

RESEARCH ARTICLE

Omitting age-dependent mosquito mortality in malaria models underestimates the effectiveness of insecticide-treated nets

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Data Availability Statement: All relevant data are within the manuscript and its [Supporting information](#) files. All code used for model fitting and plotting is available on a GitHub repository at <https://github.com/Melissa-Iacovidou/mosquito-mortality>.

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Abstract

Mathematical models of vector-borne infections, including malaria, often assume age-independent mortality rates of vectors, despite evidence that many insects senesce. In this study we present survival data on insecticide-resistant *Anopheles gambiae s.l.* from experiments in Côte d'Ivoire. We fit a constant mortality function and two age-dependent functions (logistic and Gompertz) to the data from mosquitoes exposed (treated) and not exposed (control) to insecticide-treated nets (ITNs), to establish biologically realistic survival functions. This enables us to explore the effects of insecticide exposure on mosquito mortality rates, and the extent to which insecticide resistance might impact the effectiveness of ITNs. We investigate this by calculating the expected number of infectious bites a mosquito will take in its lifetime, and by extension the vectorial capacity. Our results show that the predicted vectorial capacity is substantially lower in mosquitoes exposed to ITNs, despite the mosquitoes in the experiment being highly insecticide-resistant. The more realistic age-dependent functions provide a better fit to the experimental data compared to a constant mortality function and, hence, influence the predicted impact of ITNs on malaria transmission potential. In models with age-independent mortality, there is a great reduction for the vectorial capacity under exposure compared to no exposure. However, the two age-dependent functions predicted an even larger reduction due to exposure, highlighting the impact of incorporating age in the mortality rates. These results further show that multiple exposures to ITNs had a considerable effect on the vectorial capacity. Overall, the study highlights the importance of including age dependency in mathematical models of vector-borne disease transmission and in fully understanding the impact of interventions.

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Author summary

Interventions against malaria are most commonly targeted on the adult mosquitoes, which transmit the infection from person to person. One of the most important interventions are bed-nets, treated with insecticides. Unfortunately, extensive exposure of mosquitoes to insecticide has led to widespread evolution of insecticide resistance, which might threaten control strategies. Piecing together the overall impact of resistance on the efficacy of insecticide-treated nets is complex, but can be informed by the use of mathematical models. However, there are some assumptions that the models frequently use which are not realistic in terms of the mosquito biology. In this paper, we formulate a model that includes age-dependent mortality rates, an important parameter in vector control since control strategies most commonly aim to reduce the lifespan of the mosquitoes. By using novel data collected using field-derived insecticide-resistant mosquitoes, we explore the effects that the presence of insecticides on nets have on the mortality rates, as well as the difference incorporating age dependency in the model has on the results. We find that including age-dependent mortality greatly alters the anticipated effects of insecticide-treated nets on mosquito transmission potential, and that ignoring this realism potentially overestimates the negative impact of insecticide resistance.

Introduction

Malaria is a life-threatening vector-borne parasitic disease, which is endemic in 87 countries, mainly in the African Region [1]. The World Health Organization's (WHO) "Global Technical Strategy for Malaria 2016–2030" outlines global targets in the fight against malaria, including a 90% reduction of malaria case incidence by 2030 [2]. Significant progress towards these targets has occurred, where both the number of cases and the number of deaths due to malaria have decreased between 2010–2019, as outlined in a report published by WHO in 2020 [1]. In 2019, there were around 229 million cases of malaria globally, with 94% of them being in the African Region. Additionally, during the same year, 409,000 deaths due to malaria have been estimated worldwide.

The vectors responsible for the malaria parasite's transmission, through blood feeding, belong to the *Anopheles* genus of mosquitoes. The success of the malaria programmes to date is thanks to a range of interventions, most commonly targeted at these vectors. For example, in sub-Saharan Africa around half of the people at risk are sleeping under insecticide-treated nets (ITNs) [1] which are a way to utilise contact pesticides. Between 2000 and 2015, ITNs contributed to the averted of many cases; 68% of the cases that were prevented due to any intervention are attributed to ITNs, making them a crucial intervention [3]. Bed-nets are currently treated with pyrethroids, and there is evidence of an increase in pyrethroid resistance in malaria vectors, which threatens the elimination efforts [1, 4]. Hence, due to mutations and natural selection, mosquitoes develop the ability to resist the harmful effects of insecticides, leading to what is called "insecticide resistance" [4]. We note that there are new dual active ingredient (dual-AI) ITNs against malaria being tested [5–7], nevertheless, it is important to evaluate the potential impact that insecticide resistance actually has on the efficacy of current malaria interventions [8]. Alout *et al.* [9] claim that despite insecticide resistance, vector control is still crucial and can be effective against malaria transmission. This is in agreement with the systematic review and meta-analysis by Strode *et al.* [10], where the authors concluded that ITNs are more effective than untreated bed-nets, despite insecticide resistance. There has been a mixture of results between different studies [11–15], however, we aim to confirm and add to

the existing knowledge regarding long-term impact of insecticide on longevity by using highly resistant field-derived mosquitoes from Côte d'Ivoire in laboratory experiments.

By targeting mosquitoes, ITNs target the key point in the transmission cycle, not only reducing the opportunity for blood-feeding on humans, but increasing vector mortality and, therefore, decreasing the number of infectious blood-meals a mosquito will contribute during its life. The use of vector control to reduce or even eliminate infection is well supported by mechanistic transmission models originally developed for malaria by Ross and Macdonald and used extensively ever since [16]. A key metric linked to transmission models is the basic reproduction number, R_0 , which describes the number of secondary cases produced by a single case in an otherwise susceptible population [17]. Garret-Jones [18] took the purely entomological components of R_0 and named them *vectorial capacity*. It is defined in [19] as “the expected number of infective mosquito bites that would eventually arise from all the mosquitoes that would bite a single fully infectious person on a single day”, i.e. the average number of humans that get infected due to one infectious human per day. These metrics have been used to study the dynamics of vector-borne diseases and also quantitatively assess the possible impact of interventions to control them.

Contact pesticides were being used at the time Macdonald was researching vector control, and that is when he realised that transmission potential was affected by two important factors relating to mosquito longevity [16]:

- (a). a mosquito which is infected will only become infectious if it survives the time needed for the pathogen to develop, commonly known as the extrinsic incubation period (EIP), and
- (b). once the mosquito is infectious it must take a blood-meal in order to transmit the infection on to a host.

Hence, Macdonald concluded that the number of infectious bites taken by a mosquito will increase the longer the mosquito survives [16, 19]. The longer the EIP is, the less chance a mosquito has to survive it, therefore the younger a mosquito is when it gets infected, the more likely it is to transmit the infection. Thus, the transmission potential relies heavily on the survival of the mosquitoes [16]. Due to Macdonald's analysis, many control programmes aim to reduce the lifespan of mosquitoes [16].

The vectorial capacity depends on the mortality rate of the mosquito, which is typically assumed to be age-independent. Studies have suggested that this assumption, namely that mosquitoes do not senesce, may not be realistic enough for transmission models and can underestimate the impact of vector control strategies [20–24]. The assumption is often used to simplify, otherwise complex, mathematical models, and not because of its biological relevance [24, 25]. On the other hand, it is rare that suitable age-dependent vector mortality data are available to inform more complex models; in addition to average life expectancy with and without ITNs in place, the distribution of life expectancy and how this is impacted by intervention is also required. In the present study, we bring more complex mosquito modelling together with detailed experimental data to demonstrate how we may rethink the way the vectorial capacity is calculated.

To investigate the impact of multiple insecticidal exposures on the mosquitoes and their ability to transmit malaria, we address the following research questions:

1. Does the mortality rate of the insecticide-resistant mosquitoes change due to insecticide exposure through ITNs?
2. If it does change, how is the vectorial capacity impacted?

3. When considering age-dependent models, how different is the vectorial capacity compared to assuming mortality is age independent?

To answer these, we use data collected in Côte d'Ivoire. Laboratory experiments were conducted on field-derived female *Anopheles gambiae s.l.* mosquitoes, one of the main malaria vectors in Côte d'Ivoire [26]. The setup of the experiments allowed the comparison of the mosquitoes' survival rates when they were exposed to standard ITNs versus when they were exposed to untreated nets. We fitted various survival functions to these data to estimate biologically realistic mosquito mortality rates and used these to obtain the vectorial capacity estimates of the mosquitoes with and without exposure to ITNs. By including realistic survival in the calculation of the vectorial capacity, we can observe how the assumed effectiveness of anti-vectorial interventions is affected.

Materials and methods

Ethics statement

All experimental methods were consistent with Penn State IBC protocol no. 48219. The Pennsylvania State University Institutional Review Board determined that the experiments whereby uninfected mosquitoes were attracted to a host did not meet the criteria for human subjects research. The experimental research formed a part contribution to a larger set of studies reviewed and approved by the Côte d'Ivoire Ministry of Health ethics committee (039/MSLS/CNER-dkn), the Pennsylvania State University's Human Research Protection Program under the Office for Research Protections (STUDY00003899 and STUDY00004815).

The modelling analysis of the effect on the vectorial capacity with and without the presence of insecticides is conducted using data that were collected in Bouaké, Côte d'Ivoire. Here the focus is on the malaria parasite *Plasmodium falciparum*, which is one of the six *Plasmodium* species known to regularly infect humans, and both the most prevalent and deadly parasite in sub-Saharan Africa [27]. We present a detailed outline of the experimental setup, along with a presentation of the data, followed by a breakdown of the computation of the vectorial capacity using these data.

Experimental setup and data

The laboratory experiment was conducted on field-derived *Anopheles gambiae s.l.* at $26 \pm 1^\circ\text{C}$ and consisted of two treatments:

- (a). **control (non-exposed)**: in the presence of an untreated net
- (b). **treated (exposed)**: in the presence of an ITN

where a one-way tunnel with two cages was used; a 'holding' cage for the mosquitoes and a 'host' cage for the volunteer's foot that was covered with a treated or untreated net (see Fig 1). The mosquitoes in the experiment are considered to be extremely resistant to the pyrethroid deltamethrin, which was used to treat the ITNs. More details regarding the experimental setup are presented further on.

Mosquito populations. *Anopheles gambiae s.l.* mosquitoes were collected as larvae in natural breeding habitats around Bouaké, in central Côte d'Ivoire, and colonised at the Pierre Richet Institute. These are highly pyrethroid-resistant mosquitoes. The 1014F kdr mutation is almost fixed ($\geq 90\%$), and 1575Y, as well as upregulation of CYP6M2, CYP6P3, and CYP9K1 result in a ≥ 1500 -fold resistance to deltamethrin relative to a standard susceptible strain [28]. Larvae were reared at $27 \pm 2^\circ\text{C}$, $60 \pm 20\%$ RH and ambient light in metallic bowls of 300 larvae with 1L of deionised water. They were fed daily with TetraMin baby fish food following a

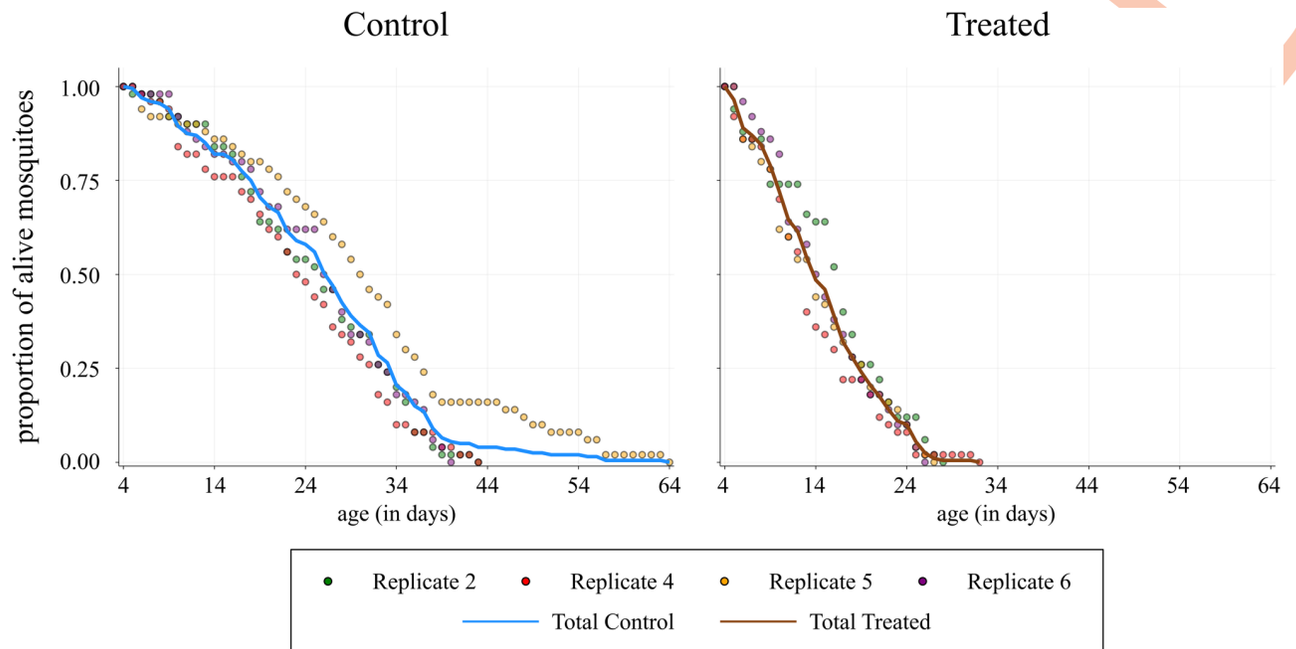


Fig 1. Setup of experiment. The figure depicts the one-way tunnel with the volunteer's foot covered in a net in the 'host' cage on the left, and the 'holding' cage on the right. Both cages are of the same size, $32.5 \times 32.5 \times 32.5$ cm. The cages are connected with a PLEXIGLAS tube ($l = 30$ cm, $d = 14.6$ cm). Depending on the chosen treatment for the experiment, the net is either an unwashed PermaNet 2.0 or an untreated net, measuring 25×25 cm in both cases. In this setup the mosquitoes have direct access to the foot for blood feeding.

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standardised 'high food' regime as described in Kulma *et al.* [29]. Adult mosquitoes were kept in $32.5 \times 32.5 \times 32.5$ cm mosquito cages, in the same environmental conditions as the larvae, and maintained on 10% sugar solution. Mosquitoes were four to five days old at their first exposure to insecticide and randomly assigned to a net treatment. Females had constant access to egg laying substrate (a wet cotton pad) and were maintained on a 10% sugar solution cotton that was renewed daily. Sugar was removed to starve mosquitoes for four hours before each experimental run.

Human host preparation. The volunteers involved in this experiment were not actively infected with malaria. They avoided the use of fragrance, repellent products, tobacco, and alcohol for 12 hours before and during testing. For the experiment, feet were washed with unscented soap and rinsed with water the day before a test. The 'host' cages were also washed with soap and rinsed with water every time after a test was conducted—to avoid the accumulation of insecticide particles. Cages were not interchanged between treatments, i.e. a cage used in a control treatment was always used for a control treatment. Note that the data were analysed anonymously.

Insecticide-treated nets (ITNs). As mentioned, two types of nets were used: an unwashed PermaNet 2.0 (Vestergaard Frandsen SA, DK) and an untreated polyester net (Coghlan's) for the control treatment. The PermaNet 2.0 is a long-lasting insecticidal net made of polyester and coated with $55^{\text{mg}}/\text{m}^2 \pm 25\%$ deltamethrin. We confirmed net efficacy by exposing sensitive mosquitoes (Kisumu strain) to WHO tubes lined with a piece of ITN; all mosquitoes died within 24 hours when exposed to the ITN, while the untreated nets killed none. For the wild-type mosquitoes, as these exhibit such a high level of resistance, there was negligible knock-down or mortality ($< 1\%$ knockdown one hour post exposure and no mortality 24 hours later) from ITN exposure in WHO cone assays [30].

Multiple exposure assay. In malaria-endemic settings with high ITN usage, mosquitoes potentially contact ITNs every time they attempt to feed. To capture this effect, we used a tunnel test in which mosquitoes had to fly a short distance between two cages to locate the host and blood-feed. The tunnel apparatus comprised a standard $32.5 \times 32.5 \times 32.5$ cm mosquito cage as a 'holding' cage, a $32.5 \times 32.5 \times 32.5$ cm mosquito cage as the 'host' cage, and a PLEXI-GLAS tube ($l = 30$ cm, $d = 14.6$ cm) forming the tunnel between cages (as shown in Fig 1). The holding cages were initially populated with 120 pupae each. After adult emergence, 50 females and 10 males were randomly selected to remain in each holding cage until their death, with the excess removed and discarded. We compared the two net treatments, where the foot of a human host was wrapped in a netting sock so the mosquitoes could land on the foot and feed if they chose. Treatments were replicated five times, giving a total of 500 female mosquitoes. Every four days at around 6pm (dusk) the mosquitoes were exposed to a human foot placed in the 'host' cage. The mosquitoes were allowed to visit the cage for 30 minutes. At the end of 30 minutes, the total number of mosquitoes that had taken a full or partial blood-meal was recorded. The tunnel was then dismantled and all mosquitoes returned to their respective holding cages. The surveillance of the mosquitoes started when they were four days old and tests were repeated every four days until all mosquitoes had died. The number of mosquito deaths was recorded daily. The net treatment was randomly allocated to one human host experimenter to ensure there were no biases due to possible differences in attraction between hosts.

During an initial inspection of the data, it was decided that Replicate 1 would be excluded from further analysis. The feeding trend (S1 Fig) of Replicate 1 suggests that the mosquitoes were not feeding and could therefore explain the mortality trend (S2 Fig). The differences in Replicate 1 highlight the fact that mosquito behaviour, for example feeding and host searching, likely depends on a lot of parameters that need further exploration in order to improve transmission models. It seems that there is no significant difference between mosquitoes tested in each treatment in Replicate 1, and this is because they almost never visit the cage with the foot, and thus were not exposed to insecticide during their life. If we were to compare Replicate 1 with the other replicates, it seems probable that blood-meals improve longevity, hence why in both treatments of Replicate 1 the mosquitoes die so early. However, the objective here is to better identify differences of the survival rates between the two treatments, the impact on vectorial capacity, and contrasts when taking age dependencies into account. Since Replicate 1 does not follow the behaviour of the mosquitoes in the rest of the replicates, it is removed from further analysis for consistency.

In the results that follow, the replicates were all combined together, as one larger, aggregated survival dataset, with data from 200 mosquitoes per treatment being examined in total. Fig 2 shows the data used for the calculations. The relevant data can be found in the supporting information (S1–S6 Tables).

Vectorial capacity

Garrett-Jones [18] introduced vectorial capacity, denoted as C , in order to estimate the risk of the introduction of malaria. C is commonly defined mathematically as:

$$C = \frac{m\alpha^2 e^{-\mu n}}{\mu} \quad (1)$$

where m is the mosquito density relative to humans, α is the biting rate, p is the survival rate, μ is the mortality rate, and n is the duration of the EIP, i.e. the number of days between the day a

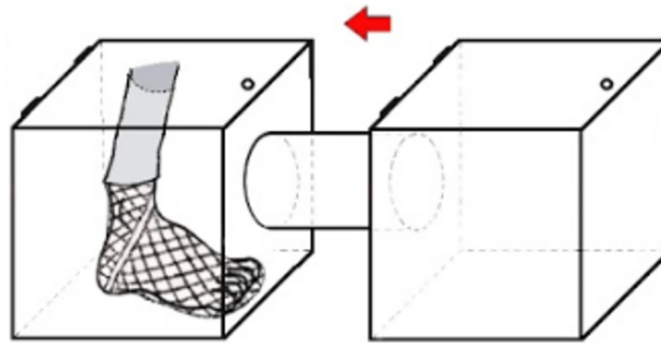


Fig 2. Proportion of alive mosquitoes. Visualisation of the data used for the calculations. The scatter plots show the proportion of alive mosquitoes at each time for 50 mosquitoes in each replicate, and the line shows the proportion of the total number of alive mosquitoes for 200 mosquitoes. Data are plotted up until the final remaining mosquito per replicate died. Replicate 1 was excluded for reasons outlined in the text.

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mosquito gets infected until its bites become infectious, and is able to transmit the infection [22].

It is important to note that this form of the vectorial capacity is predicated on some key assumptions. The first is assuming perfect transmission, and it can be overcome by including the product cb , denoting the vector competence [25, 31, 32]. However, for the purposes of this study, we will also assume perfect transmission. A second assumption is that the bite rate is fixed and constant with age. It is often calculated as the reciprocal of the average length of the gonotrophic cycle [33]. Hence, assuming a gonotrophic cycle of length four, we have $\alpha = 0.25 \text{ days}^{-1}$, i.e. the mosquitoes feed once every four days, which is in line with the experimental setup, where the mosquitoes were allowed to feed every four days. In our calculations, we will use $\alpha = 0.25$ and keep m as an unknown constant. However, we further investigate the remaining parameters.

Extrinsic incubation period (EIP), n . Another notable assumption is the use of a fixed EIP in Eq (1), with the expression $e^{-\mu n}$ representing the probability that a mosquito survives the n -day EIP [34]. However, this can be modified to account for non-fixed EIP. A popular choice is to assume the EIP follows an exponential distribution [25]. If the average EIP duration is the same as with the fixed EIP (n), then the vectorial capacity becomes:

$$C = \frac{m\alpha^2}{\mu} \frac{\sigma}{\sigma + \mu}, \quad (2)$$

where $\sigma = \frac{1}{n}$ is the incubation rate.

As mentioned, the EIP represents the time period where the malaria parasites ingested by the mosquito are developing inside it in order to be able to transmit the infection. The Centers for Disease Control and Prevention states that n is at least 9 days, but is dependent on temperature and the different kind of species of the parasites [35]. In a recently published paper by Stopard *et al.* [36], a mechanistic model fitted to data from [37–40] gave an estimate for the median (50th percentile—EIP₅₀), at 27°C, to be 10.2 days. In addition to the median, the authors provide values for the 10th and 90th percentile (EIP₁₀ and EIP₉₀, respectively), so, using these we can calculate a mean value for the EIP for a given distribution at this temperature.

Taking into account the information on the shape of the EIP, it would be more realistic to consider a different distribution which lies somewhere between the exponentially distributed or fixed EIPs. A mathematically neat choice which meets this criteria and is flexible with the addition of a single extra parameter is the Erlang distribution. The Erlang distribution is a

special case of the gamma distribution, which has been used for the incubation period in many vector-borne disease models [41–45].

The probability density function (PDF) of the Erlang distribution [46] is given by:

$$f(x; k, \lambda) = \frac{\lambda^k x^{k-1} e^{-\lambda x}}{(k-1)!}$$

for $x, \lambda \geq 0$, where k is the shape parameter and λ is the rate parameter. The mean is given by $\frac{k}{\lambda}$, which we set equal to $\frac{1}{\sigma}$. Therefore, with $\lambda = k\sigma$ we have:

$$f(x; k, \sigma) = \frac{(k\sigma)^k x^{k-1} e^{-k\sigma x}}{(k-1)!}. \quad (3)$$

We match EIP₁₀ and EIP₉₀ from [36] to our distribution to infer values for k and σ . We then take $\lfloor k \rfloor$, since the Erlang distribution requires k to be an integer, obtaining $k = 31$. Using this value for k and the value for EIP₅₀ from [36], we finally end up with $\sigma = 0.097$. Thus, we estimate the mean EIP to be 10.3 days.

Mortality rate, μ . The mortality rate is often assumed to be constant, which drastically simplifies mathematical calculations [24]. Nevertheless, it can be seen from our data that this assumption is not the most realistic. Styer *et al.* argue in [20] that ignoring mosquito senescence results in inaccurate predictions with respect to vector control effectiveness. For a more realistic approach, we consider an age-dependent mortality rate. Henceforth, we consider the following functions for μ :

$$\text{Age - Independent : } \mu_{AI}(a) = \mu_{\text{const}} \quad (4)$$

$$\text{Logistic [47] : } \mu_L(a) = \frac{\mu_1}{1 + e^{\mu_2(-a + \mu_3)}} \quad (5)$$

$$\text{Gompertz [23] : } \mu_G(a) = g_1 e^{ag_2} \quad (6)$$

where $a \geq 0$ is the age of the mosquitoes, and the (positive) parameters μ_{const} , μ_1 , μ_2 , μ_3 , g_1 , and g_2 are estimated by fitting the data. These functions are often used in demography and population models, since they describe a mortality that increases with age [22, 48]. The logistic function has an initial exponential increase and then slows down to reach a plateau, whereas the Gompertz continues to increase exponentially. In the field of survival analysis they are often referred to as ‘hazard functions’. For clarity we note that the Gompertz function (Eq (6)) is the hazard function derived from the Gompertz distribution. However, this is not the case for the logistic function (Eq (5)), here this represents a sigmoid curve often referred to as the logistic function.

Survival functions

In order to fit the data, a survival function, $S(a)$, is considered for each mortality function. Its relationship with the mortality function [49] is expressed as:

$$S(a) = \exp\left(-\int_0^a \mu(x) dx\right). \quad (7)$$

Using Eq (7), we obtain the survival function for each of the mortality functions:

$$\text{Age – Independent : } S_{AI}(a) = e^{-a\mu_{\text{const}}} \tag{8}$$

$$\text{Logistic : } S_L(a) = e^{-\mu_1 a} \left\{ \frac{e^{\mu_2(\mu_3 - a)} + 1}{e^{\mu_2\mu_3} + 1} \right\}^{\frac{\mu_1}{\mu_2}} \tag{9}$$

$$\text{Gompertz : } S_G(a) = e^{-\frac{g_1}{g_2}(e^{g_2 a} - 1)} \tag{10}$$

In what follows, we outline how we obtain estimates for the parameters found in these functions, and, additionally, how we calculate confidence intervals for them and the calculations computed in the Results section.

Maximum likelihood estimation. We use the maximum likelihood estimation method to estimate the various parameters present in the survival functions given the experimental data where the k mosquito deaths occur at times $\mathbf{t} = \{t_i; 1 \leq i \leq k\}$. To do so, we calculate the log-likelihood functions (ℓ) for each mortality type (for more information refer to the S1 File). We obtain parameter estimates by maximising the log-likelihood functions.

$$\text{Age – Independent : } \ell_{AI}(\mu_{\text{const}}; \mathbf{t}) = k \log(\mu_{\text{const}}) - \mu_{\text{const}} \sum_{i=1}^k t_i$$

$$\begin{aligned} \text{Logistic : } \ell_L(\mu_1, \mu_2, \mu_3; \mathbf{t}) &= k \log(\mu_1) + k \frac{\mu_1}{\mu_2} \log(1 + e^{\mu_2\mu_3}) - \mu_1 \sum_{i=1}^k t_i \\ &\quad - \left(\frac{\mu_1}{\mu_2} + 1 \right) \sum_{i=1}^k \log(1 + e^{\mu_2(\mu_3 - t_i)}) \end{aligned}$$

$$\text{Gompertz : } \ell_G(g_1, g_2; \mathbf{t}) = k \log(g_1) + k \frac{g_1}{g_2} + g_2 \sum_{i=1}^k t_i - \frac{g_1}{g_2} \sum_{i=1}^k e^{g_2 t_i}$$

We obtain an estimate for μ_{const} by setting $\frac{d\ell_{AI}}{d\mu_{\text{const}}} = 0$. Hence, $\hat{\mu}_{\text{const}} = \frac{k}{\sum_{i=1}^k t_i}$. For the age-dependent functions, the values of the parameters cannot be computed analytically, therefore, we use the Optim.jl [50] package in Julia [51] to minimise the negative log-likelihood.

Wald confidence intervals. The 95% Wald confidence interval of our parameter estimates are given by $\hat{\mu}_{\text{const}} \pm 1.96 \frac{1}{\sqrt{\mathcal{I}(\hat{\mu}_{\text{const}})}}$ for the age-independent case, and $\hat{\theta}_i \pm 1.96 \sqrt{(\mathcal{I}(\hat{\theta}))^{-1}}_{ii}$ for the age-dependent cases. Here, $\mathcal{I}(\hat{\mu}_{\text{const}}) = -\ell''_{AI}(\hat{\mu}_{\text{const}}; \mathbf{t})$ is the observed Fisher information, and $[\mathcal{I}(\theta)]_{ij} = -\mathbb{E} \left[\frac{\partial^2}{\partial \theta_i \partial \theta_j} \ell(\theta; \mathbf{t}) \right]$ is the Fisher information matrix (FIM).

Monte Carlo simulations. We use Monte Carlo simulations to compute 95% confidence intervals for the functions of the parameters reported in the Results section. We do so by simulating 10,000 random samples from the multivariate normal distribution fitted to the parameter estimates. In the multiple parameter cases, we use the estimated parameters as the mean and the inverse of the FIM as the covariance matrix. We then take the quantile values at 2.5% and 97.5% to obtain marginal confidence intervals of the functions of interest.

Results

Using the data and the methods outlined above, we obtain estimates for the unknown parameters in the mortality functions so that we can calculate the vectorial capacity. The parameter estimates are shown in Table 1. Using these values and our Monte Carlo simulations, we plot

Table 1. Maximum likelihood estimates for the parameters.

	μ_1	μ_2	μ_3
Control	0.193 ± 0.0854	0.132 ± 0.0331	29.5 ± 7.02
Treated	0.265 ± 0.180	0.253 ± 0.128	14.7 ± 6.78
	g_1	g_2	μ_{const}
Control	0.0094 ± 0.00275	0.0693 ± 0.00988	0.0383 ± 0.00530
Treated	0.0137 ± 0.00447	0.136 ± 0.0199	0.0671 ± 0.00930

95% Wald confidence intervals for the parameter values (3 significant figures) obtained through maximum likelihood estimation for the control and treated cases.

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the survival functions in Fig 3 and the mortality functions in Fig 4, comparing each treatment. Note that on the survival plots we add the Kaplan-Meier estimator, a non-parametric method frequently used in survival analysis, which we obtain using the Survival.jl package.

In order to investigate the differences in the vectorial capacity between the control and treated cases, and between the various mortality functions, we need to reconsider the way the vectorial capacity is calculated. In the results that follow, the uncertainty comes from the parameter estimates from the mortality functions (see Table 1). All other parameters included in the calculations have fixed values as described in the Methods section.

Rethinking the vectorial capacity

We outline the calculations required to obtain the expected number of bites a mosquito takes after being infected and show how this is linked to the vectorial capacity. This will give a more realistic value for C. To aid our calculation of the expected number of infectious bites, we pose the following four questions:

1. What is the probability of a mosquito surviving the EIP given an infectious blood-meal is taken at age a_0 ?
2. How many bites will the mosquito take if it has survived the EIP?
3. What is the expected number of infectious bites a mosquito takes in its lifetime if it has taken an infectious blood-meal at age a_0 ?
4. What is the expected number of infectious bites a mosquito takes in its lifetime?

Fig 5 depicts a timeline of these events for visualisation purposes. We attempt to answer these questions in four steps. In each step we explore the three different mortality functions: CASE (I) being the age-independent (Eq (4)), CASE (II) the logistic (Eq (5)), and CASE (III) the Gompertz (Eq (6)), and subsequently compare their results. More detailed calculations can be found in the S2 File.

Step 1: $\mathbb{P}(\text{mosquito survives EIP} \mid \text{infectious blood-meal at } a_0)$.

$$\begin{aligned} \text{CASE (I)} : \quad & \mathbb{P}(\text{surviving EIP} \mid \text{infectious blood-meal at age } a_0) \\ &= \left(\frac{k\sigma}{k\sigma + \mu_{const}} \right)^k \end{aligned} \tag{11}$$

$$\begin{aligned} \text{CASE (II)} : \quad & \mathbb{P}(\text{surviving EIP} \mid \text{infectious blood-meal at age } a_0) \\ &= \int_{t=0}^{\infty} \frac{(k\sigma)^k t^{k-1} e^{-k\sigma t}}{(k-1)!} e^{-\mu_1 t} \left\{ \frac{e^{\mu_2(\mu_3 - (a_0+t))} + 1}{e^{\mu_2(\mu_3 - a_0)} + 1} \right\}^{\frac{\mu_1}{\mu_2}} dt \end{aligned} \tag{12}$$

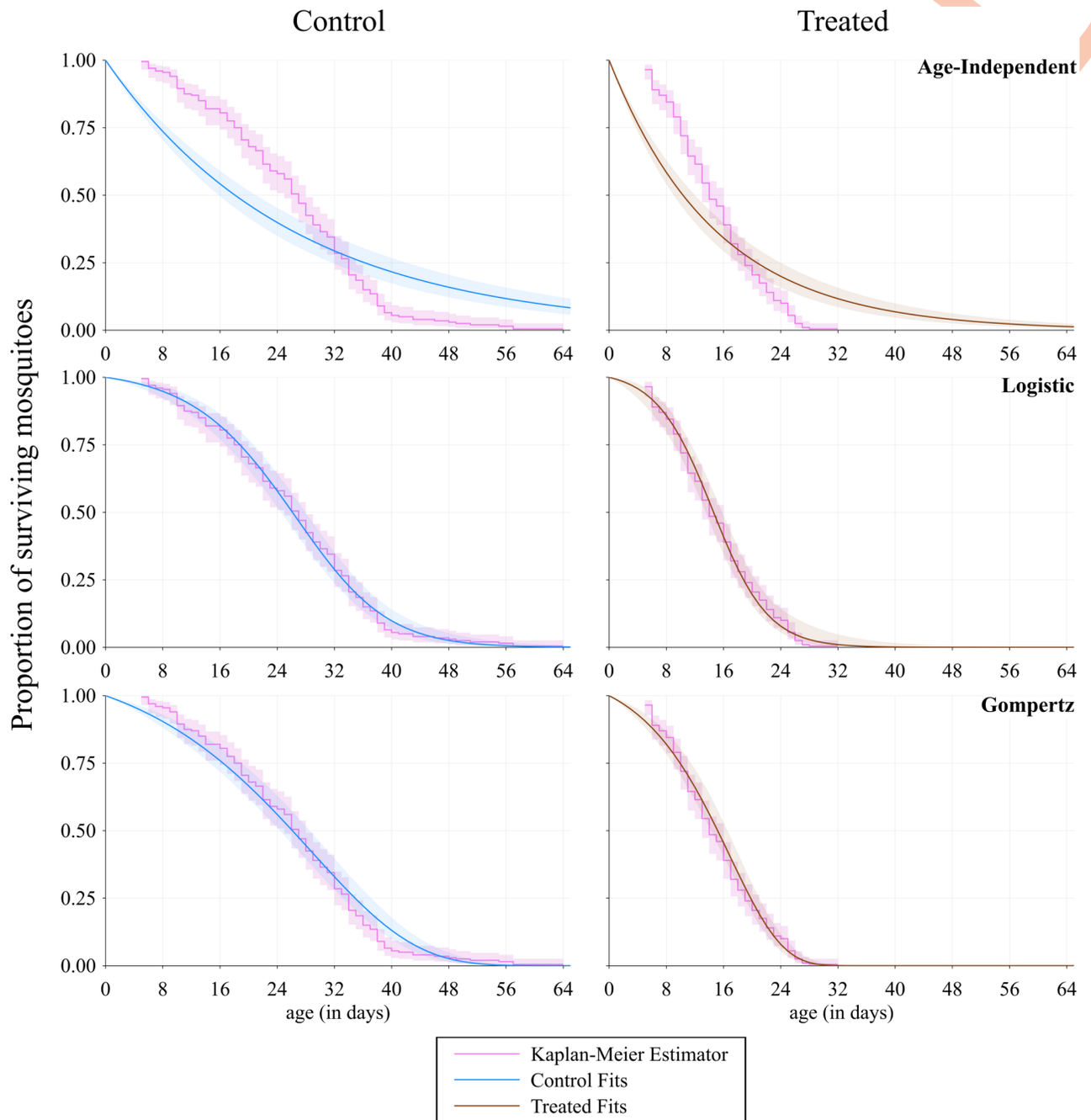


Fig 3. Fitted survival functions. Plots for the proportion of surviving mosquitoes at each day for the control (non-exposed) and treated (exposed) cases using the three different survival functions (age-independent, logistic, Gompertz). In the treated column, all mosquitoes are dead by day 33. The fits are extended to day 64 for a better comparison between the two treatments. The shaded area around the control and treated curves represents the 95% confidence interval due to the error propagated from the parameter estimates in each function using Monte Carlo simulations. The shaded area around the Kaplan-Meier estimator represents the pointwise log-log transformed 95% confidence interval.

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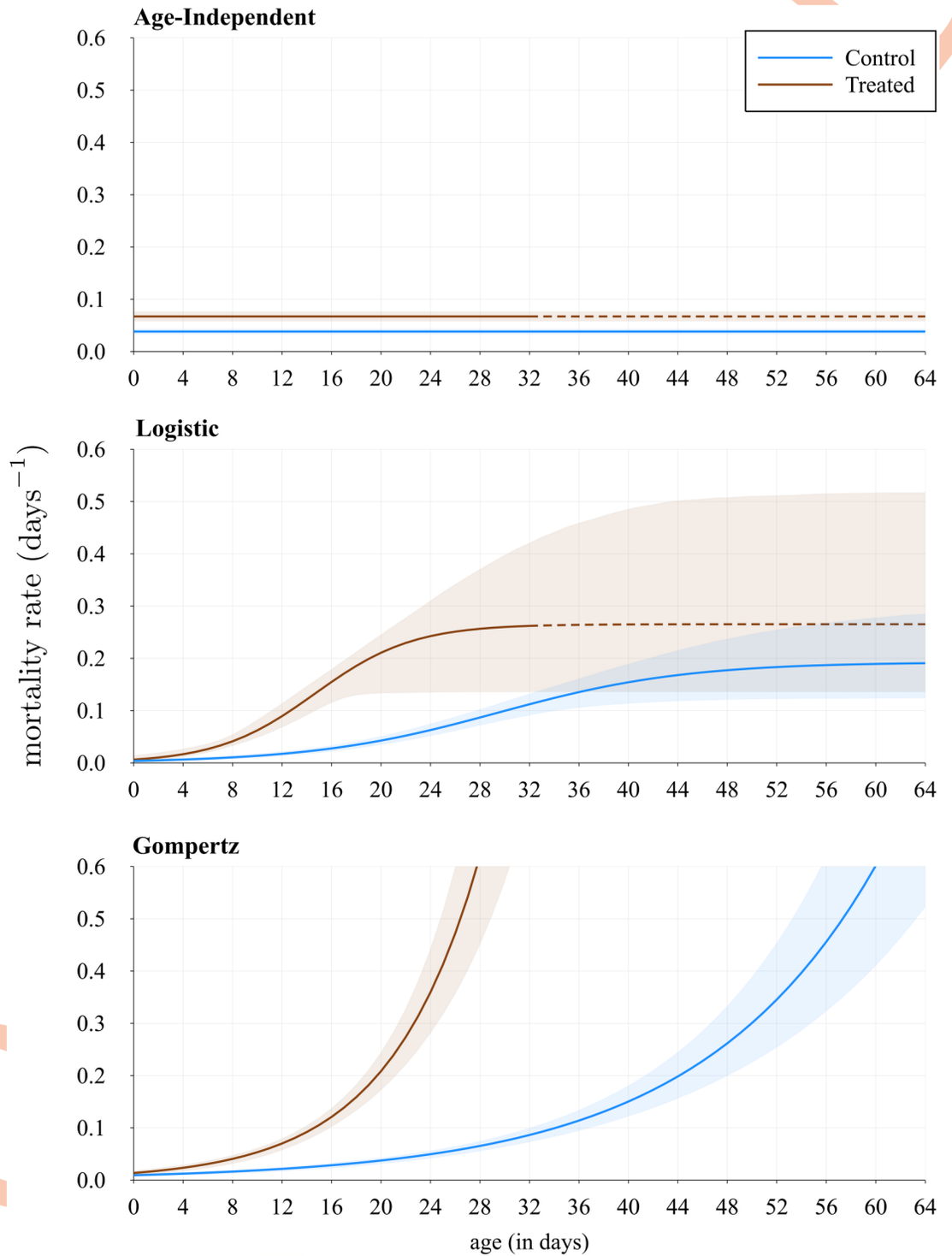


Fig 4. Comparison of mortality for the two treatments. The dashed line represents where all mosquitoes are already dead in the experiment for the treated case. The shaded area represents the 95% confidence interval due to the error propagated from the parameter estimates in each function using Monte Carlo simulations.

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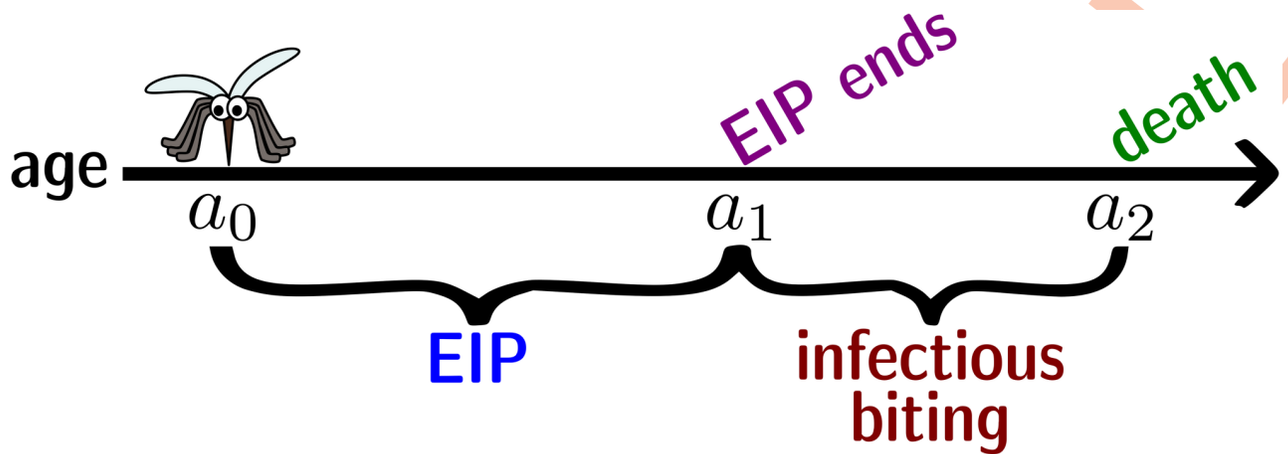


Fig 5. Mosquito timeline after taking an infectious blood-meal. We assume that a mosquito takes an infectious blood-meal at age a_0 . In order for it to become infectious, it must survive the EIP. After surviving the EIP, at age a_1 , the mosquito will take infectious blood-meals up until its death, at a_2 . [Note: the mosquito might not survive the EIP, hence, it is possible that $a_2 < a_1$.] The mosquito clip-art used to produce this figure was obtained from www.clker.com under the CC0 1.0 Universal Public Domain Dedication license.

<https://doi.org/10.1371/journal.pcbi.1009540.g005>

$$\begin{aligned} \text{CASE (III) : } & \mathbb{P}(\text{surviving EIP} \mid \text{infectious blood-meal at age } a_0) \\ &= \int_{t=0}^{\infty} \frac{(k\sigma)^k t^{k-1} e^{-k\sigma t}}{(k-1)!} e^{-\frac{\delta_1}{\delta_2} e^{a_0 \delta_2} (e^{\delta_2 t} - 1)} dt \end{aligned} \quad (13)$$

CASES (II) and (III) cannot be solved analytically, but we integrate them numerically using the QuadGK.jl package in Julia. The results for the three cases are plotted and shown in Fig 6.

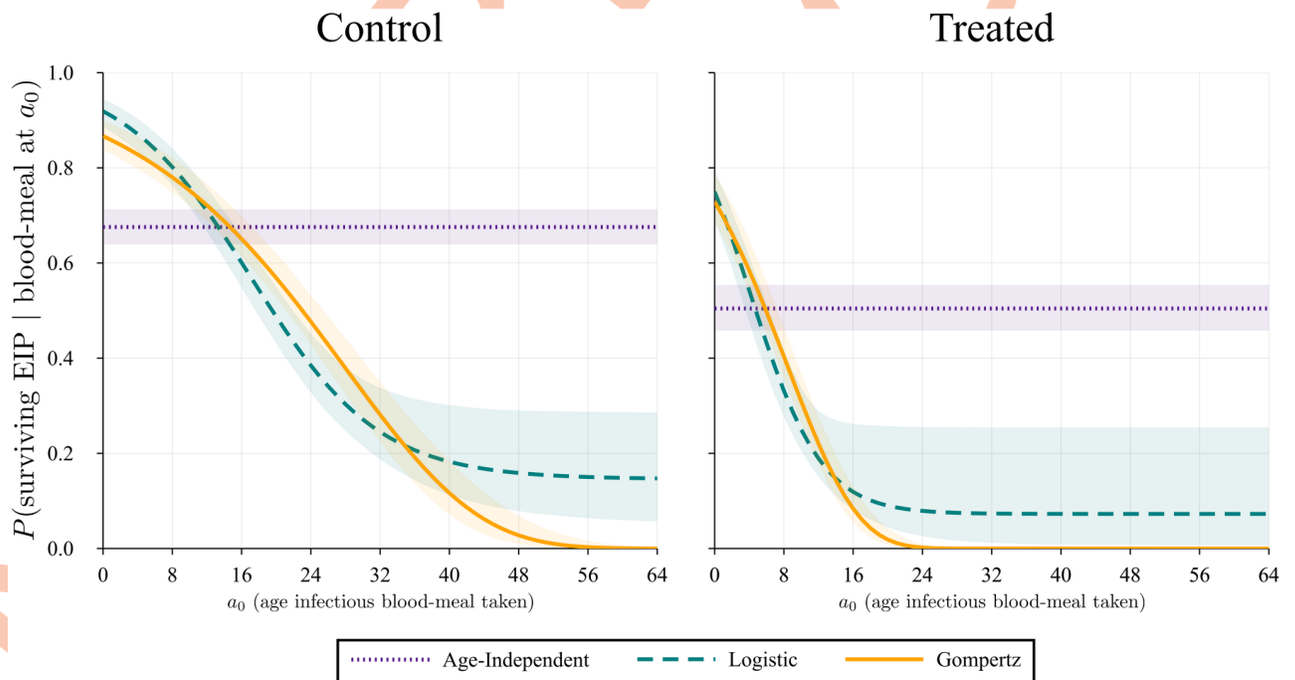


Fig 6. Probability the mosquito survives the extrinsic incubation period given that it takes an infectious blood-meal at age a_0 . The plots show the values of Eqs (11), (12) and (13) over different ages. The shaded area represents the 95% confidence interval due to the error propagated from the parameter estimates using Monte Carlo simulations.

<https://doi.org/10.1371/journal.pcbi.1009540.g006>

From this first step we notice that the age-independent case is not biologically realistic. Comparing it with the age-dependent cases, we can see that as a_0 increases, the probability a mosquito will survive the EIP decreases significantly, which makes sense, given that there is evidence mosquitoes senesce both in the data used here, but also from Styer *et al.* and Ryan *et al.* [20, 24]. We also notice that the Gompertz function’s curve approaches zero faster, which could be somewhat more realistic, whereas the logistic function reaches a plateau above zero. Comparing the two treatments, control and treated, we notice that the probability is lower initially with treatment, but also that the age-dependent functions approach zero much faster, which is in line with the mosquitoes having a lower life expectancy in the treated case (Fig 3).

Step 2: $\mathbb{P}(z = j \mid \text{mosquito exits EIP at } a_1)$. We are interested in the probability mass function (PMF) of the number of bites, z , supposing the mosquito exits the EIP at age a_1 , and dies at age a_2 .

$$\text{CASE (I) : } \mathbb{P}(z = j \mid \text{exits EIP at age } a_1) = \frac{\mu_{\text{const}} \alpha^j}{(\mu_{\text{const}} + \alpha)^{j+1}} \tag{14}$$

$$\begin{aligned} \text{CASE (II) : } \mathbb{P}(z = j \mid \text{exits EIP at age } a_1) &= \int_{a_2=a_1}^{\infty} \frac{\alpha^j (a_2 - a_1)^j}{j!} e^{-(\alpha+\mu_1)(a_2-a_1)} \\ &\quad \frac{\mu_1}{1 + e^{\mu_2(\mu_3-a_2)}} \left\{ \frac{e^{\mu_2(\mu_3-a_2)} + 1}{e^{\mu_2(\mu_3-a_1)} + 1} \right\}^{\frac{\mu_1}{\mu_2}} da_2 \end{aligned} \tag{15}$$

$$\begin{aligned} \text{CASE (III) : } \mathbb{P}(z = j \mid \text{exits EIP at age } a_1) &= \int_{a_2=a_1}^{\infty} \frac{\alpha^j (a_2 - a_1)^j}{j!} e^{a_2 g_2 - (a_2 - a_1) \alpha} g_1 e^{\frac{g_1}{g_2} (e^{a_2 g_2} - e^{a_1 g_2})} da_2 \end{aligned} \tag{16}$$

CASES (II) and (III) are again solved numerically. We plot some heatmaps to visualise the solutions of Eqs (14), (15), and (16) (Fig 7). In the plots we include the average number of bites, which is calculated using:

$$\mathbb{E}(z \mid \text{exits EIP at age } a_1) = \sum_{j=0}^{\infty} [j \times \mathbb{P}(z = j \mid \text{exits EIP at age } a_1)].$$

We also plot the PMFs for a specific age ($a_1 = 15$) which can be found in S3 Fig.

In Fig 7 we can see that, for the age-dependent cases, the older the mosquito is when it exits the EIP, the higher the probability that the number of bites it takes is small. The two age-dependent functions give similar results, but the age-independent function shows again that it is not biologically realistic, since it has a constant average and a constant probability across all ages. In the treated case, there is a higher probability that there are low, or even zero, bites compared to the control case; this happens for all choices of mortality function.

Step 3: $\mathbb{E}(z \mid \text{infectious blood-meal at } a_0)$. We can now calculate the expected number of infectious bites a mosquito takes, given that it takes an infectious blood-meal at age a_0 :

$$\mathbb{E}(z \mid \text{blood-meal at } a_0) = \sum_{j=0}^{\infty} [j \times \mathbb{P}(z = j \mid \text{blood-meal at } a_0)] \tag{17}$$

Hence, we need to calculate the PMF of the number of bites, given that an infectious blood-meal is taken at age a_0 . To do so, we use the results from Steps 1 and 2. For example, in CASE (I) we multiply the results from Eqs (11) and (14), however care must be taken for when $j = 0$,

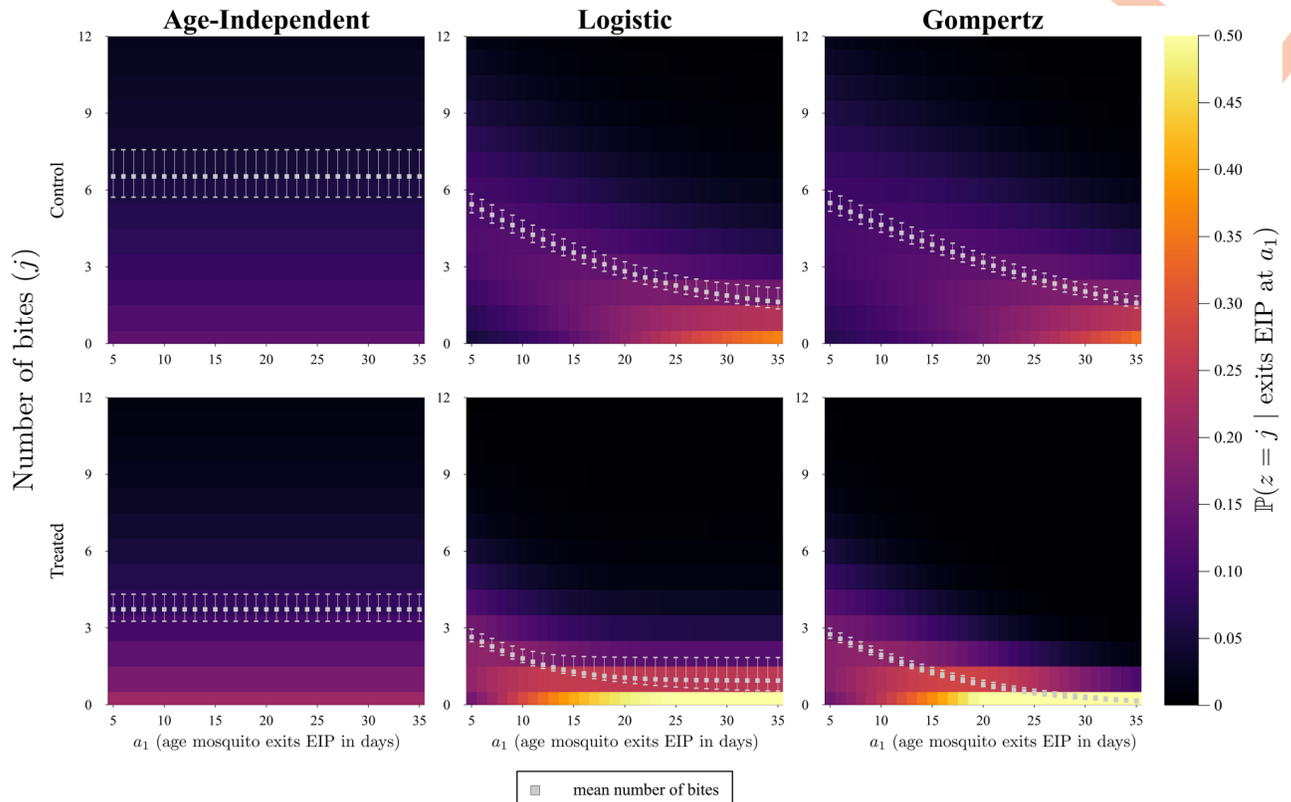


Fig 7. Heatmaps of the probability the number of bites is equal to some j given that the mosquito exits the extrinsic incubation period (EIP) at age a_1 . The heatmaps are obtained from the fitted parameters. The error bars on the mean markers represent the 95% confidence interval due to the propagated error for the fitted parameters using Monte Carlo simulations.

<https://doi.org/10.1371/journal.pcbi.1009540.g007>

where we need to consider that we definitely have zero bites if the mosquito does not survive the EIP.

CASE (1):

$$\mathbb{P}(z = j \mid \text{blood - meal at } a_0) = \begin{cases} \left(\frac{k\sigma}{k\sigma + \mu_{\text{const}}}\right)^k \frac{\mu_{\text{const}}^\alpha z^j}{(\mu_{\text{const}} + \alpha)^{j+1}} & \text{if } j \neq 0 \\ \left(\frac{k\sigma}{k\sigma + \mu_{\text{const}}}\right)^k \frac{\mu_{\text{const}}^\alpha}{\mu_{\text{const}} + \alpha} + \left(1 - \left(\frac{k\sigma}{k\sigma + \mu_{\text{const}}}\right)^k\right) & \text{if } j = 0 \end{cases}$$

Therefore

$$\mathbb{E}(z \mid \text{infectious blood - meal at } a_0) = \frac{\alpha}{\mu_{\text{const}}} \left(\frac{k\sigma}{k\sigma + \mu_{\text{const}}}\right)^k. \tag{18}$$

CASES (II) and (III):

$$\begin{aligned} &\mathbb{P}(z = j \mid \text{blood - meal at } a_0) \\ &= \int_{a_1=a_0}^{\infty} [\text{Eq (15) or Eq (16)}] \\ &\quad \frac{(k\sigma)^k (a_1 - a_0)^{k-1} e^{-k\sigma(a_1 - a_0)}}{(k-1)!} \exp\left(-\int_{a_0}^{a_1} \mu(x) dx\right) da_1 \end{aligned} \tag{19}$$

where we use Eq (15) for CASE (II) and Eq (16) for CASE (III). The above is true for $j \neq 0$. When $j = 0$, we must add $(1 - \text{Eq (12)})$ to Eq (19) for CASE (II), or $(1 - \text{Eq (13)})$ for CASE (III), following the same logic as in CASE (I). This is again integrated numerically (this time using the Cuba.jl [52, 53] package in Julia) and put into Eq (17) to obtain the required solution. The results are depicted in Fig 8 for both treatments.

The expected number of bites decreases significantly if we consider an age-dependent mortality function, as we can see in Fig 8. There is also an obvious difference between the two treatments, where in the treated case the number of bites are a lot lower to begin with. Once again the logistic function seems to be plateauing above zero.

Step 4: $\mathbb{E}(z)$. We can now use the previous steps to calculate the expected number of infectious bites a mosquito will take in its lifetime:

$$\begin{aligned} \mathbb{E}(z) &= \int_{a_0=0}^{\infty} \mathbb{E}(z \mid \text{infectious blood - meal at } a_0) \\ &\quad \times \mathbb{P}(\text{infectious blood - meal at } a_0) da_0 \end{aligned} \tag{20}$$

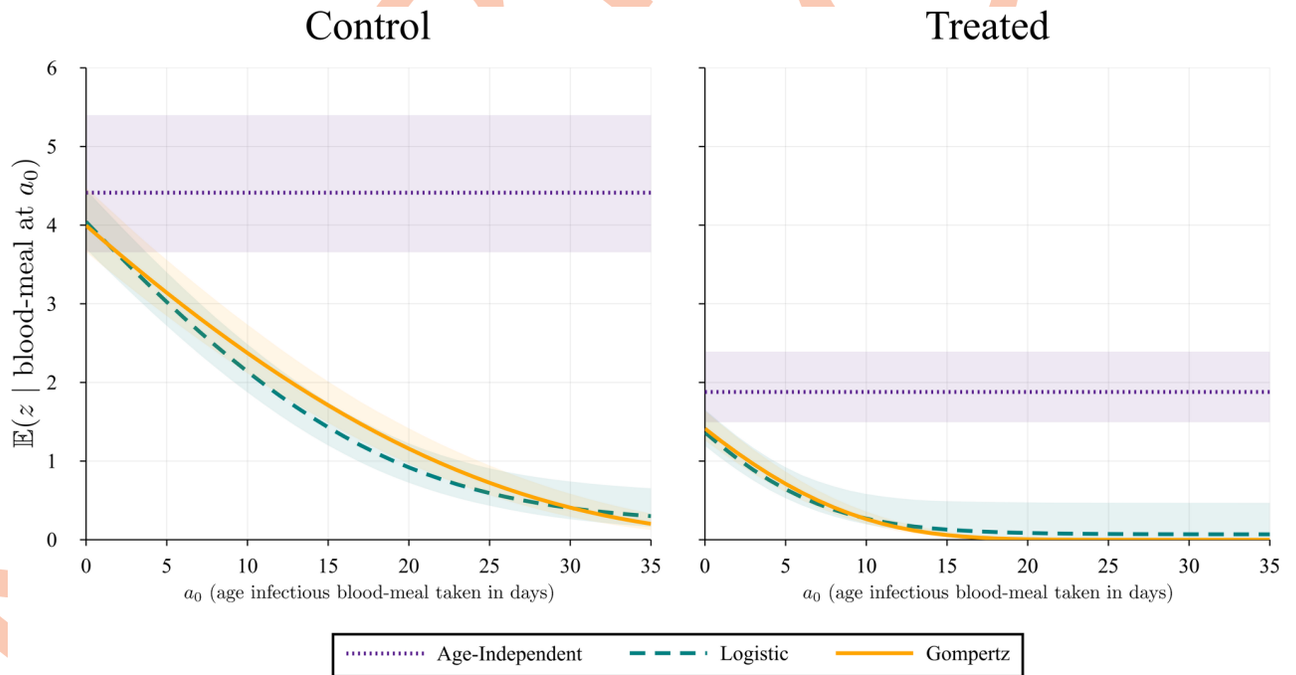


Fig 8. The expected number of bites a mosquito will take in its lifetime given it has taken an infectious blood-meal at age a_0 . The shaded area represents the 95% confidence interval due to the error propagated from the parameter estimates using Monte Carlo simulations.

<https://doi.org/10.1371/journal.pcbi.1009540.g008>

CASE (I):

$$\mathbb{E}(z) = \frac{\alpha}{\mu_{\text{const}}} \left(\frac{k\sigma}{k\sigma + \mu_{\text{const}}} \right)^k$$

which, as expected, is the same as Eq (18) as it does not depend on age. Looking back at the vectorial capacity (Eqs (1) and (2)), we can see that this is represented there by $\frac{\alpha}{\mu} e^{-\mu m}$ and $\frac{\alpha \cdot \sigma}{\mu \cdot \sigma + \mu}$ respectively. This is because the assumed EIP distribution in each case is different (fixed and exponentially distributed), whereas we have assumed an Erlang distribution.

We solve Eq (20) for CASES (II) and (III) numerically and obtain a single number for each. The results for all cases and both treatments can be found in Fig 9. We observe that the expected number of bites is lower if we consider age-dependent mortality functions, where we have 3.26 (2.96, 3.64) for the logistic function and 3.33 (3.03, 3.76) for the Gompertz, versus 4.41 (3.66, 5.40) for the age-independent. With treatment, the numbers are even lower for all cases, where for the age-dependent functions, the expected number of bites is <1.

Using all of the above, we can calculate the relative difference in the vectorial capacity between the control and treated cases for the age-dependent and age-independent mortality functions. We do so by assuming that the mosquito density, m , and the bite rate, α , are constant. Hence we can use the results from Eq (20) to make our comparisons (Fig 10).

Between the control and treated cases for each function there is a significant reduction in the vectorial capacity, where the percentage for the age-independent treated case relative to the control is at 42.6% (40.9, 44.3), or, equivalently, there is a reduction of 57.4% (55.7, 59.1). However, we also notice that the age-dependent functions have a bigger decrease: the difference in the reduction percentages between the age-independent and the logistic cases is ≈ 16.32 and between the age-independent and the Gompertz cases is ≈ 15.31 . In all previous calculations, we have seen that the age-independent function is not very realistic, whereas the logistic and

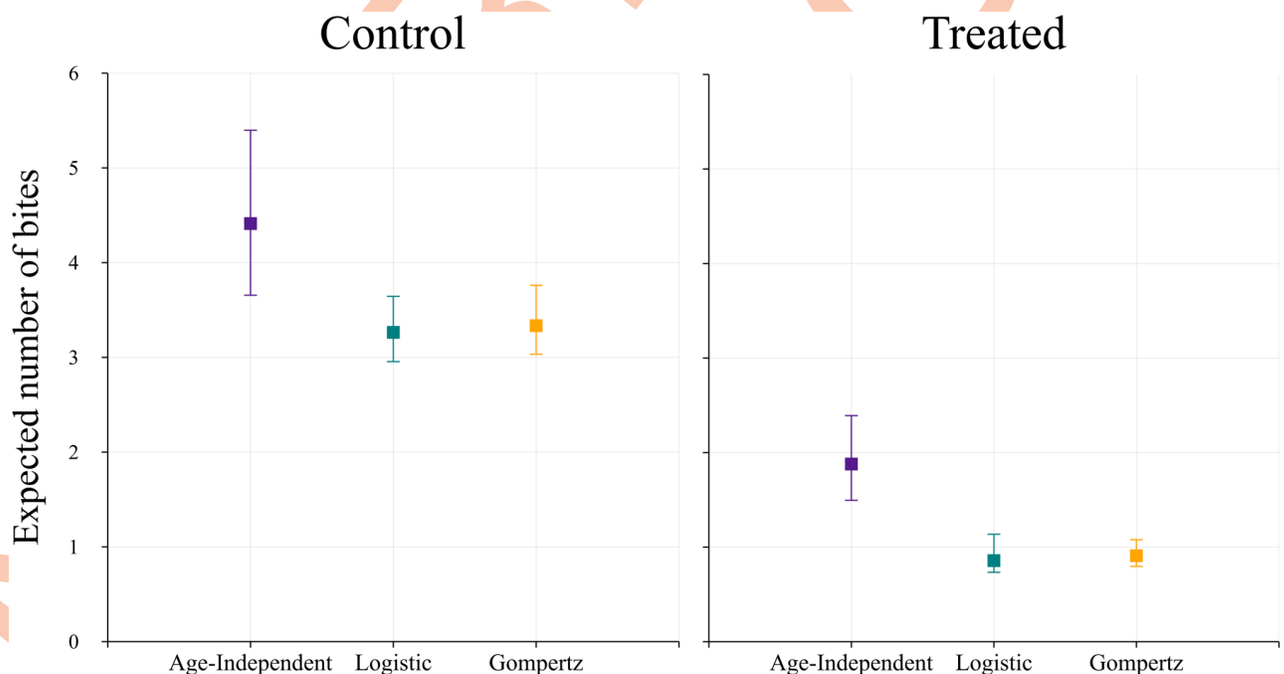


Fig 9. The expected number of infectious bites a mosquito will take in its lifetime. The error bars represent the 95% confidence interval due to the error propagated from the parameter estimates through Monte Carlo simulations.

<https://doi.org/10.1371/journal.pcbi.1009540.g009>

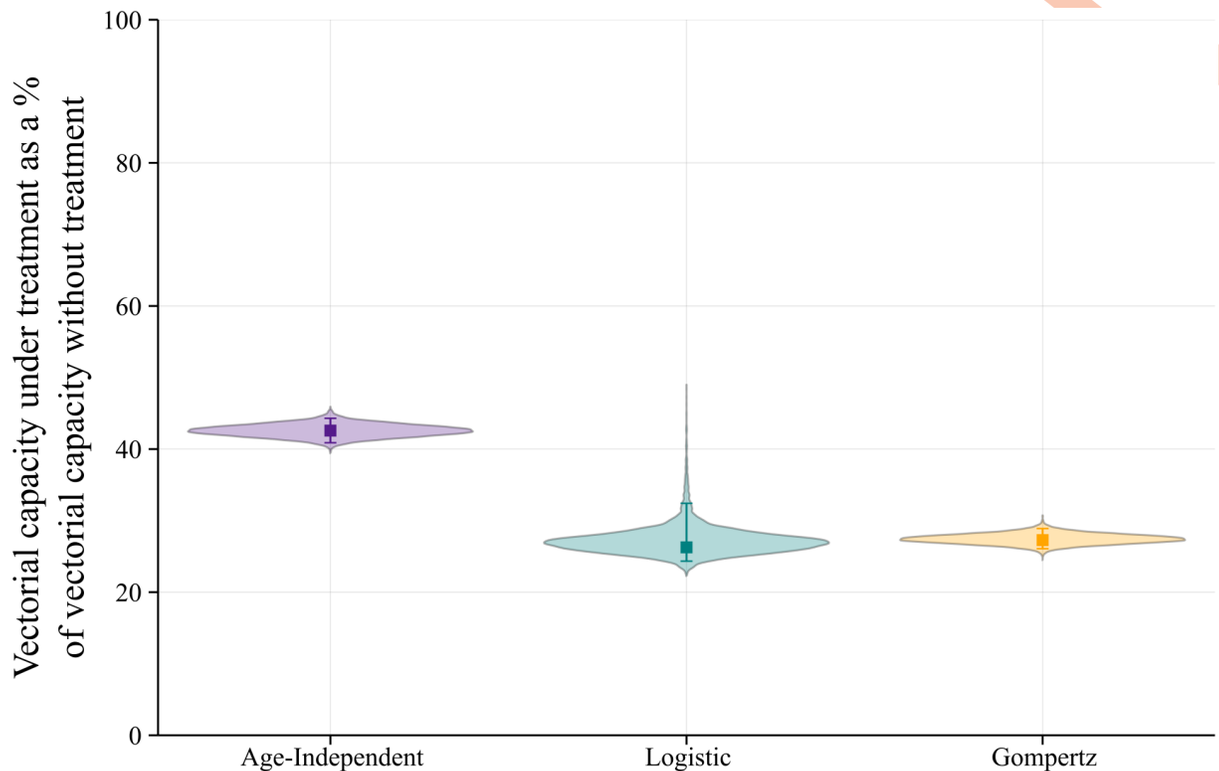


Fig 10. Violin plots showing the relative difference in the vectorial capacity with treatment. The figure shows the full distribution of the results for each function obtained from the Monte Carlo simulations. The error bars represent the 95% confidence interval due to the propagated error of the parameter estimates using Monte Carlo simulations.

<https://doi.org/10.1371/journal.pcbi.1009540.g010>

Gompertz functions behave as we might expect from our understanding of vector senescence. The Gompertz function might be slightly closer to biological realism, however the difference between the two is minimal. Both functions seem to be good candidates, though we might have a stronger preference for using the Gompertz function in future calculations, since it only requires two fitted parameters, compared to three in the logistic function.

Discussion

Mathematical modelling can contribute by recommending improved or optimised intervention strategies for various (vector-borne) diseases, but there are many challenges that modelers have to overcome to provide policy-relevant insights [54, 55]. One must find a balance between creating easy-to-use models and ensuring that biological simplifications do not alter the resultant policy recommendations by over or underestimating the impact of different intervention measures. One such challenge for modelling malaria is to accurately quantify the mortality of the mosquitoes. In the present study, we have demonstrated that using age-dependent mortality functions is important; the EIP is typically long compared to a mosquito's life-span and combined with senescence this has a substantial impact on our calculation of the vectorial capacity. Here, the age-independent assumption provides us with a conservative estimate for the impact of ITNs. This could result in less appropriate recommendations for policy decisions, especially if constant mortality is used in cost-effectiveness analyses or to optimise other policy objectives (e.g. selecting strategies to minimise costs, deaths, etc.).

This study echoes results from other studies (e.g. [22, 23, 25]), which advocate for incorporation of more biological realism in vector-borne disease modelling. Brand *et al.* [25] consider different distributions for the EIP to show how R_0 changes. On the other hand, Bellan and Novoseltsev *et al.*, [22, 23] respectively, highlight the importance of age-dependent mortality. In these studies, however, the traditionally fixed EIP is considered, and the calculations for the vectorial capacity are approached from a different perspective. Bellan [22] incorporates a fixed parameter for the impact of insecticides on the longevity of the mosquitoes, due to the lack of real-life data. Similarly, in [23], the authors consider multiple patterns for age-dependent mortality which are generalised for multiple vector species. A strength of our approach is to be able to compare and contrast the impact of vector ageing assumptions on vectorial capacity based on survival data. We note that age-dependent or age-specific mortality rates are not an uncommon addition in agent-based models of malaria, as we can see in [56–58], but are typically omitted in compartmental-type models. We have shown that without accounting for age the effectiveness of the anti-vectorial intervention against highly-pyrethroid-resistant *An. gambiae s.l.* mosquitoes would be underestimated; therefore it seems prudent to include vector senescence across agent-based and compartmental models. This highlights the clear need for better availability of real-world data for different vector-parasite systems to improve modelling predictions.

The two age-dependent functions that are explored in this paper are often used for mosquito survival analysis [20, 22, 23, 48]. Styer *et al.* investigated a large-scale mortality study using *Aedes aegypti* mosquitoes and concluded that the logistic mortality functions fit the data better on most occasions except one, where the Gompertz function was a better fit [20]. Clements and Paterson explored the survival patterns in many different mosquito species and found that their patterns are explained well by using a Gompertz function [48]. Although the logistic and Gompertz are rather flexible functions, other functional forms may be more suitable for different vector species—in this case the general framework presented in here could be readily adapted if required.

We observe an interesting feature of the mosquito survival data for the control case where suddenly the Kaplan-Meier curves flatten from day 40. One hypothesis not explored in this study is that this could arise if the mosquito population is comprised of two sub-populations with different mortality rates. A model with frailty—one sub-population with higher mortality rate and one with lower—could fit the data more closely around this 40-day mark, although our fits (see Fig 3) do capture the survival tail fairly well, especially in the case of the logistic mortality function. One argument against implementing a frailty model here is the number of parameters; partitioning the population into high and low groups has at least two extra parameters—one for the proportion of the population in the low group, and one for relative scaling of the mortality rate. Frailty seems to be the most intuitive explanation for this long tail, however there could be other plausible mechanisms or this could be attributed to slightly noisy data. Ultimately, we have chosen to avoid added extra parameters and risk “overfitting” a model. For the purposes of arguing that age-independent models fitted to lab data produce substantially different results to age-dependent models fitted to the same data, the current functions seem to be sufficient, however exploring frailty could be an interesting future avenue of research.

It is important to note that we have included the assumption of perfect transmission when considering the vectorial capacity. In reality, vector competence and host prevalence does affect the transmission potential. Here, we opted for perfect transmission as we could not estimate the vector competence from our data and field prevalence will be variable depending on factors such as the relative vector to host density. We believe that including a realistic value for vector competence and host prevalence in our calculations (as part of the step 4 calculation)

would likely show that there is a further proportional decrease in the vectorial capacity when comparing age-dependent to age-independent mortality rates, because the distribution describing the timing of the first infectious blood meal (a_0) will shift to older ages and this has a more pronounced effect when there is age-dependent mortality (see Fig 8).

The data and results show there is a significant increase in mosquito mortality when several blood-meal opportunities are offered by a host protected with an ITN vs. an untreated net. The consequence for mosquitoes is a reduced vectorial capacity induced by the presence of an insecticide. The fact that standard ITNs do not immediately kill young and unfed mosquitoes following a single and forced exposure to insecticide (which is the standard susceptibility test by WHO [30]) is not sufficient information to assess the efficacy of ITNs against the malaria vectors. In fact, when the EIP and mortality rates are taken into consideration, along with multiple exposures (which is more in line with free-flying mosquitoes that are regularly host-searching and feeding), the end result is that ITNs still retain some functionality against resistant mosquitoes and work better than untreated nets. These results are in accordance with other publications [10, 12, 13]. The experiments in this study were in a laboratory setting, so there may be an underestimation of the effect of insecticide in the field, in which case the effect on the vectorial capacity could be greater. We agree with previous authors, [8, 13, 59, 60], that encourage an update in the way ITN efficacy and resistance are measured. For example, mosquito condition (i.e. one or more blood-feeds), age (we especially care about old mosquitoes and it has been shown previously that resistance declines with age [61]), and exposure history (multiple exposures over time since mosquitoes can encounter nets at each gonotrophic cycle) are some of the factors which combined determine the overall number of mosquitoes in a cohort potentially able to transmit malaria.

There are many unanswered questions regarding the behaviour of mosquitoes. One of these relates to feeding and biting patterns. In our model, we have assumed a constant biting rate of one bite every four days on average. We could argue that we could use a different value for each treatment. However, this was not included here since additional data is required to make appropriate estimations for a biting rate. Given that the experiment here gave access to the mosquitoes for feeding every four days, it seems more appropriate to keep this assumption. Having daily access to feeding could give more insight to estimating another biting rate. Furthermore, it would be interesting to investigate feeding and biting patterns that depend on age. Furthermore, it is important to note that the nets in this experiment were new and intact to better understand the impact of maximum insecticide exposure on longevity of resistant mosquitoes. This approach follows numerous WHO lab and field bio-efficacy protocols [30]. In realistic control settings in malaria-affected regions, the ITNs encountered by mosquitoes are likely more variable and will change as the nets age and become damaged or lose insecticide due to washing. Understanding the functional impact of older, 'imperfect' nets, including the effects on age-dependent mortality as highlighted in the current manuscript, is an important area for further research [8, 60].

There have also been studies where it is suggested that there are parasite-induced behavioural changes [62–65]. This could mean that the feeding and biting patterns of the mosquitoes change significantly before, during, and after the EIP. For example, mosquitoes, having survived the EIP, could bite multiple humans to complete one blood-meal, potentially transmitting the infection to more than one person. Shaw *et al.* found evidence that the EIP can be shortened if an infected mosquito feeds an additional time [66]; this could mean that if mosquitoes feed during the EIP and the EIP shortens, the result is more infectious blood-meals and a larger vectorial capacity, hence an increase in malaria transmission. Therefore, it could be useful to explore other feeding patterns and bite rates.

For our calculations, we have used the Erlang distribution for the EIP. It is important to note that the mortality data were collected at $26\pm 1^\circ\text{C}$, whereas the values we obtained for the EIP come from data at 27°C in [36]. Data collected at various temperatures capturing the full distribution of the EIP would be extremely useful. The EIP is affected by many factors, as shown by Ohm *et al.* [67], where it is emphasised that transmission models can be improved if we have a better understanding of the EIP. Some studies focus on a temperature-dependent EIP [68–70], however, given that the experiment our data were collected from was at constant temperature conditions, we have not included this here. Nevertheless, it is important to keep in mind that some factors depend on temperature in real life, so control programmes might need adjustments depending on the time of year. Incorporating temperature dependency is something else that can be explored in the future, following in the footsteps of studies like [69] and [71].

In this study we focused on the mortality rate and how to include age dependency in the model. The data from the experiment allowed us to estimate the various parameters related to the mortality rate. However, the other parameters we included in our calculations, i.e. the bite rate and the parameterisation of the EIP distribution, were fixed values with no possibility to estimate them from our data. Hence, in our calculations, the only source of uncertainty included is coming from our mortality parameters. It is worth noting that the uncertainty in our vectorial capacity calculations is likely to be greater, had we allowed for variability in these other parameters as well.

Furthermore, to truly capture the whole picture of malaria in Côte d'Ivoire, bringing together data for human malaria cases and other on-going control strategies with this mosquito data would help calculate the human consequences for malaria transmission and control. This could be done by constructing a modified Ross–Macdonald-type host-vector disease model [72], matching it to the data, and concurrently incorporating an age-dependent mortality rate and an Erlang-distributed EIP for the vectors.

Conclusion

In this paper, we have used a modelling framework to investigate the impact of insecticide exposure on mosquitoes and their ability to transmit malaria, along with the impact of age-dependent mortality. Firstly, our results suggest that the mortality rates increase due to insecticide exposure even in mosquitoes classified as highly resistant following the WHO definition. Our analysis found that under a control (no insecticide exposure) scenario there would be a higher expected number of infectious bites by mosquitoes than under a treated scenario (with insecticide exposure). The vectorial capacity is substantially reduced when the mosquitoes are exposed to ITNs based on the experiment conducted, despite being resistant to the pyrethroids used on the nets. In addition, if age dependency is included in our model, the expected number of infectious bites is predicted to have a greater relative reduction by using insecticides than if we use constant mortality.

Without detailed vector data on survival with and without insecticides, this type of modelling analysis would not be possible. We strongly advocate for collection, not only of average mosquito life expectancies, but also distributions of survival for other vector-parasite systems where quantitative analyses of different interventions against the disease are desirable. We also suggest that modellers pay close attention to whether more could be done to factor in senescence into vector-borne disease strategy evaluations.

The above methodology could be easily used to check the insecticide resistance of mosquitoes from experiments using other pyrethroids and/or mosquito species, if similar experimental data on mosquito survival were available. The results could then be used to further examine how age dependency impacts the effectiveness of various interventions against mosquitoes.

Supporting information

S1 Table. Cumulative mortality data from the control treatment.

(XLSX)

S2 Table. Cumulative mortality data from the treated treatment.

(XLSX)

S3 Table. Feeding data from the control treatment.

(XLSX)

S4 Table. Feeding data from the treated treatment.

(XLSX)

S5 Table. Survival data from the control treatment.

(XLSX)

S6 Table. Survival data from the treated treatment.

(XLSX)

S7 Table. Data used for Kaplan Meier estimator (control).

(XLSX)

S8 Table. Data used for Kaplan Meier estimator (treated).

(XLSX)

S1 Fig. Proportion of total fed mosquitoes out of the alive ones for each replicate (access to foot is allowed every four days). The results from Replicate 1 are a lot different than the rest of the replicates. On the control plot (left), even at the beginning where many mosquitoes are still alive, the proportion of those that were fed is (close to) zero, and only at the very end a mosquito actually feeds. Similarly, on the right, almost all of the mosquitoes go through their lifetime without feeding.

(PDF)

S2 Fig. Proportion of alive mosquitoes for each replicate (deaths recorded daily). In both plots Replicate 1 seems different to the trend followed by the other replicates, especially for the control case (left). This could be explained by the trends in [S1 Fig](#).

(PDF)

S3 Fig. The probability mass function of the number of bites given the mosquito exits the extrinsic incubation period at age 15. The average number of bites for each treatment is found in the legend box of each plot. The error bars represent the propagated uncertainty of the estimated parameters. We can see that the highest probabilities are for the smaller values of j in all cases. The probability the number of bites is closer to zero is higher in the treated than in the control cases. The average number of bites is higher for the control treatment for all three cases, as expected. We also notice that for the treated case and the age-dependent functions the probability that the number of bites is equal to anything above seven is essentially zero. However, for the age-independent graph this probability goes to zero for a much higher value of j , which again shows how unrealistic an age-independent assumption is.

(PDF)

S1 File. Maximum likelihood estimation and Monte Carlo methods.

(PDF)

S2 File. Rethinking the vectorial capacity: detailed calculations.
(PDF)

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Writing – review & editing: Melissa A. Iacovidou, Priscille Barreaux, Simon E. F. Spencer, Erin E. Gorsich, Kat S. Rock.

References

1. Geneva: WHO. World malaria report 2020: 20 years of global progress and challenges. Licence: CC BY-NC-SA 3.0 IGO; 2020.
2. Geneva: WHO. Global technical strategy for malaria 2016–2030, 2021 update; 2021. Available from: <https://www.who.int/teams/global-malaria-programme/global-technical-strategy-for-malaria-2016-2030>.
3. Bhatt S, Weiss D, Cameron E, Bisanzio D, Mappin B, Dalrymple U, et al. The effect of malaria control on *Plasmodium falciparum* in Africa between 2000 and 2015. *Nature*. 2015; 526(7572):207–211. <https://doi.org/10.1038/nature15535> PMID: 26375008
4. Ranson H, N'guessan R, Lines J, Moiroux N, Nkuni Z, Corbel V. Pyrethroid resistance in African anopheline mosquitoes: what are the implications for malaria control? *Trends in parasitology*. 2011; 27(2):91–98. <https://doi.org/10.1016/j.pt.2010.08.004> PMID: 20843745
5. Gansané A, Candrinho B, Mbituyumuremyi A, Uhomoibhi P, N'Falié S, Mohammed AB, et al. Design and methods for a quasi-experimental pilot study to evaluate the impact of dual active ingredient insecticide-treated nets on malaria burden in five regions in sub-Saharan Africa. *Malaria journal*. 2022; 21(1):1–20. <https://doi.org/10.1186/s12936-021-04026-0> PMID: 35012559
6. Mosha JF, Kulkarni MA, Lukole E, Matowo NS, Pitt C, Messenger LA, et al. Effectiveness and cost-effectiveness against malaria of three types of dual-active-ingredient long-lasting insecticidal nets

- (LLINs) compared with pyrethroid-only LLINs in Tanzania: a four-arm, cluster-randomised trial. *The Lancet*. 2022; 399(10331):1227–1241. [https://doi.org/10.1016/S0140-6736\(21\)02499-5](https://doi.org/10.1016/S0140-6736(21)02499-5) PMID: 35339225
7. Accrombessi M, Cook J, Ngufor C, Sovi A, Dangbenon E, Yovogan B, et al. Assessing the efficacy of two dual-active ingredients long-lasting insecticidal nets for the control of malaria transmitted by pyrethroid-resistant vectors in Benin: study protocol for a three-arm, single-blinded, parallel, cluster-randomized controlled trial. *BMC infectious diseases*. 2021; 21(1):1–12. <https://doi.org/10.1186/s12879-021-05879-1> PMID: 33607958
 8. Lindsay SW, Thomas MB, Kleinschmidt I. Threats to the effectiveness of insecticide-treated bednets for malaria control: thinking beyond insecticide resistance. *The Lancet Global Health*. 2021. [https://doi.org/10.1016/S2214-109X\(21\)00216-3](https://doi.org/10.1016/S2214-109X(21)00216-3) PMID: 34216565
 9. Alout H, Labbé P, Chandre F, Cohuet A. Malaria vector control still matters despite insecticide resistance. *Trends in parasitology*. 2017; 33(8):610–618. <https://doi.org/10.1016/j.pt.2017.04.006> PMID: 28499699
 10. Strode C, Donegan S, Garner P, Enayati AA, Hemingway J. The impact of pyrethroid resistance on the efficacy of insecticide-treated bed nets against African anopheline mosquitoes: systematic review and meta-analysis. *PLoS Med*. 2014; 11(3):e1001619. <https://doi.org/10.1371/journal.pmed.1001619> PMID: 24642791
 11. Barreaux P, Koella JC, N'Guessan R, Thomas MB. Use of novel lab assays to examine the effect of pyrethroid-treated bed nets on blood-feeding success and longevity of highly insecticide-resistant *Anopheles gambiae* s.l. mosquitoes. *Parasites & Vectors*. 2022; 15(1):1–9.
 12. Glunt KD, Coetzee M, Huijben S, Koffi AA, Lynch PA, N'Guessan R, et al. Empirical and theoretical investigation into the potential impacts of insecticide resistance on the effectiveness of insecticide-treated bed nets. *Evolutionary applications*. 2018; 11(4):431–441. <https://doi.org/10.1111/eva.12574> PMID: 29636797
 13. Viana M, Hughes A, Matthiopoulos J, Ranson H, Ferguson HM. Delayed mortality effects cut the malaria transmission potential of insecticide-resistant mosquitoes. *Proceedings of the National Academy of Sciences*. 2016; 113(32):8975–8980. <https://doi.org/10.1073/pnas.1603431113> PMID: 27402740
 14. Hughes A, Lissenden N, Viana M, Toé KH, Ranson H. *Anopheles gambiae* populations from Burkina Faso show minimal delayed mortality after exposure to insecticide-treated nets. *Parasites & vectors*. 2020; 13(1):1–11. <https://doi.org/10.1186/s13071-019-3872-2> PMID: 31924276
 15. Toé KH, Jones CM, N'Fale S, Ismail HM, Dabiré RK, Ranson H. Increased pyrethroid resistance in malaria vectors and decreased bed net effectiveness, Burkina Faso. *Emerging infectious diseases*. 2014; 20(10):1691. <https://doi.org/10.3201/eid2010.140619> PMID: 25279965
 16. Smith DL, Perkins TA, Reiner RC Jr, Barker CM, Niu T, Chaves LF, et al. Recasting the theory of mosquito-borne pathogen transmission dynamics and control. *Transactions of the Royal Society of Tropical Medicine and Hygiene*. 2014; 108(4):185–197. <https://doi.org/10.1093/trstmh/tru026> PMID: 24591453
 17. Dietz K. The estimation of the basic reproduction number for infectious diseases. *Statistical methods in medical research*. 1993; 2(1):23–41. <https://doi.org/10.1177/096228029300200103> PMID: 8261248
 18. Garrett-Jones C. Prognosis for interruption of malaria transmission through assessment of the mosquito's vectorial capacity. *Nature*. 1964; 204(4964):1173–1175. <https://doi.org/10.1038/2041173a0> PMID: 14268587
 19. Smith DL, Battle KE, Hay SI, Barker CM, Scott TW, McKenzie FE. Ross, Macdonald, and a theory for the dynamics and control of mosquito-transmitted pathogens. *PLoS pathog*. 2012; 8(4):e1002588. <https://doi.org/10.1371/journal.ppat.1002588> PMID: 22496640
 20. Styer LM, Carey JR, Wang JL, Scott TW. Mosquitoes do senesce: departure from the paradigm of constant mortality. *The American journal of tropical medicine and hygiene*. 2007; 76(1):111–117. <https://doi.org/10.4269/ajtmh.2007.76.111> PMID: 17255238
 21. Dawes EJ, Churcher TS, Zhuang S, Sinden RE, Basáñez MG. *Anopheles* mortality is both age- and *Plasmodium*-density dependent: implications for malaria transmission. *Malaria journal*. 2009; 8(1):1–16. <https://doi.org/10.1186/1475-2875-8-228> PMID: 19822012
 22. Bellan SE. The importance of age dependent mortality and the extrinsic incubation period in models of mosquito-borne disease transmission and control. *PLoS One*. 2010; 5(4):e10165. <https://doi.org/10.1371/journal.pone.0010165> PMID: 20405010
 23. Novoseltsev VN, Michalski AI, Novoseltseva JA, Yashin AI, Carey JR, Ellis AM. An age-structured extension to the vectorial capacity model. *PLoS One*. 2012; 7(6):e39479. <https://doi.org/10.1371/journal.pone.0039479> PMID: 22724022
 24. Ryan SJ, Ben-Horin T, Johnson LR. Malaria control and senescence: the importance of accounting for the pace and shape of aging in wild mosquitoes. *Ecosphere*. 2015; 6(9):1–13.

25. Brand SP, Rock KS, Keeling MJ. The interaction between vector life history and short vector life in vector-borne disease transmission and control. *PLoS computational biology*. 2016; 12(4):e1004837. <https://doi.org/10.1371/journal.pcbi.1004837> PMID: 27128163
26. Assouho KF, Adja AM, Guindo-Coulibaly N, Tia E, Kouadio AM, Zoh DD, et al. Vectorial transmission of malaria in major districts of Côte d'Ivoire. *Journal of medical entomology*. 2020; 57(3):908–914. <https://doi.org/10.1093/jme/tjz207> PMID: 31785095
27. Ashley EA, Phyo AP, Woodrow CJ. Malaria. *The Lancet*. 2018; 391(10130):1608–1621. [https://doi.org/10.1016/S0140-6736\(18\)30324-6](https://doi.org/10.1016/S0140-6736(18)30324-6)
28. Oumbouke WA, Pignatelli P, Barreaux AM, Tia IZ, Koffi AA, Alou LPA, et al. Fine scale spatial investigation of multiple insecticide resistance and underlying target-site and metabolic mechanisms in *Anopheles gambiae* in central Côte d'Ivoire. *Scientific reports*. 2020; 10(1):1–13. <https://doi.org/10.1038/s41598-020-71933-8> PMID: 32934291
29. Kulma K, Saddler A, Koella JC. Effects of age and larval nutrition on phenotypic expression of insecticide-resistance in *Anopheles* mosquitoes. *PLoS One*. 2013; 8(3):e58322. <https://doi.org/10.1371/journal.pone.0058322> PMID: 23484017
30. WHO, et al. Test procedures for insecticide resistance monitoring in malaria vector mosquitoes; 2016.
31. Catano-Lopez A, Rojas-Diaz D, Laniado H, Arboleda-Sánchez S, Puerta-Yepes ME, Lizarralde-Bejarano DP. An alternative model to explain the vectorial capacity using as example *Aedes aegypti* case in dengue transmission. *Heliyon*. 2019; 5(10):e02577. <https://doi.org/10.1016/j.heliyon.2019.e02577> PMID: 31687486
32. Liu-Helmersson J, Stenlund H, Wilder-Smith A, Rocklöv J. Vectorial capacity of *Aedes aegypti*: effects of temperature and implications for global dengue epidemic potential. *PloS one*. 2014; 9(3):e89783. <https://doi.org/10.1371/journal.pone.0089783> PMID: 24603439
33. Moller-Jacobs LL, Murdock CC, Thomas MB. Capacity of mosquitoes to transmit malaria depends on larval environment. *Parasites & vectors*. 2014; 7(1):593. <https://doi.org/10.1186/s13071-014-0593-4> PMID: 25496502
34. Massad E, Coutinho FAB. Vectorial capacity, basic reproduction number, force of infection and all that: formal notation to complete and adjust their classical concepts and equations. *Memórias do Instituto Oswaldo Cruz*. 2012; 107(4):564–567. <https://doi.org/10.1590/S0074-02762012000400022> PMID: 22666873
35. CDC. CDC—Malaria—About Malaria—Biology; 2020. Available from: <https://www.cdc.gov/malaria/about/biology/index.html>.
36. Stopard IJ, Churcher TS, Lambert B. Estimating the extrinsic incubation period of malaria using a mechanistic model of sporogony. *PLoS Computational Biology*. 2021; 17. <https://doi.org/10.1371/journal.pcbi.1008658> PMID: 33591963
37. Shapiro LL, Whitehead SA, Thomas MB. Quantifying the effects of temperature on mosquito and parasite traits that determine the transmission potential of human malaria. *PLoS biology*. 2017; 15(10):e2003489. <https://doi.org/10.1371/journal.pbio.2003489> PMID: 29036170
38. Shapiro LL, Murdock CC, Jacobs GR, Thomas RJ, Thomas MB. Larval food quantity affects the capacity of adult mosquitoes to transmit human malaria. *Proceedings of the Royal Society B: Biological Sciences*. 2016; 283(1834):20160298. <https://doi.org/10.1098/rspb.2016.0298> PMID: 27412284
39. Bompard A, Da DF, Yerbanga SR, Morlais I, Awono-Ambéné PH, Dabiré RK, et al. High Plasmodium infection intensity in naturally infected malaria vectors in Africa. *International Journal for Parasitology*. 2020; 50(12):985–996. <https://doi.org/10.1016/j.ijpara.2020.05.012> PMID: 32681932
40. Murdock C, Sternberg E, Thomas M. Malaria transmission potential could be reduced with current and future climate change. *Sci Rep*. 6: 27771; 2016. <https://doi.org/10.1038/srep27771> PMID: 27324146
41. Gubbins S, Carpenter S, Baylis M, Wood JL, Mellor PS. Assessing the risk of bluetongue to UK livestock: uncertainty and sensitivity analyses of a temperature-dependent model for the basic reproduction number. *Journal of the Royal Society Interface*. 2008; 5(20):363–371. <https://doi.org/10.1098/rsif.2007.1110> PMID: 17638649
42. Wearing HJ, Rohani P, Keeling MJ. Appropriate models for the management of infectious diseases. *PLoS Med*. 2005; 2(7):e174. <https://doi.org/10.1371/journal.pmed.0020174> PMID: 16013892
43. Rock KS, Torr SJ, Lumbala C, Keeling MJ. Quantitative evaluation of the strategy to eliminate human African trypanosomiasis in the Democratic Republic of Congo. *Parasites & vectors*. 2015; 8(1):1–13. <https://doi.org/10.1186/s13071-015-1131-8> PMID: 26490248
44. Childs LM, Prosper OF. The impact of within-vector parasite development on the extrinsic incubation period. *Royal Society open science*. 2020; 7(10):192173. <https://doi.org/10.1098/rsos.192173> PMID: 33204441

45. Chan M, Johansson MA. The incubation periods of dengue viruses. *PLoS one*. 2012; 7(11):e50972. <https://doi.org/10.1371/journal.pone.0050972> PMID: 23226436
46. Ibe OC. 1—Basic Concepts in Probability. In: Ibe OC, editor. *Markov Processes for Stochastic Modeling (Second Edition)*. second edition ed. Oxford: Elsevier; 2013. p. 1–27. Available from: <https://www.sciencedirect.com/science/article/pii/B9780124077959000013>.
47. Rock K, Wood D, Keeling M. Age-and bite-structured models for vector-borne diseases. *Epidemics*. 2015; 12:20–29. <https://doi.org/10.1016/j.epidem.2015.02.006> PMID: 26342239
48. Clements A, Paterson G. The analysis of mortality and survival rates in wild populations of mosquitoes. *Journal of applied ecology*. 1981; p. 373–399. <https://doi.org/10.2307/2402401>
49. Rodríguez G. *Lecture Notes on Generalized Linear Models*; 2007. Available from: <https://data.princeton.edu/wws509/notes/>.
50. Mogensen PK, Riseth AN. Optim: A mathematical optimization package for Julia. *Journal of Open Source Software*. 2018; 3(24):615. <https://doi.org/10.21105/joss.00615>
51. Bezanson J, Edelman A, Karpinski S, Shah VB. Julia: A fresh approach to numerical computing. *SIAM Review*. 2017; 59(1):65–98. <https://doi.org/10.1137/141000671>
52. Hahn T. Cuba—a library for multidimensional numerical integration. *Computer Physics Communications*. 2005; 168(2):78–95.
53. Hahn T. Concurrent cuba. *Computer Physics Communications*. 2016; 207:341–349.
54. Luz PM, Struchiner CJ, Galvani AP. Modeling transmission dynamics and control of vector-borne neglected tropical diseases. *PLoS Negl Trop Dis*. 2010; 4(10):e761. <https://doi.org/10.1371/journal.pntd.0000761> PMID: 21049062
55. Hollingsworth TD, Pulliam JR, Funk S, Truscott JE, Isham V, Lloyd AL. Seven challenges for modelling indirect transmission: Vector-borne diseases, macroparasites and neglected tropical diseases. *Epidemics*. 2015; 10:16–20. <https://doi.org/10.1016/j.epidem.2014.08.007> PMID: 25843376
56. Eckhoff PA. A malaria transmission-directed model of mosquito life cycle and ecology. *Malaria journal*. 2011; 10(1):1–17. <https://doi.org/10.1186/1475-2875-10-303> PMID: 21999664
57. Arifin S, Zhou Y, Davis GJ, Gentile JE, Madey GR, Collins FH. An agent-based model of the population dynamics of *Anopheles gambiae*. *Malaria journal*. 2014; 13(1):1–20. <https://doi.org/10.1186/1475-2875-13-424> PMID: 25373418
58. Bombliès A. Agent-based modeling of malaria vectors: the importance of spatial simulation. *Parasites & vectors*. 2014; 7(1):1–10. <https://doi.org/10.1186/1756-3305-7-308> PMID: 24992942
59. Collins E, Vaselli NM, Sylla M, Beavogui AH, Orsborne J, Lawrence G, et al. The relationship between insecticide resistance, mosquito age and malaria prevalence in *Anopheles gambiae* sl from Guinea. *Scientific reports*. 2019; 9(1):1–12. <https://doi.org/10.1038/s41598-019-45261-5>
60. Thomas MB, Read AF. The threat (or not) of insecticide resistance for malaria control. *Proceedings of the National Academy of Sciences*. 2016; 113(32):8900–8902. <https://doi.org/10.1073/pnas.1609889113> PMID: 27482081
61. Glunt KD, Thomas MB, Read AF. The effects of age, exposure history and malaria infection on the susceptibility of *Anopheles* mosquitoes to low concentrations of pyrethroid. *PLoS One*. 2011; 6(9):e24968. <https://doi.org/10.1371/journal.pone.0024968> PMID: 21966392
62. Anderson RA, Koellaf J, Hurd H. The effect of *Plasmodium yoelii nigeriensis* infection on the feeding persistence of *Anopheles stephensi* Liston throughout the sporogonic cycle. *Proceedings of the Royal Society of London Series B: Biological Sciences*. 1999; 266(1430):1729–1733. <https://doi.org/10.1098/rspb.1999.0839> PMID: 10518321
63. Wekesa JW, Copeland RS, Mwangi RW. Effect of *Plasmodium falciparum* on blood feeding behavior of naturally infected *Anopheles* mosquitoes in western Kenya. *The American journal of tropical medicine and hygiene*. 1992; 47(4):484–488. <https://doi.org/10.4269/ajtmh.1992.47.484> PMID: 1443347
64. Cator LJ, George J, Blanford S, Murdock CC, Baker TC, Read AF, et al. 'Manipulation' without the parasite: altered feeding behaviour of mosquitoes is not dependent on infection with malaria parasites. *Proceedings of the Royal Society B: Biological Sciences*. 2013; 280(1763):20130711. <https://doi.org/10.1098/rspb.2013.0711> PMID: 23698008
65. Rossignol P, Ribeiro J, Spielman A. Increased intradermal probing time in sporozoite-infected mosquitoes. *The American journal of tropical medicine and hygiene*. 1984; 33(1):17–20. <https://doi.org/10.4269/ajtmh.1984.33.17> PMID: 6696175
66. Shaw WR, Holmdahl IE, Itoe MA, Werling K, Marquette M, Paton DG, et al. Multiple blood feeding in mosquitoes shortens the *Plasmodium falciparum* incubation period and increases malaria transmission potential. *PLoS Pathogens*. 2020; 16(12):e1009131. <https://doi.org/10.1371/journal.ppat.1009131> PMID: 33382824

67. Ohm JR, Baldini F, Barreaux P, Lefevre T, Lynch PA, Suh E, et al. Rethinking the extrinsic incubation period of malaria parasites. *Parasites & vectors*. 2018; 11(1):1–9. <https://doi.org/10.1186/s13071-018-2761-4> PMID: 29530073
68. Kamiya T, Greischar MA, Wadhawan K, Gilbert B, Paaajmans K, Mideo N. Temperature-dependent variation in the extrinsic incubation period elevates the risk of vector-borne disease emergence. *Epidemics*. 2020; 30:100382. <https://doi.org/10.1016/j.epidem.2019.100382>
69. Paaajmans KP, Blanford S, Chan BH, Thomas MB. Warmer temperatures reduce the vectorial capacity of malaria mosquitoes. *Biology letters*. 2012; 8(3):465–468. <https://doi.org/10.1098/rsbl.2011.1075> PMID: 22188673
70. Chua T, et al. Modelling the effect of temperature change on the extrinsic incubation period and reproductive number of *Plasmodium falciparum* in Malaysia. *Trop Biomed*. 2012; 29(1):121–8. PMID: 22543612
71. Suh E, Grossman MK, Waite JL, Dennington NL, Sherrard-Smith E, Churcher TS, et al. The influence of feeding behaviour and temperature on the capacity of mosquitoes to transmit malaria. *Nature Ecology & Evolution*. 2020; p. 1–12. <https://doi.org/10.1038/s41559-020-1182-x> PMID: 32367033
72. Reiner RC Jr, Perkins TA, Barker CM, Niu T, Chaves LF, Ellis AM, et al. A systematic review of mathematical models of mosquito-borne pathogen transmission: 1970–2010. *Journal of The Royal Society Interface*. 2013; 10(81):20120921. <https://doi.org/10.1098/rsif.2012.0921>