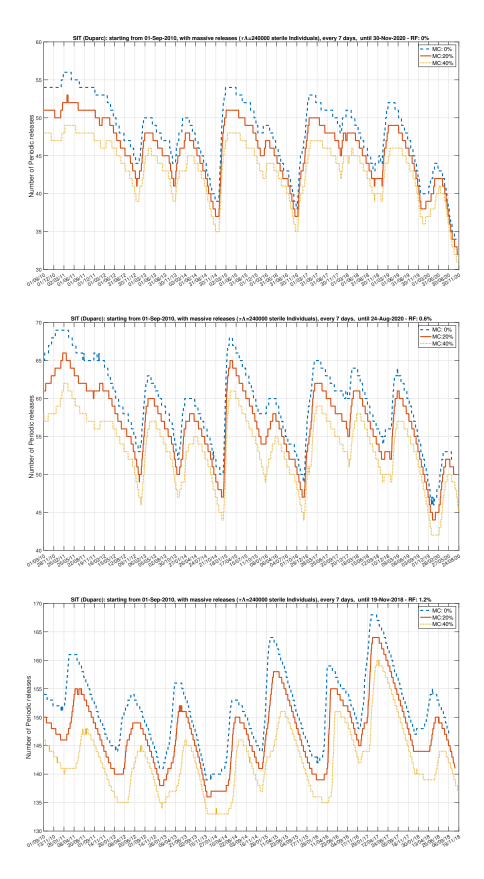


Figure 9: Temperature and rainfall dependent model - Weakly SIT control with 12000 sterile Ind/ha for various level of Mechanical control - Residual fertility: (a) 0%, (b) 0.6%, (c) 1.2%

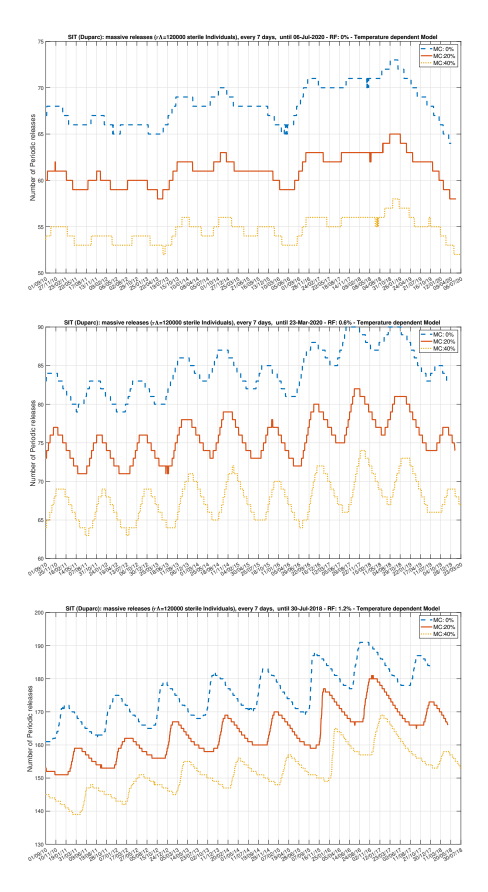


Figure 10: Temperature-dependent model - Weakly SIT control with 6000 sterile Ind/ha for various level of Mechanical control - Residual fertility: (a) 0%, (b) 0.6%, (c) 1.2%

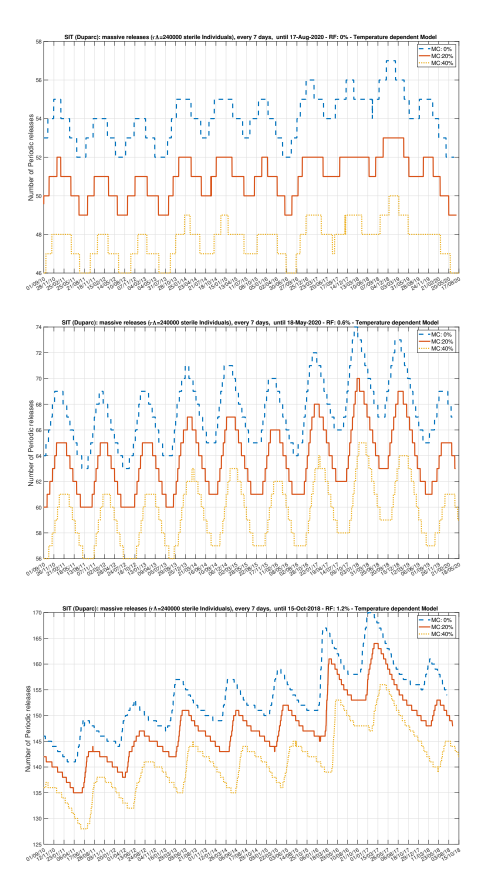


Figure 11: Temperature-dependent model - Weakly SIT control with 12000 sterile Ind/ha for various level of Mechanical control - Residual fertility: (a) 0%, (b) 0.6%, (c) 1.2%

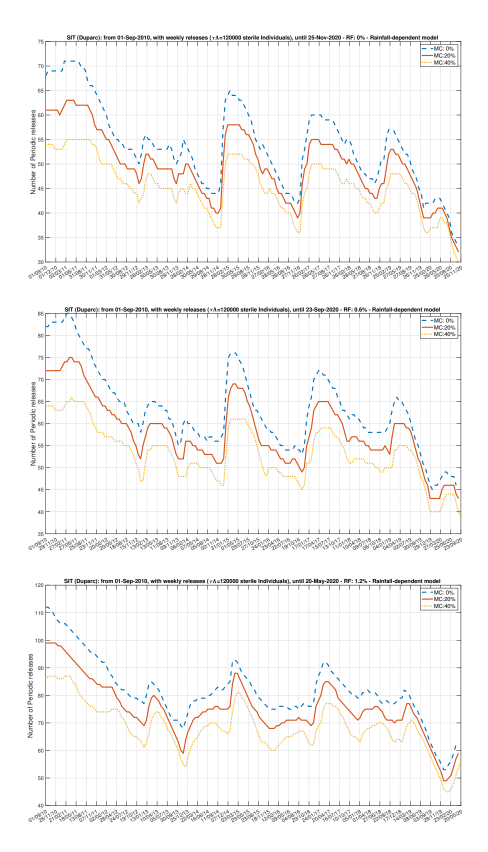


Figure 12: Rainfall-dependent model - Weakly SIT control with 6000 sterile Ind/ha for various level of Mechanical control - Residual fertility: (a) 0%, (b) 0.6%, (c) 1.2%

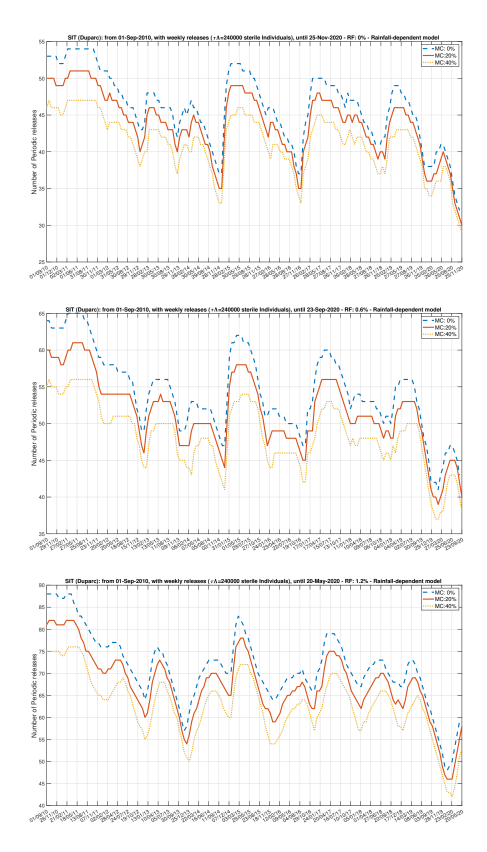


Figure 13: Rainfall-dependent model - Weakly SIT control with 12000 sterile Ind/ha for various level of Mechanical control - Residual fertility: (a) 0%, (b) 0.6%, (c) 1.2%

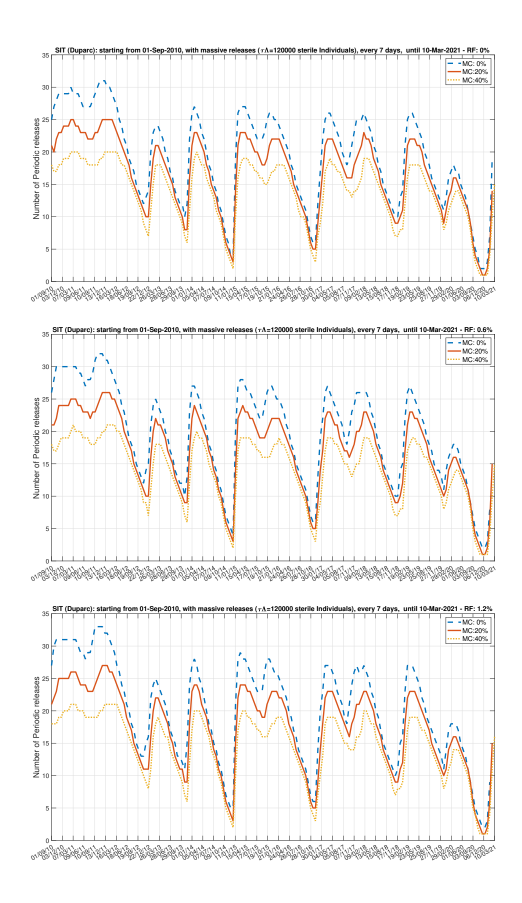


Figure 14: Temperature and rainfall dependent model - Weakly SIT control of 6000 sterile Ind/ha for various level of Mechanical control to reach  $\mathcal{R}_{eff} < 0.5$  - Residual fertility variation: (a) 0%, (b) 0.6%, and (c) 1.2%

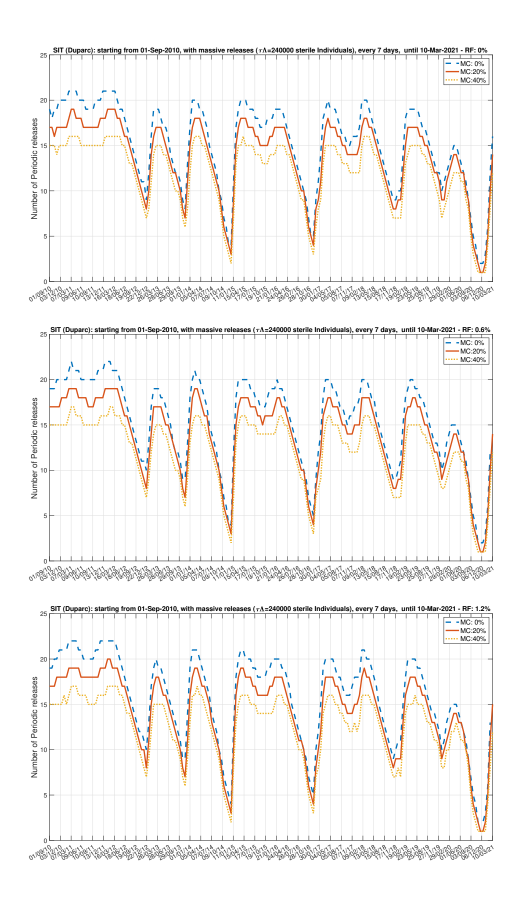


Figure 15: Temperature and rainfall dependent model - Weakly SIT control of 12000 sterile Ind/ha for various level of Mechanical control to reach  $\mathcal{R}_{eff} < 0.5$  - Residual fertility variation: (a) 0%, (b) 0.6%, and (c) 1.2%

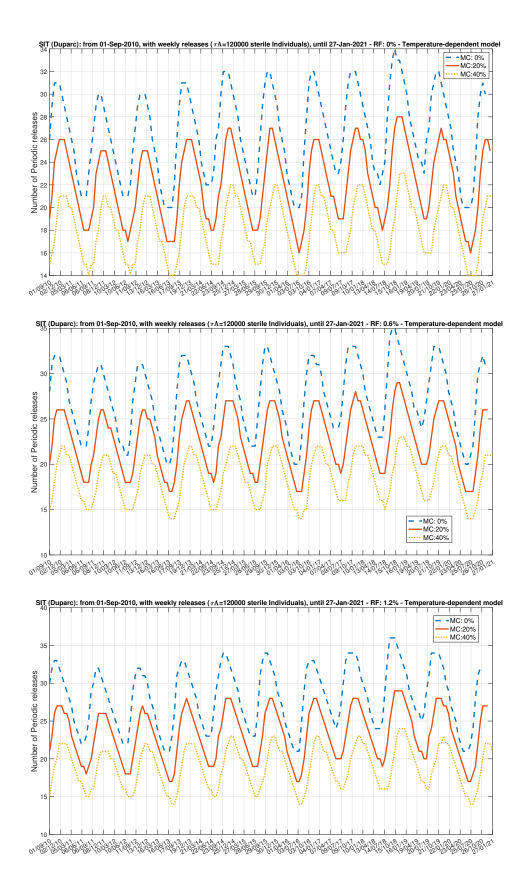


Figure 16: Temperature-dependent model. Weakly SIT control of 6000 sterile Ind/ha for various level of Mechanical control to reach  $\mathcal{R}_{eff} < 0.5$  - Residual fertility variation: (a) 0%, (b) 0.6%, and (c) 1.2%

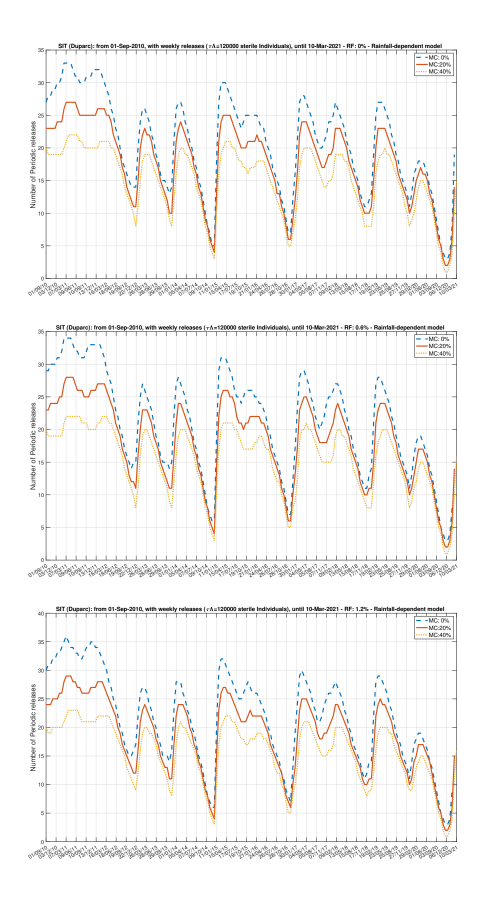


Figure 17: Rainfall-dependent model. Weakly SIT control of 6000 sterile Ind/ha for various level of Mechanical control to reach  $\mathcal{R}_{eff} < 0.5$  - Residual fertility variation: (a) 0%, (b) 0.6%, and (c) 1.2%