Unveiling Time in Dose-Response Models to Infer Host Susceptibility to Pathogens

Delphine Pessoa¹, Caetano Souto-Maior¹, Erída Gjini¹, Joao S. Lopes¹, Bruno Ceña¹, Cláudia T. Codeço², M. Gabriela M. Gomes¹*

¹ Instituto Gulbenkian de Ciência, Oeiras, Portugal, ² Programa de Computação Científica, Fundação Oswaldo Cruz, Rio de Janeiro, Brazil

Abstract

The biological effects of interventions to control infectious diseases typically depend on the intensity of pathogen challenge. As much as the levels of natural pathogen circulation vary over time and geographical location, the development of invariant efficacy measures is of major importance, even if only indirectly inferable. Here a method is introduced to assess host susceptibility to pathogens, and applied to a detailed dataset generated by challenging groups of insect hosts (Drosophila melanogaster) with a range of pathogen (Drosophila C Virus) doses and recording survival over time. The experiment was replicated for flies carrying the Wolbachia symbiont, which is known to reduce host susceptibility to viral infections. The entire dataset is fitted by a novel quantitative framework that significantly extends classical methods for microbial risk assessment and provides accurate distributions of symbiont-induced protection. More generally, our data-driven modeling procedure provides novel insights for study design and analyses to assess interventions.

Introduction

Hosts exposed to disease-causing agents respond in accordance to the challenge dose. Therefore dose-response curves contain information about disease processes that can be extracted by suitable analytic frameworks. Early examples concerning microbial risk assessment include counting lesions caused by tobacco mosaic virus on plant leaves [1], as well as human responders to experimental challenge with polio viruses [2], Vibrio cholerae [3] and Streptococcus pneumoniae [4], for escalating challenge doses. Dose-response models have been in use for analyses and extrapolation of experimental datasets [5].

Models that account for the sigmoidal shape in log-linear scale of the typical dose-response curve have been derived mechanistically, based on the assumption that each individual pathogen has a probability of infection independent of others, the so-called independent action hypothesis [6]. This results in a one-parameter exponential-function model [7]. The frequent observation of shallower-than-exponential, or overdispersed, relationships has then prompted the implementation of heterogeneity in the probability of infection of individual hosts [8–10].

In the 1960s, Furumoto and Mickey [9] developed a dose-response model that could accommodate both shallow and steep increases in the response by considering the probability of infection of individual hosts described by a Beta-distribution. Although a mechanistic justification for this specific distribution has not been given, the model has been widely applied in microbial risk assessment due to its ability to outperform the simple exponential model [5].

Susceptibility distributions other than Beta have also been considered and are more commonly used in frailty models adopted in survival analysis [11], where the data consist of survivor counts over time in host groups that are constantly subject to a hazard [12,13]. These frailty models appeared in the 1980s and have since been adapted to infection hazards, where surviving signifies remaining uninfected [14–16]. While most informative when the exposure is continued or repeated over time, these formalisms would be inadequate for estimating distributions of susceptibility to infection from instantaneous challenge protocols.

The importance of accounting for time between challenge and observable toxicity responses to pathogens or other agents has been recognized. Recent models in ecotoxicology [17,18], consider explicit kinetics within exposed organisms. Also in microbial risk analysis, previous studies [19,20] have included time postinoculation as an additional parameter in classic dose-response models, although using an approach that conceptually allows for a different susceptibility distribution at each time point. Here we present a schema to infer a distribution of host susceptibilities to infection that holds consistently across dose and time. We introduce an experimental design and inference framework that enables such inferences by analyzing simultaneously a collection of survival curves, each representing a different challenge dose. The resulting Beta distributions are compared against those obtained by classic dose-response models based on single day measurements.

Recent evidence for symbiotic interactions that reduce host susceptibility to pathogens has stimulated the development of quantitative frameworks to assess the levels of individual and population protection attributable to specific symbionts. The intracellular bacterium Wolbachia, found among many arthropod species including Drosophila melanogaster, is one such symbiont.
Author Summary

While control options for plant, animal, and human pathogens are emerging rapidly, reliable assessment of the effect of interventions in biological systems presents many challenges. A major question is how to connect laboratory experiments and measurements with the relevant process in natural settings, where hosts are subject to pathogen exposures that vary in time and geographical location. With this aim, measures of protection that are invariant under varying exposure intensity need to be developed and integrated with mathematical models. In this article, we introduce a method to assess host susceptibility to pathogens, and apply it to survival of Drosophila melanogaster challenged with different doses of Drosophila C virus. By replicating the procedure in groups of flies that carry the symbiont Wolbachia, we are able to estimate how the viral protection induced by this intracellular bacterium is distributed in the host population. Our results disentangle host infection status from observed mortality, accounting naturally for time since exposure. The multiple-dose design proposed challenges traditional study designs to assess interventions.

[21,22]. To analyze the protection conferred by Wolbachia to D. melanogaster, we apply our inference framework simultaneously to two sets of time-dependent dose-response data: in one set the flies carry the symbiont bacterium Wolbachia (Wolb+); while in the other they do not (Wolb−). In this instance we extract the Beta distribution that best describes individual protection attributable to Wolbachia, as well as population statistics valid across entire dose ranges.

Methods

Survival data

We used virus free D. melanogaster lines with DrosDel w1118 background, with or without the endogenous Wolbachia strain wMelCS [21,23,24]. Flies were reared in standard food at 25°C. To assure that potential for heterogeneities are minimized by the experimental procedure, we used fifty 3–6 days old adult males per group, 10 per replicate and 5 replicates. To study the response to viral infection, we anesthetized with CO2 and pricked flies with different doses of Drosophila C virus (DCV). We used tenfold serial dilutions – from 10¹⁰ TCID₅₀/ml to 10⁴ TCID₅₀/ml – in Tris-HCl buffer, pH 7.5. Controls were pricked with buffer solution only. We used the pricking protocol described in [24], produced and triturated virus as in [21]. After pricking, we kept flies at 18°C and checked daily survival until day 80 and twice a week until the end of the experiment. Food was changed every 5 days. We summarized the data in 16 dose-response curves (8 per group, including control) from day 0 after treatment until day 139 (Dataset S1).

Dose-response model

Starting from established models, we refine the occurrence of mortality from infection, i.e. the response, as a function of the concentration of infectious units given to hosts, i.e. the dose. We present a step-by-step derivation of descriptions that integrate dimensions that are usually treated separately as well as the motivations for doing so.

Assuming independent action of infectious units, each unit has probability p of causing an infection, while for d infectious units infection occurs with a probability described by Binomial(d, p). Given further considerations about the distribution of infectious units in a homogeneous solution (see [9] for a complete derivation of the expression), the number of units causing infection can be described by a Poisson distribution, resulting in the exponential dose-response model [7], that describes the probability of infection in a host challenged with pathogen dose d:

\[ \pi_{\text{hom}} = 1 - e^{-pd}. \]  

(1)

This most basic formulation is hereafter referred to as the homogeneous dose-response model.

Furumoto and Mickey [9] expanded this formulation by allowing the probability of infection to be described by a parametric distribution, specifically the Beta distribution. To facilitate normalization across datasets, here we maintain the probability p fixed across individual hosts (as in [23]), and introduce a multiplicative parameter, the susceptibility factor 0 < x ≤ 1, to describe any natural or induced effect that decreases susceptibility. We assume that susceptibility to infection is Beta-distributed so as to describe the variation of susceptibility in the host population. Thus, we obtain the probability \( \pi_{\text{het}} \) that a host contracts infection as

\[ \pi_{\text{het}} = 1 - \int_0^1 e^{-xpd} \beta(x) \, dx, \]  

(2)

where \( \beta(x) = x^{a-1}(1-x)^{b-1}/B(a,b) \) and B is the Beta function. We refer to this formulation as the heterogeneous dose-response model.

At last we introduce a small parameter ε to account for a small probability of ineffective challenge, such that

\[ M \sim \text{Binomial}(n, (1-\varepsilon)\pi) \]

is the random variable representing the number of infected hosts, in a group of n hosts challenged with a given dose. Assuming that an ineffectively challenged host behaves like a control host with regard to death rates, the probability that m hosts are dead a number of days after challenge is then

\[ P(M = m) = \binom{n}{m} ((1-\varepsilon)\pi)^m [1-(1-\varepsilon)\pi]^{n-m}, \]  

(3)

where π is either \( \pi_{\text{hom}} \) (1) or \( \pi_{\text{het}} \) (2) depending on which dose-response model is adopted.

The parameters to be estimated for this dose-response model are the maximum probability of infection per infectious unit (p), the shape parameters for the Beta distribution that describes the susceptibility factor (a, b), and the probability of ineffective challenge (ε).

These models require a choice of how many days post-challenge cumulative mortality should be measured, which is difficult to establish for host-pathogen systems where times to death from infection or other causes overlap significantly. To overcome this difficulty, we develop a model that integrates an explicit representation of time to death with the dose-response process for infection just described. It should, however, be noted that time is introduced with the main purpose of enabling the use of survival curves to obtain robust estimates for probabilities of infection given different challenge intensities and consistently infer susceptibility to infection. From this perspective, parameters defined from now on should be regarded as auxiliary and will be implemented as simply as possible.
Time-dependent model for control group

We first consider a survival model for a control group of flies pricked with buffer solution only (no DCV), subject to two hazards: $h_0$, an age-dependent death hazard rate; and $h_a$, a background age-independent death hazard rate. The overall death hazard rate for uninfected hosts is therefore

$$h_U(t) = h_0(t) + h_a(t).$$  \hfill (4)

Denoting $T_U$ the random variable representing time to death of control hosts, we have

$$P(T_U = t) = P(T_0 = t) P(T_0 > t) + P(T_0 = t) P(T_a > t),$$  \hfill (5)

where $T_0$ and $T_a$ are the times to death from $h_0$ and $h_a$, respectively. Their corresponding distributions are assumed to be $T_0 \sim \text{Gamma}(m_0, s_0)$ and $T_a \sim \text{Uniform}(1/k_a)$, where $k_a$ is the background mortality rate, $m_0$ is the mean time to death, and $s_0$ is the shape parameter for the Gamma distribution of day of death from aging.

Time-dependent dose-response model

Hosts challenged with pathogen can become infected or remain uninfected and this infection status is hidden. If uninfected, they are subject to the age-dependent hazard rate that affects control hosts, $h_U$; if infected, they are subject to an infection hazard rate, $h_I$, and the age-independent background mortality. Thus the overall hazard rate of infected hosts is

$$h_I(t) = h_1(t) + h_a(t).$$  \hfill (6)

Now let $I \sim \text{Binomial}(n, (1-\varepsilon)\pi)$ be the random variable representing the number of hosts infected by challenge with a given pathogen dose. Then the probability that $i$ hosts are infected after $n$ hosts were challenged is

$$P(I = i) = \binom{n}{i} [(1-\varepsilon)\pi]^i [1 - (1-\varepsilon)\pi]^{n-i},$$  \hfill (7)

where $\pi$ is either $\pi_{hom}(1)$ or $\pi_{het}(2)$ depending on which dose-response model is adopted.

Let $T$ be the random variable representing the time to death of hosts challenged by a given pathogen dose. The probability density of observing a death event at time $t$ given that $i$ hosts are infected is

$$P(T = t|I = i) = \frac{(n-i)}{n} P(T_U = t) + \frac{i}{n} P(T_I = t),$$  \hfill (8)

where $T_I$ denotes the distribution of time to death of infected hosts, given by

$$P(T_I = t) = P(T_1 = t) P(T_1 > t) + P(T_1 = t) P(T_a > t)$$  \hfill (9)

and $T_1$ is the distribution of times to death from the infection hazard rate $h_I$. This distribution is assumed to follow $T_1 \sim \text{Gamma}(m_1, s_1)$, where $m_1$ is the mean time to death of infected hosts, and $s_1$ is the shape parameter for the Gamma distribution of day of death from infection.

In setting the priors for parameter estimation we note that background mortality is small and therefore $k$ is kept small by setting $1/k$ to be much greater than the last day of the experiment. To enforce that deaths due to infection occur earlier than deaths due to aging, we constrain the mean time to infection death to be lower than old-age death, i.e. $m_1 < m_0$, and the probability of dying before the end of the study to be greater for infected hosts, i.e. $P(T_0 \leq t_{\text{max}}) \leq P(T_1 \leq t_{\text{max}})$, where $t_{\text{max}}$ is the last day of the experiment.

To construct the likelihood to be maximized by the parameter estimation procedure, we let $D$ be the random variable denoting the day fly $j$ died and $S$ the random number of survivors up to $t_{\text{max}}$. Then the likelihood of observing the actual number of survivors $s$ and the times of death $d = [d_1, \ldots, d_{n-s}], for a given dose $s$ is

$$P(S = s, D = d) = \sum_{i=1}^{n} P(S = s, D = d| I = i) P(I = i)$$  \hfill (10)

where

$$P(T > t_{\text{max}}|D = d) = \prod_{j=1}^{n-s} P(D_j > t_{\text{max}}|D_j = d_i)$$

Since the observations for each dose are independent, taking the product of the likelihoods over the different doses yields the global expression for the likelihood of the entire dataset.

In this time-dependent dose-response model, the parameters to be estimated are the maximum probability of infection per infectious unit ($\pi$) used for normalization purposes, the Beta distribution shape parameters to describe variation in susceptibility factor ($a, b$), the parameters that control death due to aging ($m_0, s_0$), infection ($m_1, s_1$), and background mortality ($k$), as well as probability of ineffective challenge ($\varepsilon$). Parameters $\pi$ and $k$ are typically small and were introduced to improve performance of the likelihood.

Parameter estimation

Model parameters were estimated using Markov chain Monte Carlo sampling implemented with the PyMC package [26] (code available from [27]). The prior distributions considered are listed in Table 1. Initial values were chosen so as to start with a non-zero likelihood. Using Metropolis-Hastings algorithm, we ran two separate chains for 252,000 iterations. The first 27,000 iterations were discarded. The recording interval was set to 250 so that the autocorrelation between samples was negligible. Convergence was assessed by inspection of the trace plots. All analyses were performed on the pooled samples from the two replicate chains.

Results

Groups of Wolbachia-negative (Wolb⁻) and positive (Wolb⁺) D. melanogaster flies were challenged with a range of DCV doses and survival curves were traced as shown in Figure 1. This dataset was analyzed by applying the models introduced in Methods.

Susceptibility distribution from selected day mortality

To emphasize the importance of day selection to infer distributions of susceptibility to infection by classic dose-response models [5] we have applied these procedures to mortality data observed by two specific days (30 and 50). Parameter estimates from these models are listed in Table 2. The model fits to the mortality data at the selected days are shown in Figure 2, as well as the associated distribution of Wolb⁺ susceptibilities and the posterior samples for the Beta distribution shape parameters. For simplicity we have adopted the homogeneous model for Wolb⁻ and focus on comparing susceptibility distributions of Wolb⁻ inferred at different days. Mean protection conferred by
Wolbachia in this illustration is estimated as 79% and 56%, based on mortality measurements at day 30 and 50, respectively. Moreover, the distributions have fundamentally different shapes, with the appearance of a high susceptibility group as time progresses. This sensitivity to the day by which mortality data are collected is a concern that raises the need to disentangle infection status from the associated time-dependent mortality. In the following sections, infection and mortality are estimated explicitly using the integrated time-dependent model described in Methods. The procedure is illustrated in Figure 3.

Aging and background mortality
Control curves from Wolb^− and Wolb^+ flies pricked with buffer solution (no DCV) were compared with the Kaplan-Meier method using the log-rank test and no significant difference was found (with a p-value of 0.47). By fitting the uninfected time-dependent model (4–6) to the control survival curves (Figure 1) we estimated the parameters describing aging (m0, s0) and background (k) mortality (Table 3).

Susceptibility distribution from survival curves
For each group of flies (Wolb^− and Wolb^+), the time-dependent dose-response model constructed in Methods was fitted simultaneously to the entire dataset of survival curves (one for each DCV challenge dose), fixing across doses the distribution of times to death from infection (mI, sI) and aging (m0, s0), while estimating the susceptibility parameters (p, a, b) that govern the dependence of response on challenge dose according to the adopted dose-response model. The estimated parameter values are listed in Table 4. The deviance information criterion (DIC) [28] favored the homogeneous model for the Wolb^− group and the heterogeneous model for Wolb^+ (Text S1). Mean time to death from infection is 9 and 14 days in the Wolb^− and Wolb^+ groups, respectively. The variance in time to death from infection is lower for Wolb^−, with a standard deviation of 2 days, compared to 6 days in the Wolb^+ group. Figure 4 compares fitted with observed survival curves.

The fitted dose-response curves that result from this analysis are shown in Figure 5A, while the inferred distribution of Wolb^+ susceptibilities normalized by the Wolb^− measure is displayed in Figure 5B and the corresponding posterior distribution of the Beta shape parameters is in Figure 5C. Given the homogeneity in the Wolb^− group, the distribution of susceptibility in Wolb^+ provides a direct indication of how antiviral protection conferred by Wolbachia is distributed among its carriers. Typically defined as 1 − RR, where RR is the risk reduction attributed to the susceptibility modifier (Wolbachia in this case), we determine the mean protection conferred by the symbiont to its host as 85% (with a 95% HPD of 60–93%).

Comparison with selected day mortality
To assess the best possible performance of classic methods [5] in the inference of susceptibility distributions (for Wolb^+ in the case)

---

**Table 1. Model parameters and their corresponding prior distributions.**

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Meaning</th>
<th>Prior</th>
</tr>
</thead>
<tbody>
<tr>
<td>m0</td>
<td>Mean time to death from aging</td>
<td>U(0, 140)</td>
</tr>
<tr>
<td>s0</td>
<td>Shape of the Gamma distribution for death from aging</td>
<td>U(0, 500)</td>
</tr>
<tr>
<td>mI , mI+</td>
<td>Mean time to death from infection (for Wolb^− and Wolb^+, respectively)</td>
<td>U(0, m0)</td>
</tr>
<tr>
<td>sI, sI+</td>
<td>Shape of the Gamma distribution for death from infection (for Wolb^− and Wolb^+)</td>
<td>U(0, 100)</td>
</tr>
<tr>
<td>p</td>
<td>Per viral particle probability of causing infection</td>
<td>U(0, 1)</td>
</tr>
<tr>
<td>a, b</td>
<td>Shape parameters of the Beta distribution for the susceptibility to infection of Wolb^+</td>
<td>U(0, 1), 10</td>
</tr>
<tr>
<td>k</td>
<td>Background mortality rate, from causes other infection or aging</td>
<td>U(10^{-6}, 10^{-2})</td>
</tr>
<tr>
<td>r</td>
<td>Probability of ineffective challenge</td>
<td>N(0.001, 0.00125)[0.1]</td>
</tr>
</tbody>
</table>

**Figure 1. Survival curves for Wolb^− (A) and Wolb^+ (B) groups of D. melanogaster.** Dots represent experimental data. Dark blue curves show the model fit to the survival of control flies. Shaded areas represents 95% CI (credible intervals). doi:10.1371/journal.pcbi.1003773.g001
we must have previously reduced the set of survival curves to a set of effectively infected proportions - one entry per challenge dose.

To search for a range of days in which absolute mortality might provide an approximate indication of infection, we compare the estimated proportions effectively infected by each challenge dose with the mortality proportion measured at each day. Using a normalized Euclidean distance between these two measures, a day-selection score is provided by the red curve in Figure 6. We identify day 30 as optimal and 17–46 as the interval of days in which the score is at least 95% of the optimal. Reassuringly, the optimal day appears to coincide with the saturation of infection-induced mortality (see position of vertical dash-dotted gray line in relation to the Gamma distributions). We now recall Figure 2 and Table 2 for the inferences based on day 30 mortality data to confirm that classic dose-response models can in principle infer susceptibility distributions that are consistent with those obtained

Table 2. Estimated parameters by applying dose-response models to selected day mortality.

<table>
<thead>
<tr>
<th>Mortality data</th>
<th>Parameter</th>
<th>Median</th>
<th>95% HPD*</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 dpc³</td>
<td>( p )</td>
<td>( 2.33 \times 10^{-6} )</td>
<td>[( 1.67 \times 10^{-6}, 3.13 \times 10^{-6} )]</td>
</tr>
<tr>
<td></td>
<td>( a )</td>
<td>0.30</td>
<td>[0.21, 0.41]</td>
</tr>
<tr>
<td></td>
<td>( b )</td>
<td>1.10</td>
<td>[0.29, 2.53]</td>
</tr>
<tr>
<td></td>
<td>( c )</td>
<td>( 1.78 \times 10^{-3} )</td>
<td>[( 4.90 \times 10^{-4}, 3.49 \times 10^{-3} )]</td>
</tr>
<tr>
<td>50 dpc³</td>
<td>( p )</td>
<td>( 2.65 \times 10^{-6} )</td>
<td>[( 1.82 \times 10^{-6}, 3.47 \times 10^{-6} )]</td>
</tr>
<tr>
<td></td>
<td>( a )</td>
<td>0.34</td>
<td>[0.24, 0.51]</td>
</tr>
<tr>
<td></td>
<td>( b )</td>
<td>0.42</td>
<td>[0.12, 0.93]</td>
</tr>
<tr>
<td></td>
<td>( c )</td>
<td>( 1.83 \times 10^{-3} )</td>
<td>[( 3.60 \times 10^{-4}, 3.32 \times 10^{-3} )]</td>
</tr>
</tbody>
</table>

*High posterior density interval.
³Days post-challenge.

Figure 2. Dose-response curves and susceptibility distributions inferred from mortality measurements 30 and 50 days post-challenge. Dose-response models adopted here are the standard formulations (1–3). A,D, Curves represent the fitted dose-response model to mortality on selected day post-challenge (dots), for Wolb⁻ (black) and Wolb⁺ (blue). Shaded areas represent the 95% CI. B,E, Distribution of susceptibility to infection in Wolb⁺. The posterior median distribution is the curve and the shaded area is the 95% CI. C,F, Posterior samples of the Beta-distribution shape parameters describing Wolb⁺ susceptibility in blue. Red dot mark the median of the respective distributions. The homogeneous model was adopted for Wolb⁻.

doi:10.1371/journal.pcbi.1003773.t002

doi:10.1371/journal.pcbi.1003773.g002
under our extended model (Figure 5). A major issue, however, is that results are sensitive to a day-selection criterion that relies on having previously carried out the entire procedure. The appearance of a high susceptibility group in distributions inferred at later days are an artifact due to the accumulation of background mortality that should be factored out. These results highlight the importance of adequately representing the time dimension in the analysis.

Discussion

Dose-response models have become standard quantitative frameworks in microbial risk assessment. Less recognized is their ability to estimate host trait distributions. Here we illustrate the concept by extracting host susceptibility distributions from mortality measured as a function of pathogen challenge dose, but similar procedures can be developed for measures of infection or infectiousness (instead of mortality), and can be made a function of other environmental variables such as temperature or humidity (instead of dose). Understanding how to detach host trait distributions from environmental variables is crucial for the

![Figure 3](https://example.com/figure3.png)

**Figure 3. Schematic illustration of the proposed experimental design and inference procedure.**
doi:10.1371/journal.pcbi.1003773.g003

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Median</th>
<th>95% HPD</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \mu_0 )</td>
<td>117.18</td>
<td>[114.99, 119.84]</td>
</tr>
<tr>
<td>( \lambda_0 )</td>
<td>118.93</td>
<td>[80.19, 166.15]</td>
</tr>
<tr>
<td>( \kappa )</td>
<td>1.14 ( 10^{-3} )</td>
<td>[5.36 ( 10^{-4} ), 1.96 ( 10^{-3} )]</td>
</tr>
</tbody>
</table>

doi:10.1371/journal.pcbi.1003773.t003

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Median</th>
<th>95% HPD</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \rho )</td>
<td>1.73 ( 10^{-6} )</td>
<td>[9.58 ( 10^{-7} ), 2.67 ( 10^{-5} )]</td>
</tr>
<tr>
<td>( a )</td>
<td>0.47</td>
<td>[0.25, 0.85]</td>
</tr>
<tr>
<td>( b )</td>
<td>3.21</td>
<td>[0.34, 8.40]</td>
</tr>
<tr>
<td>( c )</td>
<td>1.89 ( 10^{-3} )</td>
<td>[4.55 ( 10^{-4} ), 3.40 ( 10^{-3} )]</td>
</tr>
<tr>
<td>( m_0 )</td>
<td>9.34</td>
<td>[9.10, 9.58]</td>
</tr>
<tr>
<td>( s_1 )</td>
<td>35.79</td>
<td>[26.60, 47.05]</td>
</tr>
<tr>
<td>( m_1 )</td>
<td>13.79</td>
<td>[11.31, 14.94]</td>
</tr>
<tr>
<td>( s_1 )</td>
<td>5.59</td>
<td>[4.70, 11.12]</td>
</tr>
<tr>
<td>( m_0 )</td>
<td>115.20</td>
<td>[113.94, 116.45]</td>
</tr>
<tr>
<td>( s_0 )</td>
<td>140.39</td>
<td>[116.80, 166.97]</td>
</tr>
<tr>
<td>( \kappa )</td>
<td>2.15 ( 10^{-3} )</td>
<td>[1.65 ( 10^{-3} ), 2.71 ( 10^{-3} )]</td>
</tr>
</tbody>
</table>

Parameters with superscripts \( ^{-} \) and \( ^{+} \) relate to Wolb\( ^{-} \) and Wolb\( ^{+} \) groups, respectively.
doi:10.1371/journal.pcbi.1003773.t004

Table 4. Parameters governing estimated number infected per dose of DCV challenge and time to death from infection using time-dependent dose-response models described in Methods.

\[
\text{Table 3. Estimated parameters governing time to death from causes other than DCV infection.}
\]

doi:10.1371/journal.pcbi.1003773.t003
formulation of measures that can be transported between laboratory and natural conditions [29,30].

We address this problem with an experimental design and inference framework that enables the estimation of distributions of host susceptibility to infection by analyzing simultaneously a collection of survival curves, each representing a different challenge dose (Figure 3). The procedure is illustrated on a specifically collected dataset where two distinct groups of hosts (*D. melanogaster*) were experimentally challenged by viruses (*DCV*): one group consists of isogenic flies where no significant variability in susceptibility to infection is found; and another with the same genetic background but now carrying the symbiont bacterium *Wolbachia* known to reduce susceptibility to *DCV* [21,22].

Our inferences indicate that *Wolbachia* confers on average 85% *DCV* protection to *D. melanogaster* under the specified laboratory conditions, and suggest significant variability in this effect. This variance in susceptibility is induced by the symbiont, since model selection criteria did not support heterogeneity in the susceptibility of flies not carrying *Wolbachia*. Since the Drosophila and *Wolbachia* populations used in this study are isogenic, the heterogeneity in susceptibility of *Wolbachia*-carrying flies uncovered here indicates variation in the host-microorganism interaction that lacks a genetic basis. A simple hypothesis is that variance in *Wolbachia* levels at the individual host level leads to variance in resistance to viruses. Although several lines of evidence support this hypothesis [31–34], further experiments are required to discriminate whether heterogeneity in resistance is directly linked to variance in *Wolbachia* levels or, alternatively, a result of another environmental/physiological variance that is only expressed in the presence of *Wolbachia*.

Previous estimates of protection were based on survival analysis or viral titres in a dose-specific manner [21,22,24]. To our

![Figure 4. Fit of time-dependent dose-response model to survival curves.](image)

Figure 5. Dose-response curves and susceptibility distributions inferred from survival curves. A, Curves represent the estimated dose-response relationships from fitting the model described in Methods to survival over time, for Wolb⁻ (black) and Wolb⁺ (blue). Shaded areas represent the 95% CI. B, Distribution of susceptibility to infection in Wolb⁻. The posterior median distribution is the curve and the shaded area is the 95% CI. C, Posterior samples of the Beta-distribution shape parameters describing Wolb⁺ susceptibility in blue. Red dot marks the median of distribution. doi:10.1371/journal.pcbi.1003773.g005
knowledge, the experimental design and analysis presented here provides the first estimation of protection in way that is detached from challenge dose. Future developments might consider: estimation of alternative distributions to compare with the shapes suggested by the Beta family; extension of the adopted experimental design to measure responses other than mortality; and move towards host populations and environmental conditions that are closer to natural systems.

The parameters estimated here should not be seen as isolated from the relevant ecological context. On the contrary, they are intended as a first step to inform the construction of ecological and epidemiological models where Wolbachia, other symbions, or interventions that modify host susceptibility to infection, are introduced to induce desired transitions in populations. The introduction of Wolbachia into Aedes aegypti and other arthropod vectors is being considered as a promising strategy to control dengue and other infectious diseases of humans (see [35] and references therein). The inference frameworks presented can be readily adapted to provide accurate quantification of Wolbachia-induced protection and integrated in population models of public health importance.

The challenge of considering the time dependence of processes leading to observable ecotoxicity responses has also been addressed in toxicology where the so-called General Unified Model of Survival (GUTS) has been proposed [18]. These models simulate the time-course of external and internal processes leading to toxic effects on organisms to generate an output that can be fitted to mortality over time. While those studies tend prioritize the mechanistic descriptions of the toxicokinetic and toxicodynamic processes that damage the organisms, we have chosen to adopt a phenomenological approach and focus on the inference and interpretation of how susceptibility to infection is distributed in a population.

In epidemiological systems, the baseline transmission intensity is often not directly measurable but indirectly inferred in a model-based manner. Dose-response models, on the other hand, can account for experimentally controlled patterns of exposure [36,37]. Variation in host susceptibility to pathogens is one component of both classes of systems that mostly influences estimates of intervention impacts [29]. Therefore, building on the methods developed here furthers our potential to accurately evaluate the burden of infectious diseases and design effective interventions.

Supporting Information

Dataset S1 Survival data for Wolb− and Wolb+ groups of D. melanogaster challenged by various doses of DCV.

Text S1 Homogeneous vs heterogeneous model selection.

Acknowledgments

We thank Luis Teixeira for help with the experimental work, and Jorge Carneiro for valuable discussions concerning statistical models.

Author Contributions

Conceived and designed the experiments: MGMG. Performed the experiments: DP CSM BC. Analyzed the data: DP CSM EG JSL BC CTC MGMG. Wrote the paper: DP CSM EG JSL BC CTC MGMG.

References