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Modelling Parasite Transmission and Control

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DEDICATION

To our immediate families who have lived with this project, Shirin, Priya, and Patty and to numerous colleagues for their forbearance as well as their support.
Modelling parasite transmission has made enormous strides since the seminal models of Ross for describing malaria transmission developed during the early 1900s (Ross, 1911). McDonald’s use of the early malaria models to show that killing adult mosquitoes would be particularly effective in reducing infection transmission was a major advance in demonstrating the usefulness of theoretical analysis and population dynamics modelling in particular for guiding parasite control programmes, and since then parasite transmission models have also been used to guide the onchocerciasis control programme in Africa (Habbema et al, 1992), as well as for investigating best strategies for controlling a host of other parasites, including tuberculosis, trachoma and lately helminth infections, such as schistosomiasis and filariasis (Chan et al, 1995; Laing et al, 2007; Michael et al, 2004). The importance of this work is highlighted by greater understanding of threshold phenomena in transmission dynamics leading to the concept that natural “breakpoints” occur below which parasite systems will go extinct to the roles that worm mating behaviour and infection aggregation can play in both helminth transmission and control (Hairston, 1962; Macdonald, 1961, 1965). The emerging trend from this work is thus the increasing use of understanding parasite transmission dynamics via the construction and analysis of mathematical models for use in guiding the development of informed parasite control strategies, so much so that this twin objective, viz improving understanding of parasite transmission dynamics and applying models to guide parasite control, has almost become a *de facto* goal of most recent work in parasite transmission modelling.

Another growing theme in parasite transmission modelling is the evolution of modelling techniques and conceptual frameworks, from the phenomenological investigation of key factors governing transmission dynamics at large population scales, based on the generalized Anderson and May models synthesized during the 1980s and early 1990s, to the more detailed treatment of localized transmission of parasites at smaller population and spatial scales. Mathematically, this has meant that modelling techniques have increasingly moved from the application of simple dynamic differential equations-based analytic approaches to the use of simulation and other computationally intensive methods that focus on fitting and calibrating models to local data. This has raised debate about the roles of simplified a priori-derived
mechanistic models versus more complex but a posteriori models derived from data for generalizing understanding of transmission processes, and therefore how best to use models to further the theory of parasite transmission and accordingly its use for designing useful control options. This includes at its core the issue of how best to reconcile and model the impact of ecological and spatial scales as well as the effective addressing of other key, possibly scale-dependent heterogeneities of increasing complexity, such as impacts of host immunity, genetic diversity, multi-species interactions, spatial heterogeneity and between and within community connections, and external forcing factors linked to environmental variation and change.

The increasing recognition of the public health importance of parasites has led to a more recent flowering of interest to tackle and resolve these modelling issues, culminating in the convening of both the Dahlem Conference on the population biology of infectious diseases held in 1981 and the follow up Isaac Newton Institute programme on modelling infectious diseases held during 1993. These meetings summarized the then contemporary state of parasite ecology and epidemiology (Anderson and May, 1982; Grenfell and Dobson, 1995), but also underscored gaps and future research needs. This book is an attempt to present the progress made since these meetings in terms of both the conceptual and practical advances made in addressing key gaps in parasite transmission modelling (focus here restricted to the major microbial and helminth parasites), the new understanding these have yielded regarding the transmission and control of these parasites, and the future challenges they portend for population dynamics modelling.

We have organized the material in the book into two major sections, the first presenting the state of the art in models aimed at capturing complex or detailed aspects of transmission dynamics beginning with a review of the evolution of modelling malaria transmission. This first chapter, by Smith and Ruktanonchai, describes the process by which the simplifying assumptions underlying early modelling efforts were relaxed and models made more realistic (and generally more complex) over time as they became informed by new information from observational studies of vector biology and epidemiology. A key highlight is the need to consider the spatial scale in mosquito biting behaviour and pattern if more robust models of malaria transmission in real populations are to be developed. The second chapter, by Michael and Gambhir, focuses on the implications for disease transmission and control of differences in density-dependent processes regulating larval infection in the two major mosquito vectors transmitting filariasis. The authors demonstrate how a careful consideration and modeling of the occurrence and operation of multiple density-dependent processes acting on different stages of the parasite life cycle will be required to explain differential system persistence, transitions, and resilience to perturbations in different host-vector-parasite systems. In a similar spirit, but in a more general case, is the chapter by Pugliese (Chapter 3) on the dynamics of multiple parasite species infecting a host population. This might be characterized as a phenomenological investigation focusing on general questions of competition or co-existence between parasite species. Given the prevalence of co-infections commonly observed in the field, this chapter presents
a foundation from which studies of specific co-infections might be explored using models.

Chapters 4 and 5 by Gaff and Schaefer and by Cornell exemplify current advances in modeling tick-borne disease transmission and helminths, respectively, but are also of particular interest because of the modeling approaches utilized. Gaff and Schaefer use metapopulation models, which have been common in population biology for many years, because tick population dynamics are very sensitive to local environmental conditions. In addition, patch connectivity is shown to be of considerable importance, specifically with respect to control strategies. The notion of connectivity resurfaces in a similar context in the chapter by Seto and Carlton in Part II.

The chapter by Cornell is similar to that of Pugliese in its phenomenological orientation, and stems from the observation that in the case of helminths “the processes affecting parasite demographics are density dependent—i.e., depend nonlinearly on the number of worms in the host—such as the risk that a female cannot find a mate, or the modulation of infection rate by the host’s immunity.” Such questions are naturally addressed through the use of stochastic models as shown by the author. Although this chapter mainly focuses on simulation tools useful in the study of such models, Cornell also makes the case for the continued utilization of analytically tractable models in search of general principles not easily discernable from simulation studies.

In common with the metapopulation chapter of Gaff and Schaefer, Chapters 6 and 7 by Remais and by Spear and Hubbard represent a transition from phenomenological studies at an unspecified scale to a local scale where the issue is to specify effective and sustainable local control tactics. Remais (Chapter 6) describes a variant of the Anderson-May model of schistosome transmission which is adjoined to a submodel of uninfected snail density as a function of temperature and rainfall. To the extent possible, the model was parameterized to utilize both site-specific and site-invariant data for model calibration and subsequent use in short-term forecasting. Spear and Hubbard (Chapter 7) elaborate on the parameter estimation issues and residual parameter uncertainty central to model use at this scale and propose an approach tailored to deal with the heterogeneous nature of local data typically available for this purpose.

Part II of the book serves to highlight the current use of transmission models in the planning, monitoring and evaluation of parasite control programmes, and begins with Chapter 8 by Buckee and Gupta, who review advances made in using mathematical models of transmission for the design of malaria control programmes. The chapter shows how ultimately the design of appropriate control programmes will rely on better understanding of the complex dynamical interactions between vector ecology, generation and impact of pathogen diversity on host immunity and development of drug resistance, and the pathology of the disease in humans, which is only now beginning to be studied as a result of advances particularly in pathogen genomics. In a similar vein, White and Garnett (Chapter 9) highlight the advances made in modeling TB to date but also underscore how further progress in predicting the effects of control will crucially require gaining a better knowledge of the natural
history and processes involved in disease transmission, particularly rates of change in latency, and impacts of exogenous reinfection, active disease and recovery from TB disease. They suggest that a key role of models in this regard is to pinpoint existing gaps in knowledge, and in conjunction with developments in diagnostics and genetic analysis, help guide empirical research to resolve such gaps.

One study that has begun to model the impact of control programmes on both infection dynamics and disease pathogenesis (blinding sequelae) is that described by Ghambir and colleagues for trachoma in Chapter 10. A novel aspect of their modelling framework, with implications for other diseases, is the combination of a compartmental SI contact type module to model microparasite infection dynamics with a module for capturing disease prevalence based on quantifying the number of prior infections (or infection “burden”) experienced by individuals in a population.

The last two chapters in this section address the roles of parasite transmission models for decision-making in disease control. In Chapter 11, Michael and Ghambir describes the role that models can play in resolving the management questions regarding community level interventions for lymphatic filariasis elimination, including evaluating thresholds to meet various intervention endpoints, optimising control parameters for different endemic conditions, addressing the impacts of vector and spatial heterogeneities, and developing and applying rational monitoring and evaluation protocols for assessing programme success. The results indicate that instituting an adaptive management approach to parasite control, focusing on supporting the development of locally appropriate and ecologically resilient control strategies will be crucial to resolving the uncertainties surrounding parasite population dynamics and the effectiveness of presently proposed intervention options. Seto and Carlton (Chapter 12), by contrast, emphasize how site-specific epidemiologic and disease control options should and can be incorporated into modelling frameworks to support routine decision-making by public health officials in endemic regions, focusing on the model developed by their group for Schistosoma japonicum transmission in China. They demonstrate in particular how their model structure can be used to consider the spatial interconnectivity of local populations, which has profound implications for deciding not only the spatial scale of control but also the potential of the parasite to invade, persist and spread as a result of changes in host, vector and pathogen migratory patterns.

We have chosen to separate the final two chapters of the book into an Epilogue because they almost certainly point to important future applications or developments in the mathematical modelling of infectious and parasitic diseases. There are immediate challenges in addressing the implications of climate change on the extent and intensity of environmentally mediated infectious diseases in general and parasitic diseases in particular. In Chapter 13, Parham and Michael present a simple model of malaria transmission sensitive to changes in rainfall and temperature variables. Their example makes a convincing case that climate-driven transmission models will be crucial to understanding the rate at which P. falciparum and P. vivax may either infect, expand into, or go extinct in populations as local environmental conditions change. The chapter prompts two more general points, the first being that mechanistically-
based models will be centrally important tools in assessing the impact of various climate change scenarios on disease transmission because, unlike statistical models, they possess at least some credible basis for extrapolating beyond the range of current climatological experience. The second key point is that mechanistic models present a framework for assessing the relative impact of, for example, agricultural adaptation to climate change versus the narrow impact of temperature and rainfall effects in an otherwise unchanging environment.

The second dimension of future developments that is nearly upon us relates to the impact of genomics on the modelling of disease transmission and control. It is not difficult to foresee modest changes to current modelling structures to take advantage of elements of this vast new source of data. But it is also highly probable that altogether new approaches will emerge. Indeed, the utilization of genetic data relating to parasites, hosts, or vectors is only now beginning to occur in ecological or epidemiological studies of parasitic disease transmission. In Chapter 14, Tibayrenc argues for “an integrated approach to the genetic epidemiology and population genomics of Chagas disease.” Just as mathematical modelling of disease transmission originally grew from foundational studies of epidemiology and vector biology, it seems likely that we are at the beginning of a new class of modelling studies that will take advantage of the molecular epidemiology of disease transmission. While Tibayrenc argues “that Chagas disease potentially constitutes a paradigm model for the integrated genetic epidemiology and integrated population genomic approaches to understanding of disease transmission,” we suggest that his argument also points to new opportunities and challenges for the next generation of mathematical modelling studies for clarifying parasite transmission dynamics in general, as highlighted by the work of Buckee and Gupta (Chapter 8) in using information from sequence data for modelling the impact of parasite diversity on host immunity and development of resistance in malaria.

It is clear that many fascinating problems still remain to be addressed in parasite transmission modelling, from better understanding of transmission processes and natural history of infection to investigating the impact of ecological and spatial scales, climate change, host immunity and social behaviour, parasite-host evolutionary dynamics and parasite community ecology on parasite transmission. This book captures some of the advances made in recent years and provides indications of ways forward for addressing these questions by shedding light on developments in conceptual frameworks and modelling tools as well as the emergence of new data forms for aiding model construction, testing and analysis. Another important advance has been the parallel development of robust computationally-intensive statistical methods to allow model testing and parameterization by aiding the fitting of models to complex data. This is an exciting area of work, which we believe will broaden the scope of mathematical modelling in investigating parasite transmission processes. In particular, we expect this advance will now allow modellers to begin the successful development and analysis of mechanistically-rich models of parasite transmission that will facilitate better integration of the variety of mechanisms increasingly recognized as important in simultaneously affecting transmission, including abiotic processes,
trophic and evolutionary interactions, movement in space, and behaviour and even physiology of the individual. We foresee a continuing bright future for using mathematical modelling to clarify parasite transmission dynamics and address problems related to effective parasite control. Ultimately, through this improved application of models to research and management, we expect that parasite control would be an achievable goal bringing benefits to a vast number of our fellow human beings.

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Progress in Modelling Malaria Transmission
David L. Smith* and Nick Ruktanonchai

Abstract
Transmission of human malaria is a complicated dynamic process that involves populations of humans, parasites, and vectors. The first mathematical models of malaria are now more than a century old, and they are still a useful conceptual synthetic description of transmission, but they fail in some important ways. To address some of those failures, malaria transmission models have now been extended to consider malaria immunity, superinfection, and heterogeneous biting, among other factors. These extensions of the basic theory often arise from field studies in a single place, but tests of the theory come comparing standard measures of malaria taken from many places across the transmission spectrum. Several good models now exist that describe these basic patterns across the spectrum from low to high endemicity. The future of malaria modeling will involve applying these models to make decisions about real systems and finding new ways to test the underlying causes of the patterns.

Modelling Malaria Transmission, a Historical Introduction
Ronald Ross demonstrated that mosquitoes transmit malaria parasites. In the years that followed, malaria control programs focused on reducing mosquito densities, largely through larval vector control. Eleven years after Ross’s seminal discovery, he wrote about malaria transmission in Report on the Prevention of Malaria in Mauritius:2

The infection rate varies, not only in neighbouring places, but even in the same place from year to year.... What is the cause of these variations?
This question is so important as regards both the general theory of malaria and the subject of prevention that we must endeavour to obtain clear ideas about it by careful reasoning (pp. 30).

In the subsequent pages, Ross made clear that “careful reasoning” meant a quantitative description of malaria transmission and proceeded to describe the first mathematical model of malaria. Waite provided a longer description of the model and detailed analysis. He noted that the malaria rate, now called the parasite rate or the prevalence of malaria, was constantly changing due to new infections, recovery, human migration, births and deaths. He considered migration to be too complicated to treat sensibly in a simple model, so like Ross he avoided the distraction and considered the malaria rate in a closed population. He described other aspects of the model in the following way:

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I shall assume ... that both the human population and the anophelines are uniformly distributed over the area in question; that persons of all ages, whether they have previously suffered from malaria or not, are equally liable to infection; that the number of anophelines per unit of the population remains approximately constant during the period under review; and further that infected and non-infected persons are equally exposed to the risk of being bitten (pp. 422).

Waite was forthright about the limitations of his model; he considered the conclusions to be “tentative and suggestive,” but he hoped they would throw light on the factors that affected the malaria rate for “specialists who were considering remedies for the disease.” He pointed out that all of the simplifying assumptions of his model should be reconsidered and refined. Over the past century, knowledge about the parasite, vector and humans and their interactions has improved substantially and the terminology and technology has changed. Malaria models have expanded to consider most aspects of transmission, but the malaria-modelling pioneers laid out the basic malaria modelling agenda. From the beginning, malaria modelling has been motivated by malaria prevention. Models were developed to provide some useful general guidance provided the limitations were considered and soon there was progress in moving beyond the limiting assumptions of the simple models.

The next seventy years saw a part of the agenda fulfilled. Ross published a second mathematical model in 1911 in the second edition of The Prevention of Malaria and again in an article to Nature; the model retained the same simplifying assumptions, but it was formulated as a set of ordinary differential equations. This model was analyzed in great detail by Alfred J. Lotka a year after its publication and in a series of five papers, expanded on some of the earlier themes, notably the delay caused by sporogony. George Macdonald also considered the importance of this delay for control, noting that only old mosquitoes are infectious, so killing adult mosquitoes to shorten lifespan would be especially effective at reducing transmission. Macdonald’s analysis underestimated the potential benefits of killing adults since, by the assumptions of his model, the density of adult mosquitoes would also be reduced and the depletion of adults could potentially reduce recruitment of adults from larval habitat. Macdonald’s analysis also had some historical significance because it helped explain why DDT had been so effective at controlling malaria in early trials. This analysis helped to launch the Global Malaria Eradication Programme, operationalized as time-limited, intensive indoor residual spraying with DDT. Controversy over the programmatic implementation of malaria control during the Global Malaria Eradication Program and the failure to eradicate malaria continues, but the successes of malaria control during GMEP should also be remembered. This simple elaboration on the model fulfilled the mission of providing useful advice to people who do malaria control: few would argue against the great control potential of reducing the mosquito lifespan and that it should be pursued as an option if it is operationally feasible.

An important simplifying assumption from the first models was that untreated malaria infections in humans would clear at a constant per-capita rate. This is equivalent to assuming that the waiting time to clear an infection is exponentially distributed and that it would not differ in a person reinfected with one or multiple different parasites of the same species, or with multiple parasite species. In 1915, McKendrick first considered the problem of multiple infections and relapses in persons that have been infected “once, twice, thrice, etc. by the disease in question.” In 1947, Walton revisited the question of multiple infections and provided a formula to describe the distribution of the number of infections that would be carried in a population. Macdonald also considered the question of multiple infections, but as others have described in some detail, his mathematical formulation does not match the formal description in the paper: the model was supposed to describe infections that clear independently, but it actually described infections that queue up and clear sequentially. Dietz, et al derived a simple formula for the waiting time to clear parasites when infections are complex, when the system is at the steady state and when new infections arrive at a constant rate, and clear independently at a constant per-capita rate. The model that was developed for a large-scale malaria control trial in Garki, Nigeria is now called the Garki Model, which also considered some aspects of immunity. Dynamical equations describing
changes in the distribution of the number of infections were described and analyzed by Bailey and Dietz has written a synthetic review that considers all of these issues in some detail. By 1980 many of the assumptions of the simple models had thus been challenged, but the ones describing transmission remained fairly unchanged. Models up to that point had assumed that populations were closed (i.e., no migration), that human population sizes were essentially infinite and that biting was evenly distributed over space and time. Is biting even over space and time? How far do parasites move in a generation? How far do mosquitoes disperse after completing 4-6 feeding cycles, the time required to complete sporogony? How many humans live within the radius of parasite or mosquito movement? Is biting within those distances randomly or proportionally distributed or is it poorly mixed? These questions are, perhaps, most relevant if the goal of malaria control is to understand how malaria control programs work or if the goal is to contain the spread of antimalarial drug resistance. There is substantial evidence that some people are bitten more than others based on body size, body odors, proximity to larval habitat and potential alternate hosts, housing design, preferential biting of infectious hosts and other factors. Recent theory has progressively relaxed the assumptions of ideal mixing to include heterogeneous biting, preferential biting of infectious hosts, patch-based models of transmission on heterogeneous landscapes, mosquito oviposition behavior and finite host populations. These are increasingly drawing the connection between reasonably simple descriptions of heterogeneous biting in closed populations (i.e., without migration) and “realistic” descriptions of transmission that focus on the details of mosquito behavior. In this chapter, we will review theoretical and empirical studies that have moved beyond the simple assumptions that were made by Waite and Ross.

Complexity, Parsimony and Robust Descriptions of Transmission

Why should malaria be modelled and what sort of model is worth making? Again, Ross provides one of the best answers in the introduction to a seminal paper about the use of mathematics in epidemiology, in which he points out that the study of epidemics is fundamentally concerned with numbers. He also notes that different climates display striking differences in the seasonal patterns of cases and in their frequency and he asks what accounts for these differences. “Why, indeed, should epidemics occur at all?” He proposes that there are two ways to answer this question that he called the a priori and a posteriori approaches. A posteriori approaches were synonymous with statistics; “we commence with observed statistics, endeavour to fit analytical laws to them and so work backwards to the underlying cause.” To work a priori, “we assume a knowledge of the causes, construct our differential equations on that supposition, follow-up the logical consequences and finally test the calculated results by comparing them with the observed statistics.” Ross did not believe that it would be possible to answer his general question by relying on one approach or the other. He notes that many papers had investigated epidemics a posteriori, but few had done so a priori. He then proceeds to develop a purely theoretical approach to the “Theory of Happenings” and calls his theory “a priori pathometry”. The name has not caught on, but the paper established some of the foundations of epidemiology and provided rigor to ideas that are so basic in epidemiology, we almost forget they exist.

It is also worth asking what sort of model is worth making. On the subject of model building, George P. Box, a statistician, famously reminds us that “all models are wrong, but some are useful.” It is not enough, however, to merely know that models are wrong—“the practical question is how wrong do they have to be to not be useful?” Few would deny that the Ross-Macdonald model has been conceptually useful, but the model does not describe malaria well enough under field conditions to serve as an adequate quantitative guide for most purposes. Since there are many uses for models, before building a model, it is worth thinking about how the model would be used. Models serve many purposes, including conceptual understanding, prediction, statistical inference, strategic planning, persuasion and decision-making. It may not be possible to describe how a model will be used in advance, so it is often best to build a range of different models and a solid basis for understanding how the models correspond and how well they serve different purposes.
One of the most important functions of a model is to structure thought about quantitative problems. In this context, it is important to recognize the distinction between “theory” and “models.” In science, theory refers to a set of deliberately oversimplified models that have been extensively tested and that are consistent with existing evidence collected from observations and experiments. Developing and testing models form the core activities of scientists, but the word “model” applies equally well to animal models, in vitro models, conceptual models, statistical models and mathematical models. Likewise, mathematical models can have many purposes as part of basic and applied scientific processes and each model should be suited to its purpose. Theory is the unifying framework of science, but it is rarely codified into a single, grand unified model. No model can serve all purposes equally well, because models inevitably make tradeoffs between generality, realism and precision; to be useful, a model must be robust. As an example, a malaria model can strive to be “realistic,” in the sense that the model is increasingly representational, but as the model becomes more representational, it comes to describe one particular location at a certain point in time. Moreover, the more realistic a model becomes, the greater the data needs; however, the availability of data has been a limiting factor. In modelling malaria, it is useful to conduct studies that build a broad range of models and that explicitly consider the question of robustness. Some property of a model is robust if the predictions hold over a wide range of assumptions and models. When models fail to agree, we would say that a finding is not robust and careful analysis would provide guidelines about when the simpler assumptions breakdown. It may not be possible to describe malaria under every circumstance, but it may be useful to find “rules of thumb” and simple functional forms that describe malaria transmission in different contexts and at a range of spatial scales—what approximations are necessary to strike the right balance to have a model that is both simple and generally useful? It may not be possible to build a single model that would work in every place, but it may not be wise to try and build a different model for every place. A practical way forward is to consider adequate models for malaria transmission by context, including the biting habits of the vector and their bionomics and the transmission intensity.

Transmission Intensity and Its Estimations

Before reviewing mathematical models of malaria transmission, it is worth describing how malaria is measured in different contexts. What malarial indices have been developed to measure malaria and its intensity? One of the earliest indices used was the spleen rate, or the proportion of a population with a palpably enlarged spleen. The spleen rate is reasonably noninvasive, easy to measure and it is closely associated with malaria. Other causes of splenomegaly are reasonably rare, making a high spleen rate a good indicator of ongoing malaria transmission. After parasites could be reliably identified by microscopy, the parasite rate became a more commonly used index of malaria transmission intensity, a measure that addresses the proportion of humans with parasites in their blood. A third method of measuring malaria examines the prevalence of some marker of previous infection that is present in blood serum—a measure of the proportion of humans currently with antibodies that developed in response to malaria infection. Typically, seroprevalence rises with age and gives a robust measure of previous infection compared with age. The spleen rate, the parasite rate and seroprevalence all measure the fraction of the population with particular conditions at some point in time. Measures of prevalence are useful in malaria if they consider something that changes slowly—immunological responses wane slowly, untreated infections and splenomegaly last for several months.

An alternative way to measure malaria in humans is to ask how often a person gets infected. Measuring incidence (i.e., counting the number of events) is different from measures of prevalence. Prevalence measures the proportion of the population in some given state, while incidence measures the number of events—how many new infections occurred in a population over some fixed period of time. In models, the per-capita incidence rate is called the force of infection or the hazard rate for infection. Ross called this quantity the happenings rate, h. It is difficult to measure the force of infection, however, since it is impossible to observe infections directly and since many of them never present with clinical symptoms. Despite these difficulties, force of infection can be measured.
by tracking a cohort of uninfected people until they become infected. The average waiting time to infection is the inverse of the force of infection, $1/h$. Since infants are born uninfected, the waiting time to the first infection, called the infant conversion rate, is a measure of the force of infection. So, too, is a measure of the rise in seroprevalence or the parasite rate with age.

Clinical malaria is defined as fever with parasites. The definition of clinical malaria counts anyone who is infected with malaria and gets a fever from any cause, such as a viral or bacterial infection, thus, it is not always known how many of those fevers were due to malaria. An alternative index of transmission intensity would be the prevalence of clinical malaria, the population with clinical malaria at any point in time. Fevers are short compared with infection response, so clinical malaria is less common than infection with malaria. Fevers have multiple causes, moreover, so measures of disease prevalence tend to be a less specific as an index of malaria. Splenomegaly, by comparison is also a measure of disease prevalence, but it is generally more informative because the duration of fever is comparatively short and the incidence of splenomegaly from other causes is lower. Because the prevalence of clinical malaria is not reliable, clinical incidence is often measured by counting the number new cases of clinical malaria that occurred in a population over a fixed period of time. This measure is generally standardized to a fixed population and time, i.e., 200 cases of clinical malaria (or malaria-attributable clinical episodes), per thousand people, per year.

The spleen rate, the parasite rate, seroprevalence, the force of infection and clinical incidence all measure malaria transmission intensity on the human population, but none of them are perfect indices. Some infections are treated and clear before an immune response develops; this and waning immune responses can affect the accuracy of seroprevalence. Treated infections can clear rapidly and untreated infections also clear after some time, so the parasite rate is a measure of recent infection, modulo drug treatment to achieve a cure. Parasite rates tend to rise with age in young children and they decline with age throughout adolescence and adulthood, so crude parasite rate are a poor index of transmission intensity. The parasite rate in older children (aged greater than 2 but less than 10), called the standard parasite rate does provide a reliable, if imperfect, index of transmission intensity. Although complete immunity to infection may not occur in malaria, the incidence of clinical malaria does decline with age. Clinical incidence displays age-specific patterns that differ by endemicity and this makes it an unreliable index of transmission intensity.

Transmission intensity can also be measured by counting infections in the invertebrate host and entomological indices of transmission intensity can and should play a major role in defining interventions. The human biting rate is defined as the number of bites by vectors, per human, per day. The entomological inoculation rate (EIR) is the product of the human biting rate and the sporozoite rate, the proportion of mosquitoes that have sporozoites in their salivary glands. Another way to characterize EIR is the expected number of infectious bites, per person, per day (dEIR) or per year (aEIR). Just as there is a force of infection for humans, there is also a force of infection for mosquitoes. There are two ways to characterize transmission of the parasites from humans to mosquitoes—what fraction of bites on humans infects a mosquito and what fraction of bites on an infected (and presumably infectious) human infects a mosquito? The former measure is called the net infectivity of the human reservoir and the latter is called the efficiency of transmission (denoted $c$).

HBR and EIR are measures of actual transmission intensity, but there is another class of indices that measure potential transmission intensity. One of these measures is the stability index ($S$): how many human bloodmeals does a vector take over its lifetime? The stability index is a composite measure, the product of three quantities: the mosquito lifespan (denoted $1/g$; the probability a mosquito survives one day is denoted $p = e^{-g}$), the length of a feeding cycle (denoted $1/f$; the time elapsed between two bloodmeals) and the number of human bloodmeals as a proportion of all bloodmeals ($Q$). The stability index is the product ($S = fQ/g$). The parasite requires several days to complete sporogony (denoted $n$), and the probability that a mosquito survives long enough for the parasite to complete sporogony ($e^{-gn}$) also serves as a measure of potential transmission. Additionally, a measure of potential transmission is the number of vectors per human ($m$), which
can help measure potential transmission. This is directly related to the rate that new adult vectors are recruited into the mosquito population, scaled by human population density (denoted \( l \) and \( m = l/g \)). Vectorial capacity (\( V \)) is defined to be the number of infectious bites that would eventually arise from all the mosquitoes that bite a single human on a single day. It is related to the other indices by the formula \( V = \int S^2 e^{-\rho} \).

The most useful index of potential transmission is the basic reproductive number (denoted \( R_0 \)), defined as the expected number of mosquitoes that would arise from a single infected mosquito after one complete generation of the parasite. It is calculated by summing all of the infectious bites that arise from the mosquitoes that are infected by a single person (i.e., vectorial capacity) over the entire duration of a simple, untreated infection. If parasites clear from humans at the per-capita rate \( r \), then the average waiting time to clear an infection is \( 1/r \). Not all infectious bites actually produce an infection, only a fraction do (denoted \( b \)). It is defined for a population with no immunity and no malaria control. The formula for the basic reproductive number is related to vectorial capacity:

\[
R_0 = \frac{b c V}{r}
\]

This number forms a foundation for understanding malaria transmission, but its derivation relies on the simplifying assumptions made by Ross of an ideal, well-mixed population.

This list of malarialometric indices is not comprehensive, but it does give a conceptual overview of the points in the life cycle where the intensity of malaria transmission can be measured. In malaria epidemiology, it is important to measure these quantities for many reasons. These malarialometric indices provide a critical test of the models (for example, see ref. 45). How well do the models capture the variability in these indices and their relations to one another across the range of transmission intensity? The Ross-Macdonald model proposes that there is a functional relationship between these indices, but it does not incorporate immunity or a range of other factors that are known to be important. The Garki model does incorporate some aspects of superinfection immunity, but it does not correctly predict what would happen if malaria transmission were abruptly interrupted. Indeed, the frontier in malaria modelling today is to fulfil Ross’s vision of creating a model based on a set of a priori assumptions that is capable of explaining all of the patterns and much of this involves retreading old ground.

**Preferential Biting and Uneven Exposure**

The first major departure from ideal population mixing was motivated by studies from sexually transmitted diseases. The number of sexual partners and sexual activity in a population of humans are clearly uneven; many people are not sexually active, some are absolutely monogamous, some people have multiple partners and a few people in the population are highly sexually active. The idea is intuitive to most of us who suspect that everyone else is having more sex than us. If one thinks of sex as a risky behavior, it serves as a useful metaphor for malaria exposure.

It would be surprising if exposure to malaria were evenly distributed among all the humans in a population, but how large is the variance in biting heterogeneity and how is it described? One way is to characterize the proportion of bites received by a small fraction of humans. It has been proposed that 20% of the population gets 80% of the bites. Mosquito behavior and ecology affect the variability in exposure to malaria and the issue is especially complicated because malaria is transmitted by dozens of anopheline species with different biting preferences, biting habits, larval habitat and bionomics. Heterogeneous biting may also depend on other contextual details, such as the distribution of larval habitats and humans and the availability of alternative human hosts. Despite the variability, most studies of malaria have tended to focus on vectors that bite indoors and at night.

Muirhead-Thomson’s classical study of biting counted bites on family members outside their huts and concluded that adults were bitten more than children and men were bitten more than women. Muirhead-Thomson also observed that infants behaved differently from older children and adults—the infants were more likely to shake off the mosquitoes, even while asleep. Other
studies caught blood fed mosquitoes, analyzed bloodmeals and identified the host they had fed on. These studies found that the proportion of bites that a person received was roughly proportional to the proportion of skin in the household. Another curious pattern was that, relative to body size, children younger than 18 months old were bitten at a much higher rate than their siblings who were older than 18 months. However, Spencer’s study of 8 pairs of mothers and infants sleeping side by side found that the infants were never bitten, even though the mothers were bitten frequently. Pregnant women account for a larger portion of bites. These studies and many others like them (see Port and Boreham, for a review of the early studies) have found different and contradictory patterns depending on where the samples were taken and depending on whether the study evaluated landing catches or did bloodmeal analysis. The robust pattern from all these studies has been that larger individuals were bitten more than smaller individuals.

Other studies have demonstrated that mosquitoes prefer to feed on individuals who carry gametocytes, potentially through certain symptoms such as increased skin temperature and CO2 expiration. Mosquitoes with sporozoites often spend significantly more time probing blood vessels and are more likely to attempt to resume a bloodmeal if the feeding has been interrupted. This altered feeding behaviour also contributes to a higher mortality of infected mosquitoes, probably due to increased feeding activity, which increases likelihood of mortality from host defensive behaviour. Early studies suggested that multiple feeding was rare, but these studies tend to sample mosquitoes at random. Multiple feeding may be more important than it seems if it is more frequent among infectious mosquitoes.

These studies have described the proportion of bites received by individuals who live within the same household, but they don’t address the questions of how many bites are received by a household. One way to count household heterogeneity is to use insecticides to knockdown all the vectors within a household; the distribution of vectors per household is extremely heterogeneous (for example, see Fig. 2a in ref. 46). This raises the question of what accounts for the huge variability in exposure among households.

The choice of a bloodmeal host is determined by the mosquito search algorithm and chance; mosquitoes follow a set of rules, but host choice is probably opportunistic. Mosquitoes begin their search for a bloodmeal host from the larval habitat where they deposited their eggs. As mosquitoes tack across the wind, odor plumes cause mosquitoes to change direction and follow the plume upwind. As they approach the neighborhood of a potential host, different olfactory cues, CO2, movement and body temperature draw mosquitoes to a particular household or host. Mosquito behavior at the point of contact probably depends on host defensive behavior, personal protection and the mosquito’s physiological state: a mosquito that has few energy resources in reserve would be willing to take more risks than one with plentiful resources.

It follows that one would expect to find enormous variability in household risk that will be related to a large set of factors. The easiest one to characterize is proximity to larval habitat; indeed, many studies have found that proximity to larval habitat was an important risk factor for infection or clinical malaria. Other factors include the odors that emanate from a particular host—including the “limberger cheese” smell. The type of house is an important factor in determining household risk—some kinds of houses and housing designs are easier for mosquitoes to find and enter.

It is also important to note that malaria control can be a cause of heterogeneous biting. If some people in a house use an insecticide-treated bednet (ITN), the mosquitoes may be deflected onto those who do not. ITNs can have an area repelling effect and this would deflect some of the mosquitoes from houses that use ITNs onto those who do not. Since ITNs also kill some mosquitoes, delay feeding and redirect mosquitoes onto alternative hosts that would have fed on a human, the average level of risk may be reduced.

All of this discussion has focused on biting that occurs indoors and at night, but vectors bite in the forest in some areas and in the city in others. In different contexts, the cause of heterogeneous biting can will be related to the factors that predispose people to be at risk.
Although it is often useful to try and explain the variance in exposure to malaria, the total variance in exposure is, itself, an interesting quantity for transmission. When biting is heterogeneous, it means that biting is more intense on a small portion of the population and less intense on the rest. This makes it easier for malaria to invade a population when rare, essentially because person who is bitten most frequently is most likely to become infected and most likely to infect a mosquito. Thus, malaria would be able to persist at lower levels of vectorial capacity; if $a^2$ is the coefficient of variation of biting exposure, then $R_0 = \frac{bV}{(1 + a)}/r$. On the other hand, at some higher level of vectorial capacity, the average level of risk would be reduced in the population. This leads to a different relationship between EIR and PR.

Immunity and the Infectious Reservoir

The flip side of human exposure is to ask how often vectors become infected after biting a human. Another way to ask the question is which humans infect most of the vectors. Immunity to malaria is not like immunity to measles—it does not fully protect against infection or disease and it may have poor immune memory. Thus, herd immunity does not tend to develop. In highly endemic areas, immunity reduces parasite densities in older children and adults and it reduces the frequency and severity of disease. Gametocyte densities do decline and adults also naturally develop transmission blocking immunity, where human antibodies in the bloodmeal prevent infection of the mosquitoes. The infectivity of humans thus declines, but direct observations demonstrate that the adult humans in highly endemic areas remain infectious to mosquitoes.

The Garki model described some of these effects of immunity by considering semi-immune individuals that would gain and lose infections; non-immune infected individuals would pass into the semi-immune individuals at some low rate, reflecting the idea that immunity is acquired after exposure to malaria. The Garki model assumed that lower gametocyte densities in semi-immune individuals would render them uninfectious, despite being infected.

After Garki, a new trend started in malaria modelling. The Garki Model was the motivation for considering malaria as a familiar compartmental model, called an SIRS model. The infective classes are typically interpreted as susceptible (S), infected and infectious (I) and recovered and immune (R). The trick was to relabel all “semi-immune” individuals as being recovered and immune because they were not infectious, whether or not they were infected. The strength of the Garki and SIRS models were that they provided a simple conceptual way of explaining why the PR declines in older age classes, when the relationship between PR and age was far from the steady state. PR tends to rise with age in infants to a plateau in young children. At approximately age ten, it starts to decline. It then falls by about a third of the maximum by age 20 and by two-thirds of the maximum late in life. Both the Garki model and the SIRS model produce curves with this basic shape and a simple version of this model has been used to describe shifts in the age at infection when ITNs are deployed, although the models predict that the age when PR first begins to decline would rise sharply at low PR, a trend that was not observed in analysis of data. The failing of the SIRS models is that it often invites misleading comparisons with acute, immunizing childhood diseases such as measles. In malaria, immunity to severe malaria is acquired rapidly and usually early in life, but infection continues throughout life.

Following in the SIRS tradition, the first epidemiological model concluded that immunity would not affect the evolution of resistance. A subsequent model considered the dynamics of infection in semi-immunes, like the Garki model and showed that immunity would, indeed, affect the evolution of resistance. The different predictions arose because the simplifying assumption about immunity in the SIRS model forced a kind of rigidity—a person was either immune or not. When that stiffness was relaxed, the quantitative effects of immunity could play out in the models in interesting ways. Despite being less infective, they were also less likely to have clinical malaria. Since the demand for antimalarial drugs was lower, they would form a natural refuge for drug-sensitive parasites.
One of the great failures of malaria modelling has been its neglect of disease as an important epidemiological state that is distinct from infection. Indeed, there is a sharp distinction between infection and disease in malaria. Clinical malaria is important for estimating burden, for understanding the need for antimalarial drugs and their potential to control malaria and for understanding drug pressure and the evolution of antimalarial drug resistance. Most models assume that blood stage infection is a good measure of the infectious reservoir; this may not be true in places with good health systems where most clinical malaria is treated and cured. In those places, gametocytemia could persist in people who have no other signs of infection. Clinical disease and gametocytemia as important epidemiological states apart from asexual parasitemia remain largely unexplored by mathematical models (but see ref. 61). Incorporating clinical disease, the gain and loss of immunity and infectivity and identifying where these factors are important are important questions in malaria modelling.

Malaria Transmission in Real Populations

The assumption that populations are infinite and well-mixed are useful for formulating theory based on ordinary differential equations, but to apply these to real populations, it is necessary to tie the model populations at a particular place and time. For such populations, the distribution of human populations, local density and the flight distances of mosquitoes matter. Other aspects of mosquito behavior also affect the validity of the basic models. Mosquitoes can senesce, feed multiple times per life-cycle, postpone oviposition (called gonotrophic disassociation), rest in warm humid refugia in otherwise inhospitable environments, feed on nectar and engage in other behaviors that are not accounted for by the simple formulas. Understanding the factors that explain malaria transmission may require some understanding of these factors, but it is probably not necessary to describe every detail of mosquito behaviour in a mathematical model.

As a starting point, it is worth asking how the simple models get things wrong. Models that consider finite human populations and their spatial distribution provide a starting point for thinking about transmission in a more general setting. In a very large, well-mixed population, every infectious bite will land on a different person. In a small population with proportional mixing, the same individuals who make it easier for malaria to persist absorb multiple infections and this will tend to reduce the risk for everyone else and slow down the invasion. Bailey’s approach to space reformulated partial differential equation models of malaria to consider diffusive movement of mosquitoes and hosts. Later work considered diffusive movement of mosquitoes on heterogeneous landscapes by partitioning space into patches and showed that the distributions of larval habitat and humans form a template for mosquito aggregation and hence, for transmission. These extensions of the basic theory beg the question of whether it is necessary to consider local transmission in an explicitly spatial context. How far do mosquitoes move from the time when they become infected and when they become infectious? How bad are models of proportional mixing (i.e., heterogeneous biting) as an approximation to poor mixing (i.e., locally small populations)?

These ideas are particularly important for control. How far do the protective effects of ITNs extend beyond the populations where they are used? How wide does a malaria control zone have to be to minimize the chances of transmission by mosquitoes? How close do cows or other alternative hosts have to be to provide a zooprophylactic effect? At larger spatial scales, the movement of infected mosquitoes is not nearly as important as the movement of infected humans. Some infected mosquitoes do move long distances in the cargo holds of airplanes and ships and one such event sparked a major epidemic in Brazil. Perhaps the most striking example of the global spread of malaria has been the spread of antimalarial drug resistance throughout southeast Asia and South America from a few points of origin and from southeast Asia into Africa. Human movement among areas that have the potential for malaria transmission is hard to describe, but it is an important aspect of malaria spread. This calls for embedding existing models of malaria transmission into a broader metapopulation context as a way of understanding how malaria parasites persist in their vector and human host populations.
Conclusion

Mathematical models are nothing more than a tool for thinking carefully and quantitatively about malaria and they may be the only way for thinking rigorously about complex quantitative aspects of biological systems. As a scientific activity, they often engage in a different way of doing science and they provide a natural bridge to the applied sciences and public health. Models begin from a set of assumptions and work forward—if the model is correct, then the predictions of the model should be reflected in real world phenomena. This methodological difference often makes the logic of modelling seem backwards to other scientists. Ross’s vision was that mathematical modelling would provide a complementary methodology to more conventional statistical approaches. There is, however, an enormous gap between the kinds of predictions that models make and the kind and amount of data that is collected in the field. This is often a failure of modellers to engage with field biologists, but it is also a failure of field biologists to recognize how collaboration with modellers could improve their study designs to account for the real complexity of malaria. The most notable exception of this was the Garki model that was field-tested during the Garki Project. It could be argued, however, that the Garki model was the first time that the models were field tested, more than six decades after the first malaria model. In response to a new call for malaria eradication, there are many questions that arise in planning, monitoring and evaluating malaria control programs that could be informed by modelling. Models could be used in the process, at least in theory, but to be useful, models and modellers must engage with data and many different kinds of scientists to repeat the aims of the Garki project many times, although probably not on the same scale. By iteratively working from a priori to a posteriori and back, we can anticipate a race to the finish. Will we first eradicate malaria or understand it? It may not be possible to do one without the other.

References

Vector Transmission Heterogeneity and the Population Dynamics and Control of Lymphatic Filariasis

Edwin Michael* and Manoj Gambhir

Abstract

Along-standing gap in lymphatic filariasis epidemiology is quantifying the potential effect that heterogeneous infection processes occurring in the major mosquito vector genera may have on parasite transmission and control. Although previous studies have focussed on examining the forms of the density dependent mechanisms regulating larval infection in various mosquito genera, there has been little work done thus far in investigating how such differential processes might interact with density-dependent processes occurring in other stages of the parasite life cycle to influence overall transmission dynamics between areas exposed to different transmitting vector populations. Here, we explore the impact that differences in vector genus-related larval infection dynamics may have on filariasis transmission and control using newly derived parasite transmission models incorporating the forms of the density-dependent processes regulating larval infection in the two major vectors transmitting filariasis, viz. culicine and anopheline mosquitoes. The key finding in this work is that filarial infection thresholds, system resilience, transmission dynamics and parasite response to control efforts, can all be influenced by the prevailing transmitting mosquito genus. In particular, we show that infection thresholds may be raised, system resilience to perturbations lowered and effects of repeated mass treatments in eliminating infection enhanced in anopheline filariasis compared to culicine filariasis, as a direct result of the occurrence and action of multiple positive density-dependent mechanisms influencing infection in this vector-parasite system, such as the “facilitation” function regulating larval infection dynamics in the vector and the inverse probability function governing adult worm mating in the host. These findings indicate that anopheline filariasis may be easier to eradicate than culicine filariasis for a given precontrol infection level, although the actual intensity of interventions required to achieve eradication may in fact be similar to that for culicine filariasis because of the higher infection levels generated as a result of the “facilitation” process in Anopheles transmission areas.

Introduction

It has long been suggested that variable within-vector host nonlinearities, including density-dependent processes affecting parasite larval establishment and development and vector mortality, could act as major factors underlying the observed heterogeneous transmission dynamics of vector-borne diseases. Addressing the impacts of this heterogeneity has critical resonance for

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studies of the population dynamics and control of the vector-borne helminthic disease, lymphatic filariasis, in particular, given the long-held belief that the forms of the density-dependent processes acting on larval infection dynamics may differ significantly between the taxonomic group of mosquitoes involved in parasite transmission.3-8 The key empirical finding from this body of work is that the density-dependent mechanism regulating larval infection in one major group of filarial mosquito vectors, viz. culicine mosquitoes and to some extent *Aedes* mosquitoes,8 may be of the negative density dependent or “limitation” form, whereas larval infection dynamics in the other main parasite-transmitting mosquito vector, the *Anopheles* group of mosquitoes, is thought to be governed by a positive feedback or “facilitation” process.3,4,8 The latter mechanism is particularly interesting because it could lead to an unstable equilibrium population size, below which the parasite would be driven to extinction.3 As discussed previously, this outcome would clearly make anopheline-transmitted filariasis easier to eradicate through intervention efforts,5-7 a conclusion which has reinforced the impression that differences in the prevailing transmitting vector genus could largely underlie the inconsistencies observed in the success of filariasis control programmes in different endemic areas.3,9,10

In reality, however, as pointed out recently8,11 to fully understand the impact of these vector genus-specific infection processes on overall filariasis transmission and control, it is also necessary to consider how these processes interact dynamically with other density dependent processes occurring in other parts of the parasite life cycle.1,2 This is because without considering the action of such multiple density-dependent factors, it will be unclear as to what extent vector-specific infection processes alone will contribute to overall parasite transmission regulation, particularly given the likely stronger negative feedback processes, e.g., acquired immunity, which may regulate infection in human hosts.9,12 Furthermore, recent work has also underscored the crucial impact that the simultaneous presence and interactions between multiple density-dependent processes acting at different points in the life cycle of a population can have on the dynamics of parasite system persistence, including the values of parasite extinction thresholds.1,3,14 These findings suggest that a composite analysis investigating the simultaneous effects of infection processes in both the vector and human host will be crucial to efforts for determining how vector-parasite combinations occurring in different endemic regions will interact with the efforts presently being proposed to achieve the global control of this debilitating parasitic disease.

Here, we review recent work in modelling lymphatic filariasis transmission15 in order to systematically investigate the impact that vector genus-specific density-dependent processes may have on overall parasite transmission, persistence and control. Specifically, we focus on the incorporation and simultaneous analyses of the “limitation” and “facilitation” functions found to govern larval infection, in addition to density-dependent mechanisms thought to regulate adult worm infection in the human host, such as worm mating probability and acquired immunity, in the two major parasite-vector species combinations implicated in the transmission of filariasis, viz. one in which the vector intermediate host are culicine mosquitoes and the other in which they are anopheline. We begin the chapter by first providing details of the life history of the parasite, characteristics and locality of the major transmitting vectors and particulars underlying the specification and quantification of the functional forms for the “limitation” and “facilitation” mechanisms regulating larval development in culicine versus anopheline vectors respectively. This is followed by details of methods that allow successful incorporation of these functions into overall transmission models for describing culicine- and anopheline-transmitted filariasis population dynamics, highlighting in particular the key need for and derivation of expressions that describe the actions of these functions at the host population level. Model analyses are then described for investigating the influence of vector genus-specific density-dependent processes on system stability, infection/extinction thresholds and parasite population dynamics. The import of these vector genus-specific transmission dynamics for the control of lymphatic filariasis from the application of the currently-proposed mass chemotherapy intervention regimens are then examined using model simulations. We end by underscoring the usefulness of these models and future work to improve their utility in resolving remaining questions regarding filariasis transmission dynamics and control.
Lymphatic Filariasis Disease and Parasite Life Cycle

Lymphatic filariasis is a major cause of acute and chronic morbidity of humans in tropical and subtropical areas of Asia, Africa, the Western Pacific and some parts of the Americas, with over some 1.2 billion people thought to live in areas where they are at risk of infection. Of the estimated 120 million cases of people living with active filarial infection currently, 91% are caused by the filarial worm, *Wuchereria bancrofti*, while *Brugia malayi* and *B. timori* infections account for the other 9%. *Brugia timori* is only known to be endemic in Timor and the Flores islands of the Indonesian archipelago. These lymphatic-dwelling parasites cause damage to the lymphatic system which leads to lymphoedema, genital pathology (especially hydrocoele) and elephantiasis in some 41 million men, women and children. A further 76 million, most often with the transmission stage, microfilariae (Mf), in their blood, are thought to have hidden internal damage to their lymphatic and renal systems. Indeed, recent estimates made by the WHO suggest that lymphatic filariasis may be the second leading known cause of permanent and long-term disability worldwide.

The filarial parasites have biphasic life cycles involving the definitive mammalian host and various species of mosquito vectors. *W. bancrofti* seems to be exclusively a human parasite, whereas *Brugia* spp. are zoonotic in limited situations. Given the involvement of vectors, parasite transmission is indirect, with human infection initiated by the deposition of the third-stage (L3) infective larvae on the skin of the host following the bite of an infective mosquito. The deposited L3 larva penetrates the skin at the site of the bite and migrates to the lymphatic system of the host where they mature over 12 months or so into adult male and female worms. Most estimates of the life span of the adult parasite suggest a period of 5-10 years. The lymphatic-dwelling filariae are dioecious and undergo ovoviviparous reproduction resulting in the release of Mf from adult females, which circulate in the host bloodstream. The life-cycle is completed when these Mf are ingested with the bloodmeal taken by female mosquitoes of a susceptible vector species and develop over 10 to 14 days into the infective-stage L3 larvae capable of infecting a host.

Mosquito Vectors of Lymphatic Filariasis

Both the *W. bancrofti* and *B. malayi* filarial parasites are unique among the various mosquito-transmitted parasites in that larval development can take place in several genera of mosquitoes. A great diversity of mosquito spp. in the genera *Culex, Anopheles, Aedes* and *Mansonia* are capable of acquiring and supporting the development of Mf to infective L3 larvae. Indeed, based on the geographic distribution of these vectors, three main zones of filariasis transmission are recognized: the South Pacific islands and some limited areas of South East Asia, where *Aedes* vectors predominate; West Africa, Papua New Guinea, Vanuatu and Solomon islands where *Anopheles* mosquitoes are principal vectors; and China, South East Asia, Egypt, East Africa, the Caribbean and Latin America where the infection is transmitted mainly by *Culex* species, although *A. gambiae* and *A. funestus* form the predominant vectors that transmit filariasis in rural East Africa (Table 1). The variations in the biological characteristics of each vector genus listed in Table 1, including the differential vector-specific forms of the density-dependent processes regulating larval infection noted above, provide the bases for the long-held belief that the taxonomic group of mosquitoes involved in filariasis transmission may determine not only parasite transmission dynamics but also the effectiveness of any control strategy aimed at interrupting transmission in an area.

Vector-Parasite Infection Relationships

The impact that vector genera differences can have on filariasis transmission and control was first pointed out by Pichon et al, who suggested that this heterogeneous effect may arise from variations in the form of density dependent processes acting on parasite uptake and development in the different filaria-transmitting vector genera. In particular, these workers and others (see study references given in Snow et al) used experimental data on Mf uptake to L3 development obtained from mosquitoes fed on human volunteers to show that if $x$ is the number of Mf ingested by a mosquito and $y$ is the number which survive to become infective, then limitation is the
negative feedback process occurring when the parasite yield \( y/x \) decreases linearly with \( x \), whereas facilitation is the positive feedback process occurring when \( y/x \) increases with \( x \). The fits of these linearized models to data were first used to show as far back as 1974 that limitation may be most commonly associated with filarial infections of culicine mosquitoes, whilst facilitation may be associated with filarial infections of anopheline mosquitoes. Despite this, (1) variability in the results of fits to individual studies, (2) statistical problems connected with the fitting of linearized versions of essentially nonlinear functions to data\(^{25-27} \) and (3) the lack of appropriately parameterized functional forms for the Mf-L3 relationship (but see Subramanian et al\(^ {28} \)), has meant that the mathematical development of filariasis transmission models incorporating density-dependent regulatory functions reflecting variations in vector genus-related infection dynamics has thus far been hampered, with the result that current models are based on only one form of the functional relationship describing the Mf to L3 development process, viz. that related to the limitation function in culicine mosquitoes.\(^ {29,30} \)

### Quantifying the Mf-L3 Functional Response in Vector Populations

Given the above uncertainties, a major focus of recent work in modelling lymphatic filariasis transmission has been on quantifying the suggested relationships occurring between Mf uptake and L3 development in culicine and anopheline mosquitoes in the form of functional response equations that allow ready incorporation into overall filarial parasite transmission dynamics models.\(^ {8,15,28} \) Here, we describe particulars of our recent work in this area based primarily on specifying and fitting the appropriate functional response models to published mosquito feeding experiment data, as follows.\(^ {15} \)

First, for *Culex* mosquitoes, given that a limitation response occurs which results in a saturation in the production of L3 larvae as Mf loads in bloodmeals increase,\(^ 8 \) the Mf-L3 uptake and development data can be fitted to the function:\(^ {30} \)

\[
\text{(1)}
\]

where \( m \) is the Mf density in the human host (20 \( \mu \)L blood), \( \kappa_{31} \) is the maximum limiting value of L3 numbers developing in the mosquito and \( r_1 \) controls the rate at which the L3 development rises with Mf ingested. The fit of this function to data collected from all available published studies\(^ {8,31} \) is shown in Figure 1A. Note that while this is the appropriate functional form for the uptake of Mf and the development of L3 mosquitoes in a single mosquito from one bloodmeal, a fundamental
need, as remarked above, is to account for the effect of the distribution of Mf among hosts in order to quantify the average L3 output in a community if we are to model population-level effects. In order to do this in a manner consistent with field observations, the average L3 level may be calculated in the mosquito populations by assuming that they take bloodmeals from a human host community in whom the mean Mf level is $M$ and is negative binomially distributed with shape parameter $k$. This population-averaged uptake function ($L_{pop}$) can be derived as (see Gambhir and Michael):

$$L_{pop} = \kappa_s (1 - f(M))$$

where:

$$f(M) = \left(1 + \frac{M}{k} \left(1 - e^{-\frac{M}{\lambda_s}}\right)^{-k}\right)$$

This function is very similar to the individual uptake curve, but there is now a dependence on the shape parameter $k$, which determines the level of parasite aggregation in the human hosts.

For anopheline mosquitoes, a closely-related function to that used to model the Culex uptake response, but describing behaviour consistent with facilitation in the part of their Mf-L3 uptake/development curves corresponding to low Mf densities, can be given by the equation:

$$L = \kappa_{s2} \left(1 - e^{-\frac{M}{\lambda_s}}\right)^{-r_s}$$

As for the Culex function, $\kappa_{s2}$ is the maximum limiting value of L3 numbers developing in the mosquito and $r_s$ controls the rate at which L3 development rises with Mf ingested. This function has two features that distinguish it from the Culex function above. The first is the fact that it is raised to the power of two, an operation that introduces a concavity into the shape of the function prior to its limitation-associated saturation at very high Mf loads. The second is the offset parameter ($T$) appearing in the exponent of the exponential function, which ensures that the uptake value rises smoothly from its zero point only when the Mf density in a bloodmeal is

Figure 1. Functional forms relating microfilaria (Mf) uptake (from 20 $\mu$L host blood) and L3 development per mosquito in the two vector genus studied in this work. The squares denote observed data for each vector respectively (sources in Gambhir and Michael). The curve fitted to the (A) Culex data is a limiting function of Mf (equation 1 in the text) whereas the Anopheles curve in (B) describes a development response that begins with a facilitation phase which then approaches an upper limit at higher Mf uptakes (equation 4 in the text).
greater than a threshold density given by the value described by $T$. Biologically, this functional behaviour may be associated with the action of the anopheline cibarial armature or teeth, which prevents Mf from passing undamaged into the mosquito gut at intakes lower than the threshold $T_{52}$ but which allows the undamaged passage of the majority of ingested Mf at higher ingestion loads owing to the protection afforded due to the entanglement of the first passing Mf about the cibarial teeth. The best-fit of the model to published data is shown in Figure 1B. Again, this function needs to be averaged over the host population leading to the derivation of the following population-averaged function:15

$$f(M) = \sum_{n=0}^{\infty} P_{nb}(m; k, M)$$

$$\times \left[ \frac{2e^{-\gamma T/k_{52}}}{e^{-\gamma T/k_{52}} + 1} \right] \left[ \frac{1}{1 + \frac{M}{k} \left( 1 - e^{-\gamma T/k_{52}} \right)} \right] \left[ \frac{1}{1 + \frac{M}{k} \left( 1 - e^{-2\gamma T/k_{52}} \right)} \right]$$

(5)

where $P_{nb}(m; k, M)$ is the negative binomial probability mass function, with mean $M$ and aggregation parameter $k$. Table 2A provides the best-fit parameter values for both the Culex and Anopheles functions given in Equations 1 and 4.

**Derivation of Vector-Specific Models of Lymphatic Filariasis Transmission**

The basic deterministic dynamical model for lymphatic filariasis transmission primarily consists of a series of coupled differential and partial differential equations for three state variables describing the changes in numbers over time of three key parasite life stages—worm burden ($W$), Mf count ($M$) and stage L3 larvae ($L$)—and one state variable describing the acquisition and loss of immunity to parasites in human hosts ($I$).30,33,34 Structurally, these equations are divided into those which describe changes in parasitic infection in the human definitive hosts ($W, M, I$) and the L3 larval equation ($L$), which is used to calculate the change in larval density in the mosquito intermediate hosts. The equations pertaining to changes in parasite stages in the human host population describe change as a function of host age as well as time, since model inputs—such as exposure to mosquito biting—vary over age.33,35 By contrast, because the processes governing the rate of change of the average L3 larval density within the mosquito population operate on much shorter timescales than those in the human host population, the model assumes that the mosquito larval density comes to equilibrium as soon as the corresponding human host parasite and immunity levels are computed. The coupled partial differential and differential equations comprising the basic dynamical model are thus:

$$\frac{\partial W}{\partial t} + \frac{\partial W}{\partial a} = \lambda \frac{V}{H} \psi \psi_{22}^{(a)} b(a) L e^{-\phi t} - \mu W$$

(6)

$$\frac{\partial M}{\partial t} + \frac{\partial M}{\partial a} = \alpha \phi(W, k) W - \gamma M$$

(7)

$$\frac{\partial I}{\partial t} + \frac{\partial I}{\partial a} = W$$

(8)

$$\frac{\partial L}{\partial t} = \lambda_k g \int \pi(a)(1 - f(M)) da - \sigma L - \lambda \psi L$$

(9)
**Table 2A.** Parameter values used in the model for the vector uptake functions. These values were obtained through nonlinear least squares fits the data collated by Snow and Michael\(^3\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Standard Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>(k_{s1})</td>
<td>4.406</td>
<td>0.362</td>
</tr>
<tr>
<td>(r_1)</td>
<td>0.019</td>
<td>0.058</td>
</tr>
<tr>
<td>(k_{s2})</td>
<td>4.395</td>
<td>0.332</td>
</tr>
<tr>
<td>(r_2)</td>
<td>0.055</td>
<td>0.004</td>
</tr>
<tr>
<td>(T)</td>
<td>0</td>
<td>N/A</td>
</tr>
</tbody>
</table>

**Table 2B.** Descriptions and values of the parameters of the vector-specific filariasis models. Typical values of some of the parameters cannot be given due to their multi-valued or dynamic nature\(^{15}\)

<table>
<thead>
<tr>
<th>Parameter Symbol</th>
<th>Definition</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\lambda)</td>
<td>Number of bites per mosquito</td>
<td>10 per month</td>
</tr>
<tr>
<td>(V/H)</td>
<td>Ratio of number of vectors to hosts</td>
<td>Various</td>
</tr>
<tr>
<td>(\psi_1)</td>
<td>Proportion of L3 leaving mosquito per bite</td>
<td>0.414</td>
</tr>
<tr>
<td>(\psi_2)</td>
<td>Proportion of L3 leaving mosquito that enter host</td>
<td>0.32</td>
</tr>
<tr>
<td>(s_2)</td>
<td>Proportion of L3 entering host that develop into adult worms</td>
<td>0.2</td>
</tr>
<tr>
<td>(\beta)</td>
<td>Strength of acquired immunity</td>
<td>0.112</td>
</tr>
<tr>
<td>(\mu)</td>
<td>Death rate of adult worms</td>
<td>0.0104 per month</td>
</tr>
<tr>
<td>(\alpha)</td>
<td>Production rate of Mf per worm</td>
<td>0.33 per month</td>
</tr>
<tr>
<td>(\gamma)</td>
<td>Death rate of Mf</td>
<td>0.1 per month</td>
</tr>
<tr>
<td>(g)</td>
<td>Proportion of mosquitoes which pick up infection when biting an infected host</td>
<td>0.37</td>
</tr>
<tr>
<td>(\sigma)</td>
<td>Death rate of mosquitoes</td>
<td>5 per month</td>
</tr>
<tr>
<td>(k(M))</td>
<td>Aggregation parameter from negative binomial distribution</td>
<td>(k_0 + k_1M (0.0029 + 0.0236M))</td>
</tr>
<tr>
<td>(h(a))</td>
<td>Parameter adjusting rate at which individuals of age (a) are bitten</td>
<td>Linear rise from 0 at age 0 to 1 at 10 years</td>
</tr>
<tr>
<td>(L^*)</td>
<td>Equilibrium value of the larval density (see Equation 5)</td>
<td>—</td>
</tr>
<tr>
<td>(\phi(W, k))</td>
<td>Mating probability as a function of worm burden (W) and aggregation parameter (k)</td>
<td>—</td>
</tr>
<tr>
<td>(\pi(a))</td>
<td>Probability that an individual is of age (a)</td>
<td>—</td>
</tr>
<tr>
<td>(f(M))</td>
<td>Variable component of the population-averaged Mf uptake and L3-larval development function. This is a function of the average Mf level (M) (see Equations 8 and 10 below for specific functional forms)</td>
<td>—</td>
</tr>
</tbody>
</table>
The above equations are similar to those previously published for the lymphatic filarial transmission model described by Norman et al., but with a major difference in that here a mating probability \( \phi(W, k) = 1 \left(1 + W/2k\right)\) has been incorporated into the mean Mf count \( \langle M \rangle \) equation to account for the production of Mf as a function of density-dependent worm mating within the human hosts. This probability function is dependent upon the worm burden \( W \) and the shape parameter \( k \) of the negative binomial distribution describing parasite dispersion among hosts, with the form selected to account for the dioecious and polygamous nature of \( W. bancrofti \) worms. The dependence of the shape parameter of the negative binomial distribution on the average worm burden is considered to be identical to that used to describe the relationship between Mf prevalence and intensity, i.e., it is a linear function of worm burden with gradient \( k_1 \) and with zero intercept \( k_0 \). Since the rates associated with the vectors are considerably faster than the time taken in the development of worm and Mf burdens, the L3 density is assumed to instantaneously equilibrate.

By setting the right hand side of Equation 9 to zero and rearranging, the following expression is obtained for the instantaneous value of the L3 larval density, \( L^* \):

\[
L^* = \frac{\lambda k \int \pi(a)(1 - f(M)) da}{\sigma + \lambda \psi_1}
\]

here, the parameter \( \kappa \) denotes the saturation value of the vector Mf uptake function as detailed above (equations 2 and 4, i.e., \( K_1 \) denoting the saturation value for the \( Culex \) uptake/development function and \( K_2 \) the corresponding value for the \( Anopheles \) function). This expression thus readily allows the incorporation of the culicine and anopheline larval infection dynamics into the basic filariasis model in order to mimic vector-specific transmission dynamics via both substitutions for \( \kappa \) as described above and by substituting the \( f(M) \) term in equation 10 with equations 3 and 5. Table 2B provides a description of the equation parameters along with the values used in these models (details in Gambhir and Michael).15

**Impact of Vector-Specific Infection Processes on Parasite System Stability, Persistence and Extinction**

**System Equilibria, Transitions and Stable States**

The impacts of the combined vector and host density-dependent processes investigated in this study on the persistence and extinction dynamics of the filarial parasite system can be examined by conducting a numerical analysis of the stability of the solutions of the model equations based on varying initial values of \( L^* \) (details in Gambhir and Michael). The results from these analyses for both the culicine and anopheline models are shown in Figure 2. The graphs depict the steady states of host Mf infection prevalence (converted from modeled Mf loads using the expression: \( P(M, k) = 1 \left(1 + M/k\right)^k \), where \( P \) is the prevalence, \( M \) is the Mf density and \( k \) is the aggregation parameter of the negative binomial distribution15,30), as well as the nature of their transitions that appear along the vector biting rate gradient with and without the inclusion of worm mating probabilities in the respective models. As noted previously, these results, i.e., the existence of multiple parasite system regimes or states demonstrate the likely occurrence of complex dynamics in the transmission of filariasis. Here, however, we focus on the impact of vector genus transmission heterogeneity in giving rise to this complexity. Thus, in the culicine case, when worm mating probabilities were not included and where the L3 uptake and development function shows only limiting behaviour, the results show that the system gives rise to just one threshold—a vector threshold biting rate (TBR) below which the only stable equilibrium is at the zero parasite level (Fig. 2A). Above this transmission threshold, a transcritical system bifurcation appears to occur leading to the existence of endemic stable infection levels that increase in magnitude smoothly and reversibly as biting rates increase. Thus, no worm breakpoints and hence existence of alternate infection states are possible, i.e., only positive endemic states occur above the TBR, the system losing its stable endemic infection state smoothly to converge into a zero-infection state as the vector biting rate is reduced below the TBR (Fig. 2A). The effects
of introducing the worm mating function into the culicine model is shown in Figure 2B. In contrast to the situation when the function was excluded, it is clear that this can bring about a discontinuous jump in the Mf prevalence at the TBR giving rise to the appearance of three equilibria, two stable ones separated by an unstable one, mimicking thus the occurrence of a subcritical bifurcation in the system dynamics. The unstable equilibrium points of the model comprise a set of positive-valued worm breakpoints—the largest of which was found at the TBR—producing an unstable dynamic boundary or separatrix with increasing biting rates across which the system converge to either a zero or endemic stable equilibrium state. Thus, at vector biting rates above the TBR, bistable infection states, one endemic and the other zero-infection, can exist depending upon whether initial L* load values can give rise to infection levels above or below the unstable breakpoint values (Fig. 2B). By contrast, in the anopheline-mediated system in which the L3 development function was of a facilitation form, a subcritical bifurcation was observed in which a discontinuous jump in the Mf prevalence at the TBR occurs regardless of whether the worm mating function is included or not (Figs. 2C,D). The values of the vector biting rate at which the nonzero worm breakpoints exist were also far greater in this case than for the culicine system (Figs. 2B vs C,D). In addition, both the TBR and the maximum
worm breakpoint values were found to be raised by the inclusion of the mating probability function. Note that the values of these infection thresholds represent the first such model-based estimates obtained for anopheline filariasis. Their values compared to those obtained for these thresholds in culicine filariasis (maximal breakpoint prevalence of 1.01% versus 0.53% and TBR of 271 versus just 9 per month) also confirm that parasite elimination thresholds are likely to be significantly higher in anopheline filariasis.

System Resilience to Perturbations

The results of perturbing the *Culex* and *Anopheles* models by varying initial L3 values from the vicinity of the respective unstable equilibria are illustrated in Figure 3. Each age-dependent Mf prevalence curve in these graphs represents the solution of the model equations for a given initial L3 seed value set to a value either above or beneath the breakpoint level for a vector biting rate above the estimated TBR. The results depict that an increasing perturbation below the breakpoint curve leads to an eventual age-Mf prevalence of zero across all ages, whereas a similar perturbation above the breakpoint leads to stabilisation at the equilibrium endemic level (Fig. 3). The sizes of the regions between the unstable and either stable equilibrium state are also depicted in the figure for each model and clearly provide a qualitative measure of the extents of the respective basins of

![Figure 3](image-url)

Figure 3. The direction of change of the Mf prevalence age-profile among human hosts when the intermediate host was culicine and the initial Mf prevalence was (A) above and (B) below the breakpoint value (0.21%) with a vector biting rate of 11 per month; and when the intermediate host was anopheline with initial Mf prevalence (C) above and (D) below the breakpoint value (0.82%) with a vector biting rate of 280 per month. Each age-dependent curve represents a perturbation around the initial unstable equilibrium curve with the black arrows indicating the direction in which these curves are likely to travel on the way to the stable equilibrium. From Gambhir and Michael.15
attraction to either stable state. Given that they in turn provide a measure of the maximum amount each system can be changed before losing its ability to recover to either stable state, these results also graphically offer an insight into the likely resilience of the two parasite systems to each of the stable state. For example, the results show that in culicine filariasis, the basin of attraction to the endemic equilibrium state is much larger in size compared to the case in anopheline filariasis, while the opposite is true in the case of the basins of attraction to the zero infection state (Fig. 3A,B versus C,D). The larger basin of attraction to the endemic state but the smaller basin of attraction to the zero state thus suggests that culicine filariasis may be more resilient to perturbations of the system away from the endemic attractor compared to anopheline filariasis, i.e., the former may be more resilient to extinction compared to the latter.

Figure 4 portrays the effects of perturbing the anopheline system near the vicinity of the unstable worm breakpoint values (by varying initial L3 values) on the corresponding Mf age-profiles for three increasing values of the vector biting rate. For each value of the biting rate, the system was initialised as before, above and below the breakpoint Mf prevalence level (given by the red curves in Fig. 4) and the direction in which the profiles move was then tracked (indicated by the arrows in the figure). The results show that as the biting rate increases, the relative sizes of the basins of attraction for the zero-parasite and endemic equilibrium can change considerably. In particular, the range of Mf prevalence values that led to the zero equilibrium diminished while those that led to the endemic equilibrium (upper curve) grew, indicating that system resilience to perturbations even for anopheline filariasis will be greater at higher vector biting rates.

**System Hysteresis**

The existence of multistable states separated by an unstable boundary when the TBR has been exceeded introduces the possibility of the occurrence of bistability or hysteresis in the transmission dynamics of lymphatic filariasis. Figure 5A illustrates this phenomenon in terms of the equilibrium Mf prevalence versus vector biting rate relationship for the model in which the vector is culicine. The bottom black arrow shows the path taken by the equilibrium Mf prevalence when the biting rate is increased to the point on the far right (the 0.1% Mf breakpoint). Until this point is reached, the Mf steady state remains at zero. There is a jump (a subcritical bifurcation) at this point to the endemic equilibrium stage (upward pointing red arrow) and the steady state remains on this stable branch as the biting rate is increased further. Decreasing the biting rate to a point at
which the endemic state appeared, however, will not return the system to the zero Mf prevalence state. For this to happen, the biting rate needs to be reduced further backward to the TBR occurring at the maximal breakpoint (see top black arrow), at which point the endemic state loses its stability and jumps back to the origin or zero state (downward pointing red arrow) and remains there as the biting rate is decreased further. These different paths followed by the parasite system for increasing versus decreasing the vector biting rate constitute the hysteresis loop effect. Figure 5B shows the equivalent loop for the system in which the vector is anopheline. Here, the loop has a considerably greater range in the space, indicating both that the emergence or re-invasion

Figure 5. Hysteresis loops in the Mf prevalence/vector biting rate plane for (a) culicine and (b) anopheline intermediate hosts, showing the two asymmetrical ways by which a shift between alternative Mf stable states can occur with varying vector biting rates. If the parasite system is on the lower zero state but at high vector biting rates and thus close to the worm breakpoint bifurcation boundary, a slight incremental change in Mf levels may bring it beyond the bifurcation (say at 0.1% Mf prevalence) and induce a drastic shift of the system to its endemic equilibrium (rightmost red arrow). If one attempts to restore the parasite-free equilibrium state by reducing the vector biting rate (black leftward arrow), the system shows hysteresis. A backward shift to the parasite-free equilibrium (leftmost red arrow) will occur only if the vector biting rate is reduced far enough to reach the TBR bifurcation point. The hysteresis loop is wider in extent for the anopheline model compared to culicine-transmitted filariasis. From Gambhir and Michael.15 A color version of this image is available at www.landesbioscience.com/curie.
of the filarial endemic state for a similar input of infected hosts (0.1% Mf prevalence) will occur at a significantly larger vector biting rate and the subsequent state transition to the zero state will require a greater reduction of vector numbers compared to the culicine case (Fig. 5A).

**Impact of Vector-Specific Infection Processes on Age Patterns of Infection**

The results depicted in Figure 6 show that apart from effects on parasite system characteristics, including infection thresholds and dynamics of system resilience to perturbations, vector genus-specific infection processes may also play an important role in contributing to the differential community age-patterns of filarial infection observed between the culicine versus anopheline vectored filariasis endemic areas.9,10 The curves in the figure represent the equilibrium age-Mf prevalence patterns generated by the culicine and anopheline-specific filariasis transmission models for proportionately increasing rates of the monthly vector biting intensity and show that largely as a result of the greater levels of \(L^*\) produced in the case of anopheline filariasis, for proportional increases in vector biting higher age-prevalences of infection will be produced in *Anopheles*-mediated filariasis transmission areas compared to culicine areas. They also indicate the likely greater occurrence of convexity in the age-patterns of infection in anopheline compared to culicine filariasis for proportionately similar vector biting rates. This is because of the greater cumulative experience of L3 infection levels in the anopheline transmission communities, which would give rise to the earlier generation of significant amounts of host acquired immunity in these populations compared to populations exposed to culicine mosquitoes. Field support for these differential infection patterns have come partially from work carried out by Michael and Bundy12 and Michael et al,9 who clearly showed that infection prevalences could be significantly higher for a given vector biting rate in communities exposed to *Anopheles* compared to *Culex* vectors.

![Figure 6. The equilibrium age-Mf prevalence curves predicted by the culicine (upper panel) and anopheline (lower panel) filariasis models for proportionally increasing monthly vector biting rates. Results are shown at just above the TBR and at 1.5 and 2 times the TBR rates estimated for each vector (see text).](image-url)
The Impact of Vector Genus on the Dynamics of Filariasis Control

The impact of the interaction between the culicine- and anopheline-vector infection processes with a key presently proposed mass drug treatment regimen on the dynamics of filariasis infection elimination is modeled and portrayed in Figure 7. The curves in the figures show the simulated yearly decline in mean Mf prevalence in the cases of anopheline and culicine filariasis respectively as a result of annual combined diethylcarbamazine (DEC)/albendazole (ALB) mass drug treatments (efficacy values as listed in the label to the figure) applied at 80% drug coverage to communities with different precontrol infection prevalences. Two features of these results are immediately apparent with respect to the impact of vector-specific population dynamics on filariasis control by mass chemotherapy. First, a notable finding is that for a similar precontrol infection prevalence, mean community Mf prevalences will drop much more steeply to meet parasite elimination thresholds in the case of anopheline filariasis compared to the more gradual decline in infection trends generated for the case of culicine filariasis. This will clearly reduce the number of treatment cycles required to achieve infection elimination in the case of anopheline filariasis for a given precontrol infection level (compare curves in Fig. 7A versus 7B) and is a direct outcome of the two different forms of density-dependent regulation of L3 output occurring in culicine versus anopheline vectors. Essentially, this result reflects the fact that as Mf load drops progressively over time in the host community as a result of repeated drug treatment, the severe negative density-dependent regulation of L3 output will ease in the case of culicine filariasis leading to a proportionately higher output of L3 at lower community Mf loads whereas the opposite result, viz. a lower L3 output as Mf load drops, would occur in the case of anopheline filariasis.

The second notable feature of the results in Figure 7 is that this effect on the duration of annual mass treatments required to achieve parasite elimination will be further enhanced given the likely higher worm breakpoint value that may occur in anopheline (the 1.01% Mf prevalence breakpoint threshold depicted by the dot-dashed horizontal line in Fig. 7B) compared to culicine filariasis (the 0.53% Mf prevalence threshold depicted by the dot-dashed line in Fig. 7A). The dramatic effect that this difference in parasite elimination thresholds and the rates of decline induced in mean community Mf prevalences as a result of drug treatment in anopheline versus culicine filariasis is highlighted by the simulation results shown in the graphs for the high precontrol community prevalence of 20%. These show that while at least 8 annual combined DEC/ALB mass treatments at 80% population coverage will be required to reduce Mf prevalence from an initial level of 20% to under the breakpoint or elimination threshold of 0.5 Mf % in the case of culicine filariasis, the corresponding duration of annual mass treatments in the case of anopheline filariasis to reach the elimination target of 1.01 Mf % will only be in the region of 6 years. The absolute number of annual mass treatments required will be even lower than this for eliminating anopheline filariasis compared to filariasis transmitted by culicine vectors when the precontrol community infection levels are lower than a prevalence of 20% (Fig. 7).

Conclusion

A recurring theme in this volume is the potential complexity occurring in the transmission dynamics of parasitic infections, the key processes of which require careful elaboration from data and the impact of which needs the development of suitable models that are able to incorporate and address interactions between such processes. In this chapter, we have illustrated the importance of both these requirements for investigating a long-standing topic in the transmission of lymphatic filariasis, viz. the impact that vector-specific larval infection dynamics can have on the overall transmission dynamics of this macroparasitic disease. This has not only allowed us for the first time to successfully derive and analyze dynamic deterministic models for describing filariasis transmission by the two major mosquito vectors implicated in the transmission of this disease, viz. culicine and anopheline mosquitoes respectively; it has also enabled the first reliable study of the differential impact that having one or the other of these mosquitoes as the dominant transmitting vector can have on overall filariasis transmission dynamics and parasite control in host populations.
Figure 7. Annual changes in overall community Mf prevalence (scaled to 1 ml blood sampling volume) as predicted by the deterministic models for (A) culicine and (B) anopheline filariasis transmission following an annual intervention program with the DEC/ALB drug regimen. Predictions in terms of the number of annual cycles of treatment required to reach endpoints of 0.5% Mf prevalence for culicine (horizontal dashed line in (A)) and 1.0% Mf prevalence for anopheline filariasis (horizontal dashed line in (B)) are shown for a treatment coverage 80% for various baseline community Mf infection prevalences. All figures are given at the scale of 1 ml blood sampling volume. Drug efficacy values are as follows: percentage worm kill (55%), percentage Mf cured (95%) and months Mf reduced (6 months).
The key technical progress we were able to achieve in this work include (1) the successful specification and estimation of the “limitation” and “facilitation” functions acting to regulate larval uptake and development in culicine and anopheline vectors respectively and (2) the efficient incorporation of these contrasting negative (“limitation”) and positive (“facilitation”) density-dependent functions into existing basic models that contain the other major within-host density-dependent factors acting to regulate adult worm infection for describing the transmission of filariasis in host populations. This has thus allowed us to satisfactorily resolve a long-standing methodological gap that has impeded quantifying the heterogeneous effect of vector-specific density-dependent processes on overall filariasis transmission and control, viz. the effective modelling of the simultaneous effects of the major density-dependent mechanisms acting on infection at various points or stages of the parasite life-cycle.

The results from the analyses of these new models have clearly demonstrated the considerable effect that vector-related diversity in larval infection dynamics can have on filarial system dynamics, worm infection patterns and the dynamics of parasite control in host populations due to repeated mass chemotherapy. The fundamental result that emerges is that this differential vector-related impact is primarily due to variations occurring between each vector-parasite-host combination in the presence and number of positive density-dependent mechanisms regulating parasite numbers at different points in the life cycle. Our analysis of the parasite system dynamics indicate that while in culicine filariasis, in which only one positive density-dependent mechanism—the adult worm mating probability—is likely to operate significantly, relatively low infection thresholds may exist, such thresholds may be significantly higher in the case of anopheline filariasis chiefly as a result of the occurrence of at least two positive density-dependences, one facilitating larval output in the vector and the other governing the mating probability of adult worms in the host. Indeed, we show that not only may this difference in the number of positive density-dependences between the two vector-parasite systems have a differential impact on parasite extinction thresholds, but also that possibly as a result of the weaker effect of mating probability (as opposed to the combined effect of the multiple positive density-dependent factors acting in anopheline filariasis) compared to a comparatively stronger negative effect of acquired immunity, culicine filariasis may be more resilient to control perturbations than anopheline filariasis (see Fig. 3). We show that this is essentially an outcome of both the likely occurrence of a larger extent of endemic infection relative to the worm elimination threshold and the action of strong negative density-dependent feedback mechanisms—which by causing state variables to return towards their original values can produce greater stability in parasite transmission—occurring in the culicine system. Similarly, another important finding of relevance to enhancing parasite control or eradication and sustaining such states is that while hysteresis dynamics, whereby infection emergence and elimination may occur at different points along the vector biting rate (Fig. 5), could occur for both culicine and anopheline filariasis, the vector biting range over which these dynamics may occur would be longer for the latter filarial system making including vector control into intervention strategies to maintain the controlled state more important for this system compared to the culicine case.15

Our modelling of the impact that culicine and anopheline vector-specific infection dynamics could have on filariasis control by annual mass chemotherapy with the combined DEC/ALB drug regimen, a key antifilarial intervention, has supported previous more empirically-dependent conclusions that anopheline filariasis may be easier to eradicate compared to culicine filariasis. However, our analysis based on the explicit vector-specific filariasis transmission models described here, has revealed that the greater effect of this control method in achieving the elimination of anopheline filariasis is due to both the likely higher worm breakpoint or elimination threshold that may exist for this system as well as the steeper fall in infection expected over time with repeated mass treatments as infection levels in the community fall. We have shown here that this steeper fall in particular is an outcome of the lower output of L3 larvae expected with low Mf burdens as a result of the operation of the facilitation function governing larval uptake and development in these mosquitoes. However, it is important to recognize that while these results apply for a given
precontrol infection level, in reality the actual overall duration of required mass treatments in anopheline areas compared to *Culex* transmission areas will depend crucially on the prevailing levels of infection prevalence observed between the two areas. In particular, if on average higher levels of infection are likely to be generated and observed in *Anopheles* areas (as a result of higher L3 output facilitated in anopheline mosquitoes (Fig. 6)), then it is possible that despite the faster rate of decline induced on infection by mass chemotherapy, the higher precontrol levels observed for anopheline filariasis would make the overall duration of treatments required for achieving parasite elimination for this system to be similar to that required in a *Culex* region. This result indicates that a careful consideration of both transmission and control dynamics due to vector infection heterogeneity needs to be taken into account when calculating the desired magnitude of control required to eliminate either filariasis.

Although the present work has demonstrated how addressing vector-specific transmission dynamics via the derivation and analysis of appropriate mathematical models incorporating infection processes in both specific vectors and in the human host can aide the dissection of the complex nature of the transmission and control of filarial disease, the results have also indicated several lines of future research. First, it is clear that a more complete investigation of parasite transmission dynamics will require an examination of how stochastic effects influencing transmission processes would affect the findings presented here. We believe that the dynamical aspects of the current results would still hold broadly in a stochastic context but a quantification of the probabilities associated with extinction events—calculated by examining the outcomes of a large ensemble of model runs—would, however, perhaps be more realistic in guiding control programme design and management. Second, our research has also highlighted the crucial need for improved detection and determination of parameter values for all relevant components and processes occurring in parasite transmission ecology, especially those associated critically with density-dependent mechanisms governing infection transmission in different host-parasite-vector systems, if we are to more fully understand the stability, resilience and extinction dynamics of parasitic systems. In particular, we indicate here a critical need to identify and quantify more reliable mating probability functions for macroparasites, perhaps via the application of novel molecular ecological tools to parasite samples in order to reveal patterns in worm mating behaviours. We also indicate a pressing need to identify and quantify the effects of other possible positive density-dependent mechanisms that may mediate filariasis transmission, such as the role that infection tolerance processes could play in regulating infection at high vector biting rates. Although our focus here was on clarifying the dynamics and control of lymphatic filarial infection, we further also suggest that similar studies will play important roles in improving fundamental understanding of the invasion, growth, persistence and extinction dynamics of other parasitic diseases. Given the current global interest to achieve the eradication of helminth infections, it is clear that undertaking realistic parasite transmission model development and analyses, particularly from the perspective of quantifying the impact of complex population dynamics, in close conjunction with empirical studies has now become a pressing research priority in quantitative parasitology.

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**References**

Modelling Multi-Species Parasite Transmission

Andrea Pugliese*

Abstract

Some models are presented for the dynamics of a host population with two parasite species. The models differ in two main aspects: whether they include direct competition among parasites and whether the analysis is based on some approximation and which one. If the analysis is not constrained by a priori assumptions about parasite distributions, it is found that species coexistence is very unlikely without some kind of direct competition among parasites; on the other hand, coexistence generally occurs when inter-specific competition is lower than intraspecific, similarly to standard theory for free-living species. If hosts differ in their predisposition to infection, but not in an identical way towards the two parasite species, then species coexistence becomes feasible even if inter-specific competition is as strong as intraspecific; in this case, coexistence becomes easier as the variance in predisposition increases. These models do not yield universal predictions for patterns of parasite distributions; an analysis of the mechanisms of interaction in each specific system is necessary for that.

Introduction

Models for host-macroparasite interaction have a relatively long history, starting from the pioneering work of Kostizin1 and with the two seminal papers by Anderson and May2 making a strong impact also on empirical research.3,4 On the other hand, very few authors have studied models with several species of parasites, despite the fact that parasite communities are routinely found and examined in empirical research. This is presumably due to the much higher complexity of the resulting mathematical models (see below) and the difficulties in extending to multi-species the approach (approximation via the negative binomial assumption) that has been so fruitful in the analysis of single species models.

None the less, several interesting models have been developed over the years. In this chapter, I will give a personal review of the subject, mainly focused on the subject of coexistence: what are the factors that lead to species coexistence? In so doing, I will quickly review some examples of dynamic models for two parasite species competition. In the final section, I briefly discuss whether these models give any general insights to understanding for parasite community ecology.

The models studied assume that parasites have only one host (i.e., they are monoxenic) and that infections occur through free-living larvae. I believe that most results would apply to more complex systems as well.

Simple Models for Multispecies Parasite Dynamics

The main difficulty in modelling parasite dynamics is that one cannot simply divide the host population into infected and not, but one has to describe and predict the distribution of para-

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sites among hosts, since the effect of parasites on hosts and on other parasite species depend on the number present in that host and perhaps also in features of their establishment. The typical approach used in one-species macroparasite models has been to choose a priori that the parasite distribution is negative binomial, generally (following Anderson and May\(^2\)) with fixed aggregation parameter \(k\), but also with a varying aggregation parameter\(^4,6\) and to then obtain an equation for the temporal dynamics of the mean parasite burden and, depending on the model, of other variables, such as host and/or free-living larvae density.

To my knowledge, the first model on two macroparasite species interacting with one host species has been proposed by Dobson.\(^7\) His model followed this approach, assuming that each parasite species follows the negative binomial distribution with fixed aggregation parameter (\(k_1\) and \(k_2\)) and that the two distributions are independent; he then derived a model for the dynamics of the two parasite densities, assuming that the two species (and indeed all individual parasites) do not interact except in that each parasite contributes to the death of a host harboring both species. His main result is that there is an ample parameter region in which both species will coexist: the smaller the parameters \(k_i\) are (meaning the more aggregated their distributions), the larger the coexistence region will be.

That model has been extended to communities of parasites,\(^8\) to allow for interference or facilitation between parasites\(^9,10\) and to allow for logistic host growth.\(^11\) The result about parasite coexistence has proved to be robust with respect to all these changes.

Even two species identical in all parameters and differing in just one (for instance, the rate of egg production) can coexist; thus a completely inferior competitor (the one with a lower egg production) can survive. This result is actually rather puzzling and seems to be in contrast with all the theory of competitive exclusion.\(^12\) Competition theory does allow for several species to coexist on a single resource, for instance because of competitive balance shifts along a temporal cycle, or because of a colonization-competition trade-off in a metapopulation setting. In all cases, coexistence of competitors seems always to require the existence of some trade-off between traits.

The problem seems to lie in the a priori assumptions made. It will be addressed here by examining conditions for coexistence in a model based only on explicit assumptions about the interaction mechanisms.

**Structure and Parameters of Models**

The models analyzed here concern the interaction of a host population with two species of monoxenic parasites with infections occurring through free-living larvae. In contrast to the models outlined in the previous section, the model considered here does not contain a priori assumptions about parasite distributions and allows for many types of competition among parasites in the same host. The model is deterministic, with main variables the density of hosts carrying \(i\) 1-parasites and \(j\) 2-parasites, denoted as \(p_{ij}(t)\). Other authors\(^13,14\) start from a stochastic model, but then, to obtain analytic results, use some approximations leading to so-called hybrid models\(^15\) similar to those discussed here.

The system of differential equations satisfied by \(p_{ij}(t)\) is rather cumbersome and can easily be obtained with some book-keeping: it is written explicitly in the Appendix. It can be derived by noting that \(p_{ij}(t)\) may increase because some hosts that were carrying a different number of parasites switch to having exactly \(i\) 1-parasites and \(j\) 2-parasites (\(p_{ij}(t)\) increases also because of new births) and vice versa may decrease because some hosts carrying \(i\) 1-parasites and \(j\) 2-parasites switch to a different number of parasites (or die). Listing the possible transitions and their rates, as in the following Table 1, is then enough to specify the system.

In words, it is assumed here that adult parasites affect (additively) hosts’ mortality and (multiplicatively) hosts’ fertility. Moreover, parasites within a host interact directly by increasing (additively) the mortalities (according to the matrix \(\tau\) that differentiates intraspecies and inter-species effects), decreasing (multiplicatively) the fertilities (according to the matrix \(\gamma\)) and decreasing the probability of establishment of an infecting larva (through the matrix \(\gamma\)). Finally, new infections occur through encounters (at rate \(\beta\)) with free-living larvae.
### Table 1. List of the variables of the model and of the events being modeled together with their rates

<table>
<thead>
<tr>
<th>Variable or Event</th>
<th>Host Transition</th>
<th>Rate or Symbol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Host density</td>
<td></td>
<td>$N = \sum_i p_i$</td>
</tr>
<tr>
<td>Density of hosts carrying $i$ 1-parasites and $j$ 2-parasites</td>
<td></td>
<td>$p_i$</td>
</tr>
<tr>
<td>Density of free-living stages of species 1 [2]</td>
<td></td>
<td>$L_1$</td>
</tr>
<tr>
<td>Birth from a host carrying $i$ 1-parasites and $j$ 2-parasites</td>
<td>$\emptyset \rightarrow (0,0)$</td>
<td>$b(N)(1 - \xi_1)(1 - \xi_2)$</td>
</tr>
<tr>
<td>Death of a host carrying $i$ 1-parasites and $j$ 2-parasites</td>
<td>$(i, j) \rightarrow \emptyset$</td>
<td>$d + \alpha_i i + \alpha_j j$</td>
</tr>
<tr>
<td>Density-dependence in host fertility ($K$ carrying capacity of hosts)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Death of adult 1- [2-] parasites</td>
<td>$(i, j) \rightarrow (i - 1, j)$</td>
<td>$\alpha_1 i + j \tau_{12}$</td>
</tr>
<tr>
<td></td>
<td>$[i, j] \rightarrow (i, j - 1)$</td>
<td>$[\alpha_2 + i \tau_{21} + j \tau_{22}]$</td>
</tr>
<tr>
<td>Rate of parasite establishment</td>
<td>$(i, j) \rightarrow (i + 1, j)$</td>
<td>$b_1 L_1 \psi_1(I - \gamma_{11})(I - \gamma_{12})^j$</td>
</tr>
<tr>
<td></td>
<td>$[i, j] \rightarrow (i, j + 1)$</td>
<td>$[b_2 L_2 \psi_2(I - \gamma_{21})(I - \gamma_{22})^j]$</td>
</tr>
<tr>
<td>Birth from an adult 1- [2-] parasites</td>
<td>$\frac{d}{dt} L_1 = + \left[ \frac{d}{dt} L_2 = + \right]$</td>
<td>$h_1 (I - r_{11})^{i-1}(I - r_{12})^j$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$[h_2 (I - r_{21})^{i-1}(I - r_{22})^j]$</td>
</tr>
</tbody>
</table>
The dynamics of free-living larvae has to be specified; new ones are produced from adult parasites (see Table 1), while they are removed through deaths (at rate $\delta_i$ [or $\delta_j$]) or encounters with hosts. One has then the differential equation

$$\frac{dL_i}{dt} = h_i \sum_{j \neq i} p_{ij}(1 - r_{1i})^{i+1}(1 - r_{1j})^{j} - \delta_i L_i - \beta_i L_i N$$

(1)

and analogously for $L_j$. A usual simplification$^2$ is the assumption of fast dynamics of larvae, so that they are at quasi-equilibrium with adult parasites. From (1), one obtains

$$L_i = \frac{h_i \sum_{j \neq i} p_{ij}(1 - r_{1i})^{i+1}(1 - r_{1j})^{j}}{\delta_i + \beta_i N}$$

(2)

Finally, substituting (2) in the expression (Table 1) for the rate of adult parasite establishment, one obtains that, for a host carrying $i$ 1-parasites and $j$ 2-parasites, this is equal to

$$\varphi_i (1 - \gamma_{1i})^{i}(1 - \gamma_{1j})^{j}$$

with $\varphi_i = \frac{h_i \psi_i \sum_{j \neq i} \theta(1 - \gamma_{1i})^{i+1}(1 - \gamma_{1j})^{j} p_{ij}}{\epsilon_i + N}$ and $\epsilon_i = \delta_i / \beta_i$.

(3)

In the transitions listed, several relevant phenomena of host-parasite interactions have been neglected, such as nonlinear effects of parasite abundance on host mortality, facilitation of parasite establishment through parasite-caused impairment of immune response,$^{16}$ context-dependent (through the sequence of infection by different species) parasite competition.$^{17}$ The system is already complicated enough as it is and indeed it will be simplified to allow analysis; moreover, the main interest of the chapter lies in parasite competition rather than in facilitation.

The resulting system of differential equations is clearly very difficult to study, not least for its size: since the number of adult parasites in a host, $i$ and $j$, can in principle be any number, it is a doubly infinite system; even if we restrict these numbers to a maximum, say 100, we would still have a system of 10,000 differential equations. The idea of simplifying the system by some kind of moment closure, either through a negative binomial assumption,$^7$ or through a normal approximation,$^{14}$ is clearly very appealing, although it is then necessary to understand whether the results are an artifact of the approximation.

A different approach is to limit the study to the computation of the invasion criteria for each species; these are sometimes possible to compute without any approximation.$^{18}$ In this way it is not possible to infer the overall dynamics of the system, but at least one can compute exactly the parameter region that allow for species coexistence.

**Invasion Criteria**

While analyzing the dynamics of a complex nonlinear model, such as that including hosts and two parasite species is very difficult, it is often possible to study the (linearized) dynamics close to an equilibrium. In particular, the computation of the invasion coefficient (i.e., the growth rate of one parasite species in a population close to the equilibrium where hosts coexist with a first population species) is sometimes feasible. Basically, this is an extension of the basic reproduction ratio ($R_0$) of a parasite in a parasite-free population, a quantity fundamental in models for microparasites,$^{19}$ but that can be adapted for macroparasites as well.$^{11,20}$

In some cases it is then possible to characterize a quantity $R_2^1$ representing the basic reproductive ratio of parasite 2 invading a population at the equilibrium $E_1$, where hosts coexist with parasite 1. If $R_2^1$, the population of parasite 2 is able to initially increase and will establish itself; on the other hand, if $R_2^1$, that population will decrease and will eventually get extinct.

Coexistence will be deemed to occur when both $R_2^1$ and $R_2^2$ are greater than 1, i.e. when each parasite species is able to invade an equilibrium with only the other species present, together with
the host. This principle is well established in theoretical population biology and can be justified, under some technical assumptions, through persistence theory. One could extend the same principle to more than 2 species, through the computation of the invasion coefficient of parasite species $n$ into an equilibrium with the first $n-1$ present; however, this is generally very difficult because it requires finding explicitly an equilibrium with more than 1 parasite species.

Generally the basic reproduction ratio $R_1^2$ can be defined of the average number of established adult 2 parasites produced by a newly established parasite 2 during its expected life time. We can split this number in at least three components:

$$R_1^2 = \text{(average number of larvae produced over an adult parasite life time)}$$

* (probability that larva is ingested by a host) * (probability of a successful establishment) 

All these quantities will depend, according to the model used, on the features of the equilibrium $E_1$ since parasite survival and fertility may depend on how many 1 parasites are present in the same host (and perhaps also on host density); the probability that a larva is ingested by a host will depend on host density; the probability of a successful establishment may depend on host immune response, hence on the burden (as a surrogate of the host’s previous exposure) of 1 parasites.

**The Model without Direct Interactions**

This has been the case mainly analyzed in the literature, mainly because it is the simplest, so that it can also serve as a reference for studying the effect of direct interactions. All the terms in Table 1 referring to parasite interactions, the matrices $r$, $r$ and $\tau$ are set equal to 0 after tau.

Pugliese computes the invasion coefficient for that case. As discussed above, this requires to obtain some features of the equilibrium with only 1 parasite species present and then to compute the component of formula (4) for the basic reproductive ratio $R_1^2$.

Without repeating the technical steps presented there, we sketch the main ideas. First of all, at an equilibrium, if parasite interactions are neglected, the probability distribution of parasite burden follows some simple law, depending on hosts’ age: precisely, if parasite infections occur one at the time and hosts do not differ in their resistance to infection, parasite distribution for each age is Poisson with mean

$$X(a) = \frac{\phi}{\sigma + \alpha}(1 - e^{-(\sigma + \alpha) a}),$$

where $\sigma$ and $\alpha$ are parasite mortality and parasite-induced host mortality (see Table 1) and $\phi$ is the equilibrium level of the parasite establishment rate (see Eqn. 3). Alternatively (generally data do not support the assumption of a Poisson distribution of parasite load, even when accounting for host’s age), one can allow for multiple infections or heterogeneity in hosts’ susceptibility to infection, obtaining mixtures of Poisson with levels of aggregation comparable to observed ones.

One can then compute the components of (4). Since $\gamma = 0$, the probability of a successful establishment is equal to $\psi_1$. Since larvae may either die (at rate $\delta$) or be ingested by hosts (at rate $\beta N$), the probability of being ingested is $\frac{\beta N}{\beta + \delta} = \frac{N}{N + \phi}$ with $\epsilon = \phi/\beta$ (see Eqn. 3) and $N$ is host population size at equilibrium; when the equilibrium includes parasite species 1, this will be denoted as $\tilde{N}_1$.

The only part of (4) that requires lengthy computations is the average number of larvae produced over an adult parasite life time. First of all, one notes that in this model ($\gamma = 0$), fertility is constant ($h_2$); then one has

$$R_1^2 = \frac{b\psi_1 \tilde{N}_1}{\epsilon_2 + \tilde{N}_1} T_1^2$$

where $T_1^2$ represents the expected lifetime of a 2-parasite that has just infected an average host in a population at the equilibrium $E_1$, where hosts coexist with 1-parasites. This can be computed as the average of the expected lifetime of a 2-parasite establishing itself in a host of a given age and parasite
burden; the average will be weighed using the probability \( \bar{N}_1(s) / \bar{N}_1 \) of surviving to age \( s \), (according to the stationary age density) and the Poisson assumption with \( (5) \) as its mean, or the somewhat more complicated distributions following from multiple infections, for heterogeneous hosts.

From the Poisson assumption, one obtains, after some algebra,

\[
R^*_1 = \frac{b_2 \psi_2}{c_2 + \bar{N}_1} \frac{1}{\alpha_2 + \sigma_2} \int_0^\infty \left(1 - e^{-(\alpha_2 + \sigma_2)s}\right) \bar{N}_1(s) ds. \tag{7}
\]

Formula (7) shows that the reproduction ratio of parasite 2 increases with its fertility \( h_2 \), with its probability of establishment \( \psi_2 \), decreases with larval death rate \( b_2 \), with its adult death rate \( \alpha_2 \) and its induced host death rate \( \alpha_1 \). All this is rather intuitive and would not require modelling.

More interesting is the transformation of (7) into an expression containing the corresponding parameters for parasite 1; the expression is particularly simple if \( c_1 = c_2 \), which means that the parameters relative to the larval stages are the same for the two parasite species. One then obtains

\[
R^*_1 = \frac{b_1 \psi_1}{\alpha_1 + \sigma_1} \frac{1}{\alpha_1 + \sigma_1} \int_0^\infty \left(1 - e^{-(\alpha_1 + \sigma_1)s}\right) \tilde{\bar{N}}_1(s) ds \tag{8}
\]

where \( \tilde{\bar{N}}_1(s) \) represents the probability of surviving to age \( s \), at the equilibrium \( E_1 \) of coexistence with parasite 1.

Formula (8) shows clearly a principle: a parasite that is superior to the resident one (higher fertility and lower death rate and induced host death rate) will always be able to invade and will never be invaded by the other. Mutual invasibility can occur only if there exists a trade-off between traits, namely one parasite has a higher fertility (say \( b_2 > b_1 \)), but suffers also from a higher mortality or higher damages caused to its host \( (\alpha_2 + \sigma_2 > \alpha_1 + \sigma_1) \). When a trade-off occurs, Pugliese\(^{18}\) shows that coexistence is possible but only when the parameters are very precisely balanced: for given values of \( \alpha_2 + \sigma_2 > \alpha_1 + \sigma_1 \), there exists a very narrow interval of values \( b_2 \) (with \( b_2 > b_1 \)) that gives rise to coexistence.

This conclusion is in strong contrast with that obtained by Dobson\(^7\) and others,\(^9-11\) that parasite coexistence is easy and is especially facilitated by aggregation in parasite distribution. Given that the model with single infections and homogeneous hosts gives rise to very little aggregation (which is caused by the mixture of Poisson with different mean, because of hosts’ age), it may be not surprising that coexistence is very unlikely in this model.

In Pugliese\(^{18}\) the issue was tackled by analyzing models including mechanisms that built more aggregation into parasite distributions. Precisely, two models were analyzed: one with multiple infections of larvae (with a Poisson distribution of mean \( \lambda \)), the other with heterogeneity in hosts’ susceptibility to infection, i.e., \( \psi_i \) is not constant among hosts, but has a distribution with means \( \bar{\psi}_i \) and \( \bar{\psi}_i \) for the two parasite species.

In both models, the invasion coefficient can be computed from (4) along lines similar to those leading to (7) and (8). The computations are more involved and are not reported here. Eventually, one arrives at an expression of the type

\[
R^*_1 = \frac{b_2 \bar{\psi}_2}{b_1 \bar{\psi}_1} F(\alpha_1, \sigma_1, \alpha_2, \sigma_2) \tag{9}
\]

where the function \( F \) depends on all details of the model, but has two fundamental properties:

a. \( F \) equals 1 if all parameters are the same \( (F(\alpha, \sigma, \alpha, \sigma) = 1) \), which means that if two parasite species differ only in their fertility, the one with higher fertility outcompetes the other;
b. $F$ is decreasing with $\alpha_2$ and $\sigma_2$ (the reproductive ratio decreases, if a parasite suffers higher mortality, or induces a higher death rate on the host) and is increasing with $\alpha_1$ and $\sigma_1$ (it is easier for a parasite to invade, if the resident parasite suffers higher mortality, or induces a higher death rate on the host). These two properties imply that a trade-off between fertility, on the one hand and parasite-or host-induced mortality, on the other hand, is necessary for two parasites to coexist.

The coexistence region could be examined only by numerically computing the function $F$ in (9). It was found (see Fig. 1) that the width of the coexistence region was basically independent of the degree of aggregation induced by the model: one could assume a very large heterogeneity in host susceptibility and thus a very aggregated parasite distribution, but still the potential for parasite coexistence was very limited, basically the same as with homogeneous hosts and little aggregation.

The rationale for the difference in this result from what obtained by Dobson is discussed at length in Pugliese. Basically, Dobson\textsuperscript{7} assumes that parasite distributions are aggregated and independent. On the other hand, if aggregation arises from the fact that some hosts are more susceptible to infection (from whichever parasite species), it is clear that both species of parasites will be found in the most susceptible hosts, so that parasite distribution will be positively correlated; the stronger is host heterogeneity (and thus parasite aggregation), the more positive the correlation will be. Indeed, Dobson and Roberts,\textsuperscript{9} studying a negative binomial approximation with fixed correlation coefficient, show that positive correlation coefficients hinder coexistence.

Clearly, things would be different if some hosts were more susceptible to parasites of species 1 and others to parasites of species 2; this is explored in references 13-14 (see below), but it is a different explanation: coexistence arises because of differential host susceptibility, not because of aggregation per se.
Competition among Parasites

The results obtained in the previous section are very elegant mathematically, providing a subtle reason for coexistence among parasites without direct interactions: the shift in the age-dependence of host mortality when at equilibrium with either parasite species. On the other hand, they are somewhat disconcerting biologically, because coexistence is very unlikely, while routinely parasites of several species are found in the same host populations and individuals. Moreover, it refers to any kind of parasites: even parasites colonizing different organs could coexist only under very restrictive conditions, which seems plain nonsense.

The point is that in the model the only density-dependence mechanism, that keeps parasite load from growing to infinity, is the induced host mortality. Hence, parasite density at equilibrium will be at a sufficient high level to overall induce a significant host mortality. In turn, the high level of host mortality will make it very difficult for a second parasite species to invade, unless it has a higher reproduction number than the resident one and then displaces it.

In short, to make a realistic multiparasite model, it is essential to introduce competition between parasites, whether they are of the same species or of different ones. In Table 1, three levels of competition are considered, translated as rates depending on the number of parasites in a host: parasite mortality will increase, while parasite fertility and probability of successful establishment will decrease.

Empirical evidence exist for all these facts: the number of parasites in a host may be relevant because of its impact on host resources available for other parasites, or because of the immune response induced in the host; in the latter case, it might be better modelling immune response, as depending on the history of infection of individual hosts, rather than on the current parasite load, but this would be rather more complex and current parasite load may be a reasonable proxy for history of infection.

Explicit mechanisms of parasite competition have been introduced in models for competition between two parasite species by Bottomley et al., using the methods developed by Isham and coworkers. They consider separately the effect of parasite load on probability of establishment and on parasite fertility; here I consider the same cases, neglecting the effect on parasite mortality, mainly because of the mathematical complications of this. First, I use the same approach as in previous Section, that leads to an exact computation of the invasion coefficient, hence to finding the conditions for coexistence as mutual invasibility, only for the case of competition acting on parasite fertility. Then, I show the approach by Bottomley et al. that involves the computation of approximate equations for the first two moments of parasite distribution, thus yielding conditions for coexistence but also the dynamic pattern of parasite densities.

Parasite Fertility Depending on Available Resources

Let measure the effect of the load of parasite species on the fertility of parasite species ; precisely, I assume (see Table 1) that the fertility of one parasite of species 1, living in a host that harbors parasites of species 1 and parasites of species 2, is . Note that I use a multiplicative effect (and not an additive one) of parasites, to avoid the fertility becoming negative at high parasites load; as long as and are not too large, that expression can be approximated as , yielding a more usual expression.

Most of the analysis outlined for the model without direct interactions still applies. One can use (4) to find the invasion coefficient for species 2 in a population where the host coexists with species 1. In the equilibrium with only species 1, equation (5) still holds; in this case, however, the average number of larvae produced by an adult parasite is not simply the product of the fertility rate times its expected lifetime, because the fertility depends on the number of other parasites present. The computations (that will be presented elsewhere) can be performed easily using functional-analytic methods. The final result can be written as

\[ R_i = \frac{b_i \psi_i}{\epsilon + \tilde{N}_1} \frac{1}{\alpha_2 + \sigma_2} \int_0^\infty \left(1 - e^{-\left(a_2 + \sigma_2\right)}\right) \exp\left(-\frac{q_1}{\alpha_1 + \sigma_1} \left(1 - e^{-\left(a_1 + \sigma_1\right)}\right) r_{21}\tilde{N}_1(s)\right) ds, \]
or, if $c_1 = c_2$, as

$$R^2_i = \frac{b_i \psi_i}{\alpha_2 + \sigma_2} \frac{\int_0^\infty (1 - e^{-(\alpha_1 + \sigma_1) s}) \exp\left(-\frac{\psi_i}{\alpha_1 + \sigma_1} (1 - e^{-(\alpha_1 + \sigma_1) s}) r_{12} \right) \pi_1(s) ds}{\int_0^\infty (1 - e^{-(\alpha_1 + \sigma_1) s}) \exp\left(-\frac{\psi_i}{\alpha_1 + \sigma_1} (1 - e^{-(\alpha_1 + \sigma_1) s}) r_{11} \right) \pi_1(s) ds},$$

(11)

where $\psi_i$ is the rate at which (at equilibrium) new $i$-parasites establish themselves (see Eqn. 3), while $\pi_1(s)$ is as in (8).

Expressions (10) and (11), that extend (7) and (8), are rather cumbersome, but one can easily use it on a computer to find the parameter values that allow for coexistence, i.e., those for which both $R^2_i > 1$ and $R^2_j > 1$ (see Figs. 2 and 3).

Moreover, one can easily understand some particular cases:

- if all parameters of the two parameter species are identical, then $R^2_i = 1$; hence, if $r_{21} < r_{11}$ (i.e., interspecific is lower than intraspecific competition), $R^2_i > 1$. One then obtains in this context the classical result that two competing species that have the same demographic parameters but use somewhat different resources can always coexist.

- In the extreme case where $r_{21} = 0$ (i.e., the presence of species 1 has no effect on parasites of species 2) and $\sigma_1 = 0$ (hence population density is at its carrying capacity, $K$, independently of the presence of parasites), expression (10) simplifies to

$$R^2_i = \frac{b_i \psi_i}{\alpha_2 + \sigma_2} \frac{K / L}{\int_0^\infty (1 - e^{-(\alpha_1 + \sigma_1) s}) \pi(s) ds}$$

where $L$ is the average length of host life. In this case, $R^2_i > 1$ is simply the condition for species 2 to be able to persist with the host at its carrying capacity. Then two species that do not interact and do not increase host mortality can always coexist, provided each can persist with the host.

Figure 2. The region in the parameter space ($r_{21} = r_{12}$, $h_1$) that allows species coexistence (to the left of the chevron-shaped curves) for $\alpha_1 = \alpha_2 = 0$ (solid curve) or $\alpha_1 = \alpha_2 = 0.1$ (dashed curve). Other parameter values are $r_{11} = r_{22} = 0.8$, $\sigma_1 = \sigma_2 = 2$, $h_1 = 4$, $c_1 = c_2 = 1$, $d = 0.5$, $b = 1$, $K = 1000$. 
Modelling Multi-Species Parasite Transmission

Generally, if $r_{12} = r_{21} = 0.5$ (solid curve) or $r_{12} = r_{21} = 0$ (dotted curve). Other parameter values are $r_{11} = r_{22} = 0.8$, $a_1 = 0$, $\alpha_1 = \alpha_2 = 2$, $h_1 = 4$, $c_1 = c_2 = 1$, $d = 0.5$, $b = 1$, $K = 1000$.

Figure 2 shows how the coexistence region depends on the ratios of parasite fertilities and the strength of their interspecific competition. In the figure, the two parasite species have the same value of all parameters except the fertility $h_2$ and have a common inter-specific competition coefficient $r_{21} = r_{12}$. It can be seen that, when inter-specific competition is very low, the two species will coexist for almost all feasible fertility values; on the other hand, the ratio of fertilities must become very close to 1 for the two species to coexist, as the inter-specific competition coefficient approaches 0.8, the value of the intraspecific competition coefficient. When $r_{21} = r_{12} = r_{11} = r_{22}$, coexistence is impossible, as seen in Figure 2 and shown by (11). It can also be seen from Figure 2 that the coexistence regions are similar for $\alpha_t = 0$ and $\alpha_t = 0.1$ (a rather strong parasite-induced mortality, comparing with the other parameter values).

Figure 2 compares parasite species differing in only one demographic parameter; in Figure 3, the case of a trade-off between fertility and parasite-induced mortality is analyzed, as in ref. 18. One can again see the strong influence of the interspecific competition coefficient $r_{21} = r_{12}$ on the width of the coexistence region, although coexistence becomes anyway more difficult when the competitor is extremely lethal.

The trade-off between fertility and parasite-induced mortality was shown\textsuperscript{18} to be sufficient for species coexistence, in absence of intra- or inter-specific competition, although the rates had to be balanced very carefully. One may wonder whether this can happen also with inter-specific competition equal to the intraspecific one. However, repeating the computations shown in Figure 3 with $r_{12} = r_{21} = r_{11} = r_{22} = 0.8$, one sees that coexistence is impossible for any values of $\alpha_t$ or $h_2/h_1$. Vice versa, one finds a region where both $R^*_1 < 1$ and $R^*_2 < 1$, i.e., both monospecific equilibria are uninvadable. Namely, a (narrow) parameter region may exist where, even if inter-specific
competition is exactly equal to intraspecific one (and could even be slightly lower), the system exhibits competitive exclusion dependent on initial densities, as it happens for Lotka-Volterra systems when inter-specific competition is higher than the intraspecific one.

In Figure 4, two contrasting examples are shown of simulations with inter-specific competition exactly equal to intraspecific one; in the left panel, competition is rather strong \( r_{11}/g_{27} = r_{22}/g_{27} = r_{12}/g_{27} = r_{21}/g_{27} = 0.5 \) and values of \( \alpha_{i} \) and \( \beta_{i} \) have been found, for which both exclusion equilibria are attractive, as discussed above; in the right panel, competition is very low \( r_{11}/g_{27} = r_{22}/g_{27} = r_{12}/g_{27} = r_{21}/g_{27} = 0.01 \) and values of \( \alpha_{i} \) and \( \beta_{i} \) have been found that allow for species coexistence, like in the case without competition.

The simulations in the left panel of Figure 4 show a very simple behavior, typical of two-dimensional Lotka-Volterra competition systems: fast convergence to a one-dimensional manifold connecting the equilibria and then slow convergence to an equilibrium, along the manifold. In this case then parasite competition follows the standard patterns of competition theory and it becomes reasonable the search for some simple approximating system.

In the simulation shown in the right panel of Figure 4, after some initial oscillations host population and parasite loads approach a stable coexistence equilibrium. It may be noted that the population density reached (around 4) is extremely lower than the carrying capacity (1,000); this is not meant to be realistic and is due to the choice of very high induced mortalities and the very low level of parasite competition.

The previous analysis can be applied, in a relatively simple way, when parasite resource competition affects only parasite birth rate. Judging from preliminary simulations, it seems likely, that similar results will hold also when parasite competition increases death rates (through parameters \( r \) of Table 1) or decreases establishment probability (through parameters \( \gamma \) of Table 1). The computations outlined in this Section do not easily extend however, since parasite distribution will not be Poisson for fixed age (as in (5)), so that no explicit formulae can be found, although the ideas of invasion criteria still apply.

Instead, normal approximations can be obtained, as shown in the following Section.

**Normal Approximations**

Bottomley et al. have studied parasite competition through normal approximations. The idea is very simple: from the equations in the variables \( p_{ij} \), or directly from computations of the
possible instantaneous changes in parasite load, one can find equations satisfied by the first moments of the parasite distributions (means, variances, covariances...). Unfortunately, as usual in most complex models in ecology, the equations for lower moments have terms including higher moments; to obtain a closed low-dimensional system, one then needs some form of “moment closure.” A simple approach is the normal approximation, i.e., to assume that higher moments can be expressed in terms of the first and second moments, according to the same relations that hold for a normal distribution. The assumption may not seem very adequate for parasites, since normal distributions are continuous and include a negative part; the negative binomial distribution used since Anderson and May does not have these problems, but cannot be easily generalized to two or more variables; moreover, Bottomley et al. show, through simulations, that the normal approximations works reasonably well, as long as parameter values are not extreme.

The use of the normal approximation requires a correction in the laws used for density-dependence. Precisely, it becomes more convenient assuming that parasite fertility decreases with the number of parasites according to an additive law $h_i(1 - \gamma_{ij} x_i, 1 - \gamma_{ij} x_j)$ and similarly the probability of establishment is $\psi_i(1 - \gamma_{ij} x_i, 1 - \gamma_{ij} x_j)$. For the rule to be reasonable, it should be assumed that fertilities (or probability of establishment) are 0 when the quantities are negative. The following analysis is feasible only without that restriction; however, as long as the parameters $r$ and $\gamma$ are not too large, ignoring the restriction does not make a big difference.

The equation (1) for the larvae still hold, together with the quasi equilibrium approximation (2) that, with the change to the additive law, changes (3) into

$$\varphi_i = \frac{h_i \psi_i N <x_i> - \gamma_{ij} (<x_i^2>-<x_i>)- \gamma_{ij} <x_i x_j>}{\epsilon_i + N} \quad (12)$$

where $x_i$ is the $i$-parasite load and $< >$ represents the average.

It is also easy writing an equation for the total host density

$$N' = N(b(N) - d - \alpha_1 <x_1>-\alpha_2 <x_2>) \quad (13)$$

One can then write equations for $<x_i>$ obtaining

$$<x_i>' = \varphi_i(1 - \gamma_{1i} <x_i> - \gamma_{12} <x_2>) - \sigma_1 <x_i> - \rho_{1i} <x_i x_j> - \alpha_1 <x_i^2>$$

$$- (b(N) - \alpha_1 <x_i>-\alpha_2 <x_2>)<x_i>$$

$$<x_2>' = \varphi_2(1 - \gamma_{2i} <x_i> - \gamma_{22} <x_2>) - \sigma_2 <x_2> - \rho_{2i} <x_i x_j> - \alpha_2 <x_2^2>$$

$$- (b(N) - \alpha_1 <x_i>-\alpha_2 <x_2>)<x_2>. \quad (14)$$

**Competition at Establishment. No Induced Mortality**

In the case where the parasites do not induce mortality ($\alpha_i = 0$) and parasite competition acts only by reducing the probability of establishment ($r_{ij} = \tau_{ij} = 0$), equations (14) become a closed system. In fact, $N$ is then fixed at the carrying capacity $K$ with $b(K) = d$ and (14) reduces (dropping for ease of notation the brackets) to

$$X_1' = \frac{h_i \psi_i K X_i}{\epsilon_i + K}(1 - \gamma_{1i} X_1 - \gamma_{12} X_2) - (\sigma_1 + d) X_1 \quad (15)$$

$$X_2' = \frac{h_2 \psi_2 K X_2}{\epsilon_2 + K}(1 - \gamma_{21} X_1 - \gamma_{22} X_2) - (\sigma_2 + d) X_2$$

Equations (15) have exactly the form of a Lotka-Volterra competition system. Hence, according to the values of the coefficients, one can have the four possible outcomes: equilibrium coexistence of the two species, competitive exclusion of species 1, competitive exclusion of species 2, contingent competitive exclusion (either species may be excluded depending on initial densities).
First of all, it is necessary to assume that each parasite species alone is able to persist with the hosts at the carrying capacity \( K \). This condition can be written in terms of basic reproductive numbers as

\[
R_h^1 \frac{h \psi_i K}{(c_1 + K)(\sigma_1 + d)} > 1 \quad \text{and} \quad R_h^2 \frac{h \psi_i K}{(c_2 + K)(\sigma_2 + d)} > 1.
\]

Then, the conditions for coexistence are

\[
\gamma_{11} \frac{1 - (R_h^1)^{-1}}{1 - (R_h^2)^{-1}} \quad \text{and} \quad \gamma_{22} \frac{1 - (R_h^2)^{-1}}{1 - (R_h^2)^{-1}}
\]

In case \( R_h^1 \) and \( R_h^2 \) are much larger than 1, conditions (17) simply mean that inter-specific competition is lower than intraspecific one.

If both conditions are reversed (intraspecific competition lower than inter-specific one), one obtains contingent competitive exclusion.

Finally, if one of (17) holds and the other not, strict competitive exclusion of one species occurs.

Note that, while (15) are obtained from (14) without approximations, still they are not exact for the complete system because they have been derived neglecting the constraints (see above) that only the positive values of \( h_i(1 - \psi_i) \) and \( \psi_i(1 - \psi_i) \) had to be considered.

### Competition Acting on Parasite Fertility

One has to add equations for the second moments, since they appear in right hand sides of (14) unless \( r_{ij} = 0 \).

Using as variables the variances and covariance \( V = \langle x^2 \rangle - \langle x \rangle^2 \), \( C = \langle x_1 x_2 \rangle - \langle x_1 \rangle \langle x_2 \rangle \), one obtains, after some lengthy calculations, equations satisfied by them, involving the third moments. The normal approximation allows to write the third moments in terms of the first two, using the relation

\[
\langle x^3 \rangle = \langle x \rangle \langle x^2 \rangle - \langle x \rangle^3 + 3 \langle x \rangle^2 \langle x \rangle - 3 \langle x \rangle \langle x \rangle^2 \langle x \rangle \langle x \rangle - 3 \langle x \rangle^2 \langle x \rangle^2 \langle x \rangle \langle x \rangle - 3 \langle x \rangle^2 \langle x \rangle^2 \langle x \rangle^2 \langle x \rangle^2 \text{...}
\]

exact if \( (U, V, W) \) follows a multivariate normal distribution.

The expressions in the general model are very complex and add little insight, though they could be used for numerical computation. I restrict to the case of competition acting only on parasite fertility, the case analyzed above through invasion coefficient and also studied through normal approximations.

First of all, \( \theta_i \) and \( \theta_2 \) are given by (12) that is rewritten now, using \( V_1, V_2 \) and \( C_{12} \) as

\[
\theta_i = \frac{h_i \psi_i N}{c_i + N} (x_i(1 + r_i) - r_{i1}(V'_1 + x_1^2) - r_{i2}(C_{12} + x_1 x_2))
\]

and analogously for \( \theta_2 \). Then equations (14) can be rewritten for this case as

\[
X'_1 = \theta_i - \alpha_i X_i - \alpha_1 V'_1 - b(N)X_1 + \alpha_2 X_1 X_2
\]

Finally, writing the equations for the second moments and applying also (18), one obtains:

\[
V'_1 = \theta_i - \sigma_i(2V'_1 - X_1) + \alpha_2 X'_1 X_2 - (V'_1 - X_1^2)b(N)
\]

\[
C_{12} = -(\sigma_1 + \sigma_2)C_{12} - \alpha_1 X_1(C_{12} + X_1 X_2) - \alpha_2 X_2(C_{12} + X_1 X_2) - (C_{12} - X_1 X_2)b(N)
\]

Equations (13) (20) (21) (22) (with the analogous ones for \( x_2 \) and \( V_2 \)) are a closed system describing parasite competition. One can use them to find the equilibrium with only one species present and then find the invasion conditions for the second species; mutual invasibility could then be considered as denoting coexistence. Unfortunately, analytical computations are still rather difficult and it is generally necessary to resort to numerical computations.
Modelling Multi-Species Parasite Transmission

The normal approximation works effectively only when host death rates are small.14 In the limiting case where \( \alpha_1 = \alpha_2 = 0 \) and also \( b(N) = d \), at equilibrium \( x_1 = V_1 \) and \( C_{12} = 0 \); then, using these relations in (20), the equations for \( x_1 \) and \( x_2 \) take the Lotka-Volterra form (15) and one can easily analyze them.

One can also compare numerically system (13) (20) (21) (22) with the exact system presented in the Appendix; two examples of this comparison are shown in Figures 5 and 6. From the figures it can be seen, first of all, that, in both cases, simulations converge quickly to a coexistence equilibrium (inter-specific competition is half the intraspecific one). Moreover, the same qualitative trends exist for both exact and approximate models; increasing host death and birth rate (while maintaining all other parameters) results in decreased parasite loads, hence in higher host density. Similarly, parasite aggregation and correlation increase with host death and birth rate, since in these models parasite aggregation (and correlation) results from the hidden variable age: for each given age, parasite distributions are Poisson (no aggregation) and independent, but differ in their means; the mixture of these distributions results into a (little) aggregated distribution.

On the other hand, the absolute values predicted for parasite loads are rather different, especially for the more abundant species (2) and so is host density, since it will suffer from parasite-induced mortality. It must be remembered, however, that a multiplicative law for parasite fertility is used in the exact model and an additive law in the normal approximation; the latter results in a lower fertility when there are at least 3 parasites in a host, so it is no wonder that they yield quantitative different results.

**Competition and Host Heterogeneity**

As can be seen from Figures 5 and 6 (right panels), the previous models yield a very low correlation between parasite distribution and a very low aggregation in each. Then, on the one hand, one could feel justified in assuming a priori that parasite distributions are independent; on the other hand, it seems necessary to allow for some aggregation in the distributions, to have a model closer to reality. The approach by Dobson and others15 is to use independent negative binomial distributions for each species, or with a fixed correlation coefficient.9,10

Bottomley et al13,14 have instead modeled a mechanism that produces parasite aggregation, studying its effects on species coexistence, according to the detailed assumptions used. Precisely, they assume that the parameter \( \psi \) (establishment probability) is not a constant, but varies among

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**Figure 5.** Host density and parasite loads vs time (left panel); correlation between the distribution of the two species and aggregation (= variance/mean) of each vs time (right panel) in two simulations of the infinite system. Simulation “A” has \( b = 0.2, d = 0.1 \); simulation “B” has \( b = 1, d = 0.5 \). All other parameter values are the same: \( r_{11} = r_{22} = 0.8, r_{12} = r_{21} = 0.4, \alpha_1 = \alpha_2 = 0.01, \alpha_1 - \alpha_2 = 2, h_1 = 4, h_2 = 5, c_1 = c_2 = 1, K = 1000 \). Initial conditions are \( N(0) = 800 \) and Poisson independent distribution for each parasite of means 0.25 and 0.75, respectively.
hosts according to some given distribution (like in Fig. 1); the equations for the second moments will then involve mean and variance of this distribution. Note that, since $\psi$ is a probability, it must lie between 0 and 1, so that its variance cannot be large; however, it is also possible to assume that the encounter rates of hosts with larvae (parameter $\beta$ in Table 1) is variable among hosts; by a redefinition of parameters one can include this variation in $\psi$ and then its variance can take any value, as will be assumed here.

If some hosts have a higher predisposition to infection (measured by their value of $\psi$), they are more likely to get infected, so that a correlation will build up between $\psi$ and parasite load $x$. The covariance between $\psi$ and $x$, $C_{\psi x}$, will then be a variable of the system.

To keep things simple, I will restrict the analysis to the case where parasites do not induce mortality ($r_i = 0$) and parasite competition acts only by reducing the probability of establishment ($rij = 0$). Then, host population density is fixed at $K$ and equations for parasite loads $x_i$ do not depend on variances (see (15)).

The resulting competition system will assume given (different distributions) for $\psi_1$ and $\psi_2$, summarized by their means $\overline{\psi}_1$ and $\overline{\psi}_2$, variances $V_1^\psi$ and $V_2^\psi$ and covariance ($C_1^{\psi \psi}$). Variables of the system will be parasite loads ($x_1$ and $x_2$) and the covariances between $\psi_i$ and $x_j$ ($C_{\psi x}$). Through some steps,14 one arrives at the following system of equations

\[
\frac{d}{dt}x_1 = \frac{b_1 K}{c_1 + K} X_1 \left( \overline{\psi}_1 - \gamma_1 (C_{\psi x}^{\psi} + \overline{\psi}_1 x_1) - \gamma_2 (C_{\psi x}^{\psi} + \overline{\psi}_1 x_2) \right) - (\sigma_1 + d) x_1
\]

\[
\frac{d}{dt}C_{\psi x}^{\psi} = \frac{b_1 K}{c_1 + K} X_1 \left( V_1^\psi - \gamma_1 (\overline{\psi}_1 C_{\psi x}^{\psi} + V_1^\psi x_1) - \gamma_2 (\overline{\psi}_1 C_{\psi x}^{\psi} + V_1^\psi x_2) \right) - (\sigma_1 + d) C_{\psi x}^{\psi}
\]

\[
\frac{d}{dt}C_{\psi x}^{\psi} = \frac{b_2 K}{c_2 + K} X_2 \left( C_2^\psi - \gamma_2 (\overline{\psi}_2 C_{\psi x}^{\psi} + C_2^\psi x_1) - \gamma_2 (\overline{\psi}_2 C_{\psi x}^{\psi} + C_2^\psi x_2) \right) - (\sigma_2 + d) C_{\psi x}^{\psi}
\]

with analogous equations for $x_2$, $C_{\psi x}^{\psi}$ and $C_{\psi x}^{\psi}$.

In order to study coexistence in model (23), one can, as discussed above, compute the equilibrium $E_i$ with only one species present and find the conditions for invasion from the other species. As shown in Bottomley et al.,14 this can be written as reproduction numbers

\[
R^*_i = \left. \frac{b_1 K}{c_1 + K} \frac{\overline{\psi}_2 - \gamma_2 (C_{\psi x}^{\psi} + \overline{\psi}_2 x_2)}{\sigma_2 + d} \right|_{E_i} = \left. R^*_b \left[ 1 - \gamma_1 (\overline{\psi}_1 C_{\psi x}^{\psi} + \overline{\psi}_1 x_1) \right] \right|_{E_i}
\]

(24)
where all quantities are computed at the equilibrium $E_1$. Moreover, $R_0^2$ and $R_0^1$ are the basic reproduction number given in (16), using the means of $\psi_1$ and $\psi_2$. $\rho$ is the correlation coefficient between $\psi_1$ and $\psi_2$ and $cv_{1}$ and $cv_{2}$ are the coefficients of variation (standard deviation/mean) of $\psi$. The interest of the analysis of this model lies mainly in understanding the role of $\rho$, $cv_{1}$ and $cv_{2}$. In fact, high values of $cv_{1}$ and $cv_{2}$ correspond to highly aggregated parasite distributions at equilibrium; indeed, Dobson\textsuperscript{7} had found that aggregation promotes parasite coexistence.

One can start the analysis by a simple case: when $\rho = 1$, $cv_{1} = cv_{2}$ the conditions for coexistence $R_0^2 > 1$ and $R_0^1 > 1$ become identical to the conditions (17) found without heterogeneity. Hence, when there is perfect correlation among hosts between predispositions to each parasite species, a high aggregation (as long as it is the same in both species) has no effect on coexistence. This conclusion is similar to what found by Pugliese\textsuperscript{18} in the exact invasion analysis of a model without direct interactions (see Fig. 1). When $cv_{1}$ and $cv_{2}$ are different, the conclusion is not as straightforward; one can see that the invasion of a parasite is hampered, if its variation in predisposition is much larger than that of the resident parasite, while is facilitated if its much lower; on the whole, still aggregation does not promote parasite coexistence.

Decreasing the correlation $\rho$ among host predisposition to parasites makes coexistence easier; this can be seen from (24), since the coefficient of $\rho$ is negative. Intuitively, it is clear that, if some hosts are more predisposed to parasite 1 and others to parasite 2, parasite coexistence becomes easier.

Less intuitive is the fact that there is an interaction between the effects of the two parameters. When $\rho < 1$ and $cv_{1} = cv_{2}$, increasing the coefficients of variations (still keeping $cv_{1} = cv_{2}$) makes it easier satisfying the invasion conditions. This follows indirectly from (24), since higher $cv_{1}$ results in lower $x_i$ at equilibrium. Hence, in these circumstances the conclusion that aggregation promotes coexistence may be justified. A quantitative example is shown in Figure 7; it can be seen that an imperfect correlation in predisposition ($\rho < 1$) and high coefficients of variation allow for the coexistence of parasite species that are identical in all demographic parameters, but with inter-specific competition higher than intraspecific one; the effect is not very large, though, unless correlation is rather low.

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure7}
\caption{The maximum value of inter-specific competition $\gamma_{12} = \gamma_{21}$ that allows for coexistence, vs $cv_{1} = cv_{2}$, for different values of $\rho$. Other parameter values are $\gamma_{11} = \gamma_{22} = 0.5$, $\alpha_1 = \alpha_2 = 0$, $\sigma_1 = \sigma_2 = 2$, $h_1 = h_2 = 15$, $c_1 = c_2 = 1$, $K = 1000$, $d = 0.5$.}
\end{figure}
### Conclusion

Several models for parasite competition have been presented. All are rather complex and do not allow for an easy analysis. For this reason, only some special cases have been considered in this chapter and more extensive analyses would be necessary before drawing clear conclusions. In Table 2 I present a summary of the models considered and the mechanisms yielding coexistence in each.

Still, it is possible to state some general, though preliminary, results. Parasite coexistence generally requires intraspecific competition; if parasite populations are controlled only indirectly through their effects on hosts (through mortality or fertility) and do not compete directly for host resources, then at equilibrium host density will be severely reduced and parasite coexistence would require precise balances of demographic parameters that appear rather unlikely. Such a trade-off between parasite fertility and survival (including the host's) would result in a coexistence of metapopulation type, with one species quicker at reaching new hosts and another surviving longer on the colonized ones.

On the other hand, if parasite populations are controlled by their competition for host resources, or other types of interactions among them, then coexistence occurs as long as inter-specific competition is lower than intraspecific one, following a dynamics reminiscent of Lotka-Volterra equations.

A specific feature of host-parasite interactions is host predisposition for infection, that is generally considered the most relevant mechanism generating aggregated parasite distributions, a feature universally observed in empirical surveys. If there exists only a generic predisposition for all parasite species, then this has no effects on the feasibility of parasite coexistence. On the other hand, if the predisposition to different parasite species differs among hosts, then coexistence becomes more likely, as quite obvious intuitively; in this case, a higher variance in predisposition generates both a higher aggregation in parasite distribution and a larger parameter region for coexistence; in this limited sense, it may stated that aggregation promotes coexistence.
Are there general predictions about patterns that can be observed in natural population? At this level of generality, it seems very difficult; for instance, predictions about correlations between different parasite species depend on the mechanisms allowing for coexistence. Positive (but low) correlation coefficients are produced simply by the mixture of hosts of different age (and other groups); strong intraspecific competition, coupled with a low inter-specific one, will make them more positive, as well as a strong variance in generic predisposition to infection. On the other hand, predisposition to different parasite species differing among hosts will force towards negative correlation.

Finally, the dynamics exhibited by the models displayed in this chapter is very simple: quick convergence towards an equilibrium. This indeed is a typical behavior of competition systems and one may wonder whether interactions with hosts can modify this. The examples shown in this chapter have considered a rather limited parameter range, that allowed for a low-dimensional truncation of the infinite system and/or for a reasonable normal approximation. It is known, on the other hand, that host-parasite interactions may induce cycles, especially when parasites affect host fertility, but although variance in host predisposition to infection and parasite competition decrease the likelihood of cyclic behavior. Simple dynamics with quick convergence to equilibrium has been the rule in the examples examined for this chapter; however, it seems likely that more extensive numerical explorations will uncover examples of complex dynamics in host-parasite systems.

Acknowledgements
I thank Roberto Rosà for reading a first draft of this chapter and suggesting several improvements; Michael Edwin for inviting me to write this chapter and being always very supportive.

References
Appendix. The Model Analyzed

The system in the variables $p_{ij}(t)$ arising from the assumptions shown in Table 1 is:

$$
p_g'(t) = q_1(1-\gamma_{11})^{i-1}(1-\gamma_{12})^j p_{g,i-1,j}(t) + q_2(1-\gamma_{21})^{i-1}(1-\gamma_{22})^j p_{g,i,j-1}(t) + (i+1)(\sigma_1 + \tau_{11} + \tau_{12} + \tau_{13}) p_{g,i,j}(t)$$

$$+ j(\sigma_2 + \tau_{22} + \tau_{23} + \tau_{24}) p_{g,i,j}(t) - [q_1(1-\gamma_{11})^{i-1}(1-\gamma_{12})^j + q_2(1-\gamma_{21})^{i-1}(1-\gamma_{22})^j]$$

$$+ j(\sigma_2 + \tau_{22} + \tau_{23} + \tau_{24}) + j(d) p_{g,i,j}(t)$$

$$p_0'(t) = (\sigma_1 + \tau_{11}) p_{0,i}(t) + (\sigma_2 + \tau_{22})(p_{0,i}(t) - (q_1 + q_2 + d)) p_{0,i}(t) + b(N) \sum_{i,j} p_{1,i,j}(1-\xi_1)^i(1-\xi_2)^j$$

where $q_1 = \beta_1 L_1 \psi_1$ and, by convention, $p_{1,i-1,j}(t) = p_{0,i}(t) = 0$.

The system is completed by equations (1) for $L_1$ and $L_2$.

In the quasi-equilibrium approximation, $L_1$ and $L_2$ are given by (2) and $q_1$, $q_2$ by (3).
Metapopulation Models in Tick-Borne Disease Transmission Modelling

Holly Gaff* and Elsa Schaefer

Abstract

Human monocytic ehrlichiosis (Ehrlichia chaffeensis), or HME, is a tick-transmitted, rickettsial disease with growing impact in the United States. Risk of a tick-borne disease such as HME to humans can be estimated using the prevalence of that disease in the tick population. A deterministic model for HME is explored to investigate the underlying dynamics of prevalence in tick populations, particularly when spatial considerations are allowed. The dynamics of HME in a single spatial patch are considered first to determine which model components are most important to predicting disease dynamics in a local ecology. The model is then expanded to spatially-explicit patches on which patch connectivity, the surrounding environment and boundary effects are studied. The results of this investigation show that predicting risk of this disease to humans is determined by many complicated interactions. Areas that would be endemic in isolation may or may not sustain the disease depending on the surrounding habitat. Similarly, control efforts are shown to be far more effective when applied in wooded habitats than in neighboring grassy habitats. Boundary assumptions which describe the reality of increasing habitat fragmentation additionally play a large role in predicting the endemicity of an HME outbreak. Overall, HME and all tick-borne diseases are complex, nonlinear systems that have just begun to be explored.

Introduction

Tick-borne diseases are increasingly affecting human health throughout the world. These diseases include Lyme disease, Rocky Mountain spotted fever, human babesiosis, ehrlichiosis, anaplasmosis, tick-borne relapsing fever, Colorado tick fever and tick paralysis. Each of these tick-borne diseases has a unique life history that includes a combination of three elements. First, one or more tick species serves as a competent vector of the disease, i.e., the tick can transmit the disease. Second, the given tick species has a preferred host or hosts that are reservoir hosts for the pathogen, i.e., the host can transmit the disease back to ticks but is not affected by the infection. Finally, the disease is caused by a pathogen with its own dynamics and these pathogens consist of viruses, protozoa and bacteria including many rickettsia. While tick-borne diseases have great variety in their structure, one feature all of these diseases share is the erratic nature of human outbreaks, which makes the prediction and prevention of future outbreaks quite difficult. Recent literature displays increasing research efforts in this area and there is a growing understanding of the underlying dynamics of tick-borne diseases. This increased comprehension has in turn supported the development of mathematical models to further explore tick-borne disease.

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Human monocytic ehrlichiosis (causative agent: *Ehrlichia chaffeensis*) is a tick-transmitted, rickettsial disease that has recently increased substantially in the USA from 142 reported cases in 2001 to 506 reported cases in 2005.\(^2,3\) HME, as first reported in reference 4, produces a variety of clinical symptoms ranging from a mild illness to a severe, life-threatening disease with a case fatality rate of approximately 3%. In the United States, HME cases are most frequently reported in the Southeastern and south-central States,\(^5\) but the range is rapidly expanding northward.\(^6\)

The lone star tick (*Amblyomma americanum*) is suspected to be the major species that transmits HME\(^7,8\) and the white-tailed deer (*Odocoileus virginianus*) has been identified as a major reservoir host for *E. chaffeensis*.\(^9,10\) The lone star tick is a hard-bodied Ixodid tick with four life stages: egg, larva, nymph and adult. With the exception of the transition from egg to larva, the tick is obligated to acquire a bloodmeal to molt from each life stage to the next. Unlike many Ixodid tick species, the lone star tick uses the same hosts for each life stage and in many areas, the majority of their blood meals are from the white-tailed deer.\(^11\) This simplified life history allows the exploration of dynamics between a single tick species and a single host species. The system may allow us to gain insights into tick-borne diseases that might otherwise be missed because of the additional complications of multiple host species.

This chapter focuses on the analysis of a mathematical model for HME transmission, which is based on a previously published model by Gaff and Gross.\(^12\) We do not repeat their work, but rather, after a brief introduction to the model, we focus our attention on exploring the dynamics resulting from various spatial configurations of that model. These investigations reveal the importance of metapopulation structures and local environmental parameters in predicting the dynamics of HME and in evaluating the efficacy of control efforts, such as the reduction of the tick population through acaricide use. In particular, we will illustrate the importance of correctly modelling local environmental parameters, patch connectivity, the surrounding environment and boundary assumptions in HME analysis efforts.

**Methods**

**Metapopulation Modelling**

Population biologists have been using metapopulation models in some form for nearly one hundred years,\(^13\) with the word “metapopulation” first appearing in 1969.\(^13\) In such models, a patch of land is marked as either inhabitable or uninhabitable. Generally, inhabitable patches have limited connectivity to other inhabitable patches and the focus is on the probability of extinction or survival across a given set of patches. Landscape ecology has emerged more recently\(^14\) and supports the approach of viewing a landscape in continuous patches with differing ecologies. A model using landscape ecology will have four major components: variations in patches, variations in the surrounding environment, boundary effects and patch connectivity.\(^14\) Because tick population dynamics are very sensitive to the local environment, this latter modelling approach is quite useful. Within landscape ecology, there is the so-called patch-matrix theory in which a spatial region is divided into patches and the dynamics in the model occur within each patch while interaction such as migration or colonization is allowed between patches.\(^14\) In the patch-matrix theory, the patch structure is static throughout the time that the model is considered; however, other landscape ecology models may actually consider variable patch structures in which the spatial patterns and patch relationship change over time.\(^15\) In either setting, patches may be considered to have different sizes and elaborately-shaped boundaries. Recent studies have considered the effect of forest fragmentation on tick-host or other species interactions and determined that patch sizes, patch boundaries, patch interactions and direct and indirect consequences of patch placement at the edge of a grid have complex dynamics that must be considered carefully in future studies.\(^16,17\)

In this chapter, we consider a temporally static rectangular grid of uniformly-shaped patches. Each patch will have unique parameter values that reflect its local ecology. Migration will be allowed to each patch’s nearest neighbors based on the gradient in population levels. Because our
patch and habitat assumptions will be spatially explicit but not necessarily spatially realistic, we may want to consider our model to be an example of metapopulation ecology rather than landscape ecology.\textsuperscript{18}

Let’s return for a moment to the four components of metapopulation ecology and discuss their role in our investigations.

**Variations within Patches**

We will investigate the importance of the choice of parameter values to the dynamics of HME outbreaks within a single patch. In general, tick populations fare much better in wooded areas than in grassy areas as a result of exposure to heat and desiccation in open areas such as grasslands. Many of our investigations will explore the importance of the effect of division between grassy and wooded areas in our spatial models on the endemicity of disease.

**Patch Connectivity**

The amount of patch connectivity is critical in our explorations. It isn’t surprising that if migration between neighboring patches is high, then the individual spatial characteristics are quickly lost to a population with converging characteristics. It is difficult and important to determine an appropriate and realistic scheme for patch migration to capture the essential dynamics of a study.

**Variations in the Surrounding Environment**

With migration between patches in our model, we explore the importance of the surrounding environment to the sustainability of disease. For example, if an endemic area is surrounded by nonendemic areas, under what conditions can the disease be sustained when migration to surrounding nonendemic patches is allowed?

**Boundary Effects**

It has been demonstrated that patch edges are critical in spatial dynamics.\textsuperscript{17,22} We illustrate some boundary effects and contrast several choices of boundary conditions.

**Description of the Mathematical Model**

As we explained in the introduction, our model is a simplified representation of the tick-host-HME system using a single host population, a single life stage and a single pathogen. The dynamics are modeled using differential equations on a discrete spatial grid consisting of environmental patches. White-tailed deer (\textit{O. virginianus}) serve as our generic host population and lone-star ticks (\textit{A. americanum}) as the tick population. The population densities of the host and tick population in each patch are denoted by \(N_i\) and \(V_i\) respectively and the densities of the HME-infected and infectious populations are denoted by \(Y_i\) (hosts) and \(X_i\) (ticks). Both hosts and ticks are considered susceptible when not infected. Once infected with HME ticks are assumed to be infected for life.

\[
\begin{align*}
\frac{dN_i}{dt} &= \beta_i \left( \frac{K_i - N_i}{K_i} \right) N_i - b_i N_i + \sum_{j} m_{ij}(N_j - N_i) \\
\frac{dV_i}{dt} &= \beta_i V_i \left( \frac{M_i N_j}{M_i N_i} - V_i \right) - (\delta_i + b_i) V_i + \sum_{j} m_{ij}(V_j - V_i) \\
\frac{dY_i}{dt} &= A_i \left( \frac{N_i Y_i}{N_i} \right) X_i - \beta_i \frac{N_i Y_i}{K_i} X_i - (\delta_i + c_i) Y_i + \sum_{j} m_{ij} Y_i (N_j - N_i) + \sum_{j} m_{ij} \frac{Y_j}{N_j} (N_j - N_i) \\
\frac{dX_i}{dt} &= A_i \left( \frac{V_i X_i}{N_i} \right) (V_j - X_i) - \beta_i \frac{V_i X_i}{M_i N_j} (V_j - V_i) - (\delta_i + b_i) X_i + \sum_{j} m_{ij} \frac{X_j}{V_j} (V_j - V_i) + \sum_{j} m_{ij} \frac{X_j}{V_j} (V_j - V_i)
\end{align*}
\]
Parameter and function names and values are summarized in Table 1. The first equation describes the population dynamics of the host population in patch $i$, for which there is logistic growth with carrying capacity, $K_i$, as well as an external death rate, $b_i$, caused by hunting or removal. The last term in the equation models the migration from the given patch to all other patches at a rate dependent on the population differential between patches and the connectivity between patches. The second equation describes the tick population dynamics, which are similar to the host dynamics with the carrying capacity for the ticks as the product of the maximum number of ticks per host, $M_i$, and the number of hosts, $N_i$. In addition, the external death rate for ticks includes both weather, $\hat{b}_i$, and acaricide, $\delta_i$, components. The first terms in the third and fourth equations describe the new infections for the host and tick populations respectively, with transmission rates given by $A_i$ and $\hat{A}_i$. The next terms reflect the logistic growth and external deaths of the infected host and tick populations and the last terms are the migration terms, which show the proportion of infected hosts and ticks that migrate as given with the rates in the first two equations. The third equation also includes a host recovery rate, $v_i$.

Variations within Patches
Prior to performing our metapopulation experiments, we assess the relative importance of each input parameter’s influence on the results of the experiment. To accomplish this task, we use the Latin hypercube sampling (LHS) technique that is described in detail by references 19 and 20.

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
<th>Baseline Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N_i$</td>
<td>Host population density</td>
<td>Initially 10 hosts/patch</td>
</tr>
<tr>
<td>$V_i$</td>
<td>Tick population density</td>
<td>Initially 3000 ticks/patch</td>
</tr>
<tr>
<td>$Y_i$</td>
<td>Density of hosts infected with disease</td>
<td>Initially 8 hosts/patch</td>
</tr>
<tr>
<td>$X_i$</td>
<td>Density of ticks infected with disease</td>
<td>Initially 80 ticks/patch</td>
</tr>
<tr>
<td>$\beta_i$</td>
<td>Growth rate for hosts</td>
<td>0.20/month</td>
</tr>
<tr>
<td>$\hat{\beta}_i$</td>
<td>Growth rate for ticks</td>
<td>0.75/month</td>
</tr>
<tr>
<td>$K_i$</td>
<td>Carrying capacity for hosts</td>
<td>20 ticks/patch (woods)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>15 ticks/patch (transition)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>10 ticks/patch (grass)</td>
</tr>
<tr>
<td>$M_i$</td>
<td>Maximum number of ticks per host</td>
<td>200/host</td>
</tr>
<tr>
<td>$b_i$</td>
<td>External death rate of hosts</td>
<td>0.01/month</td>
</tr>
<tr>
<td>$\hat{b}_i$</td>
<td>External death rates of ticks</td>
<td>0.05/month (woods)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.05/month (transition)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.1/month (grass)</td>
</tr>
<tr>
<td>$\delta_i$</td>
<td>Acaracide-induced death rate for ticks</td>
<td>0/month</td>
</tr>
<tr>
<td>$A_i$</td>
<td>Transmission rate from ticks to hosts</td>
<td>0.02/month</td>
</tr>
<tr>
<td>$\hat{A}_i$</td>
<td>Transmission rate from hosts to ticks</td>
<td>0.07/month</td>
</tr>
<tr>
<td>$v_i$</td>
<td>Recovery rate of hosts</td>
<td>0/month</td>
</tr>
<tr>
<td>$m_i$</td>
<td>Migration rate between neighboring patches</td>
<td>0.01</td>
</tr>
</tbody>
</table>

Parameter values are taken from reference 12.
We begin by choosing the parameters in Table 1 that we believe are most likely to vary within individual ecologies and we also choose biologically feasible ranges for the parameters. Note that in our LHS runs we combine the natural and acaricide-related tick death rate into a single term, \( \hat{b} \). We then randomly sample each of the nine chosen parameters uniformly across their feasible ranges to create 600 parameter sets. We select numerical outputs that can quantify the behavior of the model in each 30-year simulation: (1) the maximum percent of infected ticks during all time steps, a measure of the intensity of an outbreak and (2) \( R_0 \), the basic reproduction number, which is a measure of the endemicity of an outbreak.\(^\text{12}\) If \( R_0 \) is greater than one, a disease is endemic and if \( R_0 \) is smaller than one, the disease will die out.

After verifying a monotone relationship between each input parameter and each output measure, we compute the partial rank correlation coefficients (PRCC) to determine how much each input parameter contributes to changes in our output measures. In Figure 1 we show the graphs of the residues from multiple linear regressions performed on the input parameter ranks compared to the reproductive number ranks. By ranks we mean that actual values have been replaced with the integer values that correspond to their relative ascending orders. The corresponding PRCC values are shown above each graph with respective statistical significances. The illustration shows the correlation between a strong PRCC value and a linear relationship between input and output ranks. In this analysis, 600 runs with random parameter samplings on a uniform distribution are performed and the PRCC values for runs with several hundred more or fewer samplings return similar values. The strength of the measured PRCC correlation depends strongly on the chosen parameter distributions; in particular, a distribution chosen with a large range will have a stronger PRCC result. The ranges chosen for this analysis are considered to be the most biologically feasible.

![Figure 1. LHS for basic reproductive number on a single patch. Each input parameter is sampled 600 times and subsequently 600 randomly-mixed parameter sets are created. The basic reproductive number \( R_0 \) is computed for each of the 600 parameter sets. Then each parameter set and the resulting \( R_0 \) values are replaced with their integer rank and multiple linear regression is performed on the ranks of each input parameter versus \( R_0 \). The residues are pictured along with PRCC and p-values which indicates the significance of the nonzero PRCC. We see that a stronger linear trend between input and output ranks corresponds to a PRCC-value near 1 (direct correlation) or -1 (inverse correlation). These results are summarized along with other PRCC data in Table 2.](image)
Table 2. PRCC values for selected parameters and outcome measures

<table>
<thead>
<tr>
<th>Parameters Varied for a Single Patch Model</th>
<th>Parameter</th>
<th>β</th>
<th>̂β</th>
<th>K</th>
<th>M</th>
<th>b</th>
<th>̂b</th>
<th>A</th>
<th>̂A</th>
<th>v</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minimum, Maximum</td>
<td>0.05, 0.3</td>
<td>0.25, 0.9</td>
<td>5, 50</td>
<td>150, 500</td>
<td>0.01, 0.1</td>
<td>0.001, 0.5</td>
<td>0.013, 0.03</td>
<td>0.045, 0.1</td>
<td>0.001, 0.12</td>
<td></td>
</tr>
<tr>
<td>Maximum Percent of Infected Ticks in any Month Over 30 Years</td>
<td>PRCC</td>
<td>-0.73*</td>
<td>-0.77*</td>
<td>0.08</td>
<td>0.80*</td>
<td>0.59*</td>
<td>0.80*</td>
<td>-0.45*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basic Reproductive Number $R_0$</td>
<td>PRCC</td>
<td>-0.62*</td>
<td>0.18*</td>
<td>0.054</td>
<td>0.82*</td>
<td>0.45*</td>
<td>0.45*</td>
<td>0.37*</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameters Varied for a Two Patch Model</th>
<th>Parameter</th>
<th>̂β</th>
<th>̂β</th>
<th>$M_i$</th>
<th>̂$b_i$</th>
<th>̂$b_2$</th>
<th>$A_i$</th>
<th>̂$A_i$</th>
<th>v</th>
<th>$m_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minimum, Maximum</td>
<td>0.05, 0.3</td>
<td>0.25, 0.9</td>
<td>150, 500</td>
<td>0.001, 0.5</td>
<td>0.001, 0.5</td>
<td>0.013, 0.03</td>
<td>0.045, 0.1</td>
<td>0.001, 0.12</td>
<td>0.25</td>
<td></td>
</tr>
<tr>
<td>Maximum Percent of Infected Ticks in Patch 1 in Any Month Over 30 Years</td>
<td>PRCC</td>
<td>-0.74*</td>
<td>-0.82*</td>
<td>-0.67*</td>
<td>-0.70*</td>
<td>-0.35*</td>
<td>0.55*</td>
<td>0.83*</td>
<td>-0.18*</td>
<td>0.0061</td>
</tr>
<tr>
<td>Maximum Percent of Infected Ticks in Combined Patches in Any Month Over 30 Years</td>
<td>PRCC</td>
<td>-0.65*</td>
<td>-0.81*</td>
<td>0.56*</td>
<td>-0.36*</td>
<td>-0.35*</td>
<td>0.47*</td>
<td>0.74*</td>
<td>-0.18*</td>
<td>0.032</td>
</tr>
<tr>
<td>Basic Reproductive Number $R_0$ in Patch 1</td>
<td>PRCC</td>
<td>-0.70*</td>
<td>0.13</td>
<td>0.61*</td>
<td>-0.75*</td>
<td>-0.40*</td>
<td>0.53*</td>
<td>0.50*</td>
<td>-0.19*</td>
<td>0.099</td>
</tr>
<tr>
<td>Basic Reproductive Number $R_0$ in Combined Patches</td>
<td>PRCC</td>
<td>-0.67*</td>
<td>-0.03</td>
<td>0.58*</td>
<td>-0.57*</td>
<td>-0.58*</td>
<td>0.53*</td>
<td>0.49*</td>
<td>-0.15*</td>
<td>-0.0069</td>
</tr>
</tbody>
</table>

The PRCC measures close to 1 indicate a strong direct correlation and those near -1 indicate strong inverse correlation. The significances of the nonzero PRCC values are checked with the student T-distribution. Superscript *notations denote PRCC values that are significant at the $\alpha = 0.001$ level.
Referring to the PRCC measurements for the one-patch disease model shown in Table 2, we see that all parameters with the exception of host carrying capacity and host external death contribute significantly to both the intensity and endemicity of HME outbreaks. We note that $\tilde{\lambda}$, the transmission rate from hosts to ticks, is much more important to the intensity than to the endemicity. Of particular interest is the tick growth rate, $\tilde{\beta}$, which contributes weakly, yet directly, to the basic reproductive number and contributes inversely to the outbreak intensity. Biologically this result suggests that a large number of ticks would cause a larger epidemic immediately, but would dilute the infection in the long term. Likewise, host recovery, $v$, decreases the immediate intensity of an outbreak but encourages endemicity; however, the impact of $v$ is fairly weak in comparison to the other parameters and we assume no host recovery as our baseline value.

We note additionally that the one parameter with extremely strong PRCC values for both the intensity and endemicity of an HME outbreak is $\tilde{\alpha}$, the external death rate of ticks. The intensity, but not the endemicity, is also strongly influenced by host and tick birth rates, $\tilde{\beta}$ and $\tilde{\beta}$, as well as by the disease transmission rate, $\tilde{\lambda}$, from hosts to ticks. Looking forward to our simulations with multiple patches, we note that because the value of $\tilde{\alpha}$ will vary in the wooded and grassy patches, we would expect patch dynamics to be important in predicting long-term disease behavior while the short-term intensity of an HME outbreak may not show as much sensitivity to patch dynamics if three of the four the most influential parameters are constant along our patches.

We presume the variables that are biologically the most sensitive to changing in various geographic locations are $\tilde{\beta}$, the external death rate for ticks and $\tilde{\alpha}$, the transmission rate from ticks to hosts. Both of these parameters are found to be significant both to the intensity and endemicity of HME outbreaks in the LHS sensitivity analysis. Differing weather patterns would influence the tick death rate, $\tilde{\alpha}$ and to a lesser extent the tick growth rate, $\tilde{\beta}$. Variability in species diversity of hosts, with some species more susceptible to the disease than others, would influence the value of $\tilde{\alpha}$ and, to a lesser extent, $\tilde{\lambda}$. In Figure 2 we illustrate the effects of varying these two parameters, which are most likely to change within a single patch.

![Figure 2. Importance of the tick death rate and host susceptibility.](image)

Figure 2. Importance of the tick death rate and host susceptibility. The graph on the left shows the long-term annual average of the percent of infected ticks. The vertical axis is the average percent of infected ticks over the last eight years of a 30-year HME simulation. We see that as the environmentally induced death rate of ticks, $\tilde{\alpha}$, increases the percentage of hosts and ticks with HME diminishes. The graph on the right shows the long-term annual average of the percent of infected ticks. We see that as the host susceptibility, $\tilde{\alpha}$, increases the percentage of hosts and ticks with HME grows as well. This is an important parameter to consider when measuring the influence of host diversity.
Patch Connectivity

Following the set-up of reference 12, we began with two patch types: grassy patches and wooded patches. In our baseline setup, both grassy and wooded areas are endemic for HME. However, the parameters for the wooded areas are more favorable to growth of both the tick population and the infection. In particular, two parameters differ from grassy to wooded areas, the host carrying capacity, \( K_i \) and the tick death rate, \( \hat{b}_i \). In our one-patch sensitivity analyses, we found that the value for the carrying capacity was not particularly important in predicting disease dynamics but that the tick death rate did have strong influence. We begin this section with a simple consideration of two connected patches that share all parameter values except for the tick death rate, \( \hat{b}_i \) and we explore how their connectedness influences disease levels.

We apply the LHS sensitivity analysis to the two-patch model and we allow our significant parameters from the one-patch model to vary in the same ranges. We have two significant new assumptions in this experiment. First, the tick death rate, \( \hat{b}_i \), is allowed to vary separately in the individual patches, while the other parameters vary uniformly in both patches. Secondly, we introduce migration as given in the mathematical model and we vary the value of \( m_{ij} \) between 0 and 0.25. The results of the analysis are presented in Table 2. We first observe that the parameters with strong PRCC values in the one-patch simulations have similarly strong values in the two-patch simulations and that the variations in migration levels were insignificant compared to the variations in these parameters. Secondly, we observe that by allowing the tick death rate, \( \hat{b}_i \), to vary individually in each patch, we effectively cut its PRCC value in half as a contributor to the overall dynamics of the infection. Of course, this makes intuitive sense because the parameter is only contributing to half of the area. The parameter’s contribution to the dynamics in its patch of origin is cut only slightly by its interaction with neighboring patches. We can reconfirm in the sensitivity analysis for the two-patch model that the effect of the tick birth rate, \( \beta_1 \), on the disease endemicity is not significant and the value of host recovery \( \nu \) is statistically significant but small. We also confirm that the dynamics of changing the tick death rate, \( \hat{b}_i \), between patches is more influential on the value of the basic reproductive number than to the maximum percent of infected ticks overall.

Though we note above that the PRCC values for the migration rates are insignificant in the LHS analysis, we caution against the conclusion that migration is not important. We showed, for example, in reference 21 that when two patches interact with migration the patches very quickly lose the individual characteristics that were caused by their local ecology. Thus, the results of the sensitivity analysis merely indicate that global changes in the landscape ecology are more important than the connectivity of the individual patches; however, in a given ecology, the patch connectivity is indeed extremely important.

To continue our exploration of the importance of patch connectivity, we now move to our consideration of the interaction of a wooded patch and a grassy patch with variable birth and death rates oscillating near the baseline values given in Table 1. The oscillating values are explained in reference 12 and are intended to mimic tick life history as well as seasonal climatic effects. We vary the patch connectivity between the grassy patch and the wooded patch. Figure 3 graphs the migration rate versus the long-term annual average proportion of infected of hosts and ticks in the combined patches. The graph indicates that as migration between patches increases, the individual characteristics at each patch become diminished. Note that Table 1 records our baseline migration level of \( m_{ij} = 0.01 \) and at this level metapopulation effects created by environmentally unique patches can be observed.

One unexpected result from this study is the importance of location for control efforts. One method frequently used to control HME in a given area is through the use of acaricides, chemical agents applied topically or systemically to hosts that kill ticks. In our model, the application of acaricide is equivalent to raising the value of the parameter \( b \), or equivalently increasing \( \hat{b}_i \). At the bottom of Figure 3, we consider two scenarios for a two-patch system. With a solid lines, we indicate the longtime average annual percent of infected ticks in the grassy (blue) and the wooded (red) patch if the acaricide control is used only in the grassy patch and with a dotted line we show the same outputs but with the assumption that the acaricide control is used only in the wooded patch. The figure shows
that without regard to migration rates, control applied in the wooded patch effectively reduces the prevalence in both patches to near zero. On the other hand, if the control is applied only in the grassy patch the prevalence remains high in both patches. It is perhaps counterintuitive that increases in migration rates contribute significantly to the reduction of HME prevalence in both patches; i.e., the patch in which the acaracide control is applied can benefit more from the acaracide if migration is allowed. Patch connectivity can play a significant role in the disease dynamics.

Figure 3. The effect of increased patch connectivity. In the top two graphs, the vertical axis is the average percent of infected hosts (left) and ticks (right) over the last eight years of a 30-year HME simulation. We observe that the individual effects resulting from a local ecology are lost with increased with large inter-patch migration. In the bottom graph, the vertical axis is the average percent of infected ticks over the last eight years of a 30-year HME simulation. We observe that when tick control is applied only in wooded areas, the control effort is effective regardless of migration levels. However, when the control is applied only in the grassy patch then the level of patch connectivity plays a large role in the effectiveness of the control.
The Surrounding Environment

In our remaining investigations, we consider more complex metapopulation ecologies. Future simulations will rely on a larger $5 \times 5$ spatial grid of patches that are connected to the nearest neighbors. Although the patches are established in a rectangular fashion, individuals move away from the center of a patch in radial directions and therefore any of the six patches that surround an interior patch would be considered equally connected to the given patch. In addition, we consider the boundaries as being reflecting boundaries, in which hosts and ticks located in edge patches are unable to cross the boundaries and thus remain in their patch rather than migrate in the boundary direction. We will discuss the importance of this boundary assumption in the following section.

As noted above, the initial work on this model discussed only wooded and grassy habitat types for patches. In reality, the grassy areas would not instantly transform to wooded areas, but rather there would be some intermediate area of land with intermediate parameter values. Additionally, transition areas may be of importance to other features such as the likelihood of host-tick encounters because ticks may be more likely to use the vegetation of intermediate height to attack hosts. We simply define transition patches as having parameter values halfway between grassy and wooded patches for the host carrying capacity, $K$, and the tick death rate, $b$. In multiple simulations, we seek to check the importance of the inclusion of such areas in the use of the model to predict the spread of HME in various settings.

Figure 4 illustrates this point with one set of simulations. In the figure, we show side-by-side the number of infected ticks that populate a $5 \times 5$ spatial grid in three scenarios that are described visually in the figure. We observe there that there is predictable sensitivity to replacing grassy areas with transition zones and perhaps greater sensitivity to replacement of the wooded areas that are highly endemic. In fact, when the graphs of the percent of infected ticks in each grid are compared, there is virtually no difference between the first two scenarios and only slight variation to the third. This suggests that in applying the model to a specific scenario, transition areas could be included as grassy areas with little loss of accuracy in predictions. Inclusion of other aspects of transition zones such as increased host-finding rates will be explored in future work.

Boundary Effects

Edge effects and habitat fragmentation have long been studied to understand the impact on various processes. We use a variety of scenarios for our metapopulation model to assess these effects on the predicted prevalence of the infected ticks. Recall that our initial simulations assume that our boundary is a reflecting boundary, which is a good assumption for an environment that is adjacent to a river or urban area. This assumption contributes to the high numbers of infected ticks in the corners of the graphs in Figure 4.

Continuing with the assumption of a reflecting boundary, we now consider several scenarios in which a small area of endemicity is placed in a nonendemic environment. In Figure 5, we show where endemic (yellow E) patches are located and include a graph of the percent of infected ticks in all patches each month over a time period of 30 years.

The first observation we see is that an isolated endemic patch cannot remain endemic when surrounded by nonendemic patches with sufficient migration ($m_e = 0.01$) between the patches; however, a single endemic patch in the corner of our grid is sufficient to create an endemic environment for multiple surrounding patches. On the other hand, even if we double the size of the endemic area and place it near, but not on, our spatial boundary, we do not maintain an endemic HME outbreak, though we see that the outbreak takes longer to diminish. Thus, patches along a reflecting boundary will experience different dynamics than patches that are in the interior of our grid.

There are two other commonly used boundary conditions in the ecology literature. One is a toroidal, or cyclic, boundary condition in which the vertical and horizontal edges of the grid are considered to wrap around to the opposite boundary. The other is an absorbing boundary in which individuals who would migrate across the edge boundary are lost. Generally for these latter two boundary choices edge effects will be lost, though some symmetries would appear for toroidal
Figure 4. Exploring the effect of transition patches. The top illustrations show the patch configurations that are assumed in each simulation and the figures below show the number of infected ticks within each of the 25 patches after 1000 months with parameter sets as given in Table 1. We conclude that the inclusion of transition zones is less important than correctly categorizing wooded patches.
Figure 5. Reflecting boundary effects and endemicity. The simulations are run with reflecting boundaries, periodic host and death birth rates and other parameters as in Table 1. A single endemic patch (denoted by E and using the wooded patch parameters) or pair of endemic patches is surrounded by patches for which the disease is not endemic (N and using the grass patch parameters with $b$ increased to 0.25). The percent of infected ticks in each month and in each patch is plotted. In the figure to the left, we see that the endemicity of a patch that is surrounded by nonendemic patches is quickly diffused and to the right we see that in time a pair of patches near but not on the boundary will also have endemicity diffused by migration. However, the center figure illustrates the effect of a reflecting boundary as would be seen with plots of land backing to a river or to urban development. In this case, the endemicity of the patch spreads to neighboring patches.
boundaries as an expected result of the structure. As an illustration of lost boundary effects, we modify our model from our standard reflecting boundaries to a toroidal boundary condition. In Figure 6, we see that an endemic patch on the boundary of our grid will no longer remain endemic when surrounded by nonendemic patches. This result shows the importance of identifying boundary conditions. Many geographic areas may be modeled as having reflecting boundaries if there are conditions such as a river or man-made buildings and structures that would preclude the ticks and hosts from moving in that direction. On the other hand, if the studied land area is surrounded by similar ecologies then toroidal conditions would most appropriately be chosen. These model results imply that the impact of these boundaries can be significant on the predicted risk of disease in a given area.

**Conclusion**

Tick-borne diseases are of increasing concern for human health worldwide as the incidence of known diseases increases and new diseases are identified. The outbreaks are notoriously sporadic in space and time making them difficult to predict or control. In order to provide some insights into the complex dynamics of tick-borne diseases, we present the results from a spatially explicit, mathematical model of human monocytic ehrlichiosis.

Using a metapopulation ecology framework, the model predictions show that spatial arrangement/location plays a significant role. Within each individual patch, the values chosen for $\hat{\beta}_i$, the growth rate of ticks and $\hat{\lambda}$, the transmission rate from hosts to ticks, were shown to have the most influence on the resulting outbreak. This indicates that favorable weather conditions that increase the tick growth rate will significantly increase the likelihood of an outbreak. Similarly, the species diversity of a given area could increase the transmission rate from hosts to ticks because many species are not susceptible to *E. chaffeensis*. This would in turn increase the likelihood of an outbreak.

Across a spatial framework, the model predictions showed that the location of a potentially endemic region determines if an outbreak will emanate out from that location or be washed out by surrounding nonendemic areas. Outbreaks are predicted to occur more commonly along areas where there is a reflecting boundary next to an endemic area. As human development continues to encroach into previously undisturbed areas, new regions that effectively have reflecting boundaries therefore create the potential for higher risk of outbreaks at the edges of development and wilderness. While transition areas were shown to be effectively modeled as grassy areas, the importance of transition into human developed areas has yet to be explored.
Similarly, the model showed that the location of the application of control measures could significantly impact the integrity of control efforts. With a two-patch system, one wooded and one grassy, the disease can be eliminated if the acaricide is applied in the wooded area but not if applied to the grassy area. This result suggests the need for future work that expands this model and seeks optimal solutions for the application of acaricide for a given spatial framework. We will use the mathematical technique of optimal control to accomplish this goal in a future work.

The predicted model results are limited by the nature of mathematical models. A mathematical model is an abstraction of reality and as such, it cannot capture all of the dynamics of any system. The model is limited by the knowledge and understandings of the biological system and thus will only be useful if we recognize its limitations and use those limitations as a benefit to point out key data gaps.

A second limitation of these results is due in turn to the limited amount of research in HME. While enough information is known about HME to construct a rudimentary model and gain many insights, the majority of tick-borne disease research has been focused on Lyme disease which has higher incidence levels. As stated previously, the advantage for working with HME is the simplifying one-host structure, but given the aggressive and seemingly indiscriminate feeding habits of *A. americanum*, additional host reservoirs may need to be added to the model system to fully capture the dynamics and risks of HME.

Our metapopulation model provides a solid basis for exploring the initial dynamics of HME across a spatial landscape. The results provide some interesting clues into the sporadic nature of outbreaks. Thus, in addition to future work in the optimal control of a potential HME outbreak, future work will also continue to expand on this model to include more information about the life history of the lone star tick as well as the impact of additional host species. As the incidence of tick-borne diseases continues to increase, insights from models such as the one presented here will provide invaluable tools for prevention of future human morbidity and mortality.

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**References**

Chapter 5

Modelling Stochastic Transmission Processes in Helminth Infections

Stephen J. Cornell*

Abstract

The number of helminths within a host can only increase by the host encountering additional infectious stages, so it is important to consider not only whether a host is infected, but also the severity of its infection. Stochastic models consider explicitly the number of parasites within the host and treat infection, death and other demographic events as random processes. I discuss stochastic helminth population models of increasing degrees of complexity, starting with the infection dynamics within a single host and finishing with the full parasite lifecycle among a population of hosts. I demonstrate the mathematical techniques that can help to analyse these models and discuss the insights into parasite population biology that these methods can bring.

Introduction

The number of helminth parasites infecting a particular host matters to both the host and the parasite. Hosts with a high worm burden suffer much stronger pathology than those with a milder infection and in many cases mount an immune response that strengthens with exposure to infectious stages of the parasite. In the terminology of Anderson and May,1 macroparasites (and helminths in particular) cannot reproduce directly within the host, so macroparasite infections develop more gradually than microparasites do because the worm burden can only increase if the host encounters additional infectious stages in its environment. Many of the processes affecting parasite demographics are density dependent—i.e., depend nonlinearly on the number of worms in the host—such as the risk that a female cannot find a mate,2 or the modulation of infection rate by the host’s immunity.3 This means that population dynamics of helminth parasites depend on the distribution of worms among hosts and cannot be adequately modelled by considering the mean worm burden alone.

A key feature of many helminth infections is that the distribution of worm burdens is aggregated, i.e., the variance in worm burden among hosts is greater than the mean.4 Much effort has been spent on building mathematical models to understand the causes and consequences of these patterns of infection. Aggregated distributions can be generated by heterogeneities in hosts’ encounters with infectious stages, or heterogeneities in the dynamics of parasites within hosts. Some of this heterogeneity is systematic—older hosts may be better at avoiding or resisting parasites—but much heterogeneity has no obvious correlation with the host’s age or other attributes.

In this chapter I shall describe stochastic models for helminth infections, where heterogeneity in host exposure and response are modelled by random processes. The most common form of randomness in these models is demographic stochasticity, so that parasite subpopulations in two different hosts may be different even when they are subject to the same underlying rates of

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infection, reproduction and death. In addition, some models have allowed the parameters determining parasite population dynamics to differ between hosts, again as a random process. Random fluctuations in environmental conditions can also play an important role in parasite dynamics,\textsuperscript{5,6} though I shall not discuss this here. I shall describe models with increasing degrees of complexity, from the infection dynamics of a single host to parasite proliferation within a host population. I shall demonstrate mathematical techniques which can be used to approximate and understand these stochastic models, using as an example a parasite which causes increased mortality in its host. This model is an exemplar of the host-parasite interaction and has been studied in different forms by several authors.\textsuperscript{7-12}

**Infection in a Single Host**

The earliest stochastic models of macroparasite transmission described the dynamics within a single host only.\textsuperscript{13} The infection rate of the host is treated as a known quantity, which is assumed to be unaffected by the reproductive output of the worms in the host under consideration. This simplifies the analysis to the extent that explicit mathematical solutions can be obtained for models with many biological components, such as acquired immunity and parasite-induced mortality.

These models give valuable insights into how different mechanisms increase or decrease parasite aggregation and how this aggregation affects nonlinear quantities such as mating probabilities.\textsuperscript{2} The main appeal of this technique is the possibility for detailed mathematical analysis, though as models become more biologically complicated it is sometimes necessary to use approximations. I shall illustrate the method by discussing a model that was studied by Valerie Isham.\textsuperscript{10}

**Clumped Infection and Parasite-Induced Mortality**

Isham\textsuperscript{10} studied a model for helminth infections, which was not aimed at describing any particular parasite but had two notable features:

- Hosts do not encounter infective parasite stages individually, but rather in ‘clumps’. This is a reasonable assumption for many helminth infections: a biting insect may transmit many infective larvae to the unfortunate host and eggs in faeces will develop into larvae that are highly clumped in space.
- The morbidity effects of the parasite causes a higher rate of mortality in hosts with high worm burdens.

Clumped infection was already a component of the earliest stochastic parasite models\textsuperscript{13} and is expected to lead to more aggregated parasite distributions, while density-dependent host death rate is expected to decrease aggregation.\textsuperscript{9}

In this section, I shall reproduce Isham’s analysis of the model, in order to illustrate the technique and to discuss its strengths and limitations. In the model, a host of age \( a \) is assumed to harbour \( m \) parasites. The processes by which \( m \) changes are

- Infection: \( m \to m + c \) at rate \( \phi(a)h_c \). This represents an infection event taking place at rate \( \phi(a) \) (which may depend on host age \( a \)) and when infected the host gains a clump containing \( c \) parasites. The distribution of clump sizes is, where \( \sum_{c=0}^{\infty} h_c = 1 \).
- Parasite death: \( m \to m - 1 \) at rate \( \mu_M m \), representing independent parasite death events at rate \( \mu_M \).
- Host death: The host dies at rate \( \mu_H(a) + \alpha m \). This represents the host’s intrinsic death rate \( \mu_H \) which may depend on its age \( a \) and an increased death rate due to the number of parasites in the host.

Under these assumptions, the following Forward equation for the joint probability \( P_m(a) \) that a host reaches age \( a \) and contains \( m \) parasites can be written:

\[
\frac{dp_m(a)}{dt} = -\{\mu_M(a) + \alpha m + (1 - h_c)\phi(a) + \mu_M m\} p_m(a) \\
+ \mu_M (m + 1) p_{m+1}(a) + \phi(a) \sum_{c=0}^{\infty} p_{m-c}(a)h_c.
\]  

(1)
The right-hand-side term on the first line of this equation represents all the processes which change the worm burden out of state \( m \); the terms on the second line represent processes where the worm burden changes into state \( m \).

This equation can be analysed with the use of the probability generating function (PGF) \( R(a;z) = \sum_{m=0}^{\infty} z^m p_m(a) \). Multiplying Eqn. (1) by \( z^m \) and summing, we note that we can simplify terms like \( \sum_{m=0}^{\infty} m z^m p_m(a) = z \frac{dR(a;z)}{dz} \) to give

\[
\frac{dR(a;z)}{dz} = - \left\{ \left[ \mu_H(a) + \phi(a) [ h(z) - 1 ] \right] R(a;z) \right. \\
\left. - (\alpha + \mu_M) z - \mu_M \right\} \frac{dR(a;z)}{dz}. 
\] (2)

This equation could be solved directly, but to study age-intensity profiles we are instead interested in the probability \( p(m|a) \) of having \( m \) parasites conditional on reaching age \( a \). If the probability of surviving to age \( a \) is \( S(a) = \sum_{m=0}^{\infty} p_m(a) = R(a;1) \), then the standard relationship between joint and conditional probabilities gives \( p_m(a) = p(m|a) S(a) \). An equation for the survival probability is obtained by substituting \( z = 1 \) into Eqn. (2):

\[
\frac{dS(a)}{da} = - \mu_H(a) S(a) - \alpha m_M(a) S(a), 
\] (3)

where we have written

\[
\frac{dR(a;z)}{dz} \bigg|_{z=1} = \sum_{m=0}^{\infty} m p_m(a) \\
= S(a) \sum_{m=0}^{\infty} m p(m|a) \\
= S(a) m_M(a),
\]

\( m_M \) being the mean of \( m \) conditional on the host reaching age \( a \).

Let \( Q(a;z) = \sum_{m=0}^{\infty} z^m p(m|a) = R(a;z) / S(a) \) be the PGF for \( p(m|a) \). Using Eqns. (2) and (3), we can derive the following equation for \( Q(a;z) \):

\[
\frac{dQ(a;z)}{da} = \{ \phi(a) [ h(z) - 1 ] + \alpha m_M(a) \} Q(a;z) \\
- \left\{ (\alpha + \mu_M) z - \mu_M \right\} \frac{dQ(a;z)}{dz}. 
\] (4)

**Explicit Solution of the Model**

Equation (4) can be solved explicitly by first defining \( \zeta(z) = \frac{\log(\phi(a) h(z) - 1)}{\alpha + \mu_M} \), and then making the change of variables \( u = \frac{a + \zeta}{z}, \quad v = \frac{a - \zeta}{z}, \) \( W(u, v) = \log Q(a; z) \). Eqn. (4) can then be written in the form

\[
\frac{\partial W(u, v)}{\partial u} = f(u + v, u - v) + g(u + v),
\]

where \( f(\zeta, \zeta(z)) = \phi(a) [ h(z) - 1], \ g(a) = \alpha m_M(a) \). This equation can be integrated directly to give

\[
W(u, v) = \int_{\zeta}^{u} [ f(w + v, w - v) + g(w + v) ] dw + F(v), 
\] (5)

where \( u_0 \) is an arbitrary constant and \( F(v) \) is an arbitrary function of its argument.

The desired solution is obtained by imposing the following two boundary conditions: (i) \( Q(0, z) = p(m = 0|a = 0) = 1 \), which states that the host is uninfected at birth, constrains the form for \( F \); and (ii) \( Q(a, 1) = m_M(a) = \sum_{m=0}^{\infty} p(m|a) = 1 \), which states that the probabilities of different worm
burdens sum to unity. While there is only one arbitrary function in (5), the second boundary condition is required because $g = c w m_{z}(a)$ needs to satisfy the self-consistency condition with $m_{z}(a) = \partial Q(a; z)/\partial z$. The final solution is

$$Q(a; z) = \exp \left[ \int_{\xi(z)-a}^{\xi(z)} f(y + a - \xi(z), y) dy - \int_{\xi(z)+a}^{\xi(z)} f(y + a - \xi(z), y) dy \right],$$

where $f$ is a general function and for the specific form $f(a, \xi(z)) = \phi(a)[b(z) - 1]$ we get

$$Q(a; z) = \exp \left[ \int_{0}^{\theta(a; z)} \frac{\phi(a + \xi(x) - \xi(z))[1 - b(x)]}{(\alpha + \mu_{M})x - \mu_{M}} dx + \int_{\theta(a; z)}^{1} \frac{\phi(a + \xi(x) - \xi(z))[1 - b(x)]}{(\alpha + \mu_{M})x - \mu_{M}} dx \right],$$

where

$$\theta(a; z) = \frac{\mu_{M} + [(\alpha + \mu_{M})z - \mu_{M}]e^{-(\alpha + \mu_{M})a}}{\alpha + \mu_{M}}.$$  

Note that the equivalent solution in ref. 10 was derived by assuming that $\phi(a)$ is independent of $a$, though as shown here a solution can be obtained without this assumption.

**When Can This Type of Model Be Solved Explicitly?**

We are now in a position to discuss the type of models which can be solved by this method.

- Eqn. (2) is a first-order partial differential equation because both host and parasite death are linear functions of $m$. If some processes took place at a rate proportional to $m^{2}$, then there would be terms of the type $\alpha^{2} P/\partial z^{2}$ and there is a much smaller chance that an explicit solution to the PDE would be possible. The problem would become completely intractable if rates had a still more complex dependence (e.g., exponential) on $m$. This severely limits the degree of biological realism of models for which explicit solutions are available.

- The component of the host’s death rate that is independent of $m$ does not bias survivorship as a function of $m$ and as a result the expression (6) does not depend on $\mu_{t}(a)$. This means that the results can be applied to organisms with any type of age-death relationship.

- Even though an explicit solution is possible for this model, the compactness of the solution belies its usefulness. In principle, the value of $p(m|a)$ can be obtained by expanding Eqn. (6) as a series in powers of $z$, though as noted by Isham this does not give very convenient expressions. Moreover, the expression is still more cumbersome when the infection rate $\phi(a)$ depends on age $a$. Nevertheless, some low moments of the distribution can be obtained in a useful form, as discussed in the next section.

**Effect of Clumping and Parasite-Induced Mortality on Aggregation**

The mean $m_{z}(a) = \Sigma_{m=0}^{\infty} pm(m|a)$ and variance $V_{z}(a) = \Sigma_{m=0}^{\infty} m^{2} p(m|a) - m_{z}^{2}(a)$ of the worm burden, conditional on the host reaching age $a$, can be straightforwardly obtained from derivatives of $Q(a; z)$ evaluated at $z = 1$. For the case $\phi = \text{constant}$, using Eqn. (6) we have

$$m_{z}(a) = \frac{\phi}{\alpha} \left[ 1 - h(\theta(a;1)) \right]$$

$$V_{z}(a) = \frac{\phi}{\alpha} b'(1) \left[ 1 - \frac{\mu_{M}}{\alpha} m_{z}(a) - \frac{\phi}{\alpha} b'(\theta(a;1)) e^{-(\alpha + \mu_{M})a} \right].$$

The mean $m_{z}(a)$ increases with age $a$ but approaches a finite limit as $a$ increases. The variance to mean ratio $I_{z}(a) = V_{z}(a)/m_{z}(a)$ is a measure of the degree of aggregation in parasite burdens, being unity for a Poisson distribution. As discussed in ref. 10, $I_{z}$ is unity when infecting clumps contain only one parasite, but increases when infecting clumps are bigger. Parasite-induced mortality has a
more complicated effect on $I_M$, however. The limiting value of $I_M(a)$ for large $a$ can either decrease or increase with $\alpha/\beta_M$ depending on the details of the distribution of clump sizes.

The model was extended by Herbert and Isham to allow for more than one life history stage of the parasite and also for non-Markovian processes in life history progression (i.e., the time spent in different stages was not assumed to be exponentially distributed). The influence of mortality on parasite aggregation was further explored by Barbour and Pugliese, who assumed nonclumped infection but considered both density-dependent parasite mortality and a range of scenarios for density-dependent parasite-induced host mortality.

**Moment Closure**

As discussed above in 'Explicit solution of the model', for many models it is not possible to obtain an exact solution for the distribution of worm burdens. However, it is possible to obtain an approximate solution by using a technique called 'moment closure'. While this is not necessary for the model described above, it is instructive to see how the approximation works in this case.

The mean $m_M$ and the variance $V_M$ of the worm burden can be expressed as derivatives of $Q(a;z)$ at $z = 1$, so from Eqn. (4) these quantities satisfy

$$\frac{dm_M}{da} = \phi(a)b'(1) - \alpha V_M(a) - \mu_Mm_M(a)$$

$$\frac{dV_M}{da} = \phi(a)\left[b''(1) + b'(1)\right] + \mu_Mm_M(a) - 2\mu_MV_M(a)$$

$$+ \alpha \left\{3m_M(a)V_M(a) + m_M^3(a) - T_M(a)\right\},$$

where $T_M(a) = \Sigma m^3 \phi(m|a)$ is the expectation of the cube of the worm burden. We find in general that the rate of change of the $n$th moment of the distribution depends on moments of order higher than $n$. This hierarchy of equations cannot be solved, because the solution to any equation depends on the solution of an equation that has not yet been solved.

The technique of moment closure gets around this problem by approximating a higher-order moment in terms of a combination of lower-order ones. For instance, Isham used the normal approximation for the third moment, in which case the term multiplying $\alpha$ on the right-hand side of Eqn. (7) (the third order cumulant) is zero. The above equations are now 'closed' as they depend only on $m_M$ and $V_M$ and can be solved. Reference 10 gives an explicit solution as a function of $a$ and compares the results of the moment closure to the exact solution.

Note that the moment equations form an unclosed hierarchy, even in a case such as this where an explicit exact solution for the moments themselves is available. However, the moment closure approximation can give approximate analytical or numerical solutions for a much wider class of problems. In particular, it is not restricted to processes which depend linearly on the internal variables as discussed in 'When can this type of model be solved explicitly?'

**Acquired Immunity**

Immunity to many helminth infections is thought to be acquired progressively in response to exposure to infection challenge. Deterministic age-structured models have been used to investigate the importance of the host’s immune response in shaping patterns of helminth infection as a function of age. Immunity has also been investigated in single-host stochastic models.

Chan and Isham studied a model similar to the one described above, with the additional assumption that the host has a stochastic integer-valued ‘immunity’ variable $I$ which increases at a rate proportional to the current infection level. In order to aid tractability, they assumed that the infection rate was modulated by a linearly decreasing function of $l$. This leads to a first order PDE for the PGF of the number of parasites and immunity variable, for the reasons discussed in 'Explicit solution of the model'. This form for the infection rate is actually negative when $l$ is high enough, but that was not found to lead to mathematical problems for the parameters studied by the authors. It was not possible to obtain explicit solutions for the PGF, because of the particular nonlinear coefficients of the first-order partial derivatives, but the equations for the first and second
moments of the distribution formed a closed hierarchy so could be solved without requiring a closure assumption.

**Host Heterogeneity**

A further cause of parasite aggregation is intrinsic differences between hosts. Hosts may differ in the rate at which they encounter parasites or in their ability to mount an immune response. The host represents the parasite’s habitat, so any differences in hosts, e.g., size and/or age, may affect their development, reproduction and mortality.

If mathematical solutions for the distribution of worm burdens is possible for a host with given parameters, then results for a heterogeneous host population can be obtained by averaging these results (either analytically or numerically) over the host parameters. Isham noted that analytical results obtained in this may be unwieldy and not helpful intuitively. It is also possible to incorporate host heterogeneity at the level of moment equations. Chan and Isham followed this approach in a model of schistosomiasis and discovered that, while the equations for the first and second moments formed a closed system for a single host, the moment hierarchy was not closed when host heterogeneity was taken into account. Two types of host heterogeneity were considered—either differences in parasite encounter rates, or differences in immune response. The former was found to have a strong effect on parasite aggregation, but the latter had only a very weak effect. Further work using the same model showed that stochastic variability may obscure the expected age-intensity relationship generated by acquired immunity.

**Mating Probability and Population Genetics**

One of the advantages of exact solutions of stochastic single-host infection-death models is that the behaviour of nonlinear functions of the worm burden may be investigated explicitly. Cornell et al. investigated a model of early-season dynamics of nematodiasis in sheep. The study focused on the reproductive output of female parasites and did not consider nonlinear processes which might affect the dynamics at high parasite density. Helminths are generally assumed to be promiscuous, but at low density there is a risk that a female may find no male to mate with in the same host. The resulting depression in average fecundity is known as the mating probability. Mating probabilities can be estimated by assuming a given distribution of worms among hosts, such as Poisson or negative binomial and when calculated in this way they decrease to zero at low density. However, Cornell et al. found that clumped infection led to the mating probability being greater than zero in this limit. This shows the importance of considering the mechanisms that generate aggregation when modelling parasite population dynamics.

Cornell et al. also considered the population genetics of the parasites’ offspring and found that if infecting clumps have a multinomial distribution of genotypes that follow Hardy-Weinberg equilibrium then the offspring show small deviations from Hardy-Weinberg equilibrium. However, if the infective clumps were more heterogeneous, then the offspring distributions showed strong deviations from Hardy-Weinberg equilibrium and in particular there was a strong promotion of rare homozygotes.

**Competition between Parasite Species**

Many hosts will harbour more than a single parasite species and each species may strongly affect the dynamics of other parasite species within the host. Bottomley et al. have extended the single-host framework to allow for more than one parasite species. They assumed that the density of adult worms affected the mortality of larval stages of either species and solved a stochastic model approximately by moment closure. When interactions between individuals of the same species were antagonistic, worm burdens were found to be underdispersed when interactions between species were weaker than interactions within species and overdispersed otherwise. However, when interactions within species were mutualistic, the variance to mean ratio was found to be close to unity.
Infection among Multiple Hosts

The models considered above in ‘Infection in a single host’ assume that the infection rate is independent of the parasite dynamics within the host. In the absence of any feedback from the parasite’s reproduction to the abundance of infective stages of the parasite, this method is not able to investigate the evolution of the force of infection over time, or the potential of control strategies to eradicate the parasite. In order to model the ecology of the parasite, it is necessary to consider the dynamics of the parasite stages that live outside the definitive host. This makes the model considerably more complicated, because there may be many hosts of different ages and infection status. The sub-populations of parasites in each of the hosts become coupled and the host themselves may also undergo complicated population dynamics.

Nevertheless, in many cases there are suitable approximations which simplify the models and make it possible to analyse their behaviour. In the present section I shall describe these techniques and again illustrate them with a specific example.

Hybrid Models

The problem of multiple hosts is simplified if one assumes that the host population is large and that the infectious stages generated by the parasite in one host subpopulation is equally likely to infect any of the other hosts (whether they are transmitted directly or via a vector organism). Each host now only sees an average force of infection (though the infection process could still be in the form of clumps), so a host’s infection rate is statistically independent of its worm burden. This means that the single host models described in ‘Infection in a single host’ can be used to describe the dynamics within each host, while the abundance of the infectious stage is a deterministic quantity related to the mean reproductive output of the parasites infecting a host. This type of hybrid model is discussed in detail by Nåsell.23 Hybrid models are often cited as a motivation for studying single-host stochastic models, though in fact few such studies do actually relate the infection process to the mean reproductive output of the parasite.

Hybrid models assume that the dynamics of the host and/or the parasite’s infectious stages are deterministic, but allow full stochastic dynamics within the host. A moment closure assumption is often used to obtain an approximate solution of the within-host dynamics. I shall illustrate this method using a host-parasite model that was studied by Anderson and May7 and later by Kretzschmar and Adler.24 This model is very similar to the single-host model studied by Isham10 and discussed above in ‘Clumped infection and parasite-induced mortality’, though without clumped infection and where the infection rate is related to the parasite population size. Using the parameter definitions in Table 1, the model takes the following form:

- Assume there are \( n_m(t) \) hosts with \( m \) parasites at time \( t \), so the number of hosts in the system is \( H(t) = \sum_{n=0}^{\infty} n_n(t) \) and the total number of parasites is \( P(t) = \sum_{n=0}^{\infty} mn_n(t) \).
- Each host is infected at a rate \( \phi(t) \). Under the assumption that infective stages develop rapidly outside the host and are much shorter lived that the adult parasites in the hosts, the force of infection is related directly to the current parasite population in the form \( \phi(t) = \frac{\lambda}{H(t) + H(t)/\gamma} \) (see ref. 24).
- Parasites die at the per capita rate \( \mu_M \).
- Hosts are born at per capita rate \( \beta \) and a host with \( m \) parasites dies at a per capita rate \( \mu_H + \mu_m \). When a host dies, its parasites die too.

Given these processes, the equations for the rate of change of \( n_m \) are:

\[
\frac{dn_0}{dt} = -(\mu_H + \phi)n_0 + \mu_M n_1 + \beta H(t) \tag{8}
\]

\[
\frac{dn_m}{dt} = -\left[ (\mu_H + \phi + m(\alpha + \mu_M))n_m + \mu_M (m + 1)n_{m+1} + \phi n_{m-1} \right]. \tag{9}
\]

Note the similarity between these equations and Eqn. (1) for the case \( b(c) = \delta_{c,1} \).
Modelling Stochastic Transmission Processes in Helminth Infections

Deterministic Models with Parasite Aggregation

Some of the earliest models of helminth infection dynamics were deterministic, but nevertheless allowed for the fact that the distribution of parasites between hosts was aggregated. Anderson and May used this approach (see also ref. 8) to discuss processes that stabilise or destabilise host-helminth interactions. An equation for \( \frac{dH}{dt} \) is obtained by summing Eqns. (8,9) over \( m \):

\[
\frac{dH}{dt} = (\beta - \mu_H)H - \alpha P,
\]

and an equation for \( P \) is obtained by multiplying Eqns. (8,9) by \( m \) and summing:

\[
\frac{dP}{dt} = \frac{\lambda PH}{H_0 + H} - (\mu_H + \mu_p)P - \alpha \sum_{m=0}^{\infty} m^2 n_m(t).
\]

The term \( \sum_{m=0}^{\infty} m^2 n_m(t) \) represents the sum of square numbers of parasites in hosts, which depends on the details of the distribution of parasites among hosts. Anderson and May related this to the mean number of parasites per host by assuming a negative binomial distribution with mean \( \frac{P}{P/k} \) and aggregation parameter \( k \). The generating function for this distribution is \( \sum_{m=0}^{\infty} z^m n_m = H(t)[1+z(x/k)(1-x)]^{-k} \) and the sum of square numbers of parasites is \( \sum_{m=0}^{\infty} m^2 n_m = P + \frac{P^2(x/k)}{k} \). Since the second moment is approximated in terms of the first by assuming a particular distribution, this is actually a form of moment closure.

Under this approximation, the system is reduced to two ordinary differential equations which can be analysed with standard methods. As shown in reference 7, a positive value of \( k \) leads to a stable equilibrium between host and parasite, while a negative value of \( k \) (corresponding to a conventional binomial distribution, which is underdispersed) destabilises the equilibrium. They concluded that parasite aggregation can stabilise host-parasite interactions.

While this approach is not able to quantify the degree of aggregation which is expected to arise from a given stochastic model, it makes it possible to explore the consequences of aggregation on

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Table 1. Definition of model parameters used throughout this chapter

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>( m )</td>
<td>Number of parasites in host</td>
</tr>
<tr>
<td>( a )</td>
<td>Host age</td>
</tr>
<tr>
<td>( t )</td>
<td>Time</td>
</tr>
<tr>
<td>( c )</td>
<td>Number of infecting parasites in a clump</td>
</tr>
<tr>
<td>( b )</td>
<td>Probability of ( c ) parasites in a clump</td>
</tr>
<tr>
<td>( p_m(a) )</td>
<td>Probability that a host reaches age ( a ) and has ( m ) parasites</td>
</tr>
<tr>
<td>( p(m</td>
<td>a) )</td>
</tr>
<tr>
<td>( R(a;z) )</td>
<td>Probability generating function of ( p_m(a) )</td>
</tr>
<tr>
<td>( Q(a;z) )</td>
<td>Probability generating function of ( p(m</td>
</tr>
<tr>
<td>( P )</td>
<td>Mean number of parasites per host</td>
</tr>
<tr>
<td>( H )</td>
<td>Mean number of hosts</td>
</tr>
<tr>
<td>( \beta )</td>
<td>Host birth rate</td>
</tr>
<tr>
<td>( \mu_i )</td>
<td>Host death rate in absence of parasites</td>
</tr>
<tr>
<td>( \alpha )</td>
<td>Additional host death rate per parasite</td>
</tr>
<tr>
<td>( \mu_{st} )</td>
<td>Parasite death rate</td>
</tr>
<tr>
<td>( \phi )</td>
<td>Host infection rate</td>
</tr>
<tr>
<td>( \lambda )</td>
<td>Asymptotic infection rate per parasite</td>
</tr>
<tr>
<td>( H_0 )</td>
<td>Host infection efficiency</td>
</tr>
<tr>
<td>( n_m(t) )</td>
<td>Number of hosts with ( m ) parasites</td>
</tr>
</tbody>
</table>
nonlinear processes such as mating probabilities\textsuperscript{2} and the evolution of drug resistance.\textsuperscript{25} It has recently been used to discuss in some detail the importance of different density dependent processes on transmission,\textsuperscript{26} control\textsuperscript{27} and drug resistance\textsuperscript{28} in helminth infections.

**Moment Closure for the Variance**

Rather than assuming a fixed degree of aggregation of parasites among hosts, Kretzschmar and Adler\textsuperscript{24} instead assumed that the distribution of parasites will itself evolve due to the stochastic dynamics. They first wrote an equation for $Q = \sum_{m=0}^{\infty} m^2 n_m$ by multiplying (8,9) by $m^2$ and summing:

$$\frac{dQ}{dt} = -(2\mu_M + \mu_H)Q + (2\phi + \mu_M)P + \phi H - \alpha \sum_{m=0}^{\infty} m^3 n_m.$$ 

To approximate the final term, they again assumed a negative binomial distribution of worms between hosts with mean $\frac{m}{H}$ and shape parameter $k$ so that $\sum_{m=0}^{\infty} m^3 n_m = \frac{(k + 1)(k + 2)p^2}{k^2} + 3Q - 2P$. However, rather than assume a fixed aggregation parameter $k$ they constrained it by stipulating that the first and second moments of $n_m$ be $P$ and $Q$. This means that $k = \frac{P}{H(Q-P)P}$, so finally we have

$$\sum_{m=0}^{\infty} m^3 n_m = \frac{P(Q - P)}{H} \left( 1 + \frac{H(Q - P)}{p^2} \right) + 3Q - 2P$$

and the equations for $H$, $P$ and $Q$ form a closed hierarchy (note that Kretzschmar and Adler\textsuperscript{24} used a slightly different formalism in terms of the variance to mean ratio, but their presentation is equivalent to that shown here).

While the formalism is more complicated than that of Anderson and May\textsuperscript{7} (three differential equations rather than two), it can still be analysed with standard methods. Kretzschmar and Adler\textsuperscript{24} also studied the case where parasites were distributed according to a Neyman Type-A distribution and found parameter values where assuming one class of distribution predicted a stable host-parasite equilibrium and the other predicted instability. More specifically, they found that aggregated distributions would only produce a stable equilibrium if the variance to mean relationship increased with increasing mean worm burden. Their study underlines the sensitivity of the predictions of moment closure models to the type of closure that is assumed.

Rosà and Pugliese\textsuperscript{11} extended Kretzschmar and Adler's model to incorporate clumped infection and free-living stages of the parasite, as well as host heterogeneity. They found the latter to be a much more strongly stabilising effect than clumped infection. Bottomley et al\textsuperscript{29} used a related hybrid approximation (though assuming normal rather than negative binomial parasite distributions) to study the coexistence of parasite species within a constant population of hosts, deriving equations related to the Lotka-Volterra equations.

**Fully Stochastic Models**

If the host population is not so large as to be essentially infinite, then the abundance of infective stages is no longer statistically independent of the worm burden of any particular host and the 'hybrid model' approximation cannot be applied. Moreover, stochastic dynamics of the host population might also need to be taken into account. When this is the case, analytical approximations are typically unavailable and results are usually obtained by individual-based computer simulations. These are relatively straightforward to program: one takes account of each host’s parasite burden, as well as other attributes such as host age and immune status as necessary. The free-living stages may be modelled as explicit populations, or as a 'force of infection' variable (which is appropriate when the numbers of infectious stages are extremely large). Parasite infection and death events can be simulated as stochastic processes, where the time to the next event is an exponentially distributed pseudorandom variable and the type of event is determined pseudorandomly by weighting each possibility by its rate.\textsuperscript{9} Alternatively, a discrete time step can be assumed and the number of
births, deaths, etc. per timestep can be generated as pseudorandom numbers with appropriate distributions (see e.g., ref. 30).

Explicit stochastic simulation models of this type are commonly used as tools to inform control strategies, such as the LYMFASIM program to model lymphatic filariasis31 and SCHISTOSIM for schistosomiasis.32 It is not possible to analyse such models in the same way as analytical models, so optimal control strategies can only be found by trial and error. However, this might not be a serious handicap given modern computing power. Simulation models are capable of handling intricate, biologically realistic processes where an analytical approach would be insurmountably difficult.

While inevitably more complicated, fully stochastic models can nevertheless sometimes be understood by comparing their behaviour to single-host or hybrid models. Since hybrid models assume that the host population size is infinite, the fully stochastic model will be similar to a hybrid model when the host population is large enough. It should be stressed that, in this context, ‘population size’ means the number of other host that can be infected by the offspring of the parasite population in one host; in this sense, an infinite population where each individual can only infect nearby neighbours has a very small effective population size. The hybrid approximation assumes that the abundance of infective stages is effectively a deterministic variable, but when the host population size (and therefore the adult population size) is finite the abundance of infective stages will fluctuate. These fluctuations will be larger larger when the host population is smaller. This idea that host population size ‘tunes’ the fluctuations in infection rate has been explored by Cornell et al.33,34 There follows a discussion of some effects of finite host population sizes on helminth population dynamics.

Mating Probability

As discussed in the section ‘Mating probability and population genetics’, worm fecundity is depressed at low density if the worms reproduce sexually. As a result, there is an Allee effect35—i.e., the parasite population will tend to die out if the density of infectious stages is below a certain threshold. In an infinite host population, where the dynamics of the force of infection are deterministic, the parasite will always go extinct if it starts below this threshold. However, the force of infection can fluctuate stochastically above this threshold when the host population is finite. This means that epidemics may be more likely in small populations than in large ones if the initial force of infection is low.33 If infection is clumped, however, the mating probability is increased and the Allee effect may be removed.34

Population Genetics and Antihelminthic Resistance

When the host population is small, the offspring of the parasites within one particular host are more likely to reinfect the same host than if host population is large. As a result, smaller host populations increase inbreeding in the parasite population, which increases homozygosity and therefore will promote antihelminthic resistance if this is a recessive trait.36 If infection is clumped and clumps represent collections of larvae whose parents all live in the same host, then these hosts will be more genetically homogeneous than the parasite population as a whole.37 As discussed in ‘Mating probability and population genetics’, this can also lead to inbreeding and thus significantly enhance the production of rare homozygous offspring.

Stability and Population Cycles

Rosà et al12 used individual based simulations to extend their study of host-parasite interactions (as discussed in ‘Moment closure for the variance’) beyond the hybrid approximation used in reference 11. They found that, when host populations were of the order of 100 or more, the mean population sizes were similar to the predictions of the hybrid model. They also found that the processes which stabilised population dynamics in the deterministic approximation would also tend to stabilise stochastic populations. However, they found that the populations would often exhibit sustained cycles even when the deterministic approximation predicted damped cycles.
Conclusion

In this chapter I have focused on the mathematical techniques which can be used to analyse stochastic models of helminth population dynamics. In fact, stochastic population modelling is increasingly performed by computer simulation, a trend which will no doubt continue. The strength of simulation models is that they can be made arbitrarily flexible and complicated. For example, spatial variation in the force of infection can straightforwardly be incorporated into a simulation model. Even when an analytical approximation would be appropriate (e.g., a hybrid approximation if there are many hosts), it may give few advantages over a simulation model if one only needs numerical results for specific parameter values and indeed the mathematical method may be slower at producing results. Stochastic simulations are very useful for making predictions provided that the parameters are known. However, it can be very tricky to fit these parameters to data and the noise in simulation output can give the illusion of biological realism. As with any model, it is important that simulation models are validated against the real world before trusting their predictions.

Nevertheless, I believe it would be a mistake to dismiss mathematical analysis as impractical or unnecessary. Mathematical analysis can elucidate robust principles and generalities which are opaque from black-box simulation models. An example of the sort of insight which can be obtained from analysis is the condition for aggregating processes to stabilise host-parasite interactions (see sections ‘Deterministic models with parasite aggregation’ and ‘Moment closure for the variance’).

Immunity and host heterogeneity are two phenomena which are important determinants of patterns of infection, yet which are difficult to measure directly. In many cases, acquired immunity is thought to increase with the host’s cumulative exposure to infection challenge. However, there are many qualitatively similar mathematical relationships between the determinants of immunity and its effects, while in a model one has to make a specific choice. Very few of these possibilities have been explored and—while they can give good qualitative and even quantitative agreement with data—the ‘correct’ or ‘best’ formalisms are unknown. There is further ambiguity when formulating stochastic models of immunity—for example, in ref. 18 immunity is assumed to be integer valued and the size of this ‘unit’ determines the strength of stochasticity in immunity. Intrinsic host heterogeneity and demographic stochasticity are additional confounding factors and bring additional parameters which need to be fitted to data. These uncertainties reduce the power of models to make quantitative predictions.

Analytical methods have another advantage over individual-based simulation models, beyond their ability to bring insight to a problem rather than just results. Helminth parasites can live in very large populations (tens of thousands of gastrointestinal nematodes can live in one sheep) and host populations can also be very large. A brute-force simulation, where every worm’s birth and death is accounted for, could be beyond currently available computer power. However, mathematically derived approximations can provide tools to make simulations feasible for very large systems. For example, a diffusion approximation for the number of parasites can allow many birth and death events at once but still model the demographic noise correctly. Similarly, mathematical expansions about the hybrid approximation can take account of a large but finite number of interacting hosts and even model spatially explicit populations of hosts. Tools of this sort are under active development and promise to help us apply stochastic models with greater biological realism to understand and manage parasite populations.

References
Modelling Environmentally-Mediated Infectious Diseases of Humans: Transmission Dynamics of Schistosomiasis in China

Justin Remais*

Abstract

Macroparasites of humans are sensitive to a variety of environmental variables, including temperature, rainfall and hydrology, yet current comprehension of these relationships is limited. Given the incomplete mechanistic understanding of environment-disease interactions, mathematical models that describe them have seldom included the effects of time-varying environmental processes on transmission dynamics and where they have been included, simple generic, periodic functions are usually used. Few examples exist where seasonal forcing functions describe the actual processes underlying the environmental drivers of disease dynamics. Transmission of human schistosomes, which involves multiple environmental stages, offers a model for applying our understanding of the environmental determinants of the viability, longevity, infectivity and mobility of these stages to controlling disease in diverse environments. Here, a mathematical model of schistosomiasis transmission is presented which incorporates the effects of environmental variables on transmission. Model dynamics are explored and several key extensions to the model are proposed.

Introduction

A common feature of many of the most debilitating macroparasites of humans is their dependence on environmental life-stages subject to dynamic climactic, ecological, hydrological and other conditions. This phase can be wholly environmental, where for example infected humans excrete parasite eggs in feces and others are exposed via contaminated food or, as in the case of hookworm, where contact with contaminated soil can result in penetration of the parasite through the intact skin. Alternatively, the environmental phase may consist of time spent in an intermediate host, such as a snail or fish, itself subject to heterogeneous environments. Transmission of human schistosomes involves environmental phases of both types and thus understanding the environmental determinants of the viability, longevity, infectivity and mobility of these phases is key to conceptualizing disease transmission and ultimately controlling disease in diverse environments.

Schistosomes enter the environment as eggs that hatch in water into a free-swimming miracidium that seeks a snail of the appropriate species to infect. Asexual reproduction in the snail produces cercariae, another free-swimming aquatic stage with a lifespan on the order of a day, which penetrate the intact skin of a definitive host and mature into adult worms. The worms sexually pair and the female lays copious numbers of eggs that are the source of
pathogenic response in the host. Some of these eggs find their way into the feces or urine, are excreted and the cycle begins again. The intermediate host, a freshwater snail and the two free-living aquatic stages are known to be subject to environmental stresses such as temperature and shear forces present in the water column. For *S. japonicum*, the species that causes schistosomiasis in east and southeast Asia, transmission is further complicated by the fact that a variety of mammals can serve as the definitive host, including rodents, dogs, cats, pigs and water buffalo, the latter of which is particularly important to sustaining transmission in the lower Yangtze environment.

In China, considerable progress has been made since the 1950s controlling transmission of *S. japonicum* in humans and domestic animals. From a total of 433 endemic counties in 1959, the disease has been eliminated from 260 counties leaving approximately 800,000 infected people and another 60 million at risk. However, these represent only a small fraction of the worldwide total of schistosomiasis cases that the World Health Organization estimates at 200 million, 85% of which are in Africa. Most of these infections are suffered by poor people, particularly children and most are preventable and treatable, although effective vaccines remain a hope for the future.

Where schistosomiasis transmission has been eliminated, targeted environmental modifications have often played an important role. Conversely, major environmental changes such as water development projects have often led to a sustained elevating effect on schistosomiasis prevalence. The underlying mechanisms shaping this relationship are poorly understood. An expansion in the preferred habitat of intermediate host snails is often implicated in these prevalence increases, yet few data exist to fortify this claim. In China, recent evidence points to the influence of changing water levels on intermediate host populations. Yet a clear mechanistic understanding of the processes that lead to increased disease is lacking and therefore opportunities to mitigate the disease impact of water projects using engineering or design principles is limited.

Schistosomes are not alone among disease systems where mechanisms bridging environmental factors and epidemiological parameters have been poorly characterized. For example, although it has been well established that meningococcal meningitis in western Africa exhibits seasonal patterns, the particular causes remain uncertain and could range from low humidity to wind speed. Similarly, multiple drivers have been proposed for the seasonal nature of cholera, including rainfall, temperature and planktonic blooms. Yet the specific roles of these drivers have not been resolved and well-established dynamic features, such as the second cholera peak experienced in endemic regions in south Asia, have gone largely unexplained.

Given the limited mechanistic understanding of environment-disease interactions, mathematical models that describe them have seldom included the effects of time-varying environmental processes on transmission dynamics. Where they have been included, seasonality is commonly incorporated phenomenologically, using mathematical functions that are periodic in time and therefore describe in a generic way the seasonal variation in a parameter—a simple sinusoidal function is a common example. Few examples exist where seasonal forcing functions describe the actual processes underlying the environmental drivers of disease dynamics. Because models that incorporate seasonality are sensitive to which parameters are externally forced as well as the shape of their forcing, there is a pressing need to identify the actual mechanisms at play. These mechanisms can include seasonal behaviors of definitive hosts, environmental forcing of vectors and intermediate hosts, sensitivities of parasite survival in the environment and annual variation in host births and deaths.

Understanding the mechanisms that tie environmental change to changes in disease dynamics is crucial for the development of comprehensive control strategies that may be more sustainable and cost effective in the long run. For the case of West Nile virus, for example, simulation studies have suggested that concentrating pesticide spraying efforts during the spring, when most transmission occurs among birds, could be more effective than the current practice of spraying in response to human cases in the late summer and early fall when mosquito numbers
are already in decline. Ultimately, models and management practices that incorporate the timing of key events such as intermediate host reproduction and parasite development are essential to developing more successful control strategies. Understanding these mechanisms is also vital for estimating the long-term impact of impending climate change at global and regional scales on environmentally mediated diseases, whereas current projections are, to a large extent, empirically-based. Indeed, it has been argued in the case of malaria, for example, that models which are mechanistic, based on plausible underlying drivers of the system and basic biology, rather than empirical relationships, are more useful for predicting and responding to, the influence of climate change.

Table 1 summarizes the evolution from simple deterministic models, to complex spatially-explicit, individually-based models appropriate for studying re-emergent scenarios in the Sichuan environment, where connectivity and environmental heterogeneity structure the dynamics of transmission. Iterative evaluation of alternative models in the light of field data is the essence of the modelling process in the application presented here. Below I summarize a model of schistosomiasis transmission in western China which aims to incorporate mechanistic environment-parasite relationships, in the hopes of understanding the local determinants of transmission and its control in endemic settings. Spatial extensions to the model are discussed and an alternative, stochastic framework is proposed for application to re-emergent disease.

Modelling Schistosome Transmission

The use of mathematical models in the study of schistosomiasis dates back to the 1960s, when a four-parameter model was first proposed and used to explain the dynamics of endemic disease. Since then, a number of models have been developed and used to explore the biological and epidemiological characteristics of schistosome species and their hosts, with a majority of them focused on *S. mansoni* and *S. haematobium* and a few on *S. japonicum*. This literature has three notable characteristics, it is explanatory rather than predictive, it is focused on phenomenological and, thereby, generalizable aspects of disease transmission and, for the most part, it has relied on analytical rather than computational methods of analysis. Koopman has written of the successes and limitations of these models in general and where they fit into a more comprehensive mathematical approach. Thus far, these models have had a very limited impact on field studies and control programs. One reason for this is the difficulty in adapting models to site-specific conditions, such as local climatic factors and intermediate host dynamics.

To date, we have used a model of schistosomiasis transmission for our work in China with tactical rather than strategic objectives. Our focus is on site-specific transmission and the issue of selecting from the limited array of feasible control modalities that are effective and sustainable in a particular village. This is because Chinese experience, as well as our recent investigations, has clearly shown considerable variability in the prevalence and intensity of human infection in villages with similar agriculture but that are geographically proximate. Hence, we regard the model as a platform for the synthesis of general knowledge of the mechanisms of disease transmission, quantitative estimates of biological parameter values and the local factors influencing transmission. To that end, the model has been extended to incorporate additional phenomena and additional data. Here, I build on the underlying model structure and parametrization described elsewhere, incorporating the influence of additional environmental phenomena.

The Model

The structure of the delay-differential equation model is shown schematically in Figure 1. Three state variables are tracked in the model, worm burden in each risk group, the density of susceptible snails in each environment and the density of infected snails in that environment. Here risk group refers to occupational subgroups known in this region to exhibit pronounced differences in the timing, intensity and location of water contact and corresponding infection levels, including farmers, students and others, the latter including domestic workers, teachers,
Table 1. The evolution of modelling frameworks for *Schistosoma japonicum* transmission in Sichuan, China

<table>
<thead>
<tr>
<th>Model Complexity</th>
<th>Deterministic</th>
<th>Stochastic</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Single-Population Model</td>
<td>Multi-Risk Group Model</td>
</tr>
<tr>
<td>Heterogeneous populations</td>
<td>No</td>
<td>Somewhat</td>
</tr>
<tr>
<td>Assumption-free worm aggregation process</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Stochastic parasite introduction</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Spatial heterogeneity(^a)</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Model complexity</td>
<td>Simple</td>
<td>Moderate</td>
</tr>
<tr>
<td>Suitable for studying control in endemic areas</td>
<td>Fair</td>
<td>Good</td>
</tr>
<tr>
<td>Suitable for studying emergence</td>
<td>Fair</td>
<td>Fair</td>
</tr>
</tbody>
</table>

\(^a\)While both deterministic and stochastic models have the potential to be spatially-explicit, stochastic models of individuals interacting in space provide additional fidelity.
etc. Environment refers to the land in which a risk group lives and farms. Hence, for each occupational group, $i$, living in environment $k$, the mean worm burden is given by the solution of the state equation:

$$\frac{dw_i}{dt} = \alpha e^{-\mu_{w} t} S_i(t - \tau_w) \gamma \frac{C_k(t - \tau_w) f(w_{ik})}{g_1 g_2 g_3 g_4} - \mu_{w} w_i(t)$$ \hspace{1cm} (1.1)$$

where

- $S_i(t)$ is the water exposure index of occupation group $i$ and reflects the seasonal variation in water contact;
- $e^{-\mu_{w} t}$ is the fraction of worms surviving the development time in humans;
- $\alpha$ is the number of parasites acquired per cercaria per m$^2$ skin surface contact;
- $f(w_{ik})$ is the density dependent worm establishment function which describes a process in which the likelihood of developing into an adult worm is assumed to be reduced when the worm burden is high due to a 'crowding effect', to concomitant immunity, or both;
- $\mu_{w}$ is the worm mortality rate; and
- $C_k(t - \tau_w)$ is the mean spatial density of cercariae in irrigation system $k$ at time $t$. The time delay is due to the fact that the rate of change in the number of adult worms at time $t$ is due to exposure to cercariae at time $t - \tau_w$, where $\tau_w$ is the worm development period in human hosts.

**Modelling Cercariae-Environment Interactions**

Cercariae are the free-living aquatic stage of the parasite which can infect humans and other mammals. They are negatively geotropic and positively phototropic, thus cercariae accumulate at the surface of water where they seek an appropriate mammalian host. They are highly susceptible to environmental stressors, including desiccation, turbulence in the water column, water temperature, aquatic chemistry and light. Water temperature and flow are key determinants.
of cercarial viability and thus $C_k(t)$ is dependent on the infected snail population as modified by these environmental factors:

$$C_k(t) = I_c(T_1) \frac{r_c(t)}{A_s} \alpha A_k z_k(t)$$

where

$I_c(T_1)$ is the temperature-dependent infectivity of cercariae, described below;

$T_1$ is the surface water temperature, measured directly using an automated logger, or estimated from air temperature using a model described below; and

$A_s$ is the nominal surface water area of the village irrigation system;

$r_c(t)$ the precipitation-and/or irrigation-dependent modulation of the average daily cercarial production $[\alpha A_k z_k(t)]$ which enters the aquatic environment, defined briefly below and in detail elsewhere;

$\sigma$ is the cercarial production per infected snail per day;

$A_k$ is the area of snail habitat;

$z_k(t)$ is the infected snail density.

**Temperature-Dependent Cercarial Activity**

Cercarial activity, including host-seeking, surface seeking, host penetration and survival are known to be temperature sensitive. Experiments that examine the influence of temperature on successful penetration and establishment in animal hosts reveal the combined effect of temperature on multiple activities. Cercariae exposed to temperatures between 15 and 30 degrees C show the highest worm recovery rates from mouse hosts. Above and below this range, recovery rates decrease, resulting in the annual infectivity cycle depicted in Figure 2 using temperature data for the Shian 5 study village in 2004. This relationship is incorporated in the model as $I_c(T_1)$, the temperature-dependent infectivity of cercariae, serving as one source of seasonal limitation of transmission in the framework presented here.

**Flow-Dependent Cercarial Activity**

Cercarial production is modulated by the availability of water in channels, $r_c(t)$, which can be predicted from precipitation and temperature using a conceptual rainfall-runoff model, IHACRES, described elsewhere and modeled following the simple binary formulation:
where $q_t$ is the normalized, IHACRES-predicted channel discharge at time step $t$; and $\Omega_c$ is the discharge threshold for cercarial release. If flow falls below the threshold for cercarial release, then $r_c(t) = 0$, effectively prohibiting cercarial penetration of hosts. When the flow threshold is met or exceeded, $r_c(t) = 1$ and transmission proceeds unimpeded. Thus, during and after rain events, when flowing water is available, cercarial dispersion and penetration can occur. This formulation is consistent with the ecology of *Oncomelania* snails, which reside above the waterline but are submerged and shed cercariae when channel flows rise. A sample classification of daily $r_c(t)$ in one study site for the year 2003 is given in Figure 3.

**Modelling Snail-Environment Interactions**

Models of schistosome intermediate hosts have typically explored a limited number of functional forms and environmental variables, such as Woolhouse and Chandiwana and Woolhouse, who selected simple nonlinear models relating water temperature and *B. globosus* recruitment and linear models relating mortality and water temperature. Woolhouse and Chandiwana adapted their previous model for flowing water environments, adding the effect of high rainfall. In contrast to the intermediate hosts of African schistosomes, modelling of the *Oncomelania hupensis* host of *S. japonicum* is rare.
O. hupensis snails are amphibious, inhabiting irrigation canals, riparian zones and littoral environments. The vegetation in these sites serves to maintain a suitable microenvironment, including temperature and humidity, as well as providing food and refuge resources. Juveniles are submerged during early stages of development, while adults are often found above the water line on vegetation and on shaded moist soil. Adults persist under environmental stress by closing their shell opening with a maneuverable operculum, allowing for aestivation and making them somewhat resistant to dry conditions.47,48

Liang et al33 previously used a temperature-dependent recruitment model coupled with constant annual mortality to model seasonal abundance fluctuations of O. hupensis, but no direct measurements of recruitment, mortality or environmental variables were made to construct this model. Others have shown that O. hupensis is highly sensitive to seasonal weather conditions including flooding, temperature and humidity.5,43 In response to these sensitivities, another study 49 used a mark-recapture technique to directly measure birth and mortality processes under changing environmental conditions, finding temperature and heavy precipitation to be most influential in determining abundance. A validated population model for O. hupensis was presented, suitable for predicting snail abundance in changing environments. In this model, the susceptible snail state equation is defined as:

\[
\frac{dx_k}{dt} = g(t - t_x, \bar{E}_{t-x}) x_k(t - t_x) - b(t, \bar{E}_t) x_k(t)
\]

(1.4)

where population gains in environment \(k\) are accomplished by the recruitment term \(g(t - t_x, \bar{E}_{t-x})\), lagged by a temperature-dependent development time, \(t_x\), required to reach mature size (estimated for O. hupensis elsewhere50) and losses are accounted for by mortality term \(b(t, \bar{E}_t)\), where \(\bar{E}_t\) is a vector of environmental variables at time \(t\). Submodels \(g\) and \(b\) are defined as follows, with model fitting and parameters described in detail elsewhere:49

\[
g(t - t_x, \bar{E}_{t-x}) = \beta_1 e^{\frac{-0.5}{\beta_2} \left( \log \frac{T(t - t_x)}{\beta_3} \right)^2} + \beta_4 e^{\frac{-0.5}{\beta_6} \left( \frac{R(t - t_x) - \beta_5}{\beta_7} \right)^2}
\]

(1.5)

where \(T(t - t_x)\) is air temperature and \(R(t - t_x)\) is a count of rain events >15 mm per month at time \(t - t_x\) and \(\beta_1, \beta_2\) are fit parameters; and

\[
b(t, \bar{E}_t) = \delta_1 + 4\delta_2 \frac{e^{\frac{T(t) - \delta_3}{\delta_4}}}{\left(1 + e^{\frac{T(t) - \delta_3}{\delta_4}}\right)^2} + \delta_5 R(t)
\]

(1.6)

where \(T(t)\) is air temperature, \(R(t)\) is a count of rain events >15 mm per month and \(\delta_1, \delta_2\) are fit parameters.

Fits of submodels \(g\) and \(b\) to environmental data are shown in Figure 4. Notice that the susceptible snail state equation is not dependent on other, endogenous transmission model state variables. Consequently, the susceptible snail model can be calibrated independent of the full transmission model, thus economizing the computation required for calibration, described further in the chapter by Spear and Hubbard in this volume.

State variable \(z_k(t)\), the density of infected snails in environment \(k\), is given by the solution to:

\[
\frac{dz_k}{dt} = \rho e^{-\mu \tau_z } \xi(T_1) M_k(t - \tau_z) x_k(t - \tau_z) - \mu_z z_k
\]

(1.7)
Figure 4. A) Relationship between instantaneous per capita recruitment rate for *O. hupensis robertsoni* and mean air temperature and mean number of rainfall events >15 mm (month⁻¹). Climate data are lagged by *t*, as discussed elsewhere. B) Relationship between instantaneous per capita mortality rate for *O. hupensis robertsoni* and mean air temperature and mean number of rainfall events >15 mm (month⁻¹). Reprinted with permission from: Remais J, Hubbard A, Wu Z, Spear R. J Appl Ecol 2007; 44(4):781-791. © 2007 Blackwell Publishing Ltd.
where

\( x_k(t) \) is the density of susceptible snails in environment \( k \);

\( \rho \) is the fraction of those miracidia which successfully infect snails;

\( \xi \) is a parameter representing the degree of spatial convergence of the distribution of snail hosts and miracidia;

\( \mu_c \) is the mortality rate of infected snails; and

\( M_k(t) \) is the mean density of miracidia in the irrigation system in environment \( k \), derived from hatched eggs, a process described below.

The implications of an environmentally-driven snail model are shown in Figure 5 using a transmission model previously calibrated for Shian 5, described in detail elsewhere and in the chapter by Spear and Hubbard in this volume. The spring snail population peak generated by the model leads to significant infected snail numbers earlier in the year when compared to the model used by Liang. As a consequence, the onset of peak cercarial release into waterways is shifted back by more than a month, from late September to mid-August, a prediction that is in line with available cercarial concentration data from field studies in Shian 5. Comparisons to field data of this sort can highlight how environmentally-driven intermediate host models can bring model performance into better agreement with real world observations.

**Modelling Ova-Environment and Miracidia-Environment Interactions**

Total egg production from all risk groups is modeled as:

\[
E_k(t) = \frac{1}{2} b \sum g_i n_i w_{i,k} \Phi(w_{i,k}, k_{w_i})
\]

where

\( b \) is eggs per gram stool (EPG) per worm pair based on Hubbard et al;

\( g_i \) is the average stool production of a member of the \( i \)th group;

\( n_i \) is the number of people in \( i \)th group whose stool is used as fertilizer;

\( \Phi(w_{i,k}, k_{w_i}) \) is worm mating probability following May and described elsewhere.

The factor \( \frac{1}{2} \) converts mean worm burden to worm pairs.
Before hatching into miracidia, excreted eggs are subject to environmental stress. They are resilient, however and can persist for days on fields before being washed into irrigation channels by a precipitation event. A composite parameter representing on-field inactivation of eggs can be calculated from literature values of egg resilience and a simple first-order inactivation process can be used to express viable eggs, $E^*(t)$, as a function of the sum of decaying eggs contributed since the last flow event:

$$E^*_k(t) = \sum_{T - T_{\tau}}^{T} E_k(t)e^{-\epsilon_d(T-t)} \quad (1.9)$$

where

- $E_k(t)$ is the sum of viable eggs shed by infected humans in environment $k$ since the last flow event;
- $E(t)$ represents eggs excreted into environment, defined above;
- $\epsilon_d$ is the decay constant governing inactivation of eggs lying dormant on fields between flow events;
- $T - T_{\tau}$ is the time since last flow event.

Miracidia are short-lived, free-swimming and are drawn to light, accumulating near the surface of water where they seek an appropriate snail host. They are sensitive to water temperature and aquatic chemistry, with the former exerting a pronounced influence on viability. Experimental data of the influence of temperature on miracidial infectivity have shown optimal activity between 15 and 30 degrees C, a relationship incorporated into the model of the net effective density of miracidia in environment $k$, $M_k(t)$, a function of viable eggs in the environment, $E_k^*$:

$$M_k(t) = I_m(T_1) \frac{r_s(t)}{A_s} \beta E_k^*(t) \quad (1.10)$$

where

- $I_m(T_1)$ is the surface water temperature dependent miracidial infectivity to snails analogous to $I_c(T_1)$ discussed above for cercariae;
- $A_s$ is the nominal surface water area of the village irrigation system;
- $r_s(t)$ is the precipitation-and/or irrigation-dependent modulation of the average daily miracidial production [$\beta E(t)$] which enters the aquatic environment, defined briefly below and in detail elsewhere;\n
$\beta$ is the fraction of the total daily egg production of infected villagers returned into the environment as fertilizer, adjusted for the presence of sanitation.

Flow events provide opportunities for viable eggs to hatch and I therefore define $r_s(t)$ analogous to the cercarial equation at time $t$ as:

$$r_s(t) = \begin{cases} 0 & q_t < \Omega_e \\ 1 & q_t \geq \Omega_e \end{cases} \quad (1.11)$$

where $q_t$ is the normalized, IHACRES-predicted discharge at time step $t$, as above. Here, if water flow falls below the threshold for egg hatching, $\Omega_e$, $r_s(t) = 0$ and eggs lie dormant. When the flow threshold is met or exceeded, $r_s(t) = 1$ and viable eggs on fields are washed into the irrigation system, where they hatch and can infect snails.

**Model Parameters**

The model was structured and parameterized to allow the use of as much of the field data as can be feasibly collected with the methods available in rural China. This includes environmental data (described below), cross-sectional data on snail population density, seasonally varying water contact patterns by group and survey data on the intensity of human infection. Some of these data are inputs to the model and some are used for parameter estimation. The issue of parameter
estimation is complex but central to our approach. Table 2 lists parameter values for the model and their literature sources. When used to study interventions (discussed in detail in the chapter by Seto and Carlton in this volume), a fundamental challenge is to reduce the residual uncertainty

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Interpretation and Unit</th>
<th>Ranges</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Biological</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\tau_w$</td>
<td>Development time of worms in human host (day)</td>
<td>20-40</td>
<td>59</td>
</tr>
<tr>
<td>$\mu_w$</td>
<td>Worm natural mortality (/day)</td>
<td>0.000183-0.0014</td>
<td>59</td>
</tr>
<tr>
<td>$h$</td>
<td>Eggs excreted (/worm pair/gram feces)</td>
<td>0.768-2.72</td>
<td>53</td>
</tr>
<tr>
<td>$\mu_z$</td>
<td>Patent and latent snail death rate (/day)</td>
<td>0.0063-0.033</td>
<td>60</td>
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<tr>
<td>$\sigma$</td>
<td>Cercarial production (/sporocyst/day)</td>
<td>20-50</td>
<td>61,62</td>
</tr>
<tr>
<td>$h_{PZQ}$</td>
<td>Efficacy of praziquantel</td>
<td>0.8-0.95</td>
<td>63,64</td>
</tr>
<tr>
<td>$D_{D1}$</td>
<td>Degree-days for sporocyst development</td>
<td>1550-1950</td>
<td>65</td>
</tr>
<tr>
<td>$T_{D1}$</td>
<td>Threshold temperature for sporocyst development (°C)</td>
<td>12-15</td>
<td>62</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>Schistosome acquired (/cercaria/m² contact)</td>
<td>0.0001-0.5</td>
<td>-</td>
</tr>
<tr>
<td>$\rho$</td>
<td>Intermediate host infection (/miracidium/m² surface water)</td>
<td>0.000001-0.00005</td>
<td>-</td>
</tr>
<tr>
<td>$\beta_{i6}$</td>
<td>Intermediate host recruitment parameters</td>
<td>See ref</td>
<td>49</td>
</tr>
<tr>
<td>$\delta_{i5}$</td>
<td>Intermediate host mortality parameters</td>
<td>See ref</td>
<td>49</td>
</tr>
<tr>
<td><strong>Site-specific</strong></td>
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<td></td>
</tr>
<tr>
<td>$w_{i0}$</td>
<td>Initial worm burden in the $i$th group</td>
<td>Data estimate*</td>
<td>Local data</td>
</tr>
<tr>
<td>$Z_0$</td>
<td>Initial density of infected snails</td>
<td>Data estimate*</td>
<td>Local data</td>
</tr>
<tr>
<td>$S_i(t)$</td>
<td>Water contact index</td>
<td>Data estimate*</td>
<td>Local data</td>
</tr>
<tr>
<td>$x_0$</td>
<td>Initial density of susceptible snails</td>
<td>Data estimate*</td>
<td>Local data</td>
</tr>
<tr>
<td>$\kappa_{i0}$</td>
<td>Initial worm aggregation parameter</td>
<td>Data estimate*</td>
<td>Local data</td>
</tr>
<tr>
<td>$x_0$</td>
<td>Initial mean snail density</td>
<td>17-35</td>
<td>66</td>
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<tr>
<td>$\gamma_i$</td>
<td>Spatial index for the distribution and interaction between exposure and cercariae for $i$th group</td>
<td>Data estimate*</td>
<td>Local data</td>
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<tr>
<td>$\xi$</td>
<td>Spatial index for the distribution and interaction between snails and miracidia</td>
<td>1</td>
<td>-</td>
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<td><strong>Inputs</strong></td>
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<td></td>
<td></td>
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<tr>
<td>$T_1$</td>
<td>Water temperature (°C)</td>
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<td>Local data</td>
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<tr>
<td>$T_2$</td>
<td>Snail microenvironment temperature (°C)</td>
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</tr>
<tr>
<td>$C_{chemo}$</td>
<td>Chemotherapy coverage</td>
<td>Data estimate*</td>
<td>Local data</td>
</tr>
<tr>
<td>$P$</td>
<td>Rainfall (mm/day)</td>
<td>No constraint</td>
<td>Local data</td>
</tr>
<tr>
<td>$r(t)$</td>
<td>Precipitation-driven channel flow (binary)</td>
<td>Data estimate</td>
<td>40</td>
</tr>
</tbody>
</table>
in model output, or its behavior more broadly, after as much of the local data as possible has been utilized to narrow the posterior distributions of the parameter values (see chapter by Spear and Hubbard in this volume).

To that end, we have conducted a variety of field studies to better understand the importance of certain elements of the model, or to obtain parameter estimates relevant to the biology of the snail or parasite specific to the region in which we work. Examples are the value of the parameter describing the production of parasite eggs per mated worm pair per gram of stool,\textsuperscript{53} the importance of rainfall in determining infected snail densities and the concentration of cercariae in irrigation water.\textsuperscript{52} There is no question that the modelling approach, with the ultimate objective of designing effective intervention strategies to meet public health objectives, has led us to seek quantitative estimates of factors controlling disease transmission that have not been of great interest to Asian parasitologists since the work of Pesigan.\textsuperscript{67}

Environmental Data

Modelling the environmental drivers of seasonality requires an accurate dataset of environmental variables, acquired by measurement where possible and prediction where not. As in all environmental monitoring, strict quality assurance measures need be taken, including instrument calibration/certification, statistically valid sampling designs, reference sites and data verification.\textsuperscript{68,69} In the work described here, air temperature, barometric pressure and relative humidity are collected relatively easily throughout the study region using continuous loggers (Hobo Onset H21-002) sampling at 12 minute intervals, validated with regional data available from the National Climatic Data Center.\textsuperscript{70} Likewise, water temperature and water column height (stage) are logged at the same interval using similar equipment (Hobo Onset U20-001-01, U22-01) in a representative sample of irrigation channels. To estimate flow (m$^3$ s$^{-1}$) from stage (m) in these channels, flow measurements must be made at multiple flow volumes in order to construct a simple rating curve. Daily precipitation is collected using a combination of tipping gauges (Hobo Onset RG3-M) and manually read rain gauges. Where data were missing due to equipment or staff error (typically accounting for «1 percent of data points in the study), data were obtained from the NOAA weather station located at the Xichang municipal airport (World Meteorological Organization ID 56571), approximately 13 km from the study sites. Where water temperature was not directly measured, it was estimated from air temperature using a standard, simple linear model:\textsuperscript{71}

\begin{equation}
T_w(t) = \alpha + \beta T_a(t)
\end{equation}

where $T_w$ = water temperature, $T_a$ = air temperature and $\alpha$ and $\beta$ are fit parameters. Time lags were excluded from the model as the observed lags (<4 hours) were much shorter than the averaging period (1 day), as is typical for temperature predictions in shallow channels.\textsuperscript{72}

Model Dynamics

The model described above, when parameterized as described in the chapter by Spear and Hubbard, generates predictions of the sort depicted in Figure 6. Mean worm burdens for the three risk groups in Shian 5 are summarized for 1000 simulations over the five year period 2001-2005. The impact of two chemotherapies, modeled as described in the chapter by Seto and Carlton in this volume, is shown in the Figure. One characteristic that can be explored using this simulation environment is the time-to-return for worm burdens after chemotherapy. As is evident in the plot, worm burden returns to precontrol levels in the farmer group in less than 3 years, while the student and other group require considerably more time to rebound, owing to their differing exposure profiles.

The seasonal rise in worm burden following the second simulated chemotherapy can be seen in Figure 7 which plots the acquisition (or loss by mortality) of worms in the three risk groups. Note that the acquisition of new worms in the Figure represents exposures to cercariae that took place as many as 6 weeks prior. Notable is the bimodal farmer pattern which results from exposures during the spring planting season. While similar activities occur during the late winter and early spring during the harvest of the winter crop, cercarial shedding and infectivity is limited in this period due to
low temperatures and limited precipitation. During the spring planting, however, temperatures just exceed the limits for cercarial activity and spring rains provide opportunities for the coincidence of cercariae and water contact. The timing and nature of interventions can be selected based on these seasonal patterns, as described in the chapter by Seto and Carleton in this volume.

All three risk groups experience a ‘shut-down’ of worm acquisition in the late fall related to the coincidence of temperature decreases, lower snail numbers and reduced water contact activities. The time-varying $I(T)$, $r(t)$ and $s(t)$ terms are at their minimum values in the late fall through winter and ‘turn on’ again in the spring, remaining ‘on’ through the transmission season. Of great interest is to explore the effects of these gating functions on the transmission process and their sensitivity...
to local and regional environmental changes. Furthermore, efforts are needed to extend the model to account not just for environmental variables, but for their spatial distribution.

**Modelling Spatial Connectivity**

Human schistosomes are model organisms for the study of and response to, the spread of disease in space and time, as their transport through the environment takes place along discrete pathways. Parasites are carried in advective flows along canals and streams as both larvae and ova. Within intermediate snail hosts, parasites are conveyed among and between aquatic and riparian habitats and as adult worms, human and animal hosts serve as the transport mode. With respect to the *S. japonicum* parasite, I term these flows parasite diffusion, using the phrase to encompass all diffusive pathways along which parasites are transported into new and existing locales. The presence of suitable pathways can affect the probability of emergence of transmission, the level of worm burden within a community once transmission is established and how transmission spreads to neighboring areas. What is more, the degree to which an endemic or emergent community is connected can have important implications for the efficacy and sustainability of various control strategies.

In a preliminary exploration of parasite diffusion, the travel time of the free-swimming forms of the parasite or snail larvae due to advective transport in typical irrigation systems was estimated,74 showing empirically that there is significant transport of viable parasite larvae within irrigation channels and that transport of larval stages occurs over considerable distance, with viable organisms detectable as far as 400 m from source snails.38 Using these key transport parameters in a follow-up project, the impact of larval transport on endemic disease transmission was assessed using a spatial-temporal model of networked villages,75 showing that diffusion of larvae via the surface water pathway, on its own, influences not just the intensity of transmission in a village, but also the effectiveness of standard interventions. Such a model allows us to better understand a number of phenomena specific to the endemic situation, such as which villages serve as “sinks” in the network, villages where worm burden can accumulate because they lie at the bottom of a watershed of numerous connected upstream villages.

The implications of a connected landscape have been explored extensively in ecology, where metapopulation models76 describe the effect of migration between connected patches on population conservation. Likewise, environmental and social connections can promote the persistence of schistosomiasis and challenge efforts to control transmission. While hydrological connectivity is relatively straightforward to characterize, social connectivity is considerably more difficult to measure and express mathematically. The indirect transmission of schistosomiasis differs from recent epidemiological modelling of social connectivity and contact networks for communicable disease spread.77-83 Within the context of the endemic transmission situation, small-scale human mobility can spread parasites from village to village. This effect may be small though in comparison to other social behaviors, such as the renting or selling of water buffalo which, if infected, can potentially release much larger numbers of eggs into the environment. While these factors might be modeled much like hydrological connectivity over the small scale via inter-village flows, they differ from the hydrological situation in that these processes can occur over much larger spatial scales and much less predictably. While difficult to estimate precisely, we look to future field data to inform the probabilities that define the movements of heterogeneous hosts; much theoretical and empirical work is needed in this area.

**Extending the Modelling Framework**

Our current research attention has shifted from endemic disease to disease re-emergence, a phenomenon we have documented in the mountainous region of Sichuan Province.84 For studying re-emergence, human infection may be more appropriately modeled by risk groups with stochastic parasite establishment in a heterogeneous environment. The simplest form would be a stochastic compartmental model, where the risk group structure would be maintained with each compartment being comprised of a number of identical individuals. However, the
static representation of the aggregation of parasites in humans in the deterministic model, even within our risk groups, does not translate easily to the re-emergent situation where, initially, the population is parasite-free. Hence, an individually-based model is preferred.\textsuperscript{31} Individual-based micro-simulation models have been utilized in studying schistosomiasis transmission in endemic settings, but without our emphasis on environmental factors.\textsuperscript{35} In ecology there has also been considerable interest in individually-based models.\textsuperscript{86} There, the analog to our discrete population of humans is a population of animals in a heterogeneous, but continuous environment. There are particularly interesting approaches being explored which might allow us to naturally utilize our GIS data base and GPS-based maps in an individually-based model.\textsuperscript{87}
Figure 8 schematically represents the elements of a new model in which the stochastic introduction of parasites is implemented by means of migration of infected hosts and advective parasite transport. Heterogeneous ditch environments, then, serve as platforms wherein eggs are released from infected individuals who, along with uninfected individuals, traverse the waterway environments and are potentially infected by contact with cercarial contaminated water. Stochastic implementations of the egg release and cercarial exposure processes are particularly suitable for a system where transmission is strongly conditioned by both environmental and behavioral factors that defy deterministic formulation. Moreover the stochastic model proposed here provides the structure to capture the potentially large influence of chance events that have been recognized to govern early epidemic dynamics, even in relatively large populations.88-90

Conclusion

The wide array of processes discussed herein can be conceptualized at various spatial and temporal scales, evaluated for their relative abilities to capture relevant transmission dynamics, including seasonal dynamics, and incorporate available field data. Measuring the potential drivers of seasonality may be relatively straightforward in the case of climate, but measuring and formalizing patterns of human behavior, particularly in a spatially explicit context, remain formidable challenges.81 Indeed, iterative evaluation of alternative models in the light of field data is the essence of the modelling process in our application. Ultimately, quantifying and synthesizing the interaction between environmental and social determinants in transmission models offers great promise for developing novel modes of control in diverse environments.

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Parameter Estimation and Site-Specific Calibration of Disease Transmission Models

Robert C. Spear* and A. Hubbard

Abstract
The use of mathematical models for developing management options for controlling infectious diseases at a local scale requires that the structure and parameters of the model reflect the realities of transmission at that scale. Data available to inform local models are generally sparse and come from diverse sources and in diverse formats. These characteristics of the data and the complex structure of transmission models, result in many different parameter sets which mimic the local behavior of the system to within the resolution of field data, even for a model of fixed structure. A Bayesian approach is described, at both a practical and a theoretical level, which involves the assignment of prior parameter distributions and the definition of a semi-quantitative goodness of fit criteria which are essentially priors on the observable outputs. Monte Carlo simulations are used to generate samples from the posterior parameter space. This space is generally much more constrained than the prior space, but with a highly complex multivariate structure induced by the mathematical model. In applying the approach to a model of schistosomiasis transmission in a village in southwestern China, calibration of the model was found to be sensitive to the effective reproductive number, $R_{eff}$. This finding has implications both for computation time for the Monte Carlo analysis and for the specification of field data to efficiently calibrate the model for transmission control.

Introduction
The theme of this book, the use of mathematical transmission models for the management and control of parasitic disease, carries the implicit assumption that the parameters of the model, as well as its structure, are directly applicable to the environment in which transmission is to be managed. While models have been used to great effect to elucidate the social and biological phenomena central to the patterns of disease on large spatial and temporal scales,¹ this has rarely been true for developing control tactics in local environments where the devil is in the details. The challenge is to use local data to tailor the model to reflect local conditions to a degree adequate for informing control tactics at that scale. In this chapter we propose a particular approach that was developed in the context of environmental models, mainly in hydrology and recognizes that the relevant data for model calibration are diverse and come in many formats and with varying degrees of confidence in their accuracy and precision.²

In this volume it can be seen that mechanistically-based models are particularly well suited for the analysis of the transmission dynamics of infectious diseases. A major reason for their...
popularity is that the structure of the model reflects known processes and mechanisms that underlie the observable features of disease transmission. However, mathematical models often become complex as they expand to incorporate more mechanistic detail and the default is often to include more detail rather than less. It is generally a major challenge to match the appropriate level of detail of the model to the purposes of a particular study.

The complexity of mechanistically-based models is in contrast to purely statistical models which are aimed at efficiently summarizing relationships between variables in a particular set of data by utilizing more general and mathematically simpler structures. Both statistical and mechanistic mathematical models include sets of parameters whose values must be specified in order to use the model for subsequent forecasts or analyses. Parameter estimation is central to the statistical modelling process in the sense of finding parameters that result in best fits to the specific data set about which the analysis revolves. In contrast, there is often prior information on the values of at least a subset of the parameters of mechanistically-based models independent from the particular application at hand. Further, some of the parameters of mechanistic models can often be estimated by independent experiments conducted before or during the modelling project. Hence, a mechanistically-based model can be thought of as a platform for the merging three types of information on the processes being modeled:

- The set of causal hypotheses that describe the current qualitative understanding of how the different processes and variables are inter-related (the structure of the model);
- Quantitative information that may be available from the literature or from independent laboratory or field experiments (prior information on the model parameters);
- The data specific to the particular problem under analysis (generally measurements dependent on a subset of the state variables of the system).

The challenge is to merge these three types of information with the objective of identifying effective management options to diminish disease transmission. This goal immediately places some constraints on the spatial and temporal scale of the processes under analysis as well as the nature and format of data available to inform the model both qualitatively and quantitatively. For example, there is an important subset of disease transmission models that implicitly depend on, although seldom acknowledge, environmental factors like air temperature or water availability. Hence, an important determinant of spatial scale concerns the relationship between the human population subject to infection and the spatial homogeneity or lack thereof in these key environmental variables. In some cases the land of a single village may be the relevant environment and in others a set of villages may be more appropriate, for example, if they lie within a connected and relatively homogeneous patch of vector habitat. Temporal scales range from days and weeks in the case of disease outbreaks to months and years for tracking the prevalence or intensity of endemic disease.

Ideally the model structure and scale are well matched to the data set available for analysis. This may not be the case in that data collection is often dictated by political boundaries and logistical considerations rather than the realities of the transmission processes. The underlying issue is one of connectivity which is easily understood by noting that it is not effective management to suppress transmission in one village if its neighbors continue to harbor ample infected people and/or vectors and the pathways exist to reintroduce them into the controlled village at the first opportunity. In any case, the nature and quantity of the problem-specific data strongly condition the approach to parameter estimation/calibration. Therefore, we now sketch the characteristics of data we postulate to be typically available for use in the management of environmentally-mediated infectious diseases.

**Local Data**

The local data relevant to understanding the transmission dynamics of infectious diseases in human populations can be roughly divided into two categories which we shall term survey data and monitoring data. Monitoring data are defined as those data that are collected routinely at locations within the geographical area of interest and at regular time intervals. Environmental
temperature and rainfall amounts are the most commonly available data of this sort. Survey data are those acquired over particular time periods of intensive investigation or, alternately, episodically as resources become available. In the developing world, prevalence or intensity of infection in the human population or the abundance of vectors or intermediate hosts are examples of survey data whereas, for reportable diseases in the developed world, such data may be classified as monitoring data if it is reliably summarized in weekly or monthly reports.

From a modelling perspective, the essential difference between monitoring and survey data is that the former are amenable to time series methods of summarization and analysis of which a great many are available. However, when addressing the issues of management and control of disease transmission in the developing world, time series data are seldom available on the key variables and processes of interest beyond the weather. For the most part, what is known of these variables is available only via survey data. Hence, time series models may be useful submodels of larger mechanistic simulation models of the overall processes of interest, but the data to be “fit,” i.e., the outcome data that define the likelihood, are generally survey data. Moreover, survey data are often from surveys of different sorts collected at different times and places within the region of interest making it very difficult to generalize about the nature, structure, or amount of such data. Hence, approaches to parameter estimation that depend on specifically structured data, e.g., monitoring data, are of limited utility in disease management analyses typical of the developing world.

A second general characteristic of the data available to inform model parameterization and calibration relates to transferability, or lack thereof, from village to village or region to region. It is useful to distinguish between two data types in this context, site-specific data and what will be called biological data, that is, data specific to the disease, the infectious agent, its vectors, or intermediate hosts that can be assumed to apply to host and/or vector populations that reside or range over extended geographical areas. For example, the average number of eggs produced per day by an adult female schistosome within a human host should not vary locally, but might over longer distances or between genetically isolated parasite populations. The importance of the distinction between site-specific and biological parameters is that estimates of biological parameters obtained from one site can, in principal, be transferred to a second area within the same ecological zone whereas this is not often the case for site-specific parameters related to local agriculture or water availability.

Neither the biological nor the site-specific parameters of a well-developed disease transmission model will ever be known precisely. Even with the most mathematically tractable models and copious time series data, the best one can hope to achieve are a set of parameter estimates but also an estimate of the residual uncertainty in their values. The attractive possibility, however, is that as the model is applied to new situations the residual uncertainty in biological parameter values, as reflected in a statistical distribution that characterizes the current information about a parameter, will continually narrow. This possibility calls out for a Bayesian approach to the parameter estimation problem. For a number of years, we have utilized a particular approach that is well suited to the nature and amount of data available in disease transmission studies. As originally formulated the approach was admittedly ad hoc, but understandable and practical. It has been widely applied and extended, mainly in the field of hydrology. More recently we have recognized that this approach can be subsumed under a more general framework termed Bayesian Melding by Poole and Rafterty. Here the procedure will be described in its practical context and later summarized more generally using their terminology.

In short, the procedure begins with the selection of a specific model structure and the specification of statistical distributions describing the uncertainty in each constant model parameter including, perhaps, those of time variable functions like the temporal pattern of exposure as driven by agricultural activity. The model structure and parameter distributions describe an ensemble of models which are then studied via Monte Carlo methods. The third element of the approach is the specification of a set of criteria to assess the performance of each model realization against the field data, that is, a measure of the goodness-of-fit. However, here goodness-of-fit is defined quite generally and can be based on any observable feature of the model’s output, for example the timing between events, or by requiring that a state variable lies in some plausible range at certain times.
Because field data are often imprecise or approximate, it is sometimes unwise to overemphasize fits to point estimates. In earlier studies, this concern led to a binary criterion of fit, pass/not-pass, where passing implies that a particular simulation run was judged to mimic the essential features of the field data. In this variant of the method, each parameter vector associated with passing or not-passing simulations is saved for subsequent statistical analysis. The Bayesian nature of the approach can be seen by recognizing that the passing parameter vectors are samples from a multivariate posterior distribution which is generally very much more constrained than the original parameter distributions, the priors that were used in the initial Monte Carlo runs.

The multivariate posterior distribution of passing parameters is the concrete expression of the fact that many different parameter combinations can produce model behavior consistent with field observations. This is not an artifact of the pass/fail classification, but it implies that, in general, these complex models are not invertible, that is, it is generally not possible to back-calculate a single optimum point in parameter space that produces the best fit, at least in any traditional sense. Beven has written extensively about this characteristic of complex environmental models, extending it to incorporate different model structures and termed it equifinality. Particularly relevant to applications of complex models to disease control is the observation that “one implication of rejecting the concept of an optimal parameter set and accepting the concept of equifinality is that the uncertainty associated with the use of models in prediction might be wider than has hitherto been considered…”

A Calibration Example

To exemplify the calibration process we propose, the model of *S. japonicum* transmission presented in the chapter by Remais is utilized in the context of exploring site-specific control options as discussed in the chapter by Seto and Carlton. Structurally, the model is comprised of a set of nonlinear differential-difference equations which track mean village worm burden in three groups of villagers; farmers, students and others who live in the village but who have limited water contact often due to non-agricultural employment outside the village. The rationale for dividing the population into three groups relates to their differences in water contact, both in timing and degree. The fourth state variable is the mean village density of infected snails. The delay terms are the development time of the parasite in the human host and the prepatent interval in snails. As recounted in Chapter 12, the model is in the tradition that began with MacDonald, but adapted to the epidemiological and environmental determinants of transmission in the hilly and mountainous areas of China. In this setting there is very limited evidence of acquired immunity as a function of age or infection history and very similar patterns in disease prevalence and intensity between adult men and women within a village.

Including the degree-day model of snail development, the model has 10 biological parameters, all of which are constant and 15 site-specific parameters of which 9 are constant and 6 are time-varying. Examples of time varying parameters are the average water contact rate, $S_i(t)$, of each group of villagers over the season and the village average density of susceptible snails, $X(t)$. The latter is treated as a time variable parameter rather than a state variable because, even in highly endemic conditions, the prevalence of infection in snails does not exceed 2%. The density of susceptible snails, however, is generated by an independent submodel which is driven by temperature and rainfall data as described in Chapter 6.

The next step in the process is to specify plausible ranges for the biological parameters from the literature as described in a paper on the calibration of an earlier version of the model. As discussed in that paper, there is no magic formula for assigning the prior parameter distributions. Some are reasonably well defined in the literature and others can only be estimated from informed guesses. However, there are technical reasons elaborated below that motivate the assumption that the individual parameter priors are uniformly distributed at least for the initial stages of the analysis that are addressed here. (Other distributional assumptions will be seen to require the full Bayesian melding procedure to properly estimate the posterior distribution as discussed below.) Similarly, survey data inform the initial ranges to be assigned to the site-specific parameters, for example, the area of snail habitat in a particular village.
In general, the ranges for the site-specific parameters are narrower than is the case for the biological parameters. However, the price paid is that collection of the necessary site-specific data is often labor intensive. Again, the ranges stipulated for both types of parameters are regarded as the limits of uniform distributions. Table 1 contains the ranges used in the main model and identifies the time variable parameters used in this example. More complete listing are given in Liang et al¹⁵ and in Chapter 6.

The particular village for the calibration example is Chaunxing Xinlong 7 which lies in hilly terrain in Xichang County of southwestern Sichuan Province. The village is hydrologically isolated from its neighbors and can be treated as a single unit, at least from the perspective of determinants of its endemic level. The field data on which the criteria for passing simulations are defined are based on cross-sectional infection surveys in 2000 and 2002, as detailed in Liang et al¹⁴ and supplemented by a more recent data set from 2003-2004.¹⁵ These are survey data as defined above and are comprised of village-average infected snail densities, generally in the spring of the year, average egg excretion data among farmers and students in the winter and estimates of village average cercarial density from mouse bioassays once per month from June through August of 2001.¹³

### Table 1. Parameter and initial condition limits

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Definition and Units</th>
<th>Limits</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Biological</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\tau_w$</td>
<td>Development delay of worms in humans (days)</td>
<td>20-40</td>
</tr>
<tr>
<td>$\mu_w$</td>
<td>Worm mortality rate (/day)</td>
<td>0.000183-0.0014</td>
</tr>
<tr>
<td>$\mu_z$</td>
<td>Infected snail mortality rate (/day)</td>
<td>0.0063-0.033</td>
</tr>
<tr>
<td>$h$</td>
<td>Parasite egg excreted (/worm pair/gm feces)</td>
<td>0.768-2.72</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>Cercarial production (/sporocyst/day)</td>
<td>0.1-5</td>
</tr>
<tr>
<td>$h_{PZQ}$</td>
<td>Efficacy of Praziquantel</td>
<td>0.8-0.95</td>
</tr>
<tr>
<td>$D_{DD}$</td>
<td>Degree-days for sporocyst development</td>
<td>1550-1950</td>
</tr>
<tr>
<td>$T_{DD}$</td>
<td>Threshold temp for sporocyst development °C</td>
<td>12-15</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>Schistosome acquisition (/cercaria/m² contact)</td>
<td>0.0001-0.5</td>
</tr>
<tr>
<td>$\rho$</td>
<td>Snail infection (/miracidium/m² surface water)</td>
<td>0.000001-0.0005</td>
</tr>
<tr>
<td><strong>Site specific</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\omega_i$</td>
<td>Initial worm burden in i-th group</td>
<td>90-140, 60-110, 30-60</td>
</tr>
<tr>
<td>$z_i$</td>
<td>Initial density of infected snails</td>
<td>0.401-0.773</td>
</tr>
<tr>
<td>$k_i$</td>
<td>Worm aggregation parameter in i-th group</td>
<td>0.4-1.1, 0.15-0.72, 0.5-0.8</td>
</tr>
<tr>
<td>$A_h$</td>
<td>Area of snail habitat (m²)</td>
<td>8004</td>
</tr>
<tr>
<td>$A_v$</td>
<td>Area of village surface water (m²)</td>
<td>2208</td>
</tr>
<tr>
<td><strong>Time-varying</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\tau(t)$</td>
<td>Parasite development in snail (days)</td>
<td>Submodel</td>
</tr>
<tr>
<td>$T_1(t)$</td>
<td>Air temperature (°C)</td>
<td>Monitoring data</td>
</tr>
<tr>
<td>$P(t)$</td>
<td>Rainfall (mm/day)</td>
<td>Monitoring data</td>
</tr>
<tr>
<td>$X(t)$</td>
<td>Uninfected snail density (snails/m²)</td>
<td>Submodel</td>
</tr>
<tr>
<td>$S(t)$</td>
<td>Daily water contact i-th group (m²/day)</td>
<td>Survey data</td>
</tr>
<tr>
<td>$r(t)$</td>
<td>Water availability (fraction of full)</td>
<td>Set to unity</td>
</tr>
</tbody>
</table>
Table 2 contains the ranges of the passing criteria. Local daily air temperature and rainfall data are used as inputs to each simulation for the uninfected snail population submodel, the temperature dependent infectivity parameters, $I_c(T)$, $I_m(T)$ and for the degree-day calculation on which the developmental delay of the parasite in the snail is based.

The simulations include environmental interventions in the 2003-04 period, notably focused snail control and the introduction of a number of household anerobic digesters that accept both human and pig wastes and subsequently provide biogas for cooking. These units also inactivate schistosome eggs with high efficiency, thereby decreasing the egg input into the environment, although the digested material is still used as fertilizer. Details are given in Liang et al.15

Because the timing of the collection of the various types of survey data is seldom coincident, the initial conditions of the state variables also need to be assigned and these, too, are typically generated by sampling from prior distributions the ranges of which are also included in Table 1. This can be tricky in models which contain developmental delays, such as the current example. A delay \( \tau \) necessitates an initial function, rather than a single initial value, in the interval \((-\tau, 0)\).

Further complications arise from a temperature-variable delay as in the present case. However, these are not complications introduced by this particular approach to parameter estimation, but are generic to any simulation application.

With the model structure given (Chapter 6), the prior parameter distributions and the distributions of the initial conditions specified (Table 1) and the pass/fail criterion defined (Table 2), Monte Carlo simulation trials can begin. In each simulation the parameters and initial conditions are randomly drawn from the uniform prior distributions, the state equations integrated over the specified interval and the calibration conditions checked at the appropriate times. The run is classed as a pass or a fail, the parameter vector stored for subsequent analysis and the process repeated.

### Table 2. Passing criteria

<table>
<thead>
<tr>
<th>Dates</th>
<th>Risk Group</th>
<th>Lower Limit</th>
<th>Upper Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Human infection</strong></td>
<td></td>
<td>epg</td>
<td></td>
</tr>
<tr>
<td>15-30 Nov 2000</td>
<td>Farmer</td>
<td>70</td>
<td>200</td>
</tr>
<tr>
<td></td>
<td>Student</td>
<td>20</td>
<td>120</td>
</tr>
<tr>
<td>19-24 Oct 2002</td>
<td>Farmer</td>
<td>11</td>
<td>37</td>
</tr>
<tr>
<td></td>
<td>Student</td>
<td>1</td>
<td>8</td>
</tr>
<tr>
<td>28 Nov-10 Dec 2004</td>
<td>Farmer</td>
<td>0.6</td>
<td>21</td>
</tr>
<tr>
<td></td>
<td>Student</td>
<td>0</td>
<td>12</td>
</tr>
<tr>
<td><strong>Infected snail density</strong></td>
<td></td>
<td><strong>Snails/m²</strong></td>
<td></td>
</tr>
<tr>
<td>20-30 Jun 2000</td>
<td></td>
<td>0.43</td>
<td>0.85</td>
</tr>
<tr>
<td>23 Jun-1 Jul 2001</td>
<td></td>
<td>0.12</td>
<td>0.38</td>
</tr>
<tr>
<td>15 June-5 July 2003</td>
<td></td>
<td>0.041</td>
<td>0.218</td>
</tr>
<tr>
<td><strong>Mouse bioassay</strong></td>
<td></td>
<td>Worms/mouse</td>
<td></td>
</tr>
<tr>
<td>30-31 July 2001</td>
<td></td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>30-31 Aug 2001</td>
<td></td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>26-27 Sept 2001</td>
<td></td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>30-31 Oct 2001</td>
<td></td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>30 Jun-1 July 2003</td>
<td></td>
<td>0</td>
<td>1</td>
</tr>
</tbody>
</table>
As has been recounted elsewhere, the fraction of simulations which meet the passing criteria is generally very small, 1% would be excellent, but not to be expected in the initial runs.\textsuperscript{4} The difficulty of achieving reasonable numbers of passes for subsequent analysis is generally regarded as the principal weakness of the approach. As a result, it is seldom feasible to use statistical analyses that rely on numbers of parameter vectors for passing simulations in excess of a few thousand, at least for complex models. In any case, there is a good reason to run the model on a fast computer with efficient simulation software. In the present case, after initial debugging, the model was run on a Dell Optiplex 755 (Intel Core 2 Duo E6850 3.00 GHz CPU) using Matlab\textsuperscript{\textregistered} version R2007a. Over the 2000-2004 calibration period, 306 passing simulations were obtained in 90,100 trials which ran overnight.

It is possible that the structure of the model, or the prior parameter ranges, are inconsistent with simultaneously meeting all calibration criteria. The result is no passing simulations, or perhaps very few. Hence, it is wise to save from each simulation run a vector indicating whether each of the components of the pass criterion was met (1) or not met (0). For a passing run, all elements of this vector are unity and, for failing runs, at least one entry will be zero. In the current example, a failing simulation produced the criterion vector keyed in order to Table 2:

\[ [1, 1, 0, 1, 1, 0, 1, 1, 1, 1, 1, 1, 1, 1, 1] \]

where the two failing sub-criteria are the mean egg excretion from farmers in October 2002 and the village average infected snail density June 2000. Calculating the correlation between each of the elements of this criterion vector is often instructive. A large negative correlation may point to either prior distributions with incompatible ranges or a structural deficiency in the model. In either case, it is an issue to be addressed before going into production runs.

The Posterior Parameter Space

An underlying reason for the complexity of the posterior parameter space is that the calibration criteria are generally dependent on functions of the basic model parameters rather than dependent on their values individually. For example, if an important rate in the model is dependent on the product of two or more parameters, as in equation 1.1 of Chapter 6, passes can occur over the entire range of each if the others assume a value yielding a product in the correct range. Similarly, for linear differential equation models of the form \( \frac{dx}{dt} = Ax \), where \( A \) is a constant matrix, the eigenvalues of the characteristic equation, which determine the dynamic behavior of the system, are complicated nonlinear functions of the elements of the \( A \) matrix. Hence, if the calibration criteria are sensitive to the eigenvalues of \( A \), the passing space of the basic parameters is geometrically complex and almost certain to be a small fraction of the volume of the space defined by their independent prior distributions. For these complex models CPU time per simulation is generally sufficiently costly to motivate the search for ways to increase the pass rate.

Recent work in our group aimed at understanding the role of the site-specific parameters in reducing or eliminating transmission suggests a strategy for increasing the pass rate that may be useful for many infectious disease analyses where the basic reproductive number \( R_0 \) is of central interest.\textsuperscript{15} The example of the eigenvalues of the \( A \) matrix discussed above motivates the speculation that the calibration criterion may depend on \( R_0 \). If so, even an approximate analytical expression of its dependence on model parameters might provide some insight into the complicated shape of the pass region and thereby suggest a means of increasing the pass rate. In the present case, a simplified model which leads to such an approximate \( R_0 \) is obtained by merging the three sub-populations in the village into one, assuming neither of the model’s nonlinearities is operational and that the time variable delay is fixed at some median value. This leads to the simplified model:

\[
\begin{align*}
\frac{dw}{dt} &= a_{11}a_{11}(t - \tau_\omega)z(t - \tau_\omega) - \mu_\omega w(t) \\
\frac{dz}{dt} &= a_{23}a_{23}(t - \tau_\omega)w(t - \tau_\omega) - \mu_z z(t)
\end{align*}
\]
where \( w(t) \) is the mean worm burden in the human population, \( z(t) \) the average infested snail density in the village environment, \( \tau_s \) the developmental delay of the parasite in humans, \( \tau_p \) the developmental delay of the parasite in snails and the \( \alpha(t - \tau) \) parameters are the time-variable exposure or environmental parameters each normalized by dividing by their annual maximum values.

The simplified model then allows the approximation \( R_p = P, P_b \) where \( P \) is a function of site-specific parameters and \( P_b \) of biological parameters. Woolhouse, in his very useful discussion of modelling and the control of schistosome transmission dynamics, used a similar approach and a two parameter description of \( R_m \) but focusing on the human-to-snail and snail-to-human portions of the transmission cycle.\(^{16}\) In the present case, the parameter aggregations are:

\[
P_s = \frac{S \gamma A \xi X \beta \delta g}{\mu_A} \quad \text{and} \quad P_b = \frac{\alpha \sigma \phi e^{-\gamma t} \mu_0}{\mu_w}
\]

There are four parameters included in \( P \) which are not listed in Table 1. \( \gamma \) and \( \xi \) relate to spatial inhomogeneities in the distribution of infected snails and of human water contact over the village irrigation system. As noted above, the parameters \( S \) and \( X \) are the annual maxima of the time variable water contact and uninfected snail density, respectively and arise from the normalization of those parameters in the simplified model. Because of this normalization and the fact that interventions affecting \( X(t) \) and the parameter \( \beta \) were implemented in the 2003-2004 period, the value of \( R \) computed from these relations for any given simulation run can be expected to overestimate the postcontrol or effective reproductive number, \( R_{ef} \). That is, it is likely that declining worm burdens post 2004 will occur for values of \( R \) based on the foregoing approximations above 1. This is essentially a scaling issue which does not alter the approach to achieving an increased pass rate. However, the value of 1 looses its traditional relevance. To underscore that fact, we redefine \( R' = P, P \) and below refer to \( R' \) rather than \( R_0 \) or \( R_{ef} \).

Since \( P_s \) and \( P_b \) are known functions of their respective parameter sets, each can be computed for each simulation run together with the value of \( R' \). The question is if this value, derived from the approximate model, is useful in discriminating between passes and fails in the full model. As noted earlier, an initial run over the full 2000-2004 calibration period yielded 306 passes from 90,100 simulations for a pass rate of 0.34%. Figure 1 shows the cumulative distribution functions of \( R', P_s \) and \( P_b \) separately for the 306 passes and 500 randomly selected fails. The passing and failing distributions differ for all three with all passing distributions shifted to the left. That is, parameters associated with simulations which meet the goodness-of-fit criteria combine to yield lower values of all three of these parameter aggregations, but \( R' \) and \( P_b \) are shifted to a much greater extent than \( P_s \). Indeed, the shift in \( R' \) is largely due to the influence of \( P_s \). None of the values of \( R' \) for the passing simulations exceeded a value of 4.0 where over 75% of the fails did so. That is, the calibration criterion are quite sensitive to \( R' \) in this particular case. It appears that for practical purposes \( R' < 4.0 \) is a necessary, but not sufficient, condition, for passing simulations. This suggests using the value of \( R' \) as a prefilter to increase the pass rate in subsequent runs. After each random draw of the parameters vector from the prior distribution, \( R' \) is calculated and the simulation conducted only if \( R' < 4.0 \). A full calibration run, from 2000-2004 utilizing the prefilter, led to 353 passes in 24,700 simulations which is about a 4-fold increase in the pass rate to 1.43%. However, there is obviously a great deal of complexity remaining to be explored in the remaining portion of the space where \( R' < 4.0 \) which accounts for the 24,200 failing simulations. Recall that the approximate model underlying the \( R' \) calculation does not take account of delay parameters \( \tau_s \) and \( \tau_p \), nor the parasite aggregation parameter \( k \) among other potential candidates potentially important to meeting the calibration criteria.

Although we have focused on the importance of parameter combinations, it is sometimes instructive to look at the univariate marginal distributions of individual parameters under passes and fails. Figure 2 shows these distributions for four parameters of the \( P_s \) aggregation, \( \alpha, \sigma, \rho \) and \( b \). (Definitions in Table 1.) For passing runs, these distributions show that somewhat higher values occur for \( \alpha \) and \( \sigma \), markedly lower values for \( \rho \) including a cutoff at about \( 6 \times 10^6 \) and that preferred values of \( b \) lie in the middle of the prior range. The most useful information in this
example relates to $\rho$ and the issue of parameter transportability. Recall that the original motivation for separating the biological parameters, $P_b$, from the site specific parameters, $P_s$, arose from the assumption that the biological parameters were transportable from one village to another in this region. With that in mind, the difference in the distribution of $P_b$ under passes and fails, shown in Figure 1b, suggests that we extend the prefilter idea to constrain $P_b$ to be less than $4 \times 10^{-5}$ in future applications of the model in the Xichang region. The univariate marginal distribution of $\rho$,
shown in Figure 2C, suggests further that the distribution of this single parameter be altered from its prior range of $1 \times 10^{-6}$ to $5 \times 10^{-4}$ to the range $1 \times 10^{-6}$ to about $6 \times 10^{-6}$.

There are clearly many other possible and potentially useful explorations of the posterior space, but the foregoing illustrate the challenge and its complexity. More generally, however, the $R_0$ strategy, both for both increasing the proportion of passing simulations and understanding the characteristics of the posterior parameter space, even in an approximate way, is a strategy of that might be of general utility in informing disease transmission modelling for management purposes. In such applications one is generally focused on the goal of assuring that the zero equilibrium state is stable and that the range of attraction of the zero state is as large as possible in the face of considerable uncertainty in many parameter values. Hence, preliminary studies using models to determine the nature and extent of calibration data sensitive to $R_0$ should be useful in setting data collection priorities for field surveys. In this context, the foregoing example leads to the speculation that there might be field data which is not cost effective to collect with respect to estimating $R_0$ as a prelude to studying management options.

Assuming that, in the end, the calibration runs result in at least several hundred passing parameter vectors, these vectors can be used directly to explore the effects of intervention strategies as recounted in Liang et al.14 In the present example, suppose that it was feasible to reduce the extent of the snail habitat by 20%, perhaps by concreting a portion of the village’s irrigation ditches. The issue is then to determine if this intervention would lead to a high probability of decreasing transmission with time. Suppose this intervention was to be implemented right after the end of the calibration runs in 2004 and the forecast was to run until 2009. The baseline run from 2004-09 would use the passing vectors unaltered. The habitat reduction run would allow the comparison by using the same vectors with the value of $A_h$ in each of the passing vectors decreased to 80% of its original value. Other control options could be studied in the same fashion.

**Bayesian Melding**

As noted earlier, the pass/fail procedure for incorporating field data into the mechanistic modelling framework is a special case of what is called Bayesian melding.8 We will first summarize the approach and then indicate how it can provide a more complete description of the posterior parameter space where the prior distributions are not uniform. The following exposition below draws heavily on an earlier description of the approach.17

In Bayesian melding, two priors on the output are compared. One prior is based on literature or field data as to what is reasonable output. These have been referred to above as the calibration criteria and are summarized in Table 2 for the example used herein. The other prior on the output is induced by running the model on prior information on input parameters, e.g., Table 1 in the example. These two output priors are “melded” together and inverted to the input parameter space, thereby refining the estimate of the input parameters. In detail and using Poole and Raftery’s terminology,8 let,

$$M : \theta \rightarrow \phi, \quad \theta \in \Theta \subseteq \mathbb{R}^n, \quad \phi \in \Phi \subseteq \mathbb{R}^p,$$

where $M$ is the deterministic model that relates a $n$-vector of input parameters, $\theta$, to a $p$-vector of outputs:

$$\phi = M(\theta).$$

From the current example, one component of $\theta$ might be, $h$, the number of eggs/worm pair/gm feces and a component of $\phi$ might be $w_{ik}(t)$, the worm burden in the $i$th village, $k$th risk group at time $t$. Define the posterior, joint model of inputs and outputs to be

$$\pi(\theta, \phi) \propto \begin{cases} p(\theta, M(\theta)) \text{ if } \phi = M(\theta) \\ 0 \text{ otherwise} \end{cases}$$

where $p(\theta, M(\theta))$ is the premodel joint distribution. One can think of $p(\theta, M(\theta))$ as containing the statistical information and relationships among the parameters and state variables before
considering how these inputs determine the outputs, \( \phi \). The postmodel joint distribution, \( \pi(\theta, \phi) \), however, only puts mass on input/output combinations consistent with the model, so it is a rescaled version of \( p(\theta, M(\theta)) \) with the mass of the impossible input/output combinations set to 0. In the schistosomiasis model, for instance, if the prior \( p(\theta, M(\theta)) \) allows positive probability on the combination \( h = a \) and \( w(\ell) = b \), whereas the structure of the model suggest such a combination could never exist (although each values are acceptable in different combinations), then the posterior puts mass 0 on \( (a, b) \). The interesting distribution with respect to the estimation of input parameters is the marginal posterior of the inputs, or,

\[
\pi(\theta) \propto p(\theta, M(\theta)).
\]

As discussed above, we will ignore the possibility of having sufficient data to construct meaningful likelihood on the outputs. Thus, one can typically write the prior as,

\[
p(\theta, \phi) \propto q_1(\theta) q_2(\phi)
\]

where \( q_1 \) and \( q_2 \) are the prior distributions for \( \theta \) and \( \phi \), respectively. If there is sufficient data on one of the outputs, so that a likelihood could be defined (say, the distribution of egg counts given the mean per village, group is negative binomial), say \( L(\phi) = p(D|\phi) \), where is the data on the outputs, now we have

\[
p(\theta, \phi) \propto q_1(\theta) q_2(\phi) L(\phi).
\]

As discussed by Poole and Raftery, adding the likelihood here removes the problem of Borel’s paradox, where the posterior distribution on the outputs is no longer dependent on the arbitrary parameterization of the model, \( M \). Though we have not defined a likelihood in our example, optimally and not surprisingly for many reasons, one would have sufficient data on the outputs in order to use something more akin to a typical Bayesian procedure.

Figure 2. Univariate marginal distributions of \( \alpha, \sigma, \rho \), and \( h \) for 306 passing and 500 failing simulations.
Because $\phi = M(\theta)$, the model, $M$ and the prior $q_1(\phi)$, induce another, independent prior on $\phi$, say $q_2(\phi)$. If one had a single prior on the outputs, there are existing Bayesian synthesis methods to define a posterior on inputs. Bayesian melding is a method for reconciling these two priors on the output. Specifically, Poole and Raftery suggest logarithmic pooling, or $\tilde{q}_2^{(\alpha)}(\phi) \propto q_1(\phi)^{\alpha} q_2(\phi)^{1-\alpha}$ resulting in a weighted (on the log-scale) average of the prior on the outputs induced by the prior on the inputs, $q_1(\phi)$ and the specified prior on the outputs, $q_2(\phi)$, resulting in a single prior, $\tilde{q}_2^{(\alpha)}(\phi)$. The choice of $\alpha$ is discussed in their paper. This prior on the outputs is then inverted to get a “melded” prior on the inputs, or $\tilde{q}_1^{(\alpha)}(\theta) = \tilde{q}_2^{(\alpha)}(\phi)$, for $\theta = M^{-1}(\phi)$. This leads to a posterior on the inputs, $\pi^{(\alpha)}(\theta) \propto \tilde{q}_1^{(\alpha)}(\theta)L(\theta)$, which itself leads to standard Markov Chain Monte Carlo approaches to estimating posterior distribution on the input parameters, $\theta$.

The distillation of the above technical discussion for our purposes, that is without a formal likelihood, is that Bayesian melding takes existing information on the input parameters in the form of a prior distribution, $q_1(\theta)$ and new information/expert opinion on the outputs, $q_2(\phi)$ and constructs a new, refined information on the parameter inputs, $\pi^{(\alpha)}(\theta)$. Note that the pass/no pass procedure simply places a uniform prior on a subspace of $\Theta^*$; $q_2(\phi) \propto I(\phi \in \Theta^*)$ where $\Theta^*$ is the acceptance region of $\Phi$. This implies that $\tilde{q}_2^{(\alpha)}(\phi) \propto q_1(\phi)$ if $q_2(\phi) > 0$ and $= 0$ otherwise. Thus, the choice of $\alpha$ does not affect the pass/no pass procedure and a random sample from the implied $\pi(\theta, \phi) \propto q_1(\theta)\tilde{q}_2^{(\alpha)}(\phi)$ if $\phi = M(\theta), 0$ otherwise, can be generated as follows: 1) generate a random $\theta$ from $q_1(\theta)$, 2) calculate $\phi = M(\theta)$, 3) keep $(\theta, \phi)$ if $q_2(\phi) > 0$ (a pass).

As suggested, most procedures that attempt to estimate the marginal posterior, $\pi^{(\alpha)}(\theta)$, do so, not by finding the distribution directly, but by methods which generate random samples from the underlying distribution of interest, one example being the pass/no pass method as discussed above. The result is repeated random draws from $\pi^{(\alpha)}(\theta)$. So, defining the posterior distribution is an exercise in multivariate density estimation. Tree-based density estimation is a nonparametric multivariate density estimation technique particularly well-suited to estimation of high-dimensional data. In addition, newly developed nonlinear principal components methods permit the discovery of strong nonlinear relationships among the parameters that conventional principal components methods would not discover. Depending on the model and the prior information on the inputs and outputs, our experience suggest that strong nonlinear relationships (sometimes deterministic nonlinear relationships suggesting that an optimal set of parameters based on $\pi^{(\alpha)}(\theta)$ is not identifiable) among the posterior distribution of the input parameters is common in these types of disease models. Thus, examination of these posterior distributions can help to reparameterize the model in ways that avoid identifiability problems.

Conclusion

The fundamental fact underlying the approach to parameter estimation presented in this chapter is that there are almost certain to be many parameter combinations that are equally consistent with the local data available to calibrate the model. Moreover, when used in a forecasting mode, these combinations can lead to different qualitative performance of the model. For example, the effective reproductive rate for some may be greater than one and less than one for others. So, the search for effective control strategies seeks the feasible control option that will maximize the probability of declining transmission over time.

It is in the context of future forecasting that the full Bayesian melding concept becomes attractive. If likelihoods can be estimated for the prior distributions, then the posterior distribution can support more informed and hopefully more reliable, inference on the outcomes of the various intervention options. This point further illustrates that, if the objective is to design effective local control strategies, there is a very intimate relationship between laboratory-generated data, field data and the modelling process. It can be argued that low esteem in which modelling is held by many
epidemiologists and infectious disease experts arises from the rarity of data-informed modelling studies of the sort we advocate. Conversely, modelers can argue that much of the available laboratory and field data is of marginal relevance to control programs. There is ample opportunity for progress at the intersection.

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References

Modelling Malaria Population Structure and Its Implications for Control

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Abstract

Mathematical models of malaria transmission have been used to inform the design of malaria control programs since the mid 20th century, and many of these models have provided useful insights into the complexity of the disease. Among developing countries, however and particularly in sub-Saharan Africa, malaria remains a major cause of morbidity and mortality. One of the main difficulties in controlling the most virulent human malaria parasite, *Plasmodium falciparum*, is its genetic diversity, which confounds attempts to design an effective vaccine. The population structure of *P. falciparum* remains poorly understood but plays a key role in determining epidemiological patterns of disease and the development of immunity. We discuss the seminal model of malaria transmission developed by Ross and MacDonald, and the modifications that have been made since to include more realism. We show that age profiles of disease and serological data support a theoretical model in which the parasite population is diverse and structured into several antigenic types and highlight the implications of this structure for controlling malaria. Lastly, we discuss the current sequence data on parasite antigen genes that are important for the acquisition of immunity, and the results of a new analysis of *P. falciparum* population structure at the genomic level.

Introduction

Malaria remains a major global public health problem, causing about 500 million cases and over a million deaths each year.1 Although the disease has been eliminated in many parts of the developed world due to intense chloroquine use and mosquito control programs during the 1950s and 1960s, the logistical difficulties associated with the implementation of these programs in many developing countries, as well as the emergence of resistance to DDT in mosquitoes and to chloroquine in the parasite, led to a discontinuation of the WHO’s Global Malaria Eradication Program in 1972. The breakdown of malaria control led to a resurgence of malaria cases worldwide and severe epidemics in areas such as Madagascar and Sri Lanka (reviewed recently in ref. 2). Renewed interest in controlling the disease in the past decade has led to significant funding for basic research and control efforts, however. The design of appropriate control programs relies on an understanding of the complex dynamical interactions between vector ecology, the lifecycle of malaria parasite and the pathology of the disease in humans, which remain poorly understood. The complexity of these interactions leads to nonlinear relationships between important epidemiological parameters...
and the use of mathematical models can provide an important tool for understanding the potential effects of control programs on malaria prevalence and disease.

Ronald Ross formulated the seminal model of malaria transmission between humans and mosquitoes in 1911, and it was subsequently modified by MacDonald in the 1950’s. Based on a simple framework which tracks the proportions of human and mosquito populations that are susceptible and infected, these models emphasized the relative importance of different aspects of the transmission system in determining malaria prevalence. They were used to successfully inform the design of control programs in many countries, by providing support for the intense efforts to kill adult mosquitoes by the global eradication program, for example. The Ross-Macdonald model is deficient, however, in capturing certain essential features of malaria transmission, such as naturally acquired immunity. Numerous models have been developed to add “realism” to the original framework, but the fact remains that we still lack a thorough understanding of the relationship between infection and immunity in the host and its relation to the biology of the malaria parasite. One of the major reasons for this is the genetic and phenotypic complexity of the parasite itself. Figure 1 illustrates the lifecycle of the most virulent human malaria parasite, *Plasmodium falciparum*, highlighting its variety of morphological forms. Disease symptoms in humans are associated with the blood stage of the parasite, when rounds of asexual replication occur within red blood cells leading to recurring fever, anemia and occasionally coma and death. It remains unclear why some

![Figure 1. The lifecycle of *Plasmodium falciparum*.](image)

**Figure 1.** The lifecycle of *Plasmodium falciparum*. A) Sporozoites are injected into the human host when an adult female mosquito takes a blood meal. They travel rapidly to the liver (B), where they invade liver cells and undergo rounds of asexual reproduction and produce hundreds of merozoites. Upon release into the bloodstream (C), merozoites invade red blood cells and begin cycles of replication, lysis of the cell and repeated invasion (D). It is during these rounds of replication that disease symptoms manifest themselves. Some merozoites differentiate into male and female gametocytes (E), the sexual forms of the parasite responsible for transmission. Mosquitoes take up gametocytes upon feeding (F). Within the mosquito gut gametocytes unite and form zygotes (G), mature, eventually cross the gut barrier and travel as newly formed sporozoites (H) to the salivary glands for transmission back to the human host.
individuals suffer these complications and others don’t. Although a range of host factors such as genetic background and immune status are undoubtedly important in determining disease outcome, heterogeneity within the parasite population also appears to play a major role.

By the late 19th century, distinct species of the malaria parasite *Plasmodium*, as well as different ‘broods’ or ‘colonies’ within these species, had been identified as producing different kinds of fever and other variable disease phenotypes such as quinine resistance and virulence (reviewed in ref. 3). This heterogeneity within the parasite population was also apparent in age profiles of infection and disease, which suggested that naturally acquired immunity builds up gradually and requires exposure to many different parasites. Furthermore, serological studies have shown that immunological responses to parasite isolates seem to be primarily isolate-specific. We will discuss this evidence that the targets of immunity are polymorphic, focusing on mathematical models exploring the mechanisms behind and the implications of a parasite population made up of independently transmitted parasite types. The epidemiological parameters of transmission are significantly affected by this assumption and this must be taken into account when designing control problems.

Sequencing projects have now given us insight into the structure of the parasite population at the genomic level. Phylogenetic analysis of these sequences is often impossible for malaria antigens, however, due to high rates of homologous and nonhomologous recombination. New techniques for understanding patterns of genetic diversity are being developed that do not rely on multiple alignments or particular evolutionary models. We discuss results from these approaches, which support the notion that there may be restricted antigenic types that could be targeted by vaccination. A solid link between genetic and phenotypic structure is needed to understand how this gene sequence data relates to expression in the host and disease phenotype, however. Mathematical models will continue to play an important role in uncovering this link and in our understanding of the relationship between the population structure of *Plasmodium falciparum* and malaria epidemiology and control.

Adding Realism to the Basic Framework of the Ross-MacDonald Models

The seminal models of malaria transmission formulated by Ross and MacDonald have remained the framework for the vast majority of theoretical models describing malaria epidemiology since they were developed during the first half of the 20th century. Although these models capture many defining features of malaria epidemiology, there are several important assumptions affecting their utility in guiding public health programs. First, mosquito biting rates in the model are assumed to be homogeneous, despite evidence that mosquitoes have systematic preferences for certain hosts. Second, it is assumed that humans do not acquire any immunity to the parasite, which we will show is incompatible with both serological evidence and age profiles of disease and infection. Third, human, mosquito and parasite populations are assumed to be homogeneous. This chapter will review all these assumptions, but will focus on recent attempts to understand parasite population structure and the implications of these studies for estimates of the basic reproduction number and the design of vaccines.

Heterogeneity in Biting Rates and Susceptibility

Many models have been developed which add various levels of complexity to the basic Ross-MacDonald framework. One of the first issues to be addressed was the assumption of homogeneous biting rates by mosquitoes. Studies measuring the proportion of “freshly fed” mosquitoes containing human blood have shown that host selection by mosquitoes is not random, as assumed in the original model. In fact, mosquitoes appear to actively select hosts harboring gametocytes, the stage of the parasite lifecycle responsible for transmission. By dividing the human population into subpopulations with different biting rates, Dye and Hasibeder showed that the assumption of uniform biting leads to an underestimation of $R_0$, the basic reproductive rate of the disease, even without assuming that mosquitoes preferentially select infected hosts. Heterogeneity in biting rates also leads to changes in the estimates for overall prevalence of infection, which under assumptions of homogeneous biting rates is overestimated in highly endemic areas and underestimated in low transmission regions. It also causes large differences between subpopulations of hosts with respect
to their contribution to transmission,\textsuperscript{10} such that control programs targeting hosts that are bitten frequently will be most successful. Furthermore, current estimates of how heterogeneous biting rates relate to the prevalence of infection in the field show that the interaction between the two may be variable and this heterogeneity must also be taken into account when modelling the effects of control programs on malaria transmission.\textsuperscript{11,12}

Extensions of Dye and Hasibeder model, in which host populations were also subdivided with respect to susceptibility and duration of infection, explored the effects of genetic heterogeneity in the host population on malaria transmission. Variability between hosts due to innate genetic differences in blood group or sickle cell type, for example, was modeled by assuming that two types of host exist with high or low susceptibility, long or short duration of infection. Calculation of $R_0$ then required the inclusion of the variance in these parameters and their covariance with respect to each other (reviewed in ref. 17).\textsuperscript{13} In this case $R_0$ may be increased or decreased, depending on whether the covariance between parameters is positive or negative. The prevalence of infection will always be increased, however, when there is significant variance in host susceptibility or the length of infection. Again, the inclusion of heterogeneity in these models highlighted the often counterintuitive effects of changes to a highly complex system and the importance of mathematical models in our understanding of these nonlinearities.

**Superinfection**

It has long been recognized that the population structure of the malaria parasite may be better represented as multiple “broods” than a single entity and that different “broods” may coinfect one host (reviewed in ref. 3). MacDonald criticized the original Ross model in which hosts cannot be reinfected until they have cleared current infections and suggested a model of superinfection in which “broods” co-infected humans independently of each other. Dietz\textsuperscript{14} incorporated superinfection into a model of malaria transmission by dividing the host population into different classes depending on the number of infections they harbor. He explored a range of assumptions about the density-dependent constraints on multiple infections, ranging from models in which infections occur completely independently of each other, to those in which only one infection is possible. With complete density-dependence (i.e., the original model) the prevalence predicted was much lower than in models that include superinfection. As in previous studies, models allowing superinfection showed that for higher $R_0$ values, changes in transmission due to control programs have little effect on prevalence. These models also highlighted the importance of the type of density-dependence occurring both in the human host and in the mosquito, which both remain poorly understood, for the estimation of $R_0$.

**Incorporating Immunity**

One of the most obvious problems with the Ross-MacDonald model is the absence of acquired immunity in humans. The relationship between infection and immunity to the malaria parasite is complex and poorly understood, although its inclusion in mathematical models is vital for an understanding of the potential effects of vaccination. Models by Aron and May\textsuperscript{15-17} incorporated observations of semi-immune adults living in endemic areas by adding an immune class of people that were infected but asymptomatic. A key assumption of this framework is that individuals are removed from the immune class and became susceptible again after some period of time in the absence of immunological boosting by new infections. Thus, the rate of acquisition of new infections determines the duration of immunity. Analysis of this model showed that in endemic regions, the suppression of transmission through drug treatment for example, may sometimes increase the prevalence of disease. Although several more recent studies have attempted to incorporate immunity into the Ross-MacDonald framework,\textsuperscript{18} few have generated qualitatively different results from Aron and May\textsuperscript{16} and until we have a better understanding of the mechanisms underlying the development of immunity, their validity remains un-testable.

Understanding how naturally acquired immunity to the malaria parasite develops remains a major challenge and one that is critical to vaccine design. Figures 2A-C show the age profiles of malaria infection and mild and severe disease in an endemic region (from Snow and Marsh,
Figure 2. Age profiles of infection and disease in a malaria-endemic region (from Snow and Marsh, 1990, Gupta et al, 1994). A) The prevalence of malaria infection, defined as the percentage of individuals of different age groups harboring blood-stage parasites, measured by microscopy. B) The incidence of mild malaria episodes defined as uncomplicated infections presenting at the out-patient clinic in West Africa. C) The incidence of severe malaria disease in the same region: severe malarial anemia (SMA) and cerebral malaria (CM).
The prevalence of infection (Fig. 2A) is extremely high among infants and young children and drops gradually in adolescence and into adulthood. Clinical disease shows a different pattern (Fig. 2B,C) and can be divided into several distinct syndromes from mild malaria to severe cerebral malaria to severe noncerebral malaria. Mild episodes (Fig. 2B) are suppressed among infants under one year, peak at approximately 1-2 years and fall gradually with age into adolescence. Severe noncerebral malaria shows a similar peak among 1-2 year olds, but falls much more rapidly among young children (Fig. 2C). Cerebral malaria also declines rapidly with age, but it shows a delayed peak in incidence (Fig. 2C) compared to severe noncerebral disease. In spite of the number of studies of parasite prevalence and disease, our understanding of naturally acquired immunity remains incomplete and recent mathematical models fitted to data from Tanzania and The Gambia have shown that the generation of immunity is likely to require at least two different mechanisms, both suppression of disease and parasite clearance, occurring on different time scales. In addition, maternal antibodies clearly play an important role in protection against disease in the first few months of life, however there may be differences in the period of postnatal protection against cerebral and noncerebral malaria. Gupta et al also identify a “strain-transcending” immunity as being important for protection against severe (noncerebral) disease after the first few infections. More basic research is required to untangle the contribution of different mechanisms of immunity to patterns of infection and disease.

Modelling the Effects of Parasite Population Structure

The observations of the gradual decline in infection and disease and the heterogeneity of disease phenotypes led to the suggestion that immunity may be directed against polymorphic parasite antigens. Under this hypothesis, individuals growing up in endemic regions require exposure to many different parasites in order to build up a repertoire of antibodies to different antigenic types. This idea was supported by a number of serological experiments showing that patients generate agglutinating antibodies specific to their own isolate during infection, but not to isolates from other patients. Furthermore, Gupta et al showed that exposure to particular isolates among children

![Figure 3. Age profiles of serological responses to different parasite isolates among children from Papua New Guinea (from Gupta et al, 1994). The proportion of people of different age groups with agglutinating antibody responses to each of four wild isolates from PNG and one lab isolate (HB3).](image-url)
from Papua New Guinea rose gradually with age compared to the rapid rise in exposure to any *P. falciparum* isolate (Fig. 3). In studies of serological responses of Kenyan children, it was shown that the frequency of recognition of particular parasite isolates was associated with both young host age and a limited antibody repertoire, suggesting that there may also be a hierarchy of infection with different antigenic types, possibly caused by their different frequencies in the parasite population.

**Implications of Antigenic Diversity for Control**

An antigenically diverse parasite population has important implications for our understanding of the basic epidemiological parameters of malaria transmission. For example, Gatton and Cheng used a stochastic simulation model to demonstrate that the time taken to develop immunity was directly proportional to parasite antigenic diversity. A model exploring the interaction between immunity to different parasite phenotypes and the rate of evolution of the parasite showed that cross-reactivity in the parental genotypes would delay the appearance of and increase the chances of extinction of, a resulting recombinant form. An extension of this framework to include population effects demonstrated that the presence of increasing numbers of independent parasite genotypes in the parasite population may significantly prolong the persistence of each one in the host population.

Assuming antibodies generated to one antigenic type will not protect a host from infection with a different antigenic type effectively removes competition for hosts between strains. Thus, the parasite population is divided into distinct antigenic systems transmitted essentially independently. Models by Gupta et al. explored the implications of this removal of competition between parasite strains. In many theoretical models exploring the relationship between virulence and *R₀*, competition between parasites with varying virulence and *R₀* values will select for a pathogen population characterized by an intermediate level of virulence that maximizes *R₀*. Under this framework, competitive exclusion of pathogens outside the range of optimum transmission will occur. In a population composed of discrete antigenic types, however, the lack of shared antigenic epitopes between strains reduces direct competition for hosts between different types, facilitating the coexistence of a wider range of *R₀* values and levels of virulence. For falciparum malaria, the separation of severe and mild isolates into distinct antigenic systems allows for the coexistence of two different sets of maximized *R₀* values. These models predicted that cerebral malaria, with its delayed peak in prevalence, may be caused by a distinct set of circulating strains, whereas severe malarial anemia represents a rare outcome of infection with any number of “mild” strains.

Another important implication of a discretely structured population relates to the calculation of epidemiological parameters from *R₀*. For simple pathogen systems like measles, the basic reproductive rate of a pathogen is inversely proportional to the average age of first infection:

$$ R₀ \propto \frac{L}{A} $$

where *L* is the average lifespan of the host and *A* is the average age of first age of infection. Estimates based on this relationship for malaria yield an extremely high *R₀* value, since infants in malaria-endemic regions usually experience their first malaria infection before they are one year-old. In a system composed of discrete antigenic types, however, *R₀* for the whole parasite population is calculated as the sum of the *R₀* values of individual strains. Thus, rather than assuming that immunity to the malaria parasite is short-lived, Gupta et al. showed that long-lived strain-specific immunity can lead to a young average age of first exposure to any of *n* circulating strains, *A*:

$$ A = \frac{1}{\left(\frac{1}{n} \sum_{i=1}^{n} R_{0i}\right)} $$

Here, relatively low *R₀* values for each antigenic Type *i* contribute additively to the probability of becoming infected with any antigenic type. In this case, immunity to each strain is assumed to be life-long. Data from Papua New Guinea showing a gradual rise in antibodies to particular parasite isolates compared to antibodies to any isolate support this hypothesis, as discussed in the
previous section. The recalculation of $R_0$ as a cumulative value also has implications for control programs. For most pathogens the proportion of the population that must be vaccinated ($p$) in order to eradicate a disease, for example, can be related to $R_0$ by the following equation:

$$
(7)
$$

Hence, to eradicate a disease with a very high reproductive rate, nearly everyone in a population must be vaccinated. Assuming a number of discrete antigenic types each with a relatively low $R_{0i}$, however, the targeting of particular types in a vaccine, those associated with cerebral malaria for example, becomes a more realistic possibility.

Recent attempts to estimate $R_0$ in the field have made painstaking efforts to parameterize a modification of the Ross-MacDonald model using data from around the world. Their models predicted the range of $R_0$ values from over a hundred African countries to vary between less than one and close to ten thousand, several orders of magnitude greater than previous estimates. They assume a system with infinite broods and independent clearance however (see ref. 18 for details) and make no attempt to model the effects of population structure on immunity. Furthermore, they include immunity only as a reduction in infectivity of both humans and mosquitoes (due to transmission-blocking and liver-stage immunity, respectively) that varies as a function of the entomological inoculation rate. These simplifications largely ignore the important aspects of population structure described above and lead to the extreme values of $R_0$ they calculate.

**Emergence and Maintenance of Strain Structure**

In order for distinct antigenic strains of the malaria parasite to persist in spite of high rates of recombination, however, some mechanism for preventing the breakdown of associations between antigenic epitopes must exist. Mathematical models have shown how discrete antigenic types may be structured and maintained in the presence of recombination, however, through immune selection. These models assume that when infected individuals become immune, they may gain cross-immunity to parasites that share antigenic variants with the infecting isolate. Hosts infected with a particular parasite gain ‘exposure’ to any other parasites expressing the same epitopes, depending on $\gamma$, the strength of cross-immunity. In the simplest hypothetical case, for example, each antigenic type is defined by two loci, each with two possible allelic variants, giving four possible types (see Box 1). This model is defined by a system of ordinary differential equations the changing proportions of the population that are either infected, immune, or exposed to different antigenic variants are modelled for different levels of cross-immunity. When cross-immunity is very weak, infected hosts will not gain protection to different parasites even if they share antigenic epitopes and all antigenic types will coexist at a prevalence determined by their intrinsic transmissibility. This situation could arise due to epitope shedding or weak immune responses, for example. At intermediate levels of cross-immunity, unstable population structure emerges, displaying cyclical or chaotic patterns of dominance of different antigenic types. When cross-immunity is strong, however, immunity to one antigenic type will confer substantial protection against isolates that share antigenic epitopes, leading to the dominance of a set of genotypes with non-overlapping antigenic repertoires (which will not be competing for susceptible hosts). Discrete, non-overlapping antigenic repertoires will persist over time in spite of recombination between parasites, because immune selection will suppress the prevalence of recombinant combinations that share epitopes with dominant antigenic types.

Testing this hypothesis remains difficult while the immuno-dominant parasite epitopes responsible for acquired immunity remain unknown, however. The serological studies described above, showing isolate-specific immune responses in patients from endemic regions, are suggestive of strong antigenic structuring. However, until a system of defining antigenic repertoires based on known epitopes is developed and we understand more about expression patterns within the host, it remains an untested hypothesis.
Despite these difficulties, now that three full genome sequences of \textit{Plasmodium falciparum} lab strains are available and sequencing technology continues to become cheaper and easier, we can now sequence-based analysis of population structure of malaria parasite antigens. Pathogen strains are defined by two loci (shown as either the circle or the triangle in the figures), each with two alleles (yellow or white, in this case). Once infected with a particular strain, hosts gained partial immunity to any other strains with shared antigenic determinants. The strength of cross-immunity was determined by the parameter gamma ($\gamma$) and all strains were assumed to have the same transmissibility, $\beta$. For each strain, the host population consisted of three overlapping compartments: the proportion that were infectious to other hosts, $x_n$, the proportion that were completely immune, $z_i$, and the proportion partially and completely immune, $w_i$. Immunity was gained immediately upon infection and infectiousness was lost at a rate of $\sigma$, such that those who were infected ($x$) were a subset of those who were completely immune ($z$), who were in turn a subset of those who were partially and completely immune ($w$). It was assumed that the duration of infectiousness ($1/\sigma$) was short compared to the average host life-span ($1/\mu$) and that immunity was life-long. The effect of recombination was not explicitly included in the model, however all possible strains were present from the start in order to investigate the competitive interactions between them. The figures show the effects of immune selection assuming (A) weak, (B) medium and (C) strong cross-immunity ($\gamma$). At low levels of cross-immunity, all strains coexist at the same prevalence. When cross-immunity is very strong, the pathogen population is dominated by two strains with non-overlapping antigenic repertoires, since these will not be competing for hosts and the remaining strains are suppressed. At intermediate levels of cross-immunity, cyclical behavior is observed and pairs of strains with non-overlapping antigenic repertoires oscillate with respect to each other. Chaotic behavior is also observed for some parameter regions. A color version of this image is available at www.landesbioscience.com/curie.

**Sequence-Based Analysis of Population Structure of Malaria Parasite Antigens**

Despite these difficulties, now that three full genome sequences of \textit{Plasmodium falciparum} lab strains are available and sequencing technology continues to become cheaper and easier, we...
have the opportunity to start exploring the genes underlying parasite population structure. One of the best candidate targets of the isolate-specific agglutinating antibodies is the highly diverse, polymorphic family of proteins *Plasmodium falciparum* erythrocyte protein 1 (PFEMP1), expressed on the surface of infected red blood cells.\(^{37}\) PFEMP1 is implicated as a virulence factor due to its role in the cytoadherence of infected cells to the endothelial lining, mediated by host cell receptors such as CD36 and ICAM1.\(^{38}\) The *P. falciparum* genome contains approximately 60 *var* genes encoding PFEMP1. These undergo clonal antigenic variation during infection; different variants in the parasite *var* repertoire are expressed sequentially in a mutually exclusive manner,\(^{39}\) prolonging infection and increasing the parasite’s chances of transmission. *Var* genes also exhibit extremely high rates of recombination, not only between parasites during the sexual stages in the mosquito but also in the human host between different genes within the same parasite.\(^{40}\) This antigenic diversity and complexity of expression during infection complicates our understanding of the structuring of these loci, both within individual parasite genomes and across the parasite population.

Despite the incredible diversity of *var* genes across the parasite population, some constrained structure is apparent. Serological studies have shown that PFEMP1 proteins from parasite isolates infecting young children tend to be recognized by most adults in an endemic region and that by the time children reach the age of about fifteen in areas of high transmission they have gained exposure to most PFEMP1 variants circulating in the local parasite population.\(^{3}\) Following the sequencing of the *Plasmodium falciparum* genome, analysis of *var* repertoires has also hinted at limits to their diversity. Attempts to classify *var* genes into meaningful groups based on fully sequenced lab strains have led to the identification of approximately 6 different *var* groups, differentiated based on protein domain structure, different upstream promoters, the direction of transcription and chromosomal position.\(^{41-44}\) This system of classification identifies three major groups (A, B and C) and two minor groups (B/A and B/C). Different groups appear to show marked differences in their cytoadherent properties, for example groups B and C bind to the host receptor CD36, whereas group A does not.\(^{45}\) They also have different immunological properties, with group A *var* genes being better recognized by antibodies from sera of children growing up in malaria-endemic regions.\(^{46}\) The diversity of the sequences themselves has made sequencing whole *var* repertoires from clinical isolates extremely difficult, however. A sequence-based approach to *var* gene classification has been developed which uses short sequence fragments.\(^{47}\) Regions of homology at the beginning and end of these tags allow for the design of primers that can amplify most *var* genes. The sequence tags are grouped according to a series of semi-conserved regions and the number of cysteine residues present between them. A study by Bull et al\(^{48}\) analyzing these sequence fragments from wild isolates showed that *var* genes belonging to the 6 different groups were represented in the same proportions within different parasite genomes, but their expression patterns varied considerably in different hosts. Figure 4 illustrates this finding, showing the genomic structure of *var* gene repertoires from 12 wild Kenyan isolates and their profiles of *var* expression. The expression of particular groups of *vars* was also found to be associated with different disease phenotypes, such as rosetting and the immune status of the host.\(^{48}\) The groups identified using these sequence-based techniques are therefore linked with phenotypic characteristics and represent a useful system of classification.

Understanding the evolutionary relationships between *var* genes and *var* gene groups remains a major challenge, however. Phylogenetic analysis of the relationship between *var* sequences is confounded by the high rates of homologous and nonhomologous recombination; most phylogenetic techniques rely on an initial alignment of sequences, which is not possible for most *var* genes. Bockhorst et al\(^{49}\) have overcome this problem for a small group of conserved *var* genes associated with malaria in pregnant women, by identifying segments of correlated nucleotides within sequences and using whole segments in a phylogenetic analysis and generating “population trees”, however this technique cannot be applied to the vast majority of *vars*. Recently, a technique for visualizing the relationships between *var* sequence fragments has been developed by Bull et al\(^{50}\) which is not reliant on an alignment. Sequence tags are represented as nodes in a network, with edges between nodes representing exact sequence matches at variable regions termed “position
specific polymorphic blocks” (PSPBs). These regions show considerable mosaicism, such that otherwise unrelated sequences may share one or more PSPB. Figure 5 illustrates an example of some of these sequence fragments, the network they form and the results of an analysis of over 1000 \textit{var} genes from clinical and lab isolates, as well as several \textit{P. reichenowi} (the primate malaria parasite) homologues. The large network is coloured according to the 6 groups previously identified. The network has two distinct lobes with loose links between them and these lobes correspond well to different groups of \textit{var} genes based on both classification systems described above. All group 1 sequences fall within the smaller lobe, for example, whereas group 4 sequences are found in the larger lobe. Furthermore, group A genes (using the Lavstsen et al\cite{44} system) are all found within the smaller lobe. Thus, the separation of groups seems to represent a recombination hierarchy, with certain types of \textit{var} gene recombining with each other more frequently than others. Interestingly, \textit{P. reichenowi} genes fall within the network, indicating that many of these polymorphic regions are relatively ancient. The diversity of the malaria parasite is therefore generated by recombination, which shuffles variable regions between \textit{var} genes.

A major field of research in the next few years will be the search for antigenic epitopes within \textit{var} gene sequences and the investigation of how the expression of different \textit{var} epitopes relates to parasite phenotype and disease patterns in the host. Without this link between genotype and phenotype, it will be difficult to directly relate the sequencing projects discussed above with the mathematical models of population structure presented in previous sections.

Conclusion

The use of mathematical models in the design of malaria control programmes dates back to the beginning of the 20th century. The Ross-MacDonald model of malaria transmission provided the basis for most modelling efforts since then and gave valuable insights into the importance of
epidemiological variables such as the mosquito vector lifespan for determining the spread of the disease. Many variations on this framework have been developed, however we still lack a thorough understanding of the acquisition of immunity to *Plasmodium falciparum* and the relationship between infection, disease and immunity in malaria-endemic populations.

The models discussed in this chapter have aimed to understand mechanisms behind malaria epidemiology and dynamics. Many studies have also explored models focusing on specific interventions, however, comparing the effects of different types of control programs and the practicalities involved in their implementation. For example, Saul developed a simulation model of vaccination in which he examined the effects of different types of vaccines—antigametocyte or antisporeozoite/asexual—and the expected changes in malaria prevalence and disease with variable levels of vaccine coverage and efficacy. He showed that antisporeozoite or asexual vaccines were more effective at lower levels of coverage and that in combination with other methods of control, vaccination could be a feasible strategy. His recent models (for example ref. 52) take into account the antibody levels needed to maintain herd immunity towards a specific mosquito-stage transmission-blocking
vaccine. Other investigations of the relative merits of different types of vaccines and vaccine strategies, as well as the effects of only partially protective or "leaky" vaccines, have been explored using age-structure models.\textsuperscript{53-55} These models included one set of equations for vaccinated hosts and one for unvaccinated hosts and highlighted the importance for successful control programs of vaccine efficacy and duration and boosting of immunity by natural infections. Another more specific model reproduced the conditions of a particular country, Sri Lanka and modeled the effect of transmission-blocking vaccines under epidemic conditions.\textsuperscript{56}

We have also not dealt with mathematical models of the spread of drug resistance through the malaria parasite population. This aspect of parasite population structure has been modeled extensively,\textsuperscript{57-61} and shows complex relationships between the number of genetic loci involved, the level and type of drug use and the level of malaria transmission in the region. These models either focus on the spread of resistant alleles through the parasite population (i.e., population genetic models) or they incorporate infected hosts and use an epidemiological framework (reviewed in ref. 60). Despite the differences in modelling approaches, most of these studies conclude that the key to preventing rapid emergence and spread of resistance is minimizing drug use and using them in combination with transmission-blocking strategies such as bednets.

Even careful use of malaria drugs will eventually result in the emergence of drug resistance however, making the development of a vaccine a priority. We have shown that the population structure of the malaria parasite at immunodominant loci such as the \textit{var} genes has important implications for the basic epidemiology of malaria and for the design of control programs like vaccination. A major goal will be to link \textit{var} gene repertoire structure and expression patterns with serological and disease profiles in endemic regions. This will enable the re-assessment and validation of the mathematical models discussed in this chapter, providing the opportunity to design appropriate control programs in the future.

References


Mathematical Modelling of the Epidemiology of Tuberculosis

Peter J. White* and Geoff P. Garnett

Abstract

Despite the infectious agent that causes tuberculosis having been discovered in 1882, many aspects of the natural history and transmission dynamics of TB are still not fully understood. This is reflected in differences in the structures of mathematical models of TB, which in turn produce differences in the predicted impacts of interventions. Gaining a greater understanding of TB transmission dynamics requires further empirical laboratory and field work, mathematical modelling and interaction between them. Modelling can be used to quantify uncertainty due to different gaps in our knowledge to help identify research priorities. Fortunately, the present moment is an exciting time for TB epidemiology, with rapid progress being made in applying new mathematical modelling techniques, new tools for TB diagnosis and genetic analysis and a growing interest in developing more-effective public-health interventions.

Introduction

Despite the availability of effective treatment tuberculosis remains a major global cause of morbidity and mortality, with around one-third of the world’s population believed to be infected. It caused an estimated 1.7 million deaths and 8.9 million new cases of infection in 2004.1 The highest incidence of disease is in sub-Saharan Africa, in part due to interactions with HIV,2-4 which has fuelled dramatic rises in incidence of the disease in many countries. (Globally, TB is the proximate cause of many HIV-related deaths, particularly in Africa.3) Even in many countries where its overall incidence is low TB remains a problem: there has been an outbreak in the New York in the recent past5,6 and incidence is currently rising in the UK.7

TB Natural History

TB’s natural history is summarized briefly here; it is discussed in more detail below with respect to its representation in mathematical models.

Tuberculosis is caused by the slowly-replicating bacterium Mycobacterium tuberculosis (M tuberculosis, Mtb). Mtb is in the same genus as M bovis, the cause of the disease of cattle bovine TB, which humans can acquire from contaminated milk; it can also infect other species such as badgers (Meles meles). Other members of the genus, such as M avis, can cause human disease, particularly in the immunocompromised host.

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Person-to-person transmission of \textit{Mtb} is via the respiratory route, which can occur both through close contact between persons and through infectious bacilli being carried throughout buildings by air currents (which makes ventilation an important preventative measure;\textsuperscript{8} this can be supplemented by ultraviolet irradiation).

It is important to distinguish between TB \textit{infection} and TB \textit{disease}. Most people infected with TB never develop disease during their lifetime, but remain latently infected—and uninfected. The progression of an infection to disease is more likely in the years shortly after infection, known as primary disease. Thereafter, there is a much lower rate of progression to disease, called endogenous reactivation, resulting in a lifetime risk of roughly 10\% of developing TB disease (in the absence of HIV—see below). Endogenous reactivation can take decades to occur\textsuperscript{9,10} and is the main cause of disease in older people in settings where the prevalence of TB disease is very low. However, those with latent TB infection are also subject to exogenous reinfection, where they acquire another TB infection; this can cause rapid progression and can be an important cause of incident disease in settings with a high prevalence of TB disease and consequently a high force of infection.\textsuperscript{11}

Active TB disease can affect other parts of the body besides the lungs, e.g., causing Pott’s disease in the spine and tuberculosis meningitis in the brain, causing morbidity and mortality. However, only those with \textit{pulmonary} (lung) disease can be infectious. (Most mathematical models of TB transmission only consider pulmonary TB, although nonpulmonary infection causes significant morbidity. Also, since the natural history of TB is different in children,\textsuperscript{12} most models consider only adults.) Pulmonary TB can be divided into smear-negative and smear-positive, according to whether a sputum sample gives a negative or positive result in microscopic observation. Smear-negativity is caused by a lower concentration of bacilli in the sputum and is associated with lower, if any, transmissibility.

Without treatment, individuals may recover from active TB disease, but there is disagreement over whether natural immune recovery clears the infection, or leads to (re-establishment of) latent infection. In some individuals, untreated active disease leads to chronic, persistent, infectious active disease.

Treatment requires long-term use of antibiotics (at least 6 months is recommended for short-course therapy), but is generally highly effective, including in those with HIV, provided the patient is adherent.\textsuperscript{4} Lack of adherence can result in the bacterium acquiring drug resistance; transmission of drug-resistant strains is a significant problem in many parts of the world: globally an estimated 4\% of patients have multi-drug resistant TB,\textsuperscript{4} but this proportion can be much higher in particular settings. Treatment of drug-resistant TB takes longer and is less effective.\textsuperscript{4} Importantly, most patients—even those with HIV—become uninfected for TB soon after treatment commencement.\textsuperscript{3}

Chest X-ray screening can identify pulmonary lesions that potentially indicate active tuberculosis. Definitive diagnosis requires sputum smear microscopy, or laboratory culture of sputum specimens (which can take several weeks), since a low bacterial load may result in a ‘false negative’ sputum smear test; nucleic-acid amplification tests also lack sensitivity for smear-negative disease.\textsuperscript{4} Diagnosis of latent TB infection previously relied on the tuberculin skin test (TST) where inflammation associated with antigen challenge was used to identify those previously exposed. Unfortunately, this test lacks sensitivity (particularly in the early stages of primary tuberculosis, in immunocompromised persons and in disseminated tuberculosis) and specificity (due to confounding by BCG vaccination or other mycobacterial infections). Fortunately, several tests with far superior sensitivity and specificity have recently become available.\textsuperscript{4,13}

The immune suppression associated with HIV infection increases the rate of progression of latent TB infection, thus increasing the incidence of TB disease.\textsuperscript{3} However, it is debatable how much this increases TB transmission, since HIV increases the proportion of active disease that is smear-negative and shortens the infectious period due to more-rapid death associated with AIDS.

Currently, there is one vaccine against TB, which uses a live attenuated strain of \textit{M bovis} called Bacillus Calmette-Guérin (BCG). Unfortunately, this offers only partial protection against acquisition
of infection and progression to active disease and a more-effective vaccine is badly needed. There is also need for new antibiotics. Fortunately, in recent years there has been renewed research interest in TB,\textsuperscript{14} in the areas of diagnostics, drug therapy and vaccines,\textsuperscript{15,16} promoted by initiatives such as the Global Alliance for TB Drug Development (GATB), which is supported by the Bill and Melinda Gates Foundation and the Rockefeller Foundation. Greatly-improved diagnostic tests have recently become available\textsuperscript{4,13,17,18} and have already offered new insight.\textsuperscript{19}

**Mathematical Models of TB Transmission Dynamics**

The first model of TB was published by Waaler et al in 1962,\textsuperscript{20} followed by others.\textsuperscript{21-29} Early models tended to treat infection incidence as a parameter, rather than modelling the transmission of infection per se. Subsequent modelling work was limited until a number of circumstances generated substantial interest in tuberculosis and modelling of the infection. Firstly, there was the realization that infectious diseases had not been defeated by effective antimicrobial treatment in developing countries, where crowded accommodation and poor nutrition facilitated transmission and progression to disease and where diagnosis and treatment were inadequate. Secondly, the emergence of AIDS and associated increases in tuberculosis disease incidence renewed concern over the disease. Thirdly, the emergence of resistance to the limited number of antituberculosis drugs available lead to the need for more-complex and expensive treatments and to concerns that new drugs were not in development.

A new generation of mathematical models have been developed and analyzed exploring many aspects of tuberculosis. Unfortunately, the lack of a clear and cheap tool to diagnose latent infection and uncertainty about the natural history of tuberculosis have led to a range of different assumptions about the infection and disease being adopted, which affects the findings of the models. These assumptions may be adopted for the sake of simplicity or alternatively may reflect the authors’ understanding of the infection and disease. It is the assumptions about natural history which distinguish many of the models rather than the mathematical structure employed. Most mathematical models of tuberculosis have been deterministic compartmental models using ordinary or partial differential equations or difference equations; these have tended to be either relatively simple models for algebraic analysis of equilibria and stability conditions, or more-realistic, more-complex models analyzed numerically. In addition, there have been some stochastic, discrete-event simulation models developed\textsuperscript{30-33} and this approach is likely to increase in popularity. The following discussion of the natural history of tuberculosis is applicable across the model types.

**Modelling the Natural History of TB**

**Latent Period**

Different approaches have been taken to modelling the latent period of TB. Some models have a single latent stage which all newly-infected individuals enter, meaning that all individuals have the same exponentially-distributed period between initial infection and progression to active disease (e.g., see refs. 32, 34-36). Other models have more-realistic heterogeneity in rates of progression: some have a proportion of newly-infected individuals immediately developing active disease with the remainder developing slower-progressing latent infection,\textsuperscript{37-52} others divide newly-infected individuals into fast- and slow-progressors,\textsuperscript{33,53-60} whilst others have all newly-infected persons passing through an early latent phase which has a relatively high risk of progression to disease before entering a late latent state which has a lower risk of progression.\textsuperscript{61-63} The models of Porco et al\textsuperscript{64} and West and Thompson\textsuperscript{53} have several latent stages of TB infection before the development of active disease.

**Exogenous Reinfection**

Exogeneous reinfection is where an individual with latent tuberculosis infection is subsequently infected with TB bacilli from another source. Models differ in the extent to which this occurs and its consequences: some models do not incorporate this process\textsuperscript{32,34,37,38,41-44,46,56,62,64} and others implement it in different ways—in some models it causes progression to active disease,\textsuperscript{3,46,45,47,48,50-52}.
The effect of exogenous reinfection in tuberculosis epidemiology can be profound. Empirical studies have found that it is an important cause of disease in not only high-incidence areas, but also medium- and even low-incidence ones. This last counter-intuitive finding is due to the presence of high-incidence foci of transmission even in low-incidence areas (see Contact patterns, below). Gomes et al found that a large range of incidence of disease in different settings is explicable by a relatively smaller range of contact rates. The greater the importance of exogenous re-infection in causing new episodes of disease in a particular setting, the more rapidly infection can be brought under control; conversely, if much disease is occurring due to endogenous reactivation of previously-acquired, latent infection then disease cases will continue to arise for some considerable time—unless latent infection can be effectively detected and treated. However, modelling work has shown that where exogenous reinfection is the dominant cause of disease, a vaccine that is only partially protective against reinfection (as is likely to be the case with vaccines developed in the near future) will be of limited benefit.

Active Disease

Individuals with active tuberculosis disease are not necessarily infectious. To be infectious, one must have pulmonary disease, which can then be sub-classified as smear-negative or smear-positive, according to whether the infection can be detected by sputum-smear microscopy (which is less sensitive than sputum culture). Some believe that smear-negative individuals are un-infectious, others that they are infectious, but much less so than smear-positive individuals. Whilst many models consider only infectious active TB, some also consider non-infectious disease (e.g., see refs. 32, 40, 45, 56, 60); in some of these models non-infectious active TB can be a precursor to infectious disease. Typically, models of TB do not consider sputum smear status (e.g., see refs. 33-35, 37, 38, 41-44, 46, 48, 51-53, 55, 57, 62, 63, 65, 69). However, the model of Salomon et al considers both smear-negative and smear-positive individuals, with the former being less infectious, having a lower detection rate and converting to smear-positivity over time.

Recovery from TB Disease

There is uncertainty over whether those who recover from active disease through natural recovery or antibiotic treatment completely clear their infection, or whether they return to a state of latent infection. In some models, there is no natural recovery, whilst in other models natural recovery from active disease returns the individual to latent infection, and in others natural recovery clears infection from the host. Of those models that incorporate exogenous reinfection, some allow it to occur to those who have recovered naturally from infection whilst others do not. Models also vary in their implementation of antibiotic treatment. In Blower et al 1996 and Ziv et al 2001 treatment is effectively lifelong, whilst in Lietman and Blower and Ziv et al 2004 treatment is effectively instantaneous and removes individuals from the model population. However, most models have a ‘treatment’ compartment that is entered when individuals commence treatment and exited when they cease treatment, due to successful completion, treatment failure, loss-to-follow-up, or mortality. Typically, models assume either that successful treatment clears infection completely, or, more often, that it returns the individual to latent infection. Some models explicitly consider those who have experienced treatment failure. Of those models that incorporate exogenous reinfection, some allow it to occur to those who have been successfully treated whilst others do not. In some models, natural recovery and successful antibiotic treatment have the same effect—e.g., both clearing infection in Dye and Williams 2008, but in others they do not. In the model of Salomon et al, ‘natural cure’ does not clear infection, but returns individuals to slow-progressing latent infection, whilst successful therapy does clear infection; exogenous reinfection can subsequently occur in both cases. In Dye and Williams 2000 and Dye and Espinal 2001, natural recovery moves the individual into a
distinct latent state, from where relapse—but not exogenous reinfection—can occur. In the models of Blower et al 1995 and Porco and Blower 1998,19 individuals who recover from disease enter a ‘Recovered’ compartment, which is a latent state from which relapse can occur (these models do not consider treatment), whilst the model of Blower et al 1996 considers treatment interventions and does not feature ‘natural recovery’ (none of these models incorporate exogenous reinfection). In two models of Gomes et al17,63 there is no natural recovery; in the former model treatment clears infection, whilst in the latter it does not (in both models exogenous reinfection can occur to treated individuals). In Dye et al 1998,40 Cohen and Murray 2004 and Cohen et al 2007,33 both natural recovery and treatment result in distinct, separate latent-infection states (Dye et al 1998 also distinguishes between “good” and “bad” treatment).

Vaccination

A key area of uncertainty in tuberculosis epidemiology concerns immunity induced by vaccination. It has long been believed that vaccination with BCG (a live attenuated strain of \( M. bovis \)), did not protect against acquisition of \( Mtb \) infection but gave partial protection against progression to active disease, thus protecting the vaccinated individual against disease and reducing onward transmission of infection. Consistent with this, Vynnycky and Fine argued, based on their modelling analysis,61 that prior tuberculosis infection does not reduce susceptibility to subsequent (re) acquisition of \( Mtb \) but does reduce the probability of progression to active disease. However, recent studies using a new TB diagnostic test that can discriminate between an immune response to BCG and to infection with \( Mtb \) have shown that BCG does offer partial protection against acquisition of \( Mtb \) infection as well as against progression.19 BCG vaccination has been found to have variable efficacy in different settings, for reasons that are unclear.73-75 One possibility is that there has been differential attenuation in the different laboratories where it is maintained for vaccine production.76 Another possibility is that in some populations there is a relatively high rate of exposure to environmental mycobacteria, which might elicit a partially-protective immune response to which BCG vaccination adds negligible additional protection.47,77,78

The assumed effects of vaccination have varied amongst models. Note that most models considered hypothetical future vaccines rather than BCG. It is likely that any new TB vaccine developed in the near future will be only partially protective against acquisition of infection and/or progression to disease.71 In the models of Lietman and Blower42 and Ziv et al 2004,48 vaccination partially protects uninfected individuals against acquisition of infection and reduces the proportion who progress instantly to active disease in the event of their becoming infected; in those who have already acquired TB and are latently infected, vaccination reduces the rate of progression to active disease. In the models of Garcia et al36 and Dye et al 1998,40 vaccination offers complete protection against acquisition of infection by naïve individuals but does not affect the course of infection once acquired, whilst in Dye and Williams 200840 vaccination protects, not only against acquisition of infection, but also prevents progression to active disease in latently-infected individuals. In Gomes et al 2004,47 vaccination is partially protective against acquisition of infection and exogenous reinfection, but does not affect progression, whilst Gomes et al 2007 consider the impact of vaccines that promote recovery from latent infection, without affecting susceptibility to (re)infection. Murray and Salomon54 and Murray 200230 consider vaccines that protect against acquisition or progression. Models also differ with regard to whether the effect of vaccination wanes over time,36,40,42,48 or not.47,60 Clearly, the modeled impact of vaccination depends upon the assumed natural history of tuberculosis (particularly whether exogenous reinfection occurs), the properties of the vaccine—whether it offers complete or only partial protection; whether it protects against initial acquisition, exogenous reinfection, or progression; whether it promotes recovery from latent infection; and whether its effects wane or are life-long—and the assumed vaccine coverage.

Population Age Structure

For simplicity, many models do not incorporate population age-structure. However, there are several reasons why incorporation of age-structure is desirable. Firstly, there is evidence that the
risk of active disease is age-dependent (see refs. 40 and 61 for references). Secondly, protection from BCG vaccination wanes with time since vaccination (and effectively with age, therefore). Thirdly, since infection can last for many decades, TB's transmission dynamics have long time-scales and incorporating age structure can be important, particularly when the annual risk of infection changes over time, resulting in a changing age-prevalence profile. Fourthly, patterns of contact between individuals are often age-dependent.79 Models that do incorporate age-structure include refs. 30, 32, 40, 60, 61 and 80.

Interactions with HIV

Papers modelling the interaction of HIV and TB have recently been reviewed by Bacaër et al.81 Modelling in detail the natural histories of both HIV and TB leads to complex models—e.g., see ref. 64. A more-common approach is to simplify the modeled natural history of HIV40,45,58,81 and/or to make HIV incidence a parameter, rather than modelling explicitly the dynamics of HIV transmission.49,58

Contact Patterns

Vynnycky and Fine developed a detailed, age-structured model of TB natural history and published a series of papers applying the model to England and Wales.61,82-85 Similar to the earlier models of TB, this model does not incorporate the dynamic transmission process per se, but rather estimates annual rates of tuberculosis acquisition by fitting to case-notification surveillance data. The authors argued that long-term declines in TB disease in the UK were primarily due to declining contact rates between individuals.84

Currently, what constitutes a potentially-infectious contact is poorly understood, which limits our understanding of relevant contact patterns. Whilst it is clear that tuberculosis is often clustered in households, schools and workplaces and much transmission occurs in populations living at high density in poorly-ventilated conditions such as prisons, homeless hostels and many hospitals, with transmission further associated with prolonged contact,86-89 there is also evidence that TB can be USBOTNJUUFE UISPVHI ADBTVBM DPOUBDUT transmitted through ‘casual’ contacts.90,92

To date, most models of TB transmission have assumed a homogeneously-mixing population, although a model considering a population divided into households has been developed93 and then extended to incorporate both household and casual contacts.94 A recent model by Cohen et al.200733 combined a more-sophisticated network structure (using the method of Read and Keeling95) with a more-realistic natural history of tuberculosis and found that exogenous reinfection can play an important role even in ‘low incidence’ settings, because tuberculosis infection tends to be clustered, meaning that those with TB infection typically experience a ‘local’ prevalence of active TB that is much higher than the overall population prevalence and hence they experience a much higher force of infection that promotes their progression to active disease, which in turn promotes a high local prevalence for others who are infected. The same authors96 found that heterogeneity in the network of contacts between individuals results in localized outbreaks occurring stochastically, without requiring a particular ‘cause’.

In developed countries with a low overall incidence of tuberculosis, the disease has been found to have a relatively high incidence in specific sub-populations such as homeless persons, problem drug users and prisoners (which are often overlapping groups).88,97-99 It is not clear at present how much transmission occurs from these high-risk populations to the rest of the population. A combination of empirical work and modelling has shown that targeted screening by mobile X-ray unit followed by confirmatory testing can be an effective strategy for tackling TB in these particular groups.100-102

The Basic and Effective Reproductive Numbers of TB

Estimation and interpretation of TB’s basic reproductive number, $R_0$, is complicated, for several reasons.83 Significant changes in the contact rate can occur over the long timescale of a tuberculosis infection, affecting the potential for a new infection to spread. Also, the proportion
of infected individuals who progress to infectious disease is also affected by age-specific rates of progression and mortality.

Furthermore, exogenous reinfection complicates the relationship between the basic reproductive number, $R_0$ and the effective reproductive number, $R(t)$. Without exogenous reinfection, in a homogeneous population at least, the relationship between $R_0$ and $R(t)$, is simple, with the latter being the product of the former and the proportion of the population that is uninfected. However, this relationship does not hold for TB because exogenous reinfection affects $R(t)$ but plays no part in $R_0$, the invasion threshold for a totally-susceptible population. Exogenous reinfection increases the proportion of infected individuals who progress to disease and hence become infectious, which tends to increase $R(t)$; however, this increases the prevalence of infection, which reduces the proportion of the population that is susceptible, which in contrast tends to reduce $R(t)$.

It was proposed that there may be a ‘backward bifurcation’ resulting in exogenous reinfection enabling TB to persist even where $R_0 < 1$, but this phenomenon is not relevant for reasonable parameter values, as it requires those already infected to have increased susceptibility to subsequent reinfection, rather than reduced susceptibility due to immunity.

Modelling Strains of TB

There are three main reasons for modelling distinct strains of TB: to understand the emergence of drug resistance at the population level, to develop methods of analysis of DNA-fingerprinting data to elucidate transmission patterns and to explore the evolution of the bacterium and its relationship with pathogenesis and transmissibility.

Inference of Tuberculosis Transmission Patterns from DNA Fingerprinting Data

Tuberculosis’s variable and usually long latent period makes investigating its transmission dynamics challenging. Recent developments in DNA fingerprinting techniques for TB offer an exciting new opportunity to gain new insights into TB transmission patterns by identifying transmission clusters. The inference of transmission patterns from identification of clusters of related isolates is not simple, however and modelling has an important role to play in developing methods. Some isolates may have the same fingerprint without being part of the same local transmission cluster. Conversely, incomplete sampling may mean that isolates from recent transmission or newly-imported strains may appear unlinked; however, independent importations of a strain endemic elsewhere may falsely appear to be a local transmission cluster. Modelling studies have shown that clustering due to recent transmission will tend to be underestimated, due to incomplete sampling. (However, a modelling study applied to the Netherlands found that in the case of older individuals—who were usually infected early in life when TB was more prevalent and who experienced reactivation—clustering overestimated the amount of recent transmission because those individuals were more likely to have been sources of infection, rather than recently-infected cases.) Murray’s thorough modelling study also found that odds ratios for risk factors associated with clustering are also typically underestimated, particularly if the odds ratios are high, ‘true’ cluster sizes are small, or the proportion of cases sampled is low. She further concluded that, due to variation in these factors between studies, comparison of studies is difficult.

Drug Resistance

A key concern of tuberculosis control is combating drug resistance, which can arise through poor adherence to treatment and can then be transmitted. Resistance to isoniazid is relatively common; multi-drug resistant tuberculosis (MDR-TB) is resistant to both isoniazid and rifampicin. (It is important to note that there are phenotypic properties—i.e., there are multiple strains of TB that are drug-resistant.) Treatment of drug-resistant infection is much more difficult and expensive than normal treatment and, with the emergence of extensively-drug-resistant (XDR) tuberculosis there is a concern that some infections may become untreatable. Understanding transmission patterns is important to the development of effective control strategies.
Unfortunately, the expense of testing for drug resistance has meant that in developing countries it has been common to place tuberculosis patients on first-line therapy unless they have certain risk factors for drug-resistant infection (e.g., likely acquisition from a known drug-resistant case or a past history of drug-resistant infection); only if therapy fails are they tested for drug resistant infection. Clearly, this promotes the evolution of further resistance and the transmission of drug-resistant strains by providing ineffective therapy for a period of months. A new diagnostic test, MODS, offers both faster diagnosis of infection and simultaneously determines its drug-sensitivity profile and is much cheaper than traditional methods. The most cost-effective way to use MODS remains to be determined and modelling has an important role to play in estimating numbers of infections averted using different protocols.

Blower et al 1996 presented the first model of the transmission dynamics of drug-sensitive and MDR TB; a later paper discussed its application to control strategies. Clearly the fitness of MDR-TB strains will determine how difficult MDR-TB is to combat and will determine whether the problem may remain localised or become more widespread. Commonly it is assumed that drug-resistant strains must ‘pay a price’ for their resistance in terms of reduced transmission efficiency. For example, the fitness of MDR-TB relative to drug-sensitive wild-type strains was varied in the range 70-100% in the model of Dye and Williams 2000 and in the range 10-70% in the model of Salomon et al, whilst Dye and Espinal 2001 assumed that MDR-TB’s fitness was 70% fitness of wild-type.

Worryingly, an in vitro study by Gagneux et al found that resistant strains may not necessarily be less fit. Furthermore, even if there is an ‘initial’ fitness cost to acquisition of drug resistance, compensatory mutations acquired over time may negate it. Cohen and Murray 2004 presented a model that incorporates heterogeneity in the fitness of MDR-TB, which can change through mutation; they found that relatively fit MDR-TB strains may emerge over time, meaning that short-term measures of the burden of drug resistance may not be indicative of the long-term scale of the problem. Blower and Chou developed a multi-strain model that allowed for heterogeneity in strain fitness and incorporated the process of ‘amplification’ of resistance, where strains can sequentially acquire resistance to multiple drugs as a consequence of suboptimal treatment. They recommended that second-line drugs should be rapidly deployed to potential ‘hot zones’ where amplification of resistance by sequential acquisition of mutations could lead to the emergence of fit MDR strains.

Rodrigues et al report that when exogenous reinfection is relatively common, resulting in individuals being infected with both drug-sensitive and drug-resistant strains, the outcome of the resulting within-host competition which determines which strain dominates can have important consequences for the population-level emergence of resistant strains.

The recent emergence of extensively drug resistant tuberculosis (XDR-TB) has prompted new modelling work by Blower and Supervie, which applied the model of Blower and Chou and a more-detailed analysis by Basu et al of the prospects for reducing the incidence of XDR-TB through measures to combat nosocomial transmission in a rural South African setting. As XDR-TB becomes better-understood it will clearly be an important concern for further modelling work.

Host Genetic Factors and Within-Host Modelling

An area of mathematical modelling that we will not consider in detail here, but which is an area of growing importance, is within-host modelling. This aims to gain a greater understanding of the interaction of the pathogen and the immune system, including how latent infection is established and maintained, how progression to active disease occurs, the process of recovery and the effect of vaccination. It has been applied to the fundamentals of the interaction of \textit{M tuberculosis} and the immune system, the impact of antibiotic therapy and the emergence of resistance and the effect of HIV co-infection. Many host-genetic factors have been associated with susceptibility or resistance to tuberculosis infection and there is growing interest in the possibility that strains of TB may be adapted to particular host populations and that this may have implications for the efficacy of vaccines and antibiotic therapies in different settings. However, there has been relatively little modelling work to date on the effect of host genetic factors on the epidemiology of
Future modelling work is likely to combine within-host and between-host (transmission) modelling to get a better understanding of TB evolution.

**TB-Control Strategies**

The first attempts to model TB were motivated by a desire to improve control programs and this has continued to inspire much work. Models have been used to set targets for (e.g.) the proportion of cases found and successfully treated and how long gaining control of TB is likely to take under different scenarios. Since resources are inevitably limited, models can be used to determine the most efficient allocation of those resources. Of course, it should be remembered that there is uncertainty in the predicted impact of interventions, due not only to uncertainty in parameter estimates but also uncertainty in model structure—i.e., how the natural history of TB is represented in the model. Improvements in our understanding of TB’s natural history and transmission dynamics will obviously reduce the uncertainty of model predictions.

Modelling has been used to inform WHO’s Stop TB Strategy guidelines for the proportion of cases of active TB that need to be diagnosed and treated successfully in order to control TB and to assess the impact of interventions by predicting numbers of cases averted (e.g., see ref. 128). Salomon et al modelling the impact of potential new, shorter DOTS regimens which may have lower rates of default and make more-efficient use of limited treatment resources. Resch et al found that providing second-line drugs for resistant TB infections in Peru was cost-effective.

Treating latent infection has been recognized to have an important role because the prevalence of latent infection is high relative to that of active disease and there is the risk of endogenous reactivation (which may be prompted by HIV), as well as by exogenous reinfection, leading to active disease. Even if new acquisition of TB infection were to cease immediately then there would continue to be cases of TB disease due to endogenous reactivation for decades to come. Currie et al 2003 modeled the interaction between HIV and TB in Uganda, Kenya and South Africa and found that the most effective way to control TB epidemics fuelled by HIV was to focus on TB control through preventive therapy and treatment of active TB disease. Cohen et al 2006 found that isoniazid preventive therapy (IPT) for HIV-positive persons with latent TB in a sub-Saharan African setting would reduce TB incidence at the cost of increased drug resistance, requiring policies for the effective detection and treatment of resistant TB, which would be costly. They also found that HAART treatment for HIV would have little short-term impact on TB transmission, although it would avert HIV-associated deaths. It is interesting to note that a modelling paper from the preHAART era also concluded that prophylaxis and treatment of active TB disease could also be highly effective in averting TB deaths in a population with high HIV prevalence.

Ziv et al 2001 modeled the impact of targeted treatment for early latent TB infection, which has the greatest hazard of progression to active disease (with these recently-infected individuals identified through contact tracing, although this is not modeled explicitly) and compared it with a strategy of treating longer-term latent infections (which would tend to be identified through screening programs) and found that the former may be more efficient since it targets those at greatest risk of progression. They found that to achieve TB elimination by combining treatment of latent infection with treatment of active disease, the proportion of cases of early latent infection that need to be treated is lower than the proportion of cases of long-term latent infection.

Typically, combining interventions is potentially much more effective than using any one alone due to synergies. For example, even if new acquisition of infection could be halted immediately, endogenous reactivation would cause new cases of active disease to arise in those who are already infected for decades, so effective treatment strategies would still be required and any new vaccine (which is unlikely to be completely protective against acquisition, in any case) will become part of an ensemble of interventions against TB.

The most cost-effective combinations of interventions are likely to vary amongst settings and are likely to change with time, as the transmission dynamics of TB change as a consequence of interventions and other factors. For example, Ziv et al 2004 found that postexposure vaccines (protecting latently-infected individuals against progression to active disease) would initially
have a greater impact than pre-exposure vaccines (protecting against acquisition of infection) on reducing the number of new cases of disease in a high-prevalence setting, but that over time the impact of the former would decline and the latter increase.

A new vaccine may have a role to play in combating the emergence of drug-resistant strains, by limiting transmission and the opportunity to acquire ‘amplified’ resistance or compensatory mutations increasing transmissibility. Modelling would be required to determine the necessary vaccine properties (minimum efficacy against acquisition or progression, duration of protection) and coverage.

**Conclusion**

Recent years have seen intense activity modelling tuberculosis, with some interesting insights about modeled behavior emerging. Unfortunately, much of this work is speculative because of our incomplete understanding of tuberculosis natural history and the slow timescales of tuberculosis epidemics. This slow spread of infection and emergence of disease hinders efforts to compare model predictions with observed patterns. Many important questions remain to be resolved regarding the natural history of TB infection and gaining a greater understanding of TB transmission dynamics requires further empirical laboratory and field work, mathematical modelling and interaction between them. Modelling can be used to quantify uncertainty due to different gaps in our knowledge to help identify research priorities. Fortunately, the present moment is an exciting time for TB epidemiology, with rapid progress being made in applying new mathematical modelling techniques, new tools for TB diagnosis and genetic analysis and a growing interest in developing more-effective public-health interventions. Since future field trials of vaccine candidates will occur against a background of BCG vaccination and TB treatment a key application of mathematical modelling will be in trial design and analysis, as has been advocated for HIV-vaccine trials.

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Modelling Trachoma for Control Programmes
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Abstract
Trachoma is a major cause of blindness in the developing world and 63 million people are currently infected. Large-scale control programmes are being implemented to clear ocular Chlamydia trachomatis infection—the causative agent of trachoma—and improve environmental conditions to reduce transmission. Chemotherapeutic intervention involves antibiotic administration and the effectiveness of this treatment is currently under investigation. A mathematical model has been developed to allow the impact of control programmes on infection and blinding disease sequelae to be predicted. The model has a structure that allows an important aspect of trachoma pathogenesis to be taken into account, namely the effect of repeated cycles of infection and recovery leading to scarring and the damaging disease sequelae. This novel model structure reproduces many age- and time-dependent epidemiological patterns observed in endemic settings and allows the dynamic effect of treatment on infection and disease sequelae to be gauged.

Introduction
Trachoma is the leading cause of infectious blindness in the world and in 1996 the World Health Organization (WHO) adopted a resolution for the Global Elimination of blinding Trachoma by the year 2020 (GET 2020). In addition, trachoma has been included among several ‘neglected tropical diseases’ affecting the developing world which are currently being targeted for very large integrated control programmes. Mass drug administration (MDA) with the antibiotic azithromycin is the cornerstone of the WHO ‘SAFE’ strategy for trachoma control and there have been several recent studies whose aim has been to investigate the effect of one or several rounds of mass treatment for populations with varying baseline levels of disease endemicity. One of the main findings of these studies has been that communities respond heterogeneously to mass treatment and in some communities a persistent reduction in infection prevalence is achieved whereas others return very quickly to pretreatment levels. The likelihood of eliminating the infection appears to be strongly related to the baseline prevalence and, therefore, the transmission environment but this relationship remains to be properly quantified. Mathematical models that include the salient features of disease pathogenesis and the mechanisms of contact and infection transmission can, when fitted to baseline data, allow control interventions to be simulated; the long term consequences of treatment can therefore be investigated.

*—Surgery for trichiasis; A—Antibiotic for infection; F—Facial cleanliness; E—Environmental improvement.

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This chapter begins with a brief review of the mechanisms behind trachoma disease pathogenesis and we explain that these mechanisms can lead to distributions of bacterial infection and disease that are observed in endemic populations. The facts pertaining to disease pathogenesis are then included in the construction of a simple mathematical model of the transmission of infection in a population and the progression of individuals to disease sequelae. The structure of the mathematical model is of interest because it combines aspects of the simple SIS approach—most often used for viral and bacterial microparasitic infections—with the infection burden approach more commonly used to model macroparasitic infections such as helminths. The simple model of disease pathogenesis is shown to reproduce the population patterns of infection, active disease, bacterial load and trichiasis (the final disease sequela before blindness) by age. Once these static patterns have been reproduced, a treatment scenario can be applied to the model, which is then examined dynamically as it rebounds from a state of low infection back to the original endemic state. Mass-antibiotic treatments are modelled by assuming that the transmission context remains essentially the same but infected individuals are returned to their susceptible states. We examine the effect of antibiotic treatment on infection and disease and its implications on programmes to control blinding trachoma and we discuss the usefulness of modelling for control programme impact projections.

**Disease Pathogenesis and Epidemiology**

The causative organism of trachoma is the infectious bacterial agent *Chlamydia trachomatis* and ocular infection with this agent can eventually lead to a chronic conjunctivitis the most severe sequelae of which are corneal opacity (CO) and blindness. A first infection with the bacteria leads to some inflammation and usually a complete clearance; however, individuals are left hypersensitive to subsequent exposure and repeated infections cause chronic inflammation and, eventually, scarring that leads to the severe disease sequelae. The WHO simplified grading scheme—shown in Table 1—outlines the five progressive disease stages for use by nonspecialists working in a field setting (omitting the final stage, blindness). These WHO disease grades will be used throughout this chapter. Until recently, with the advent of sensitive Polymerase Chain Reaction (PCR)-based tools (compared with e.g., Giemsa staining and culture-based methods) to detect infection, the connection between infection with *C. trachomatis* and progress to the various stages of disease has not been clear. However, recent qualitative and quantitative PCR studies in which it has been possible to detect the presence of infection with *C. trachomatis* and estimate the bacterial load in conjunctival swabs, have made it possible to examine the distribution of the infection load over the community and compare with disease diagnoses. These new data can, in turn, be used to construct and calibrate mathematical models of the pathogenesis and epidemiology of trachoma.

| Table 1. WHO simplified clinical grading of trachoma disease and disease sequelae, adapted from Taylor |
| --- | --- |
| **Criterion Definition (and WHO Acronym)** | **Criterion Description** |
| Trachomatous Inflammation-Follicular (TF) | At least five follicles (of at least 0.5 mm in diameter) on upper tarsal conjunctiva |
| Trachomatous Inflammation-Intense (TI) | Papillary hypertrophy and inflammation covering more than half of deep tarsal vessels |
| Trachomatous Scarring (TS) | Clear trachomatous scarring of upper tarsal conjunctiva |
| Trachomatous Trichiasis (TT) | One, or more than one, eyelash rubbing on the eyeball or signs of the eyelash’s removal |
| Corneal Opacity (CO) | Corneal opacity caused by trachoma, some of which obscures the pupil |
as will be shown in the following. These studies have also highlighted discrepancies between cases of disease and infection and these are probably brought about by the kinetics of infection in which, following clearance, disease may persist. Infection without disease, detected by PCR, may be a result of contamination with infections that are not replicating, an idea recently put forward in a new quantitative PCR study for 16s RNA expression.

The prevalence of active disease (TF, TI in the WHO grading, Table 1) peaks for children aged 1-5 years and drops to much lower levels in adults. Often, the larger number of adult cases of TF and TI are female and this is thought to be because of their more frequent contact with infectious children compared with males. The prevalence of scarring (TS), trichiasis (TT), corneal opacity (CO) and blindness in the youngest age groups is often found to be very low—though some recent studies have shown high levels in very compromised populations—with a rising trend with age. Only a small fraction of the trachomatous disease burden for older adults is generally found to be corneal opacity and blindness, but the prevalences of scarring can be high. When the prevalence of active disease is high, the prevalence of the disease sequelae is also high, which lends support to the idea that repeated episodes of inflammation lead to disease progression.

Infection and clinical disease tend to have a shorter duration in older adults from trachoma endemic communities and this may be due to an acquired immunity that involves cellular and humoral components. It is also possible that a decreased exposure to infection among adults may play a role in their observed shorter durations of infection. The acquired immunity tends to accelerate the clearing of infection and may have only a very small protective effect against re-infection, though some genital animal models show significant levels of protection.

Studies investigating infection loads have found strong age-dependences and in some cases close to half of the community bacterial load has been found in children under 1 year. The infection loads of adults are often lower than those of children which is thought to be due to the acquisition of clearing immunity with age. This imbalance between adults and children is particularly marked in communities in which active disease is hyperendemic (here considered to be where the prevalence is greater than 20%) and the infection tends to be more evenly distributed in lower prevalence populations. Acquired immunity is less stimulated in lower prevalence communities due to lower exposure levels and so the immunological reaction to infection is more similar in children and adults.

Another proposed mechanism for chronic inflammation and subsequent scarring disease progression involves persistent chlamydial infection and antigen release, though studies have found this idea very difficult to distinguish from repeated infection that is clustered in households.

**Antibiotic-Based Control Programmes**

The main weapon in the arsenal against trachoma is azithromycin and this antibiotic is currently being deployed across trachoma endemic countries. Both theoretical and empirical studies have sought to answer questions regarding, for example, the frequency with which the antibiotic should be administered, the routes of reinfection in a community and the rate of re-emergence of infection and disease and what these rates tell us about the transmission environment.

The mathematical model of Lietman et al has investigated the frequency of antibiotic treatment needed to eliminate *C. trachomatis*. In this study, the rate of re-emergence of infection leads to an estimate of the initial exponential rate of increase of the infection prevalence in the community. This rate then forms the basis of an analysis to determine the necessary treatment frequency to achieve a steady prevalence decline and the value thus obtained has been compared with data from a field trial (carried out by the same authors) in the Gurage zone of Ethiopia, where annual treatment was found to be sufficient for elimination at an 80% or higher coverage. This study has informed the WHO policy for community antibiotic treatment frequency and duration, but it does not include details pertaining to the natural history of infection that allow the number of infections experienced to be tracked. The gradual experience of infection and associated inflammation, leads to scarring and consequently the disease sequelae and these phenomena are explicitly included in the models presented in this chapter.
Field studies include those by Burton et al. in the Gambia and by West et al. and Solomon et al. in Tanzania, who all look at either one-off or multiple single dose azithromycin treatments. Burton et al. found that most treated villages cleared their infection 12 months after treatment for low initial infection levels, but that, in some, infection was reintroduced externally. Solomon et al. have reported that, at a low endemic level, a single dose at very high coverage caused infection to almost completely disappear and a second treatment 24 months later has eliminated infection, though here azithromycin coverage was supplemented by the use of tetracycline. West et al. found that, after an initially large drop in infection, for a hyperendemic community, infection was reemerging 18 months after treatment.

Methods

A First Mathematical Model of Trachoma

Here, we outline two mathematical models; the first is a basic model accounting for repeat infection and the accumulation of scarring and the disease sequelae TS and TT (Fig. 1) and the second includes an active disease compartment that corresponds to the WHO grading TF/TI but does not model sequelae (Fig. 2). The conceptual mathematical model is based on the following tenets: (1) trachoma is a disease for which progress to the severe sequelae is dependent upon multiple infections; (2) while the accumulation of scars on the way to a TS or TT diagnosis may occur gradually, with each successive infection, it is useful to think of there being a threshold number of infections, beyond which an average individual might be diagnosed as TS or TT; (3) the duration of infection and disease tends to decrease with age, which is likely due to the acquisition of immunity that more rapidly clears infection following the experience of a greater number of infections; (4) bacterial infection load also tends to decrease with age, which is also likely to be due to a stronger infection-clearing acquired immunity in those who have experienced a larger number of prior infections. These tenets, centred on the response of an individual to re-infection.

Figure 1. A compartmental diagram illustrating the model described in the text. Each susceptible and infected compartment is connected to the compartment above so that the population passes up a ‘ladder’ of infection. The subscript \( i \) corresponds to the number of prior infections experienced. Adapted from Gambhir et al. 57.
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with the bacteria, can be incorporated into a simple mathematical model. The model proposed here is of the Susceptible-Infected (SI) framework, with two basic compartments; however, here each of the compartments is indexed by the number of infections that an individual has experienced prior to the current one \( i \) (see Fig. 1).

The model illustrated in Figure 1 is constructed as successive susceptible and infected stages, indexed by suffix \( i \). Individuals pass up a ‘ladder’ of infection through susceptible (\( S_i \)) and infected (\( I_i \)) compartments, each of which is followed by the next compartment above it. On progression up the ladder, individuals’ acquired immune response is assumed to result in more rapid clearance of infection, reduced bacterial load and more severe disease, in accordance with the relevant aspects of Chlamydia immunobiology outlined above. The rate of clearance of infection is assumed to be a function of \( i \), with parameters estimated by fitting to available data on the duration of infection with age.\(^7\)\(^,\)\(^47\) The probability of transmission of infection from an infected individual by flies, fomites, or direct contact is assumed to be proportional to the bacterial load of that individual.

The following equations (adapted from Gambhir et al\(^57\)) describe the rate of change in the numbers of susceptible and infected individuals of age \( a \) and at time \( t \), experiencing their \( r \)th infection. The equations presented here are continuous partial differential equations in age and time and these were discretised for numerical solution using the Euler integration scheme in Matlab\(^6\).

The three terms on the right-hand side of each equation represent: (1) transmission of infection through contact between susceptible and infected individuals; (2) mortality of hosts; (3) recovery from infection.

\[
\frac{\partial S_i(a,t)}{\partial t} + \frac{\partial S_i(a,t)}{\partial a} = -\beta S_i(a,t) \sum_j \rho_j f_j I_j(a',t) + \mu(a) S_i(a,t) + v_{i-1} I_{i-1}(a,t) \tag{1}
\]

Figure 2. Diagram showing a second model that includes a diseased-only class, as well as an infected and diseased class. This model allows the population to clear the infection yet still show signs of disease, a phenomenon commonly observed in field-studies.
\[
\frac{\partial I_1(a,t)}{\partial t} + \frac{\partial I_1(a,t)}{\partial a} = \beta S_1(a,t) \sum_{a'} w(a,a') \int \rho_j(a',t) \sum_j I_j(a',t) \, da' - \mu(a) I_1(a,t) - \nu I_1(a,t)
\]

where \(\beta\) is the transmission parameter, which determines the rate at which individuals move from susceptible to infected states when they come into contact with one another and this parameter was estimated from fitting the model to the available data; \(w(a,a')\) is a mixing matrix (in the discretised version of the model) describing the rate at which individuals of age \(a\) and \(a'\) mix,

\[
w(a,a') = \varepsilon \delta_{a,a} + \frac{(1 - \varepsilon) N_{a'}}{\sum_{a'} N_{a'}}
\]

where \(\delta_{a,a}\) is the Kronecker Delta; \(N_{a'}\) is the number of individuals of age \(a'\) and \(\varepsilon\) is a mixing parameter ranging from 0 (to represent random mixing among age groups) to 1 (to represent assortative mixing, i.e., each age group mixing only with itself, which is set here to 0.5) for an intermediate level of random and assortative mixing; \(\rho_j\) is the infectivity of infected individuals in compartment \(I_j\); \(\mu(a)\) is the death-rate at age \(a\) obtained from the 2001 WHO Tanzanian life table; \(\nu\) is the recovery rate of individuals in compartment \(I_1\). Aside from the ‘ladder’ structure of the model, the other important features of the disease pathogenesis are the changing recovery rate and the infectivity with number of prior infections. These are outlined next.

**Recovery Rate**

The recovery rate per individual \(\nu_i\) from infection \(i\) (measured as the rate per year, so that the reciprocal of this value gives the average duration of infection), is assumed to follow a monotonically increasing function of \(i\) with lowest value \(\nu_1\) (corresponding to the recovery rate from the first infection) with saturating maximum value of \(\nu_\infty\) (corresponding to the recovery rate following a large number of prior infections),

\[
\nu_i = (\nu_1 - \nu_\infty) \exp \left( -\gamma (i - 1) \right) + \nu_\infty
\]

Parameters \(\nu_1, \nu_\infty\), and \(\gamma\) were estimated by fitting the model to the mean duration of infection for different age groups using data already reported but re-analyzed by Grassly et al. with results shown in Table 2 (\(\gamma\) represents the per infection rate of change of the recovery rate).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Parameter Definition</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1/\nu_1)</td>
<td>Mean duration of first infection</td>
<td>13.6 months</td>
</tr>
<tr>
<td>(1/\nu_\infty)</td>
<td>Mean duration of infection after multiple prior infections</td>
<td>2.8 months</td>
</tr>
<tr>
<td>(\gamma)</td>
<td>Rate of drop of duration of infection per prior infection</td>
<td>0.75 infection (^{-1})</td>
</tr>
<tr>
<td>(I_1)</td>
<td>Infection load per person at first infection</td>
<td>(1.03 \times 10^5) copies omp1 per swab</td>
</tr>
<tr>
<td>(\phi)</td>
<td>Rate of drop of infection load per prior infection</td>
<td>0.05 infection (^{-1})</td>
</tr>
<tr>
<td>(\beta)</td>
<td>Transmission coefficient: the rate of transmission (per year) of infection between individuals</td>
<td>29.6 year (^{-1})</td>
</tr>
</tbody>
</table>

Table 2. Parameter definitions and values for the model obtained through maximum likelihood using a function combining infection prevalence, bacterial load and recovery rate data from a hyperendemic setting adapted from Gambhir et al.
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Infectivity

The infectivity of an individual $\rho_i$ is assumed here to be linearly proportional to the bacterial load $l_i$ carried by that individual, such that the maximum value of the infectivity is 1 and it decays with subsequent infections experienced by an individual and we assume that the load is a function of the number of prior infections experienced by that individual, in accordance with the decline in load with age that has been observed in trachoma endemic communities.\textsuperscript{4,5,14,51} We use a simple exponential decay, which describes a decline in the load from its initial value $l_1$ according to the expression shown below,

$$l_i = l_1 \exp \left[ -\varphi \left( i - 1 \right) \right]$$ \hspace{1cm} (5)

The infectivity $\rho_i$ is then calculated by dividing this expression by $l_i$, such that the maximum value of $\rho_i$ is 1 and it decays exponentially with the decay constant $\varphi$. The decay rate of this exponential is estimated from age-dependent data on bacterial load.

Parameter Estimation

The model parameters ($\beta$, $v_1$, $v_\infty$, $\gamma$, $l_1$ and $\varphi$) were estimated by simultaneously fitting the model-generated age-structured infection prevalence, rate of recovery from infection and infection load. A fit over six parameters represents a substantial search problem, so several starting points were used for the minimisation algorithm to increase the chance that the global minimum was found.

Data

The model was fitted to the published data of West et al\textsuperscript{51} (age-dependent prevalence levels and infection load) and Grassly et al\textsuperscript{47} (age-dependent durations of infection), both of which were collected prior to intervention from study sites that were hyperendemic in active disease prevalence.

Prevalence of Disease Sequelae with Age

A simple mechanism for the accumulation of trachomatous scarring (TS) is incorporated into the basic model by assuming that scarring increases with the number of prior infections. With worse scarring comes the more severe disease sequela, trachomatous trichiasis (TT). Corneal opacity (CO), the final disease sequela prior to blindness, is not modelled here because its aetiology differs from that of the other sequelae. While TS and TT are caused by and worsen with repeated infection, CO represents something of a point of no return for individuals who have reached the TT stage and whose eyelids will continue to inflict mechanical corneal damage without surgical intervention. Since the population dynamical model here is a first attempt at capturing the relevant contact-related factors for infection and disease, this separate mechanical process is not included; however, the central importance of CO for the GET2020 goal to eliminate blinding trachoma means that progress to CO should be considered for inclusion in future models.

The simplest possible scheme is used to determine the prevalence of each of the sequelae: thresholds exist, along the ladder of infection, beyond which each of the sequelae are assumed to be present. Beyond the threshold corresponding to a specific sequela, that sequela is assumed to be detectable in the population that has reached this stage e.g., when the threshold for TT has been passed, both TS and TT are present. For simplicity, it is assumed here that the two disease sequelae can coexist with one another i.e., a person may be graded with scarring or scarring and trichiasis.

Disease Sequelae and Prior Infection Number

Once the model has been fitted to the hyperendemic infection data in the endemic setting, the threshold infection levels for the disease sequelae TS and TT are determined. These can be found by comparing the model-generated disease sequelae prevalence curves—given specific threshold
infection numbers—with the data from the hyperendemic setting. The data set used for this hyperendemic setting, while from the same district in Tanzania as the baseline set of West et al., was collected several years prior to the infection data, but it is assumed here that since there had been no treatment interventions prior to the MDA study, infection and disease in the district was in a steady state.

A Second Model Including an Active-Disease Class

The assessment of the level of trachoma in an area—a community, district, or country—is usually conducted on the basis of clinical examination. Since the primary method for the treatment of trachoma is antibiotic for the bacterial infection, the relation between infection and disease is important. In addition to the model already presented, which takes into consideration only the infected state, a second model—entirely separate from the first, but very similar in structure—has been constructed to account for active disease (TF/TI). Infection and disease are not perfectly coincident in trachoma and the age-structured model outlined implicitly assumes that they are, an assumption that allows the model to explore, with great simplicity, the epidemiological patterns of trachoma. However, the inclusion of a stage in which individuals are diseased allows the exploration of: (1) the consequences of the existence of a proportion of the population that is diseased without being infected; (2) the time-lag between the onset of infection and the manifestation of disease (as well as the lag between the cessation of infection and disease), which is of particular importance when infection in a community has been treated by antibiotic and, while the infection load may drop substantially, the signs of disease remain among many members of the population.

Due to the added complexity of investigating the effect of a diseased stage, this second model is not age-structured. The addition of the new diseased class allows the model population to include members who are diseased without at the same time being infected. However, for simplicity, individuals cannot be infected without showing clinical signs of disease. The aim here is to include a diseased class, since it is clinical disease (TF/TI) that is measured—more frequently than the infection prevalence—to determine the level of trachoma in a community.

Figure 2 illustrates the model; the main features distinguishing it from the age-structured model are: (1) the diseased-only state (with recovery rate from this state of 100) and (2) the new path that individuals can take to advance along the ladder of infection from the diseased-only state, bypassing a recovered state and straight into the next infected-and-diseased state above. The addition of a new way in which individuals can progress along the infection ladder is intended to allow for the possibility that, even after antibiotic distribution has reduced the level of infection in the community, disease remains and there may be individuals who continue to experience disease for a considerable time, possibly even progressing in the severity of the disease they experience. This addition is intended to also capture the lag between the clearance of infection and the eventual clearance of disease from a community. Figure 2 shows that this additional path along the ladder is dependent upon the prevailing force of infection in the population, but that it is attenuated by a factor $f$. The compartments shown in Figure 2 represent a particular level of the ladder of infection and should have been labelled with subscripts $i$ as for the earlier model (Fig. 1) but, for simplicity, these have been suppressed here.

Fitting the Model to Data

The second model was developed in Berkeley Madonna™ and fitted to data available from three published studies from, Upper Saloum district, Gambia; Rombo district, Tanzania and Kongwa district, Tanzania. Pre- and posttreatment data—with varying numbers of follow-up surveys following the baseline assessment—were used to determine the parameter values of the model using least squares fitting of the model-generated time-dependent infection and disease prevalence to the data.
Results

Fitting the First Model to Static Infection and Disease Sequelae Data

Once the basic model has been simultaneously fitted to the prevalence, infection load and recovery rate data from the hyperendemic setting—with parameter values found to be as in Table 2—the epidemiological model outputs can be generated as shown in Figure 3 and Figure 4. These figures show that the steady state hyperendemic model results in a decreasing prevalence with age that drops from an initially very high value (Fig. 3A), a decreasing age profile of the infectivity (proportional to the bacterial load) (Fig. 3B), a decreasing profile for the duration of infection (giving a monotonic rise of the recovery rate) beginning at a high value and asymptoting to a maximum value (Fig. 3C), a rising profile with a flattening at older ages of the TS scarring sequela (Fig. 4A) and an almost linearly rising profile of the TT sequela starting at a critical age of around 20 years (Fig. 4B). The threshold number of infections for the detection of TS and TT were found to be 88 and 130 respectively, though these numbers would be lower if, for example, age-related changes in contact rate were included in the model. This set of close fits to the epidemiological data suggests

Figure 3. Age-profiles generated by the best-fitting parameters to the hyperendemic data set of West et al.51 and—for the recovery rate—Bailey et al.,7 reanalysed by Grassly et al.47 (data displayed as square data points with 95% confidence intervals and model fits shown as solid lines): (A) the prevalence of chlamydial infection measured by PCR; B) the infectivity, which is proportional to the bacterial load, measured by quantitative PCR; C) duration of infection. The model was fitted simultaneously to all three data sets. Adapted from Gambhir et al.57

Figure 4. Age-profiles of disease sequelae generated by the best-fitting parameters to the hyperendemic data and using the disease sequelae threshold infection numbers given in the text. Adapted from Gambhir et al.57
that the current paradigm for *C. trachomatis* pathogenesis, as encapsulated in the model, is largely correct. These fits form the basis for applying one or more control perturbations to the model in its steady state and investigating how epidemiologically relevant quantities vary though time.

**Simulating Control Scenarios**

The aim of the models is to simulate control scenarios based on the parameters estimated from different prevalence settings. Here, we give an overview of the kinds of control scenario that can be investigated using the model developed. We look at the effects of antibiotic treatment on the prevalence of infection and on active disease in what follows and we confine the study to the hyperendemic setting, since it is for these data that the model has been fitted.

**The Effect of Mass Treatment on the Prevalence of Infection by Age**

At the time of treatment, a proportion of individuals (given by the fraction: efficacy × coverage) is treated so that their bacterial carriage is reduced to zero and, in the model, they are moved from their infected state \( I_i \) back into the susceptible state \( S_i \); their memory of the prior number of infections is retained. In Figure 5, the plot at time \( t = 0 \) shows that the prevalence of infection in all age groups has dropped to a much lower level than before treatment (shown by the dotted line). Figure 5 also shows the time development of the prevalence of infection from an initial time point at which mass antibiotic administration (MDA) has occurred. The efficacy, for an individual case, of a single dose of azithromycin is taken to be 95%\(^{53,54}\) and the community coverage level achieved is assumed to be 86% as given by West et al.\(^{51}\) The pattern of the spread of infection throughout the population following mass treatment illustrates the following main points: (1) because the prevalence of infection in children is higher than in the rest of the population, the same percentage reduction in the infected individuals in each age group will leave a larger proportion of children still infected posttreatment than the rest of the population; (2) the rate at which infection returns to children is greater than for the rest of the population. This can be seen in Figure 6 where, in addition to the posttreatment prevalence of infection being greater in children than adults, the return to pretreatment levels of infection is particularly rapid in those under 10 years-old. There are two reasons for this: the rate of recovery from infection in children is lower than for adults and, therefore, children retain their infections for greater lengths of time causing greater prevalence levels; the second reason is dependent on the comparatively greater infectivity of children than adults in the model. Since children are assumed to interact more frequently with those their own age (\( \varepsilon = 0.5 \) i.e., there is some assortative mixing), this greater infectivity leads to a greater probability of becoming infected and, therefore, a greater prevalence. The greater infectivity among

![Figure 5](image-url)
children is very similar to the effect of an increased frequency of exposure to infection in children compared with adults.

**Age-Targeted Treatment**

As well as the mass treatment of as large a fraction of the entire community as possible, there are treatment options that target those sections of the community most likely to harbour high infectious bacterial loads. The modelling of protocols in which household members of those who show signs of active trachoma are treated requires a model more fine-grained than the one examined here (as outlined by Blake et al\(^5\)). However, it is instructive to look at the selective treatment of children and the impact of this treatment on the prevalence of infection in the whole community. Figure 5 has illustrated the effect over time of the treatment of the whole age-range at coverage level 86% and 1 year following treatment the hyperendemic community has already returned to its pretreatment prevalence of infection. While it is likely that the treatment of 0-10 year-old children would be a less effective strategy than community-wide treatment, because much of the community infection load is concentrated among children, it is not obvious that the treatment of children alone will not show a similar impact.

The results shown in Figure 6 illustrate the impact over time of a single treatment round of the population between 0-10 years. For the purpose of this illustration, the efficacy and coverage of the round of treatment have both been set to 100%, which, while unrealistic, clarifies the dynamical picture. At the time of treatment, all individuals up to and including the age of 10 years are treated so that their bacterial carriage is reduced to zero and, in the model, they are moved from their infected state back into the susceptible state. The plot at time \( t = 0 \) shows that the prevalence of infection in all people under 10 years is zero while the prevalence of infection for those over this age remains as it was prior to treatment (dotted line). As time progresses, the prevalence of infection in the 0-10 year olds rises as these children come into contact with older infected people. Because the youngest children had, in general, experienced fewer infections than older children at the time that they were treated, they have lower levels of acquired immunity. They therefore clear infection more slowly and have higher bacterial loads when infected, resulting in a more rapid increase in the prevalence of infection among these children.

The dotted curves in Figure 6 represent the age-dependent infection prevalence prior to treatment. By comparison with the solid curve, it can be seen that a treatment round administered to children under 10 years not only reduces prevalence among these children (direct impact) but also decreases the prevalence of infection among those of all ages (indirect impact). Moreover, the size of this impact can be significant. By removing infectious children from the population, adults benefit because they are less likely to come into contact with an infectious source. Because
the younger children tend to harbour greater bacterial loads, a large portion of the community infection load has been eliminated by treating these children.

### Table 3. The parameters used for the active disease model

These parameters were obtained by fitting the model to the pre and posttreatment data of the three endemic settings analysed in this chapter: (a) Upper Saloum district, Gambia; baseline infection prevalence 3.2%, (b) Rombo district, Tanzania; baseline infection prevalence 9.5% and (c) Kongwa district, Tanzania; baseline infection prevalence 57%. Note that the rate of recovery from the diseased-only state, denoted by $\gamma$, represents the total rate at which individuals recover from this state.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Parameter Definition</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$1/\nu_1$</td>
<td>Mean duration of first infection</td>
<td>13.6 months</td>
</tr>
<tr>
<td>$1/\nu_r$</td>
<td>Mean duration of infection after multiple prior infections</td>
<td>2.8 months</td>
</tr>
<tr>
<td>$1/\eta_1$</td>
<td>Mean duration of first episode of disease</td>
<td>31 months</td>
</tr>
<tr>
<td>$1/\eta_r$</td>
<td>Mean duration of disease after multiple prior episodes</td>
<td>6 months</td>
</tr>
<tr>
<td>$f$</td>
<td>Attenuation factor of the force of infection for those in the diseased-only state</td>
<td>a) very low (&lt;0.01), b) 0.35, c) 0.99</td>
</tr>
<tr>
<td>$\beta$</td>
<td>Transmission rate: the rate of transmission (per year) of infection between individuals</td>
<td>a) 2.5 year$^{-1}$, b) 6.4 year$^{-1}$, c) 20.8 year$^{-1}$</td>
</tr>
</tbody>
</table>

The Effect of Treatment on Active Disease

The values of the parameters pertaining to the second, active disease, model, obtained through fitting to the pre and posttreatment data, are listed in Table 3. The parameter values for the fraction of the population advancing along the ladder of infection without recovery show that there is no consistent value obtained from fitting to the data. In the low-endemic setting the fraction is found to be very small, whereas for the high-endemic setting the value is very high. The model-generated

Figure 7. The development through time of the prevalence of infection and active disease for the model and for recent baseline and posttreatment data$^{4,16,31}$ in each of three endemic settings: (a) Upper Saloum district, Gambia; baseline infection prevalence 3.2%, (b) Rombo district, Tanzania; baseline infection prevalence 9.5% and (c) Kongwa district, Tanzania; baseline infection prevalence 57%. Treatment causes a sharp drop in the infection and disease prevalence followed by a rebound at differing rates.
pattern obtained with the extreme values of 0 and 1 (close to the values actually obtained through fitting) give very similar outcomes.

Using these parameter values, the model generates outcomes for the trajectory of the prevalence of active disease (TF/TI) and infection pre and posttreatment. Figure 7 shows the model-generated prevalence levels of active disease and infection and the data to which the model was fitted, for the three data sets. The model curves show the two key features of the kinetics of trachoma infection and disease as it affects trachoma epidemiology: (1) the prevalence of infection is different from the prevalence of disease; in this model, the prevalence of disease is always assumed to be higher than that of infection; and (2) once antibiotic treatment has been administered, the prevalence of infection drops to a low level immediately, while the prevalence of active disease lags behind infection at a higher level.

**Conclusion**

This chapter has detailed the development of a model for ocular infection with *C. trachomatis* in a community setting. New data from qualitative and quantitative PCR (as well as 16S RNA-based PCR), collected at the baseline and follow-up points during trachoma control programmes, are providing new high resolution data on ocular infection with *C. trachomatis* at individual and community levels. Mathematical models of trachoma transmission and control, of a kind able to incorporate this new data, are currently being developed in order to inform control policy for the achievement of the GET 2020 goals. The model outlined here is based upon the idea of a ‘ladder’ of infection in which individuals progress to greater numbers of infection when they come into contact with infected members of the population. With increasing numbers of infections, individuals in the model become progressively more scarred and develop the damaging disease sequelae. The model, therefore, combines aspects of the SI contact framework—often used for modelling microparasitic infections—and the infection burden framework where, in this case, the number of prior infections is equivalent to the infection ‘burden’. This novel approach to modelling disease pathogenesis when multiple infections are important may prove to be useful for modelling other infections, such as malaria.6 Previous models of trachoma infection have not taken into account the progress of an individual up the ladder of infection and have had no simple way of incorporating disease sequelae into the framework.

The model is shown to represent adequately the pretreatment epidemiological patterns of trachoma in a hyperendemic setting, namely: the prevalence of infection and disease sequelae at all ages and the age-profiles of the bacterial load and the recovery rate.

When the model is fitted to baseline infection prevalence data from a recent antibiotic intervention trial in a hyperendemic setting, the resulting age-dependent infection prevalence curve shows magnitudes over all ages that are very close to the observed data and the maximum point of the prevalence of infection is found for young children.

When a single mass drug treatment is administered to the whole age range, the infection prevalence drops instantaneously. Subsequently, the prevalence rises back up to its pretreatment profile, with the most rapid rise occurring in children due to their partially assortative mixing pattern and their high infectivity. In a hyperendemic setting, this infection rebound is shown to happen over the course of 1 year, after which the profile has returned to its pretreatment level. When only a subset age-range of the population is treated, the impact of the treatment is felt over all ages due to the decreased force of infection. However, the single treatment round is found to result in very little impact after a year. In a hyperendemic setting, due to the rapidity of the infection rebound, the WHO-recommended treatment schedule of three annual antibiotic treatments would therefore probably result in no lasting effect on infection or the disease sequelae. Very few cases of trichiasis and corneal opacity would therefore also be averted by these treatment programmes and the GET 2020 goals in this setting would seem to be very difficult to achieve. It is therefore necessary to implement the full SAFE strategy, which will reduce the transmission level of the population, thereby reducing the endemic infection prevalence and the rate of rebound of infection following treatment allowing the impact of repeated rounds of mass drug administration to accumulate over
time. The most recent follow-up surveys from the studies used here to fit the model have found that, in areas with very low initial endemisities one or two high coverage MDAs may be sufficient to achieve elimination, whereas in a high-endemic setting two treatments are not sufficient to result in lasting drops in transmission.

Following a single treatment, a model that includes an active disease stage generates trajectories in which disease prevalence is higher than infection and is found to lag behind it, reproducing these two key observations of field studies. This model also finds that the infection and disease rebound following treatment is very rapid for a setting in which initial endemity is very high and considerably slower in lower endemic communities. The relation between infection and active disease in this basic model can be used to inform surveys carried out by treatment programmes in which clinical status is measured but infection is treated: following treatment the relation between these two states is not simple. Easy-to-use field tests for determining the level of infection are certainly more reliable for infection measurement but while these tests are not widely available the conversion of active disease prevalence to an estimate of infection prevalence would be useful.

The models discussed here have focused on reproducing the progression of individuals through the infection ladder that appears to be so important for trachoma epidemiology and the progression to ocular disease. However, the models are very simple and the conclusions are subject to their assumptions, including: the mixing and exposure patterns with age, sensitivities to parameter assumptions and estimates and the deterministic nature of the model (which can also be adapted to a stochastic framework, taking, for example, elimination of infection into account). The age-structured model should also be fitted to lower endemic settings and posttreatment data to predict properly the frequency and duration of antibiotic and nonchemotherapeutic treatment interventions. Further refinements to the model should also focus on important heterogeneities such as risk groups which cannot easily clear infection, those who harbour particularly large infection loads or respond poorly to antibiotic treatment, as well as the household structure of transmission. These refinements will allow better modelling of targeted treatments and the prevention of infection re-emergence. As more data emerge from intervention trials and control programmes, the models presented in this chapter are capable of being fitted to these new data to allow projections to be made of the impact of treatment in trachoma-endemic communities. This may help guide programmes in achieving the elimination of blindness due to trachoma.

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Transmission Models and Management of Lymphatic Filariasis Elimination

Edwin Michael* and Manoj Gambhir

Abstract

The planning and evaluation of parasitic control programmes are complicated by the many interacting population dynamic and programmatic factors that determine infection trends under different control options. A key need is quantification about the status of the parasite system state at any one given timepoint and the dynamic change brought upon that state as an intervention program proceeds. Here, we focus on the control and elimination of the vector-borne disease, lymphatic filariasis, to show how mathematical models of parasite transmission can provide a quantitative framework for aiding the design of parasite elimination and monitoring programs by their ability to support (1) conducting rational analysis and definition of endpoints for different programmatic aims or objectives, including transmission endpoints for disease elimination, (2) undertaking strategic analysis to aid the optimal design of intervention programs to meet set endpoints under different endemic settings and (3) providing support for performing informed evaluations of ongoing programs, including aiding the formation of timely adaptive management strategies to correct for any observed deficiencies in program effectiveness. The results also highlight how the use of a model-based framework will be critical to addressing the impacts of ecological complexities, heterogeneities and uncertainties on effective parasite management and thereby guiding the development of strategies to resolve and overcome such real-world complexities. In particular, we underscore how this approach can provide a link between ecological science and policy by revealing novel tools and measures to appraise and enhance the biological controllability or eradicability of parasitic diseases. We conclude by emphasizing an urgent need to develop and apply flexible adaptive management frameworks informed by mathematical models that are based on learning and reducing uncertainty using monitoring data, apply phased or sequential decision-making to address extant uncertainty and focus on developing ecologically resilient management strategies, in ongoing efforts to control or eliminate filariasis and other parasitic diseases in resource-poor communities.

Introduction

Parasite transmission modelling is becoming increasingly recognized as a vital tool in guiding the management of parasitic disease control or elimination and for predicting the impacts and benefits of competing management options.1,3 This recognition arises largely from the difficulty of reliably predicting the long-term impact of repeated chemotherapy on the rate of transmission of parasites and hence reinfection within a community, which has resulted in inducing uncertainty surrounding operational issues, such as the required duration of treatment and determination of
infection thresholds below which transmission is controlled or ceases, for most of the current interventions against the major parasitic diseases.\textsuperscript{4-6} It also stems from increasing recognition that empirical field studies alone are unlikely to provide the required information due to constraints on resources and time and the effects of variations in local transmission conditions which make generalizations from individual studies difficult. This issue is particularly problematic for parasites with complex life cycles like the lymphatic filarial worms, which exhibit both complicated transmission dynamics—and therefore likely to demonstrate complex outcomes in response to chemotherapeutic interventions—as well as high levels of aleatory and epistemic uncertainties in their transmission processes, parameters and values.\textsuperscript{4,7-13}

Here, we focus on recent work in modelling lymphatic filariasis transmission to highlight how mathematical models of parasite transmission dynamics can play useful roles in resolving each of these questions and therefore provide the management decision support framework required for defining optimal intervention strategies and for monitoring and evaluating community-based interventions for controlling or eliminating parasitic diseases. A key focus is on how understanding the nature of transmission dynamics, particularly with regard to complex system transitions, heterogeneities and uncertainties, will be crucial to informing the design of effective parasite management. A secondary objective is also to indicate how the application of such tools can serve to reveal gaps in existing knowledge on the biological controllability or eradicability of parasite transmission by management action and hence help guide critically required research for developing a more robust theory of parasite eradication.

Transmission Models and Decisions in Parasite Management

Informed decision-making in any parasite control programme consists of four key elements: (1) the setting of objectives, (2) specification of management options or actions, (3) an understanding of the structure and dynamics of the parasite system to be managed, i.e., how the parasite population responds dynamically to perturbations, and (4) periodic monitoring of the results of management in order to inform and possibly adjust subsequent management decisions.\textsuperscript{5,6,14,15} Figure 1 summarizes how parasite transmission models can be used to address these elements and hence provide a framework for supporting the design of parasite control monitoring and evaluation plans. According to this schema, the key advantage of using dynamic transmission models (over subjective or even empirical assessment by an expert or managers) is that such models will be key to resolving the following four major components of a successful parasite control or elimination programme: (1) conducting rational analysis and definition of endpoints for different programmatic aims or objectives, including transmission endpoints for disease elimination, (2) undertaking strategic analysis to aid the optimal design of intervention programs to meet set endpoints under different endemic settings, (3) providing support for performing informed evaluations of ongoing programs, including aiding the formation of timely adaptive management strategies to correct for any observed deficiencies in program effectiveness, and (4) addressing the impacts of uncertainty and complexity in developing a coherent approach to parasite management.\textsuperscript{5,6,15}

Models and Quantifying Intervention Endpoint Targets

Quantifying Parasite Breakpoint Thresholds in the Human Population

Recently, we and other workers have shown that the derivation of parasite infection thresholds or endpoints to serve as parasite elimination targets fundamentally requires an analysis of the dynamic properties of the parasite transmission system, as these variables essentially represent stability or persistence components of (such) dynamical systems.\textsuperscript{5,8,16-18} In particular, as pointed out in the chapter by Michael and Gambhir (Chapter 2), the two extinction thresholds (one occurring in the vector-to-host transmission process referred to as the “threshold vector biting rate” and the other in the host-to-vector transmission process known as the “breakpoint worm burden”) that exist in the case of vector-borne macroparasitic diseases, such as lymphatic filariasis, arise principally as the result of the operation of positive density dependences acting to regulate infection, which gives rise
Transmission Models and Management of Lymphatic Filariasis Elimination

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Figure 1. The potential role of parasite transmission models in integrating management actions and monitoring data for providing policy- and management-relevant support to the design and monitoring of parasite control programmes. Models can help define endpoint targets for intervention goals, predict the effects of management actions, aide the selection of outcomes and indicators for monitoring and guide interpretation of monitoring data. Feedback from model-based evaluation of monitoring outcomes will be vital to adapting and refining both management actions and the model via revealing topics for research. Reproduced with permission from Michael E et al. Adv Parasitol 2007; 65:191-237; ©2007 Elsevier.

to the occurrence of multiple stable parasite states and crucially the formation of transmission system boundaries the crossing of which can cause a parasite population to either persist in an endemic state or go to an extinct state. Although our initial work has provided some estimates of the likely numerical values of these thresholds, the finding that elimination thresholds for filariasis are very sensitive to the local level of endemicity, vector species and specific population processes occurring in the local host and vector populations means that ultimately estimating reliable endpoints for a locality will require the fitting of the appropriate vector-species specific transmission model to site-specific infection data. Such data-driven model-based estimation is also necessitated by the often large uncertainties associated with model structure, parameterization (especially when such models are characterized by a relatively large number of parameters, as is typical with dynamic parasite transmission models) and prediction. 3,19-21

Fitting complex ecological models to data is not a trivial task, especially when there is corresponding uncertainty and lack of detail in the site-specific infection data. However, recently we have extended a methodological approach based on fitting dynamic parasite transmission models to data using computer simulation techniques in conjunction with a Bayesian information melding (BM) method, to show how such methods may allow the joint use of data and dynamic models to quantify and test locally relevant parasite elimination endpoint values. The BM method (see a more detailed description given by Spear and Hubbard; Chapter 7) takes all available prior information on model inputs and outputs and, where available, likelihood functions for data and generates posterior distributions of model inputs and outputs through statistical comparisons of
predictions with data. The essence of the method is to initially assign to each parameter of a model a distribution function reflecting current uncertainty of its value and to refine these estimates from new information as embodied by the data. The form implemented in our analysis used: (1) uniform or vague prior distributions for each of the model input parameters and (2) likelihood functions for the available data, which in the present case are age-prevalence of infection and therefore, are assumed to be binomially distributed. The multidimensional space defined by the set of prior distributions for each input parameter is then evenly sampled 100,000 times. For each instance of a sampled parameter vector, the model is run and likelihoods of predictions compared to observed data are calculated for each of the age prevalence curves generated. We then used the Sampling Importance Resampling (SIR) algorithm\textsuperscript{20} to resample 500 parameter sets from the original set of 100,000 parameter generates with a probability given by their output likelihood values. These can then be used to generate distributions of the model outputs, i.e., worm breakpoints, threshold biting rates (TBRs) and the basic reproductive ratios ($R_0$) estimates for any given study community. Figure 2 shows the 500 resampled age-dependent equilibrium curves obtained by the SIR algorithm plotted against actual age-prevalence data from communities in East Africa (where the anopheline LF

![Figure 2](image_url)

Figure 2. The 500 curves (blue solid lines) generated by importance resampling of the input parameter sets according to their likelihood for each of the datasets investigated via the application of the Bayesian Melding model fitting approach described in the text (black crosses with solid black lines showing upper and lower 95% CIs of the data). The Annual Biting rate is given in parentheses above each of the plots. The appropriate model corresponding to Anopheles or Culex vector transmission species was used as required. A color version of this image is available at www.landesbioscience.com/curie.
transmission model was fitted) and India (where the culicine LF transmission model was fitted).

Each of the plotted curves is generated by a different model parameter set and the range of curves produced represents the uncertainty remaining in the parameters following the BM updating procedure. Figure 3 portrays the maximum values of worm breakpoints, the corresponding TBRs and a histogram of the $R_0$ values obtained for each of the data sets. The results highlight how this approach can be used to fit, refine and estimate extant uncertainty in complex dynamic parasite transmission models and how it may be used to estimate locally relevant elimination thresholds. They also show that for both the endpoints (both TBRs and mf prevalence breakpoints), not only can there exist a wide distribution of values consistent with the data in each study community when the model is fitted to mf age prevalence, but also that these thresholds can vary significantly between the study communities.

Vector Infection Thresholds: Theory and Empirical Data

To assess local filarial transmission elimination, it is essential also to not only determine if larval infection thresholds exist in the vector population that could signify parasite transmission elimination but also, as in the case of worm breakpoints above, to quantify what the numerical values of such thresholds might be. There are two principal reasons for this interest in identifying and quantifying these thresholds. First, clearly larval infection thresholds represent the parasite elimination target for drug intervention programmes in vector-borne infections, such as filariasis. Second, information regarding the numerical value of such vector infection thresholds is required if xenomonitoring tools, such as parasite DNA-based PCR poolscreen methods, can be deemed to be feasibly applicable, on the basis of mosquito sampling sizes, in filariasis monitoring programmes compared to methods based on measurement of infection in humans.
Recently, we have shown how a joint stability analysis of the deterministic filariasis transmission model described in the chapter by Michael and Gambhir (Chapter 2) and analyses of vector infection data will be critical to address these issues. Figure 4 shows the results from a numerical stability analysis carried out using the deterministic model of lymphatic filariasis transmission. It depicts the existence of a mf prevalence breakpoint threshold of 0.53% prevalence in the human population (shown for culicine transmitted filariasis for given averaged values of each model parameter) that corresponds to a density threshold of 0.0009 L3 per mosquito in the vector population. However, since no L3-specific probes to monitor vector population transmission now exist (these would need to be based on a stage specific RNA transcript, protein or other nonDNA biomarker), control programs are left with estimating the comparable “all-stage infection” (mf, L2 and L3) prevalence figures to gainfully use the theoretical larval breakpoint results depicted in Figure 4. This requires investigating how L3 density per mosquito is related to L3 prevalence and how in turn L3 prevalence is related to prevalence of all-stage larval infection prevalence. Figures 5A,B show how we can use published data on mosquito dissections to derive both these prevalences based on estimating (1) the relationship between average L3 density recorded among dissected mosquitoes versus the prevalence of L3 infection observed in these mosquitoes in order to obtain the threshold L3 infection prevalence (Fig. 5A) and then (2) to use this threshold value in conjunction with estimation of the relationship between vector L3 infection prevalence and overall infection prevalence again observed in published data to derive the vector overall or all-stages infection threshold (Fig. 5B). These results are clearly crude (e.g., the relationships are estimated for both culicine and anopheline mosquitoes combined and the data are derived from a variety of mosquito collection methods, including a mix of resting and/or biting catches); nonetheless they provide a first estimate of the magnitudes of the infective L3 and overall larval infection threshold prevalences that may be used as preliminary transmission endpoint targets in the global LF elimination programme. More specifically, the results in Figures 5A,B suggest that in this regard if L3 infection thresholds obtained through mosquito dissections are to be used as endpoint targets then the relevant endpoint value could be preliminarily set as 0.085%, while for overall larval infection, the corresponding value would be higher at 0.65%. As noted by Michael et al, the importance of these actual threshold values is that they will play a major role in the choice of the infection indicator for monitoring. For example, if dissections are to be used for estimating infection levels in the vector population, the higher value of the overall larval

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Figure 4. Results from a numerical stability analysis of the coupled deterministic transmission model of lymphatic filariasis indicating the existence of a L3 larval density transmission threshold (value of 0.0009 per mosquito given by the dashed line crossing the X-axis) in culicine mosquitoes corresponding to the existence of a mf prevalence threshold in the human host (value of 0.53% given by the dashed line crossing the Y-axis). Details of model and analysis methods are as given in Gambhir and Michael. Reproduced with permission from Pedersen EM et al. Trends Parasitol 2009; 25:319-27; ©2009 Elsevier.
Figure 5. A) The relationship between average L3 per mosquito and prevalence of L3 infection obtained in wild-caught mosquitoes from filarial endemic communities. Each circle in the Figure denotes paired data on these variables obtained from mosquito samples within each study community \((n = 29, \text{ refs. given in Pedersen et al}^{23})\). The solid curve shows the fit of the negative binomial model relating infection intensity to prevalence given by: \(P(M,k)=1-(1+M/k)^{-k}\) where \(P\) is the L3 infection prevalence, \(M\) is the L3 density and \(k\) is the aggregation parameter of the negative binomial distribution. The model was fit to the data by maximum likelihood using binomial errors giving an estimation of \(k\) of 0.007. Application of the model to an \(M\) value of 0.0009 (the L3 transmission threshold density) gives a corresponding value of 0.085% as a potential L3 threshold prevalence for determining filariasis elimination (shown in bold on the graph). B) The relationship between overall all stages vector infection prevalence and prevalence of L3 infection obtained in wild-caught mosquitoes from filarial endemic communities. Each solid circle in the Figure denotes paired data on these variables obtained from mosquito samples within each study community \((n = 63, \text{ refs. in Pedersen et al}^{23})\). The solid curve shows the best fit quadratic logistic regression model of the form: \(\log[p(\frac{M}{M+1})]=a+b_1 X+b_2 X^2\), where \(p\) denotes the prevalence of overall all stages infection, \(X\) is the prevalence of L3 infection, \(a\) represents the intercept and \(b_1\) and \(b_2\) represent the linear and quadratic terms respectively, with estimated values of \(a = 0.239, b_1 = 0.210\) and \(b_2 = -0.008\). Application of this model to the L3 threshold prevalence value of 0.085% gives an empirical value of 0.65% as the corresponding all-stages larval infection prevalence threshold (Figures in bold on the graph). Reproduced with permission from Pedersen EM et al. Trends Parasitol 2009; 25:319-27; ©2009 Elsevier.
infection prevalence indicator, which would reduce the required sample size plus ease the effort of dissections for estimating this prevalence, would mean that this indicator would be the choice for transmission monitoring when using mosquito dissection as a method to monitor the impact of an intervention. Again, as noted for worm breakpoints, model uncertainties as well as heterogeneities in initial conditions between localities, such as in infection aggregation, host immunity and type and magnitudes of positive density dependences, will result in both a distribution of larval infection endpoints within communities as well as significant site-to-site variability in the actual values of this threshold.

Models and Design of Optimal Filariasis Intervention Strategies

Modelling for Choosing Optimal Strategies

A major utility of mathematical models of infectious disease transmission in disease control decision making is that by facilitating the integration of information on pathogen transmission dynamics with programmatic factors such as drug and other intervention efficacies, they may provide a powerful strategic tool for designing and planning community-based control programs against infectious diseases. In particular, if control targets are well defined and model structure, parameter estimates and outputs are validated, the predictions of such models can be used for finding the optimal choice among different strategies for meeting a set control-program objective. Here, we illustrate and discuss the important role that filariasis transmission models can play in guiding the design of optimal intervention strategies to eliminate culicine-mediated filariasis under different endemic and programmatic conditions, focussing in particular on the optimal design of the two currently proposed mass drug administration (MDA) regimens for filariasis—i.e., combined DEC/ALB and IVM/ALB therapies—and the added impact of including vector control to these regimens.

Figure 6A portrays the results of model predictions of the impacts of these regimens in terms of the number of years required by each option to reach a target threshold of 0.5% mf prevalence (assumed here as the threshold for parasite elimination (but see above)) for a range of precontrol community prevalence or endemicity levels. Three key features of these results illustrate how such model simulations can support the planning and optimal design of parasite elimination programs. First, the dotted droplines in the figure show the feasibility boundaries, i.e., the maximum endemicity level at which it would be feasible to achieve the target threshold of 0.5% mf prevalence within the prescribed 6 years of control, for each intervention and highlight the endemicity ranges over which a particular MDA (at 80% population drug coverage) or MDA plus vector control (incorporating a 90% reduction in vector biting) intervention strategy can successfully meet the above control criterion. The foremost finding here is that annual mass treatments using the combined IVM/ALB regimen may successfully achieve the above elimination criterion only in communities with up to a relatively low precontrol mf prevalence of 3.75% (at the scale of 1 ml blood sampling volume), while in the case of annual mass combined DEC/ALB treatments, the corresponding controllable precontrol community endemicity levels may be only moderately higher at around some 10% mf prevalence (Fig. 6A). These results clearly imply that achieving elimination of filariasis transmission by mass chemotherapy alone, whether by the application of the IVM/ALB or the DEC/ALB regimens, may not be feasible within the prescribed 6 years timeframe in areas of high endemicity, although clearly the DEC/ALB option (largely as a result of its greater macrofilarial activity) would achieve parasite elimination for a higher range of precontrol endemicity levels compared to IVM/ALB treatments.

This is highlighted by the second major feature of the results, viz that for both drug regimens, the addition of vector control will extend the range of endemicity levels over which it would still be feasible to achieve the target threshold in the prescribed 6 years compared with that afforded by drug treatments alone (Fig. 6A). Apart from extending the range of controllable endemicity levels, combining vector control with drug administration will also enable the achievement of the set target infection level faster and earlier compared with using each...
Figure 6. A) Deterministic filariasis transmission model predictions of the number of years of intervention required by various annual MDA and combined annual MDA plus vector control options to achieve the target elimination threshold of 0.5% mf prevalence (44), for a range of precontrol community endemicity (Mf %) levels. Solid curves give predictions for IVM/ALB (open squares) and DEC/ALB (closed squares) annual MDA regimens respectively, while dashed curves portray the corresponding results for each of these regimens combined with vector control (i.e., open squares-dashed line for IVM/ALB plus vector control and closed squares-dashed line for DEC/ALB plus vector control). Drug efficacy values: DEC/ALB—55% worm kill, 95% mf cured and 6 months mf suppression; IVM/ALB—35% worm kill, 99% mf cured and 9 months mf suppression; while vector control is assumed to be 90% effective in reducing vector biting. The lowest dashed curve with transparent squares denote the effects of increasing the frequency of mass treatment to the DEC/ALB plus vector control regimen. The droplines in the Figure serve to indicate the maximum endemicity level at which it would be feasible to achieve the set target threshold of 0.5% mf prevalence within the prescribed 6 years of control by each intervention. All results at 1 ml blood sampling volume. B) Model simulations of the number of years of intervention required by the mass annual DEC/ALB regimen either administered alone (solid lines) or with vector control (dashed lines) to achieve the 0.5% mf prevalence threshold for a mix of precontrol community mf prevalences (5%, 10% and 25%) and drug coverage values. 90% Vector control efficacy assumed. Vertical droplines show the optimal drug coverage required at each endemicity level by each of these options to meet the control criterion of achieving the 0.5% mf prevalence threshold in 6 years. Reproduced with permission from Michael E et al. Trends Parasitol 2006; 22:226-33;©2006 Elsever.
drug regimen alone. The number of control years thus saved by the inclusion of vector control to each of the two drug regimens is given by the areas between the respective solid and dashed lines in Figure 6A and indicates that on average between ½ to 1 year at low (<10%) and 1½ to 2 years at moderate/high (>20%) precontrol community infection levels may be saved by adding vector control to each of the DEC/ALB and IVM/ALB regimens respectively (Fig. 6A). These findings underscore the critical role that including vector control to MDA programs can play in filariasis elimination programs, especially in areas of high endemicity, where the benefits of this combined strategy would be most apparent and in areas, such as Africa, where MDA using IVM/ALB is to be implemented.

A third important insight of significance to the global filariasis control initiative gained from the model simulations depicted in Figure 6A concerns the finding that when combined with the set control criterion, i.e., reaching 0.5% mf prevalence in 6 years, the addition of vector control (at 90% efficacy) to DEC/ALB mass treatment (at a coverage of 80%) produces a rather moderate upper limit to the endemicity level (approximately 18% at the scale of 1 ml blood sampling volume) that can be feasibly controlled even by this most effective combined strategy. This result implies that we need to either extend the duration or frequency of mass treatment in areas of higher precontrol prevalence or examine other more effective options in these areas. Indeed, model simulations (lower curve in Fig. 6A) indicate that increasing treatment frequency to once in every 6 months may substantially increase the upper limit to the endemicity level (to almost 70% mf prevalence) that can be feasibly controlled by the combined DEC/ALB-vector control strategy, suggesting that this option may well need to be considered in areas with the very highest prevalences if filariasis elimination is to be achieved within a reasonable timeframe in such regions.

Predictions from transmission models can also play a key role in quantifying another important element regarding the rational design of mass treatment programs, viz the optimal drug coverage needed to meet a set control criterion for various endemic conditions.1,33-35 Figure 6B shows the predictions made by the deterministic filariasis transmission model of the number of years of intervention required using annual MDA with DEC/ALB alone versus mass annual DEC/ALB administration plus vector control (at 90% efficacy) to achieve the 0.5% mf prevalence threshold for a mix of precontrol community mf prevalences (5%, 10% and 25%) and drug coverage patterns and exemplifies how the application of models can facilitate the derivation of optimal drug coverages for drug-based intervention strategies. Several important points regarding optimal program design in this regard emerge from the simulations shown in the figure. A key finding is that for both the MDA alone and the MDA plus vector control strategies, optimal drug coverages will increase with precontrol infection endemicities and for both strategies this impact of variations in drug coverages, particularly in the case of sub-optimal coverage, in achieving a set infection control will be greater at higher precontrol endemicity levels compared to low precontrol infection prevalence levels (Fig. 6B).

The results in Figure 6B also highlight a final key point regarding the inclusion of vector control to MDA programmes, viz that for all endemic situations including vector control will lower the optimal MDA coverage required to meet a set control criterion compared to MDA alone. Given that typical coverages achieved in large-scale community treatment programmes are normally only around 65%,46 this result again clearly affirms the value and importance of considering the inclusion of vector-management options in current filariasis mass treatment programmes,1,37-39 especially in areas with high precontrol endemicities.

Table 1 shows how these results can be summarized as a type of decision table to aide the identification of the optimal coverage needed to achieve a programme target for various precontrol community mf endemicity levels. The shaded diagonal region in the table represents the present decision criterion (reaching 0.5% mf threshold in 6 years) for various combinations of endemicity levels and coverage rates. The optimal coverage required to meet the control threshold of 0.5% in 6 years for a particular precontrol prevalence are then found simply by tracing the corresponding paired figures off this diagonal. These results not only contribute to providing a decision support framework for determining the optimal coverage for the successful execution of the mass treatment programme
Transmission Models and Management of Lymphatic Filariasis Elimination

Choosing an Optimal Strategy Under Uncertainty

While the above results are pertinent in identifying optimal strategies under conditions of low uncertainty regarding both program objectives as well as biology or ecology of the parasite system, choosing the best option and management decision-making under uncertainty in either these areas presents a more complex problem in parasite management. Recently, we have shown how uncertainty in transmission endpoints behooves us to take a more flexible sequential approach to achieving filariasis elimination based initially on attaining disease control and then proceeding to the target of parasite elimination when information regarding endpoints becomes better known. This work, which combined analysis and quantification of the disease control threshold, mathematical modelling of the effects of MDA with IVM/ALB and economic analysis, was applied to the control of filariasis eradication in Tanzania, is summarized in Table 2 in terms of expected present costs (EPCs) and option values of strategies with and without a flexible sequential decision option based on exercising disease control first. The results in Table 2 depict that the EPCs of strategies that contain the flexibility of switching between disease control and eradication are lower than that of an eradication strategy in which no flexibility to either switch to disease control if eradication is not feasible or to embark upon eradication after disease control when eradication becomes favourable exists. Since the EPCs of the flexible strategies are lower than that of the inflexible eradication strategy, it is clear that such strategies should be preferable highlighting their optimality when uncertainty regarding eradication occurs. Indeed, the results show that switching to long-term disease control from year 5 (MDAs given every 10 years following an initial 5 annual MDAs to achieve disease control: Strategy 2) can be the cheapest intervention option for LF if eradication can never be achieved (Table 2), while the lower EPCs of Strategies 3a and 3b compared to Strategy 1 highlight the costs which can be saved by waiting for better information regarding the feasibility of eradication before switching to eradication from disease control. The value of including such flexibilities in the filariasis eradication program for Tanzania is given by option value figures depicted in Table 2 and indicates that including flexible decision-making based initially on disease control can yield savings of between US$ 3.3 to 5.0 million depending on the feasibility of eradication (from never (Strategy 2) to waiting to switch to eradication at different times when knowledge and technology improves (Strategies 3a and 3b)). These results also clearly highlight that when substantial biological and programmatic uncertainty exists, there is value in designing and implementing adaptive management options that will facilitate learning about the parasite system and ultimately the best long-term strategies for achieving overall management goals.

Table 1. Developing decision support for the design of filariasis elimination programs. Model predictions of the number of years required to reach a 0.5% microfilarial prevalence threshold using combination DEC-ALB mass drug treatment in relation to endemicity and coverage.

<table>
<thead>
<tr>
<th>Endemicity (Mf %)</th>
<th>Coverage</th>
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<tr>
<td></td>
<td>2.5%</td>
</tr>
<tr>
<td>60%</td>
<td>7</td>
</tr>
<tr>
<td>70%</td>
<td>6</td>
</tr>
<tr>
<td>80%</td>
<td>5</td>
</tr>
<tr>
<td>90%</td>
<td>4</td>
</tr>
<tr>
<td>95%</td>
<td>3</td>
</tr>
</tbody>
</table>
However, the finding of the existence of hysteresis loops in the mf prevalence/vector biting rate plane in filariasis transmission dynamics (see Fig. 5 in Chapter 2 by Michael and Gambhir; this volume) suggests that a second tactic to promote filariasis elimination might be to avoid focusing solely on meeting the objective of uncertain elimination, but to exploit the ability of even a relatively poor model to give fairly good guidance to promote good parasite system transitions (e.g., parasite elimination) and prevent bad transitions (e.g., infection re-emergence following control). Given that including vector control with MDA can, by increasing the worm break-point threshold value reduce the resilience of the endemic state and by raising the re-emergence infection threshold promote the resilience of the parasite-free state, it is clear that a combined MDA/vector control strategy can play this resilience-enhancing role in facilitating and sustaining filariasis elimination.

Models and Design of Monitoring Programs

Recently, it has been shown how by conceptualizing critical components and processes of parasite transmission dynamics, mathematical models of parasite transmission can provide a rational framework for undertaking monitoring and evaluation activities via their ability to unveil information on the following four specific areas: (1) determination of intervention endpoint targets to meet program objectives; (2) predictions regarding the expected magnitude of changes in the states of the parasite system under various interventions to determine if these are on track to meet set endpoint targets, including estimation of optimal frequency of monitoring and suggestion of remedial measures to get the intervention back on track; (3) determination and exploration of the roles of key indicators for monitoring effectiveness; and finally (4) guiding interpretation of monitoring results to ascertain if successful control has been achieved.

A difficulty with the above role and use of models in monitoring programs is that it is premised on the notion that it is possible to predict and anticipate the consequences of management

Table 2. Present value costs of LF eradication strategies for the Republic of Tanzania with and without sequential-decision making flexibility based on exercising chronic disease control options

| Strategy | Design | EPC | Option Value
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</thead>
<tbody>
<tr>
<td>S1</td>
<td>Inflexible: 10 year annual MDA</td>
<td>157,165,771</td>
<td></td>
</tr>
<tr>
<td>S2</td>
<td>Flexible: Switch to long-term disease control after 5 years of annual MDA when it becomes evident eradication is not possible</td>
<td>152,110,572</td>
<td>5,055,199</td>
</tr>
<tr>
<td>S3a</td>
<td>Flexible: Switch to eradication with 3 extra annual MDAs from year 10 following disease control with 5 initial annual MDAs</td>
<td>153,822,682</td>
<td>3,343,089</td>
</tr>
<tr>
<td>S3b</td>
<td>Flexible: Switch to eradication with 3 extra annual MDAs from year 15 following disease control with 5 initial annual MDAs</td>
<td>152,968,795</td>
<td>4,196,976</td>
</tr>
</tbody>
</table>

1Denotes the three different scenarios described in detail in the text. 2Expected present value cost of each strategy in US$. 3The difference between the Expected Present Costs (EPCs) of flexible strategies 2 to 3 over strategy 1. Represents the value (costs saved here) of retaining the option to switch between eradication and disease control depending on the state of eradication feasibility. From Michael E et al. PLoS One 2008; 3(8):e2936.
decisions, i.e., that once all the necessary information is gathered, a right “scientific” decision can be made. For filariasis (and for most other parasitic systems), given that major uncertainties exist in almost every area of our understanding of transmission dynamics and control, including (1) model and process uncertainty which gives rise to approximate thresholds and predictions of the effects of intervention, (2) observation uncertainty related to diagnostic tool accuracy, (3) the role of spatial and population ecologies on the sampling process, (4) intervention uncertainty in which there is little reliable information available regarding the effectiveness of the various proposed interventions to reduce or interrupt parasite transmission, and (5) that the transmission dynamics is likely to be furthermore ecologically complex, it is clear that taking such a traditional anticipatory approach to management is problematic. This highlights that a major role of monitoring programs should be to provide data that can be used to develop our knowledge of parasite system dynamics and responses to management. In particular, in the face of complex infection dynamics, there is a need to plan and use monitoring data to identify the alternate attractors accessible to the parasite infection system, the feedbacks which maintain the system at the attractors, the external ecological influences (environmental, vector species, vector and human migration) which define the context for a specific attractor and conditions, including interventions, which allow or promote “good” flips between attractors.

The above considerations suggest that there is an imperative need to modify existing command and control management models for filariasis control based on globally-set targets in order to account for the complexity and uncertainty in parasite system dynamics and resulting intervention thresholds that are likely to vary from site to site. Specifically, we identify here a need to develop an adaptive management plan that (1) acknowledges uncertainty and system complexity, (2) actively contemplates using monitoring system responses to interventions to learn about and reduce uncertainty in system models and (3) anticipates that future management interventions will be modified as we gather information and learn more about system behaviour. Such an approach must also link monitoring data with robust methods, such as Bayesian analysis, that allow ready and effective updating of new information to refine model structure or reduce parameter uncertainty. This approach will also require a change in attitudes to implementing parasite interventions for at its core it argues for the use of interventions as natural experiments to compare selected policies or practices in order to evaluate alternate hypotheses about the parasite system being managed. This is in sharp contrast to the current rigid top-down anticipatory management strategies used for parasite control, which sees such deviations from set protocols as “errors” to be avoided.

Conclusion

Informed parasite management requires information about both the parasite system state at a given time as well as the impact that potential management actions will bring about on that state. We have shown here how mathematical models of parasite transmission can play a vital role in informing and guiding the development of filariasis elimination programs by their ability to provide quantitative information for decision-making in all key areas of optimal program design and execution, ranging from (1) initial definitions of thresholds to meet various intervention endpoints, (2) optimising program parameters for different endemic conditions and (3) creation and application of rational monitoring and evaluation plans for assessing management success. The results, however, have also highlighted the continuing gaps in understanding and the urgent need to resolve such gaps in both knowledge of parasite population dynamics and hence model formulation and parameterization, and the impact of stochasticity in key infection processes as well as intervention effectiveness—for example, currently there is little reliable information available regarding both drug efficacy rates against the underlying adult worms as well as the rates of efficacy of various vector control measures in reducing vector population size—if modelling results are to be reliably integrated and applied within control programs. We show how a critical advance will also be how best to address the impact of ecological complexity in parasite transmission if we are to develop a more reliable approach to parasite control management.
Whereas new model fitting approaches based on Bayesian updating methods can be used to both reduce model uncertainty and gain an understanding of localized transmission complexities and sequential decision frameworks could be used to guide management policies in the face of uncertainty, perhaps the most important conclusion from the work described here is that this need to resolve uncertainties and address complexities means that control programs should also be sufficiently flexible to adapt to local transmission dynamics and even accommodate change in intervention methods as the program proceeds. Complexity in transmission dynamics also indicate that several intervention methods, for example adding vector control to MDA in filariasis inventions, may need to be applied simultaneously or sequentially during a program to enhance the resilience and sustainability of “good” system transitions.8,41,42

Overall, these results thus imply that rather than using a simple “one size fits all” command and control approach, as currently promulgated in parasite control programs, a more nuanced approach focusing on using intervention and monitoring programs to aid greater learning and understanding of parasite system states and the forces that govern critical system transitions and how best to manage such transitions by interventions to achieve desired objectives, will be required if we are to successfully attain the goal of controlling or elimination human parasitic infections. Transmission models will clearly play a major role in the development and application of this adaptive management framework. We indicate that if science, including mathematical models, is to effectively support the management of large-scale control or elimination of parasitic diseases, including lymphatic filariasis and other major human helminths, from endemic communities, such management methods closely linked to and aided by models should now be urgently examined and deployed at all programmatic levels.

Acknowledgements
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References
Disease Transmission Models for Public Health Decision-Making: Designing Intervention Strategies for *Schistosoma japonicum*

Edmund Y.W. Seto* and Elizabeth J. Carlton

Abstract

The purpose of infectious disease transmission modelling is often to understand the factors that are responsible for the persistence of transmission, the dynamics of the infection process and how best to control transmission. As such, there should be great potential to use mathematical models to routinely plan and evaluate disease control programs. In reality, there are many challenges that have precluded the practical use of disease models in this regard. One challenge relates to the mathematical complexity of the models, which has made it difficult for field workers and health officials to understand and use them. Another challenge is that, despite their mathematical complexity, models typically do not have sufficient structural complexity to consider many of the site-specific epidemiologic and disease control details that the practicing health official routinely considers. Moreover, most modelling studies have not been sufficiently explicit or exemplary in explaining how field data may be incorporated into the models to impact public health decision-making. In this chapter, we start with a classic model of schistosomiasis transmission and relate its key properties to the more detailed model of *Schistosoma japonicum* model presented in chapter by Remais and chapter by Spear and Hubbard. We then discuss how various controls (e.g., chemotherapy, snail control and sanitation) may be evaluated via the detailed model. We then demonstrate in a practical manner, using *S. japonicum* data from China, how field data may be incorporated to inform the practice of disease control. Finally, we present a new model structure that considers how heterogeneous populations are interconnected, which has particular relevance to understanding disease control and emergence in today’s highly mobile world.

Introduction

Nearly two decades ago, Woolhouse reviewed the modelling literature for schistosomiasis and how various disease controls may be evaluated.¹² We remark upon his rather depressing introduction, which stated that since Macdonald developed the first model in 1965 and subsequent reviews of the modelling literature in the 1977 and 1982 by Cohen and Barbour, respectively, the models had very limited impact on actual schistosomiasis control.³⁴ One of Woolhouse’s reasons for this, which we tend to agree with, is that there exists a disconnect between disease modelers who focus their work on theoretical mathematics and field workers who make real-world public health decisions related to disease control. Another reason is that, while mathematically complex,
the models still lacked the realism to capture local variations and logistical constraints that affect the implementation and efficacy of controls. Woolhouse argues that this does not need to be the case. Instead, models may be one of the tools that health planners use, in addition to field experience and epidemiology in designing control programs. With this as background, we begin where Woolhouse left off, using the same simple schistosomiasis modelling framework to consider disease control. We discuss how the \textit{S. japonicum} model presented in Chapters 5 and 10 relates to the basic modelling framework and allows us to consider various disease controls, including chemotherapy, snail control, sanitation and health education. We then demonstrate in a practical manner, using \textit{S. japonicum} data from China, how field data may be incorporated to inform the practice of disease control.

We also present a new development in schistosomiasis modelling that explicitly considers how heterogeneous populations are interconnected. These connections may occur via natural physical processes (e.g., hydrological connections via watersheds) and/or via social-economic processes (e.g., migrant labor or trading of buffalo). Such a model offers exciting new possibilities to consider how disease control may be best practiced in highly connected environments. As many areas are becoming increasingly connected due to economic development, it also provides opportunities to evaluate the impact of changes in connectivity due improvements to road and irrigation networks, for example, on disease transmission and (re)introduction.

**Model Framework**

Schistosomiasis infection persists in over 70 countries and remains a cause of great morbidity despite several opportunities in the lifecycle of the parasite to disrupt transmission. Figure 1 illustrates the lifecycle for \textit{S. japonicum} in China and points of possible intervention, which include the treatment of human hosts via chemotherapy, snail control through molluscicides or more permanent habitat removal, improvements in sanitation and reductions in exposure (e.g., via health education). We describe each of these controls in more detail below.

Woolhouse’s basic model framework is based on the simplification of the worm-snail-cercariae-miracidia 4-state system to a 2-state system of just mean number of schistosome worms per person \(m\) and the prevalence (as a proportion) of patent infections in snails \(y\). This is reasonably assumed as the lifespans of the larval stages are considerably shorter than worms and snails (hours versus months and years). Hence, it is assumed that larvae are at equilibrium and we can simply model the remaining states via the following coupled differential equations (see Table 1 for a description of the model parameters):

\[
\frac{dm}{dt} = \alpha N y - \gamma m \\
\frac{dy}{dt} = \beta H m (1 - y) - \mu y
\]

(0.1)

where each equation consists of a gain and loss term. For the worm burden equation, the gain term consists of the product of the number of snails \(N\), prevalence of patent snail infection \(y\) and the rate of infection per person per patent infected snail \(\alpha\). The loss term consists of the product of per capita worm mortality \(\gamma\) and the mean worm burden per person \(m\). For the snail infection prevalence equation, the gain term consists of the product of the number of humans \(H\), mean worm burden per person \(m\), the proportion of uninfected snails \((1 - y)\) and the per capita rate of infection of snails per schistosome \(\beta\). The loss term consists of the product of per capita infected snail mortality \(\mu\) and the patent snail infection prevalence \(y\).

The simple model above has a number of assumptions that are inherent in the underlying Macdonald model, upon which (0.1) is based. First, it is assumed that the helminthes are bisexual and that the mating probability of these worms is incorporated in the \(\beta\) term. Moreover it is generally assumed that there is equal probability of male and female larval infectivity and worms are monogamously paired. More importantly, the basic model ignores possible density dependent effects that may arise through acquired immunity. Furthermore, it is assumed that the loss of
Miracidia from the environment due to snail infection is negligible. Snail numbers are assumed to be held constant, which does not account for seasonal variations or increased morbidity imposed by larval infection. Multiple infections of snails are ignored, as is the issue of infection latency due to a prepatency period. Additionally, human population numbers are assumed to be held constant, with homogeneous exposure to larvae occurring for the population. It also assumes humans are the single definite host. We comment on various extensions of the basic Macdonald model that relax some of these assumptions later.

Table 1. Variables in the transmission models

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
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<tbody>
<tr>
<td>$m$</td>
<td>Mean number of schistosome worms per person</td>
</tr>
<tr>
<td>$y$</td>
<td>Prevalence of patent (shedding) infections in snails</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>Rate of infection per person per patent infected snail</td>
</tr>
<tr>
<td>$N$</td>
<td>Number of snails</td>
</tr>
<tr>
<td>$\gamma$</td>
<td>Worm mortality</td>
</tr>
<tr>
<td>$\beta$</td>
<td>Rate of infection of snails per schistosome</td>
</tr>
<tr>
<td>$H$</td>
<td>Number of humans</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Snail mortality</td>
</tr>
<tr>
<td>$k$</td>
<td>Aggregation parameter for schistosomes in the human host</td>
</tr>
<tr>
<td>$\phi$</td>
<td>Mating probability of schistosomes</td>
</tr>
<tr>
<td>$T_{SH}$</td>
<td>Snail to human transmission rate</td>
</tr>
<tr>
<td>$T_{HS}$</td>
<td>Human to snail transmission rate</td>
</tr>
<tr>
<td>$P$</td>
<td>Prevalence of human infection</td>
</tr>
</tbody>
</table>

Figure 1. Schistosome lifecycle with main opportunities for control shown in parentheses.
1) Humans become infected by the free-swimming cercarial form of the parasite from skin contact with contaminated surface water. 2) Once infected, adult schistosome worms will develop in the portal veins of the liver or bladder depending on the species of schistosome. Male and female worms will mate and produce eggs, which are released in either the feces or urine. 3) Inadequate sanitation results in parasitic eggs being deposited into an aquatic environment (e.g., ponds, irrigation ditches), where they hatch into a free-swimming miracidial form of the parasite. 4) Miracidia infect the snail intermediate host, which lives in these aquatic environments. 5) After a prepatency period, infected snails will begin to release cercariae, which infect humans to complete their lifecycle.
Classically, the approach to studying the Macdonald system is first to evaluate the equilibrium solutions of the model. Equilibrium for each state denoted as \( m^* \) and \( y^* \) are reached when both the mean worm and patent snail infection prevalence reach a stable unchanging level (i.e., where \( \frac{dm}{dt} = 0 \) and \( \frac{dy}{dt} = 0 \)). In discussing this equilibrium it is helpful to define the following terms:

\[
T_{SH} = \frac{\alpha N}{\gamma}
\]

\[
T_{HS} = \frac{\beta H}{\mu}
\]

where \( T_{SH} \) considers those parameters that relate to snail-to-human infection and \( T_{HS} \) considers those that relate to human-to-snail infection.

The equilibria then can be described according to a quantity known as the basic reproduction number, \( R_0 \):

\[
R_0 = T_{SH} T_{HS}
\]

When \( R_0 < 1 \) then the system does not sustain transmission and the equilibrium values are:

\[
m^* = 0
\]

\[
y^* = 0
\]

However, when \( R_0 > 1 \) there exists a positive equilibrium at:

\[
m^* = T_{SH} - \frac{1}{T_{HS}}
\]

\[
y^* = 1 - \frac{1}{T_{SH} T_{HS}} = 1 - \frac{1}{R_0}
\]

Hence, an important property of the basic schistosomiasis model is that endemic infection will exist for a population when the Basic Reproduction Number, \( R_0 > 1 \).

We note that \( R_0 \) is explicitly defined by the snail-to-human and human-to-snail infection parameters. However, these terms can be factored into more specific terms, as we saw in the Macdonald-like \( S. japonicum \) model of Chapter 7. Indeed, we note even in Macdonald’s early work, there was a distinction between biologic and site-specific variables (synonymous with the \( P_s \) and \( P_b \) variables, respectively mentioned in Chapter 7). For instance, the probability of a cercaria finding and infecting a suitable host, (Macdonald’s so-called “exposure factor”) may depend upon a biologic constant representing the efficiency of cercarial penetration and a site-specific variable that relates to the number and characteristics of water contacts made by hosts.

In Macdonald’s work, considerable detail is spent in describing the effect of adding more detailed mating probability function \( \phi \) as one of the factors that makes up the \( \beta \) term. It is assumed that male and female schistosomes infect with equal probability, but mate at random, with at least two worms, one of each sex required for mating. The results of adding the mating probably function is the establishment of break point in the system, which concerns the resulting equilibrium when the parasite is introduced into a community. As described by Woolhouse,

\[
A constant mating probability results in any parasite introduction establishing endemic transmission (positive \( m^* \) and \( y^* \)). However, when a more realistic assumption that the mating function depends upon the aggregation of worms in hosts (typically regarded as negative binomial distributed), an unstable break point occurs in the system, such that parasite introductions below a threshold do not establish transmission (i.e., the system will fall to \( m^* = 0 \) and \( y^* = 0 \) equilibrium). Conversely, when there is a sufficient introduction of parasites, then endemic transmission will be established (i.e., the system will settle on the \( m^* > 0 \) and \( y^* > 0 \) equilibrium).

From a control perspective there are thus two theoretical goals in eliminating transmission. One goal may be to reduce the longer-term potential for transmission such that \( R_0 < 1 \). Alternatively, another goal may be to perturb the system so much as to bring the system below the break point,
whereby it will fall to the 0, 0 equilibrium. We note, however, that accomplishing either says nothing of the dynamics (how long it will take) for transmission to be eliminated. We deal with this issue of system dynamics later, after we first explore how the simple Macdonald model performs for real-world data.

**A Crude Estimate of \( R_0 \) for *S. japonicum* Based on Macdonald’s Model**

Based on the simple Macdonald model’s equilibria equations (0.5) for endemic transmission, we note a rather surprising finding: the only field data required to estimate \( R_0 \) is snail infection data. Note that \( y^* \), the equilibrium patent snail infection prevalence (expressed as a proportion) is the sole estimator for \( R_0 \):

\[
y^* = 1 - \frac{1}{R_0}
\]

In China, as in many other places the proportion of snail surveyed that are patent infected typically does not exceed 0.05. In our study of 20 villages in Xichang county in Sichuan Province, we found the highest snail prevalence to be 0.03. Substituting 0.03 for \( y^* \), results in a solution for \( R_0 \) of 1.03. Because \( R_0 > 1 \), the condition for endemic transmission is satisfied.

**Control Needed to Terminate *S. japonicum* Transmission Based on Macdonald’s Model**

The above estimate of \( R_0 \approx 1.03 \) is surprising because it seems so small. One way to sustainably eliminate transmission is by reducing the potential for transmission by driving \( R_0 < 1 \). We must effectively transition from a \( R_0 \) of 1.03 to an effective reproduction number \( R \) (the common term for \( R_0 \) after control has been implemented) of <1. We note that because of the following definition of \( R_0 \), there are many possible control options to bring \( R < 1 \):

\[
R_0 = \frac{\alpha N \beta H}{\gamma \mu} = \frac{\alpha N \beta H}{\gamma \mu} = 1.03 \Rightarrow R = R_0 L = \frac{\alpha N \beta H}{\gamma \mu} L < 1
\]

where \( L \) represents a factor that reduces \( R_0 \) sufficiently to bring \( R < 1 \). In this case, to terminate transmission, \( L \) must be <0.97. It is quite shocking that only a 3% reduction in \( R_0 \) would terminate transmission. It is even more shocking when one considers that factor \( L \) could come from any of the possible controls and even combinations of controls.

Routine chemotherapy can be implemented as an increase in \( \gamma \), the per capita mortality of schistosomes:

\[
\gamma' = \gamma - \ln(1-g'h)
\]

where \( g' \) is the coverage rate and \( h \) the efficacy, both as proportions. The increased mortality essentially results in a decrease to \( T_{sh} \), thereby creating \( R < R_0 \). And our goal is to obtain:

\[
R_0 = \frac{\alpha N \beta H}{\gamma \mu} = 1.03 \Rightarrow R = R_0 \left( \frac{\gamma}{\gamma'} \right) = \frac{\alpha N \beta H L}{\gamma \mu} \left( \frac{\gamma}{\gamma'} \right) < 1
\]

Indeed, if we assume a fairly non-aggressive control strategy that consists of a coverage rate of single treatment every 2 years and 70% efficacy when the natural lifespan of worms is 3 years, working through the calculation suggests this strategy would be more than sufficient to eliminate transmission (at least eventually, at equilibrium)! Similar arguments could be made for relatively easy elimination through snail control by modifying \( N \Rightarrow N' \); improved sanitation by modifying \( \beta \Rightarrow \beta' \) and reducing exposure by modifying \( \alpha \Rightarrow \alpha' \) (see\(^2\) for examples of how these additional controls may be implemented).
Problems Inherent in the Macdonald Model Assumptions

The above analyses suggest that with such low $R_0$ (i.e., slightly above 1), it should be easy to terminate transmission. If only it were so easy in the real world! Indeed, because the Macdonald model is unrealistic in its assumptions, the equilibria conditions are thus unrealistic.

Barbour was quite vocal about the limitations of the basic Macdonald model, attempting to improve the model (or “rescue it” as he states) by accounting for various heterogeneities that may scale $R_0$ upwards.9,10 Specifically, Barbour accounts for snail latency to patency by the addition of a full snail population submodel based on *S. japonicum* data,11 spatial heterogeneity in water contacts and age-dependent immunity.9 Despite various attempts to rescue the model, the values of $R_0$ from variations on Macdonald’s model may still be unrealistically low.10

We note that in Chapter 7, for our calibrated Macdonald-like model for *S. japonicum*, we arrived at an approximate $R_0$ (or $R'$ representing the prefILTERING calibration criterion) of approximately 4. Moreover, as this is a prefILTER criterion, it is an upper limit on $R_0$! Again, using chemotherapy as an example, two treatments per year with 70% efficacy should eliminate transmission for this region of China—something that is already being conducted in many endemic areas of China. However, it is important to note that we still have not considered how long it would take to reach elimination, which we discuss later in terms of the dynamics of the system. We only note here that there may be benefits to combining chemotherapy with other more sustainable disease control strategies.

Still, there remain other heterogeneities that have not been widely considered. For one, more work is needed to consider age-acquired immunity, for which there is evidence for various *Schistosome* species, although the role of age-acquired immunity in *S. japonicum*, in particular, remains unclear.12 In addition, the presence of animal reservoirs may need to be considered for some environments particularly for *S. japonicum*, as over 40 different wild and domestic mammals have been shown to play a role in transmission.13 Stochastic effects and the transport of parasites between connected environments may also create heterogeneities that modify $R_0$, the latter discussed in greater detail below. Space and time-varying effects, such as attenuations that may occur through the synchrony of exposure may also modify $R_0$. In these respects, recent studies by various modelling groups may be considered (e.g., for *S. japonicum*, age-specific or population group-specific models in,14,15 models of parasite persistence in connected populations,16 time-varying factors (Chapter 6) and inclusion of animal hosts17).

Barbour has suggested an alternative to the Macdonald model (the so-called Ross model) based on human infection prevalence rather than worm burden.10 The rather obvious benefit of such a model is that it is based on human infection prevalence, which can be measured (or at least approximated from the currently best-available diagnostic tests), as opposed to worm burdens which cannot be directly measured in humans and are difficult to relate to egg count data. Under Barbour’s model:

$$R_0 = \frac{1}{(1 - y^*)(1 - P^*)}$$  \hspace{1cm} (0.9)

where, as before, $y^*$ is the equilibrium infected snail prevalence and now $P^*$ is the equilibrium prevalence of infection in humans. Real-world values of 3% snail infection and 50% human infection, immediately result in values of $R_0$ twice as high as the Macdonald model without accounting for any heterogeneities. Indeed, there is some acceptance of this modelling approach by those working in China and the Philippines on *S. japonicum*.15,17

Modelling Control Dynamics

As we alluded to earlier, regardless of our choice of model (e.g., Macdonald or Ross), the equilibrium analyses of the models only tell us the eventual equilibrium values of the system, but not how fast the system will arrive at those equilibria. In the real-world, there is a practical need to understand the comparative performance of competing control programs, which often depends upon what can be done in short time frames (e.g., 5-10 year control programs) and how large of an impact might be expected. Such issues ultimately lead to questions about the dynamic properties of the system.
We note from our work, that when various heterogeneities and delays are incorporated into the system, analytical solutions are essentially not possible. Thus, numerical simulations are required to understand the equilibrium and dynamics of the system. Moreover, often quite complex models are numerically simulated and calibrated to field data, which can be computationally intensive. Thus, in Chapter 7, considerable attention was paid towards optimizing simulation times via tricks, such as prefiltering the parameter space based on $R_0$. Indeed, in our work, we have spent considerable time to model the dynamic behavior of various combinations of disease controls (e.g., various combinations and levels of chemotherapy, snail control and sanitation). We note also that the dynamics of these controls may act differently. For instance, in contrast to the Anderson-May method of modelling a chemotherapy program as described above, we have modeled chemotherapy as a resetting of the worm burden state (i.e., an instantaneous drop in worm burden when chemotherapy is provided), which can lead to disease elimination when considered in terms of the break point phenomenon of the model system. In contrast, snail control may be implemented either as a instantaneous drop in snail populations, reduction in snail habitat, or additional mortality, which depends on how the snail control was implemented. Thus, some disease control strategies may be used to yield immediate effects on reducing disease morbidity (good public health benefit), while others may be aimed at long-term, sustainable disease control (moving towards disease elimination). This can be clearly seen in the time-series figures of that show sudden drops in worm burden due to chemotherapy, but gradual reductions in transmission over time due to combined control of snails and/or sanitation.

Given the above dual objectives for disease control, it is interesting to comment upon the various stages of schistosomiasis control in the world. Global schistosomiasis control policy currently focuses strongly on the distribution of praziquantel. As the drug has become increasingly affordable, it has taken a leading role in disease control efforts. In contrast, 15 years ago, the World Health Organization placed health education, access to safe water supplies and sanitation at the center of schistosomiasis control efforts. Clearly, for some countries, drug treatment to control morbidity needs to remain the focus, particularly in environments where being able to reliably deliver the drug remains a challenge. However in China, where much of our work is focused, there is already good access to drugs, yet the disease persists. There is recent evidence of disease re-emergence and a rise in prevalence following a 10-year chemotherapy campaign in some endemic areas. The situation in China and in other middle-income countries dealing with parasitic diseases is transitioning away from asking the dynamic questions of how to achieve short term reductions in morbidity, but rather how to effectively reduce and/or eliminate transmission over the long term. Moreover, a related question concerns the threat of disease (re)emergence and how might models be best-used to evaluate this threat in terms of its probability, likely time-frame and place of occurrence. Current efforts in our group are focused on exploring these latter issues, particularly in the context of using models to control and provide early warning for emerging schistosomiasis.

New Model Developments: Incorporating Population Heterogeneity and Connectivity

Quite early on it was recognized that one of the limitations of the simple schistosomiasis model was the assumption of population homogeneity, which led subsequently to adjustments such as those for worm aggregation and differences in exposure (e.g., different risk groups in our work). Still it was assumed that transmission occurred such that host populations were isolated from other hosts and parasite populations were isolated from other parasite populations. The real world, however, is made up of a complex mixture of populations that are defined by natural and social and political boundaries. Moreover, while parasites may be transported via hydrological or social networks, it is political boundaries that often govern how control is conducted in the real world. In China, for instance, each county organizes their own local control strategy against *S. japonicum*. In any given year, a county may decide to perform surveillance or mass control in some villages, but not in others.
Increasing evidence suggests that transmission actually occurs in groupings of heterogeneous populations that are in fact connected to one another in varying degrees. Connections may occur via numerous processes, most notably via hydrological channels that transport the larval stages of the parasite between environments and via socio-economic connections, such as host movement between environments. Generally, this understanding has led to the following model as described by Guarie and Seto (modifying their notation slightly to match our’s):

\[
\frac{dM}{dt} = \alpha \pi_c \hat{\Theta} \left( \Omega^T \cdot T_c^{-1} \right) \cdot \hat{N} \cdot Y - \gamma M
\]

\[
\frac{dY}{dt} = \beta \alpha \pi_M \left( \Omega \cdot \hat{\Theta} \cdot \hat{H} \right) \cdot (1 - Y) M - \mu Y
\]

in which different heterogeneous populations are represented in matrix notation as \( M \), a vector of village worm burdens and \( Y \), a vector of infected snail prevalences. The model accounts for differential social contacts through the matrices \( \Theta, \Omega, \) per capita cercarial and miracidial outputs \( \pi_c, \pi_M \) and spatially explicit transport for cercariae and miracidia that incorporate hydrological connectivity, survival and mortality in the matrices \( T_c, T_M \) and different vectors of human \( H \) and snail \( N \) populations.

Various elaborations of this system have been explored by our group for *S. japonicum*. The more recent theoretical analysis of the system by Gurarie and Seto describes both the general characteristics of the system and a few surprising aspects that have practical disease control implications. First, we define the terms:

\[
R_{SH} = \hat{N}^{-1} \left( \Omega^T \cdot T_c^{-1} \right) \cdot \hat{N}
\]

\[
R_{HS} = \left( T_M^{-1} \cdot \Omega \right)
\]

which represent the contribution of social and hydrological connectivities for snail-to-human and human-to-snail transmission, respectively. The product of these factors and the “internal potential” \( \rho \) results in the “Basic Reproduction Matrix”:

\[
R = R_{HS} \cdot \hat{\rho} \cdot R_{SH}
\]

Similar to the \( R_0 > 1 \) inequality for the simple Macdonald system (0.1), this matrix analog, \( R \) conditions sustained transmission for the distributed-connected system (0.10). Specifically, transmission is sustained when the largest eigenvalue of \( R \) satisfies the condition:

\[
\lambda_1 (R) > 1
\]

We note that (0.12) and (0.13) precisely define the role that “internal potential” and “connectivity” play in transmission. Both are important in sustaining transmission. The underlying parameters that make up internal potential \( \rho \) relate to the local potential for transmission within each village in the absence of connectivity (i.e., \( R_0 \)). In fact, in the absence of connectivity, the matrices simplify in such a manner that each individual village’s \( R_0 \) conditions transmission.

The practical implications of the system on control are large. We can show that due to connectivity, infection may be high in certain villages. Yet focusing treatment on these high-infection villages may be inefficient. Instead, careful exploration of the network of connectivity and choosing key nodes that contribute to downstream infection may be much more useful in achieving region-wide control. Indeed, it can be shown that the structure and characteristic of various social and hydrological connectivities can determine whether region-wide schistosomiasis transmission will occur optimally, or not at all (Fig. 2).

We have observed this in our work in 20 villages in Xichang County located in Southwest China: the village with the highest infection prevalence (73% of residents were infected in 2000) was located downstream from an endemic village that pertained to a different county. High levels of cercariae were detected in the irrigation channels through which water entered...
the village suggesting local infection may not only be dependent on village conditions but the influx of parasites from upstream.\textsuperscript{7} Despite treatment of all infected residents in 2000, infection prevalence was 52\% only two years later. We surmise that disease control efforts in this village that do not address the sources of parasite import will yield transient reductions in infection. The connected model confirms this hypothesis and suggests a more regional perspective to control activities may be useful, particularly in environments like those in China where rapid re-infection occurs and re-emergence is common. This understanding has led most recently to spatial analyses aimed at identifying connected watersheds that maybe used by local disease control agencies to better coordinate regional control strategies (Fig. 3).

While it has not yet been explored extensively for \textit{S. japonicum}, we note that the dynamics of such a system maybe particularly interesting from the perspective of emerging disease and disease spread. Because population connectivities are explicitly modeled in the system, it would be possible to consider the impacts of increasing population connectivity that might occur via improved infrastructure, such as road and water resource development. In some cases, as we have shown in our work, increased social dispersion may actually reduce the potential for transmission by moving people away from hotspots in the network that sustain infection.\textsuperscript{16} The practical analog to this is the increasing migration of labor away from rural environments to urban ones. Moreover, stochasticity may be incorporated into the model to consider the effects of chance perturbations to the network, including, for example, the occasional introduction of infected hosts into areas with low or no transmission. Such a model would allow for the identification of environments particularly favorable to the (re)initiation of transmission.

**Conclusion**

Given almost half a century of model development for schistosomiasis, surprisingly little use of models occurs in the planning and implementation of disease control programs. As described above, simple models and crude $R_0$-based analyses allow for easy calculations of the effect of chemotherapy and other control strategies, however, such calculations may lead to misleading results.
because of the over-simplified assumptions of these models. Accounting for various heterogeneities in the transmission of the parasite has led to more complex models, which have suggested that the disease may be more difficult to control than suggested by simpler models. Indeed, there is a trend towards increasingly complex models because simple models are not able to mimic real-world field data. Another reason for increasing model complexity may also be less justified, simply because there exists computing power to consider such complex models via numerical simulation. Unfortunately, the trend towards more complex models and numerical simulation is pushing the field of schistosomiasis modelling outside of the realm of practical field use at the local or regional level. Indeed, after over a decade in our group of developing our *S. japonicum* model for China, there has not been the knowledge transfer necessary for our complex model to be run at a provincial-level Chinese research institute.

However, our modelling work has demonstrated some encouraging applications. The most important application has been the ability to use the models as an independent tool based on field data to assess whether current control policies seem reasonable. In this regard, we have been able to confirm regional and national-level policies promoting combinations of disease control measures as having the best chance at eliminating transmission. Much of this knowledge has come about from studying the more complex dynamics of the model, rather than the equilibria of the system, which is consistent with the practical question of how long it takes to reach appreciable levels of morbidity reduction from various combinations of controls. Even more encouraging are the recent developments in modelling heterogeneous, yet connected populations that are beginning to tackle some of the most challenging aspects of schistosomiasis control, such as how to reduce the potential for re-emergence. In our increasingly connected society, these connected models also have great potential to identify the impacts of changes in connectivity on transmission of parasitic diseases such as schistosomiasis.

Figure 3. Three-dimensional rendering of watersheds (bounded by white lines) in Xichang County, Sichuan Province, China, computed in a geographic information system and displayed in Google Earth. These visualizations were created to assist the county in better planning their regional control efforts by taking into account how villages may be hydrologically connected to one another.
Although many practical field decisions regarding schistosomiasis control continue to be based on experiential knowledge, there exists a new class of problems, for which there is little to no field experience. This is the case with disease emergence and re-emergence. In our group, the current work on developing connected models may be used to design surveillance systems and examine predictors of disease emergence. Such models may prove to be very useful in exploring the possible impacts of (re)emerging disease, making full use of the ability to run computer simulations for a variety of hypothetical scenarios when very little experiential knowledge is available from the field. Our group is also continuing our work on complex site-specific models to explore the effect of various mixtures of control strategies, which is very much needed to combat endemic transmission as well as to prevent disease re-emergence.

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Abstract

The impact of climate change on human health has received increasing attention in recent years, with potential impacts due to vector-borne diseases only now beginning to be understood. As the most severe vector-borne disease, with one million deaths globally in 2006, malaria is thought most likely to be affected by changes in climate variables due to the sensitivity of its transmission dynamics to environmental conditions. While considerable research has been carried out using statistical models to better assess the relationship between changes in environmental variables and malaria incidence, less progress has been made on developing process-based climate-driven mathematical models with greater explanatory power. Here, we develop a simple model of malaria transmission linked to climate which permits useful insights into the sensitivity of disease transmission to changes in rainfall and temperature variables. Both the impact of changes in the mean values of these key external variables and importantly temporal variation in these values are explored. We show that the development and analysis of such dynamic climate-driven transmission models will be crucial to understanding the rate at which *P. falciparum* and *P. vivax* may either infect, expand into or go extinct in populations as local environmental conditions change. Malaria becomes endemic in a population when the basic reproduction number $R_0$ is greater than unity and we identify an optimum climate-driven transmission window for the disease, thus providing a useful indicator for determining how transmission risk may change as climate changes. Overall, our results indicate that considerable work is required to better understand ways in which global malaria incidence and distribution may alter with climate change. In particular, we show that the roles of seasonality, stochasticity and variability in environmental variables, as well as ultimately anthropogenic effects, require further study. The work presented here offers a theoretical framework upon which this future research may be developed.

Introduction

The potential effects of global climate change and ozone depletion on human health have received increasing attention in recent years, with those due to changes in vector-borne disease (VBD) incidence and distribution thought to represent one of a range of major direct and indirect effects. Malaria has arguably attracted the most attention of all VBDs, due to both the sensitivity of its transmission dynamics to changes in environmental variables and its status as one of the biggest causes of worldwide mortality due to infectious disease, with an estimated 247 million cases in 2006.
Climate change may affect malaria transmission via changes in the ecology and behaviour of humans, Anopheles mosquitoes and Plasmodium parasites.\textsuperscript{11} Perhaps the most important effect will be on the vector population itself. Immature stages of the lifecycle are aquatic and increasing rainfall, leading to increased availability of breeding sites, is a strong driver of abundance. The use of precipitation as an empirical predictor of incidence,\textsuperscript{12-16} besides its direct impact on vector abundance,\textsuperscript{17-22} is well-established. The role of temperature has also been considered in terms of its effects on vector abundance (as a physiological factor affecting the development rate of larvae, adult daily survival probability and biting rates due to increased processing rate of blood meals), local dispersal, parasite dynamics (such as reproduction rates as a function of temperature) and human behaviour.\textsuperscript{4,23-25}

Given the links between malaria incidence and variation in environmental variables, mathematical models incorporating climate variables have been developed in recent years.\textsuperscript{7,20,26-28} The growing importance of these models derives from the fact that while statistical models have been useful in elucidating relationships between environmental variables and transmission intensity,\textsuperscript{9} process-based mathematical models permit a more explanatory understanding of the role of internal (due to biological processes) versus external (such as those due to changes in environmental variables) drivers of transmission. Dynamical models are fundamentally important, since the biological processes driving malaria transmission are embedded within a dynamically changing environment on a range of timescales. Moreover, to meaningfully capture the emergence of new outbreaks (e.g., due to changing climate making areas previously unsuitable for transmission more favourable, human movement or mosquito dispersal to previously disease-free regions), a dynamic model is required to capture invasion dynamics.

The modelling work to date incorporating environmental variables into dynamic transmission models has almost entirely focussed on the impact of changes in temperature (although see ref. 20). Most of the attention of previous modelling work has also been on capturing equilibrium dynamics based on derivation and analysis of static quantities such as \( R_0 \), the basic reproduction number of the disease. In this chapter, we firstly consider how rainfall may be incorporated within a dynamic transmission model and secondly highlight how important disease transmission issues not examined to date can only be addressed within such a dynamic modelling framework. More specifically, we examine the impact of climate change on mosquito (and hence malaria) extinction due to changes in environmental conditions, malaria invasion dynamics in previously disease-free regions and the effects of temporal variability in climate variables on mosquito population dynamics, invasion behaviour and endemic prevalence.

**Mathematical Model Development**

In terms of developing a dynamic framework for understanding the impact of climate change on malaria transmission, a deterministic or stochastic transmission model may be developed, embedded within an environment assumed static or fluctuating over the timescale of interest. Arguably, the most realistic framework is a stochastic transmission model within dynamic environmental conditions, in which the inclusion of environmental forcing would be key to obtaining a better understanding of the effects of climate change on the spread and control of malaria. Despite this, little research to date has examined the effects of temporal forcing by climate (i.e., variability in environmental variables occurring at intra-annual (seasonal), inter-annual, decadal and longer timescales) on malaria transmission, despite receiving attention for other infectious diseases.\textsuperscript{29,30} Understanding the dynamical impact of trends and variability in climate over the next 40 years, for instance, may affect a goal of malaria eradication by 2050. Thus, although a more thorough theoretical study of the effects of temporal forcing is beyond the scope of this chapter, a key objective of the work presented here is to begin considering how seasonality effects may be incorporated into models and may impact vector population dynamics, invasion behaviour and changes in the \( R_0 \) of malaria.

We assume a deterministic model, which despite making simple biological assumptions, is expected to be sufficiently realistic to permit valuable insights into climate-driven malaria transmission dynamics. Let \( S_M, E_M \) and \( I_M \) represent the number of susceptible, exposed (but
not infectious) and infectious mosquitoes respectively, along with $S_H$ and $I_H$ representing the analogous categories in humans (where we assume a fixed latent period of duration $\tau_H$ in humans). All model parameters, functional forms and baseline values assumed in this analysis are summarised in Tables 1-3. The model is described by the ordinary differential equations

$$
\begin{align*}
    \frac{dS_M}{dt} &= b - ab_1 \left( \frac{I_H}{N} \right) S_M - \mu S_M, \\
    \frac{dE_M}{dt} &= ab_1 \left( \frac{I_H}{N} \right) S_M - \mu E_M - ab_1 \left( \frac{I_H}{N} \right) S_M (t-\tau_M) l(\tau_M), \\
    \frac{dI_M}{dt} &= ab_2 \left( \frac{I_H}{N} \right) S_M (t-\tau_M) l(\tau_M) - \mu I_M, \\
    \frac{dS_H}{dt} &= -ab_2 \left( \frac{I_H}{N} \right) S_H, \\
    \frac{dI_H}{dt} &= -ab_2 \left( \frac{I_H}{N} \right) S_H (t-\tau_H) - \gamma I_H, \\
\end{align*}
$$

where the total mosquito population $M(t) = S_M(t) + E_M(t) + I_M(t)$ and human population $N(t) = S_H(t) + I_H(t) + R_H(t)$.

**Functional Forms for Incorporating Temperature and Rainfall Effects and the Derivation of $R_0$**

We assume that the mosquito birth rate $b$ depends on rainfall (through the dependence on breeding site availability) and temperature. Let us write the birth rate as

$$
b(R,T) = \frac{B_p p_e(R) p_b(R) p_f(R)}{\tau_E + \tau_L(T) + \tau_f},
$$

Table 1. Malaria model parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Definition (and Units)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$b(R,T)$</td>
<td>Adult mosquito birth rate (per day)</td>
</tr>
<tr>
<td>$\mu(T)$</td>
<td>Adult mosquito per capita death rate (per day)</td>
</tr>
<tr>
<td>$a(T)$</td>
<td>Mosquito biting rate (per day)</td>
</tr>
<tr>
<td>$b_1$</td>
<td>Proportion of bites by susceptible mosquitoes on infected humans that produce infection</td>
</tr>
<tr>
<td>$\tau_M(T)$</td>
<td>Duration of the sporogonic cycle (days)</td>
</tr>
<tr>
<td>$l_M(T)$</td>
<td>Survival probability of infected mosquitoes over the incubation period of the parasite</td>
</tr>
<tr>
<td>$b_2$</td>
<td>Proportion of bites by infectious mosquitoes on susceptible humans that produce infection</td>
</tr>
<tr>
<td>$\tau_H$</td>
<td>Latent period of infection within humans (days)</td>
</tr>
<tr>
<td>$1/\gamma$</td>
<td>Human average duration of infectiousness (days)</td>
</tr>
<tr>
<td>$M(t)$</td>
<td>Total number of mosquitoes ($S_M(t) + E_M(t) + I_M(t)$)</td>
</tr>
<tr>
<td>$N(t)$</td>
<td>Total number of humans ($S_H(t) + I_H(t) + R_H(t)$)</td>
</tr>
</tbody>
</table>
where \( B_{E} \) is the number of eggs laid per adult per oviposition, \( p_{E}(R), p_{L}(T, R) \) and \( p_{P}(R) \) are the daily survival probabilities of eggs, larvae and pupae, \( \tau_{E}, \tau_{L}(T) \) and \( \tau_{P} \) are the duration of each development stage, \( 1/\gamma \) is the average duration of infectiousness in humans and we highlight the parameter dependence on daily temperature \( T \) (in °C) and rainfall \( R \) (in mm). The average larval duration depends on temperature as \( \tau_{L}(T) = 1/(c_{1} T + c_{2}) \), corresponding to larval daily survival probability \( p_{L}(T) = e^{- (c_{1} T + c_{2})} \), while the development rate of eggs and pupae is relatively independent of temperature.32 While rainfall has been shown to positively correlate with malaria incidence, it has also been suggested that excessive rainfall may flush out larvae and breeding sites31,33 and we assume a quadratic relationship between the daily survival probabilities of eggs, larvae and pupae and rainfall. For larvae, we assume

\[
\frac{p_{ML}}{p_{L}(R)} = \left( 1 + \frac{R - R_{L}}{R_{L}} \right) 
\]

where \( p_{ML} \) is the maximum daily survival probability (i.e., when there is optimum rainfall for mosquito breeding) and \( R_{L} \) is the rainfall limit beyond which breeding sites get flushed out and no immature stages survive. We assume an analogous expression for eggs and pupae. We also assume that temperature and rainfall act independently on the survival probability of larvae such that \( p_{L}(R, T) = p_{L}(R) p_{L}(T) \), although this is likely to be an approximation in reality. Increased evaporation of breeding sites or the melting of snow packs as temperatures increase provide two examples of this. Substituting these expressions into (2) gives the birth rate as a function of temperature and rainfall and this functional form is plotted in Figure 1.

As well as \( b(R, T) \), the biting rate \( a(T) \), mosquito mortality hazard \( \mu(T) \) and probability \( l(\tau_{M})(T) \) of a mosquito surviving the duration of the sporogonic cycle also depend on temperature.4 These are summarised in Table 2 and plotted in Figure 2, where we additionally note that \( \mu(T) = -ln p(T) \) and \( l(\tau_{M})(T) = p(T)^{e^{\tau_{M}}(T)} \). Note that the expression in ref. 4 for \( p(T) \) is true only at favourable humidities for mosquito development. Survival drops off rapidly at relative humidities below 50-60%,36 highlighting that \( p(T) \) is also indirectly dependent on rainfall through its impact on relative humidity and illustrates again the interdependence of temperature and rainfall. We assume that the proportions of bites by susceptible mosquitoes on infectious humans \( (b_{1}) \) and infectious mosquitoes on susceptible humans \( (b_{2}) \) that produce infection are independent of environmental conditions.

Arguably, one of the most important concepts in infectious disease epidemiology is that of the basic reproduction number \( R_{0} \), defined as the average number of secondary cases generated per infectious individual over their duration of infectiousness in an entirely susceptible population.37 This may be similarly defined for VBDs as the product of the number of vectors infected per person and the number of people infected per vector (over their respective infectious periods). While \( R_{0} \) may be calculated in a variety of ways for infectious disease models, calculation for models assuming homogeneous mixing is relatively straightforward. The transmission model (1) has two equilibrium
states, namely the disease-free state and the endemic equilibrium. In the latter case, it is readily shown that the prevalence in mosquitoes is given by

$$I^*_m = \frac{M(R_0 - 1)}{I(\tau_m) + \left(\frac{ab_m M}{\gamma N}\right)}$$

while for the human population

$$I^*_h = \frac{N(R_0 - 1)}{R_0 + \left(\frac{ab_1}{\mu}\right)}$$

where \((ab_1/\mu)\) is Macdonald's index of stability. Stability analysis about the endemic state demonstrates that malaria will persist when \(R_0 > 1\) where

$$R_0 = \frac{Ma^2 b_1 b_2 (\tau_m)}{\gamma \mu N}.$$

Table 3. Baseline parameter values assumed for model simulations

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Assumed Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>(B_M)</td>
<td>200</td>
<td>Dimensionless</td>
</tr>
<tr>
<td>(p_{ME})</td>
<td>0.9</td>
<td>Dimensionless</td>
</tr>
<tr>
<td>(p_{ML})</td>
<td>0.25</td>
<td>Dimensionless</td>
</tr>
<tr>
<td>(p_{MP})</td>
<td>0.75</td>
<td>Dimensionless</td>
</tr>
<tr>
<td>(R_1)</td>
<td>50</td>
<td>mm</td>
</tr>
<tr>
<td>(\tau_1)</td>
<td>1</td>
<td>days</td>
</tr>
<tr>
<td>(c_1)</td>
<td>0.00554</td>
<td>(°C days)(^{-1})</td>
</tr>
<tr>
<td>(c_2)</td>
<td>-0.06737</td>
<td>(days(^{-1}))</td>
</tr>
<tr>
<td>(e_p)</td>
<td>1</td>
<td>days</td>
</tr>
<tr>
<td>(T_1)</td>
<td>19.9</td>
<td>°C</td>
</tr>
<tr>
<td>(D_1)</td>
<td>36.5</td>
<td>°C days</td>
</tr>
<tr>
<td>(b_1)</td>
<td>0.04</td>
<td>Dimensionless</td>
</tr>
<tr>
<td>(A)</td>
<td>-0.03</td>
<td>(°C days(^{-1}))</td>
</tr>
<tr>
<td>(B)</td>
<td>1.31</td>
<td>(°C days(^{-1}))</td>
</tr>
<tr>
<td>(C)</td>
<td>-4.4</td>
<td>days(^{-1})</td>
</tr>
<tr>
<td>(b_2)</td>
<td>0.09</td>
<td>Dimensionless</td>
</tr>
<tr>
<td>(\tau_f)</td>
<td>10</td>
<td>days</td>
</tr>
<tr>
<td>(DD)</td>
<td>111 ((P. falciparum))</td>
<td>°C days</td>
</tr>
<tr>
<td></td>
<td>105 ((P. vivax))</td>
<td></td>
</tr>
<tr>
<td>(T_{min})</td>
<td>16 ((P. falciparum))</td>
<td>°C</td>
</tr>
<tr>
<td></td>
<td>14.5 ((P. vivax))</td>
<td></td>
</tr>
<tr>
<td>(\gamma)</td>
<td>1/120</td>
<td>days(^{-1})</td>
</tr>
</tbody>
</table>
Modelling Climate Change and Malaria Transmission

although note that this is conditional on assuming static environmental conditions. The expression for $R_0$ when we account for fluctuations in temperature and rainfall is considerably more complex and we postpone a more theoretical discussion of the implications of environmental variability on $R_0$ (and transmission dynamics more generally) to future work.

**Vector Population Dynamics**

The dependence on mosquito abundance in the nonlinear vector-human transmission term of (1) results in considerable sensitivity of the model to vector population dynamics. Thus, understanding the dependence of $M(t)$ on temperature and rainfall (and seasonality therein) is key.

![Figure 1. Mosquito birth rate $b(R,T)$ as a function of daily rainfall $R$ (in mm) and temperature $T$ (in °C).](image)

![Figure 2. Anopheles biting rate and survival probability and Plasmodium incubation period as a function of temperature.](image)
Adding equations in (1) related to the mosquito population gives the deterministic model for the vector population

\[ \frac{dM(t)}{dt} = b(R,T) - \mu(T)M. \]  

(7)

In reality, mosquito population dynamics behave stochastically and a full understanding of the impact of the mosquito population on transmission dynamics requires the stochastic equivalent of (7). If \( p_M(t) \) denotes the probability of having \( M(t) \) mosquitoes at time \( t \), the master equation is

\[ \frac{dp_M(t)}{dt} = b(t)p_{M-1}(t) + \mu(t)(M+1)p_{M+1}(t) - (b(t) + \mu(t)M)p_M(t) \]  

(8)

for \( M(t) = 0, 1, 2, \ldots \) and where \( p_{-1}(t) = 0 \). If we define the probability generating function to be

\[ G(z,t) = \sum_{i=0}^{\infty} p_i(t)z^i, \]

it is readily shown by successively multiplying (8) by \( z^i \) for each \( i \) and adding, we can convert (8) into a partial differential equation for \( G(z,t) \) as

\[ \frac{\partial G(z,t)}{\partial t} = b(t)(z-1)G(z,t) - \mu(t)(z-1)\frac{\partial G(z,t)}{\partial z}, \]  

(9)

which can be solved using a variety of methods. We are interested in the dependence of the mosquito population on temperature and rainfall and their seasonal variability, a full understanding of which may be obtained by solving (9). The full solution of (9), however, for the case of general \( b(t) \) and \( \mu(t) \) is complex, so we simplify the analysis by considering only the dependence of the dynamics on the mean values of these environmental factors, not their variability.

We consider the mosquito population in Tanzania where malaria is highly-endemic, but the conclusions drawn about the effects of seasonality in environmental factors are general. Using temperature and rainfall data from WorldClim\(^4\) (http://www.worldclim.org) and averaging across all regions in Tanzania, we fit the temporal profiles

\[ T(t) = T_1(1 + T_2 \cos(\omega_1 t - T_1)), \]  

(10)

and

\[ R(t) = R_1(1 + R_2 \cos(\omega_2 t - R_1)), \]  

(11)

giving the values in Table 4 (with the profiles plotted in Figure 3A. Substituting (10) and (11) into the earlier expressions for \( b(R,T) \) and \( \mu(T) \) and solving (7) gives the mean population dynamics plotted in Figure 3B, with the amplitude and frequency of vector oscillations strongly dependent on the seasonality factors \( T_2 \) and \( R_2 \). To better understand the sensitivity to variability in temperature and rainfall, Figure 3C plots the mean and standard deviation in mosquito numbers as a function of \( T_2 \) and \( R_2 \). Increasing seasonality in rainfall about a fixed mean (below the level at which flushing out occurs), broadly corresponding to more breeding sites, is found to always increase the mean size of the mosquito population, while increasing seasonality in temperature always reduces the vector population. Large amplitude variability in temperature is also found to drive mosquito populations to extinction, thus suggesting that extinction dynamics are more sensitive to changes in temperature than rainfall. This highlights that while understanding predicted changes in environmental variables from climate models is important, it is just as important to understand the impact of global warming on changes in the variability of climate variables.

Solution of (9) for general \( b(t) \) and \( \mu(t) \) would permit insight into the effects of climate variability on the stochastic population dynamics, but the complex and theoretical nature of the problem is beyond the scope of this chapter. Instead, we consider the extinction dynamics as a function of changes in mean temperature, fixing rainfall to simplify matters, but the analysis is
readily extended to include changes in rainfall at fixed temperature. When \( b(t) = b \) and \( d(t) = d \), (9) can be solved using Laplace Transforms or characteristics.\(^42\) If we assume \( M_0 \) initial mosquitoes, solution of (9) gives

\[
G(z,t) = \frac{b \cdot (z-1) e^{-zT} \cdot (1 + (z-1) e^{-\mu T}) M_0}{1 \cdot (z-1) e^{-zT} \cdot (1 + (z-1) e^{-\mu T}) M_0},
\]

whereupon substituting \( z = 0 \) gives, as a function of temperature and rainfall, the probability that the mosquito population fades out at or before time \( t \) as

\[
p(t) = e^{-\frac{b \cdot R}{\mu(T)} (1 - e^{-\mu T}) M_0}.
\]

At extreme temperatures, mosquitoes are more likely to fade-out (e.g., due to thermal death at temperatures beyond around 40˚C),\(^31\) although the risk decreases with increasing rainfall when \( R < R_L \). Beyond these early stages of mosquito invasion, the probability that the population ultimately fades out is \( e^{-\frac{b \cdot R}{\mu(T)} (1 - e^{-\mu T}) M_0} \). Thus, reliable estimates of parameters included within the birth and death models (e.g., the rainfall threshold beyond which breeding sites are washed out) will enable robust conclusions not only about when and where mosquitoes may be driven to extinction, but also the implications for malaria transmission. While a full analysis of malaria fade-out requires a stochastic transmission model, the proportionality of vector abundance and \( R_0 \) means that the above analysis yields useful insight into the impact of changes in climate variables on malaria elimination.

### Invasion Dynamics

As well as providing a useful framework for understanding how the stochastic dynamics of mosquitoes may lead to the establishment of malaria as environmental conditions change, dynamical models are vital for capturing the invasion dynamics in previously susceptible populations. In addition to the direct effects of climate change on transmission, indirect effects such as those on malnutrition, poverty or the more frequent occurrence of extreme weather events (e.g., floods or heat waves) may also contribute to changing distributions of incidence, as human and animal ecosystems and habitats become more susceptible. However, while malaria is, in general, most prevalent in the tropics and sub tropics, predictions that increases in global temperatures may lead to emergence in currently temperate regions and at higher altitudes\(^27\) are not universally accepted.\(^43\) Nonetheless, it is well-accepted that changes in climate variables influential in malaria transmission are likely to cause the global distribution of incidence to change over the next 100 years. Thus, understanding the invasion behaviour into new regions represents a key challenge and one which has considerable public health implications.
Figure 3. A) Annual temperature and rainfall profiles (10) and (11) fitted to WorldClim data (http://www.worldclim.org) averaged across Tanzania, (B) predicted mosquito population dynamics for Tanzania from the solution to (7) (with parameters from Tables 2 and 3) and (C) the solution to (7) as a function of increasing seasonality in temperature ($T_2$) and rainfall ($R_2$).
Again, a more detailed stochastic transmission model, incorporating seasonally-varying environmental variables, is required for a more thorough understanding of invasion dynamics, but considerable progress about the expected behaviour can be obtained from (1). When most of the population are initially susceptible, $S_H(t - \tau_H) = N$ and $S_M(t - \tau_M) = M$, whereupon substituting into (1), the invasion dynamics are described by

$$\frac{dI_H}{dt} = ab_H \left( \frac{I_H(t - \tau_M)M}{N} \right) I_M(t) - \mu I_M,$$  (14)

and we look to solve for the growth rate of $I_H(t)$. We approach this problem in a way that allows illustration of the method and thus how to apply the technique to more complex transmission models.

In essence, we look to rewrite the system as a single equation in $I_H(t)$ and its derivatives, which we note currently depend on factors proportional to $1, I_H(t), I_M(t - \tau_H)$ and $dI_H(t)/dt$. Here, we have a dependence on four factors, but only two independent equations. However, a third independent equation may be obtained by differentiating (14), since this will depend on some combination of the same four factors. Considering the resultant equation, together with (14), at time $t + \tau_H$ gives the three independent equations

$$\frac{dI_H(t + \tau_H)}{dt} = ab_H I_M(t) - \gamma I_H(t + \tau_H),$$

so that $I_H(t)$ and its derivatives now depend on three factors and we have three independent equations. If $J$ is the $3 \times 3$ matrix with elements containing factors proportional to $I_H(t)$ and its derivatives, we can rewrite (16) as the matrix equation

$$J \begin{pmatrix} 1 \\ I_H(t) \\ \frac{dI_H(t)}{dt} \end{pmatrix} = 0,$$  (17)

whereupon solving $\det J = 0$ leads to

$$I_H(t + \tau_H) + (\mu + \gamma)I_H(t + \tau_H) + \mu I_M(t) - R_0 I_H(t - \tau_H) = 0$$  (18)

for the number of infectious humans. Substituting the trial solution $I_H(t) = e^{rt}$ gives

$$r^2 e^{rt} + (\mu + \gamma)re^{rt} + \mu e^{rt} - R_0 e^{rt} = 0,$$  (19)

where $r$ is the real-time growth rate and the solution reduces to the standard growth rate equation for an SEIR model (with analytical solutions) when $\tau_H = \tau_M = 0$. Substituting parameters and functional forms from earlier, Tables 2 and 3 allow numerical solution of (19) and Figure 4A plots the outbreak doubling time $T_D = \ln(2)/r$ as a function of temperature at different mosquito densities. The rate of spread into a susceptible population is found to be extremely sensitive to temperature, with a clear window around 32-33°C where the doubling time is shortest. At lower
temperatures, fewer mosquitoes survive long enough for the sporogonic cycle to complete, while at higher temperatures, mosquito survival probability drops off rapidly. Invasion dynamics are also found to be strongly dependent on vector abundance, itself driven by rainfall, with doubling times around 4-6 weeks at vector densities typical of rainy seasons and increasing to the order of months at lower abundances typical of dry seasons.

A more theoretical analysis of the effects of seasonality in the vector population on the growth rate has been considered elsewhere and the applicability of such methods to more complex

![Figure 4. A) Doubling times of a *P. falciparum* outbreak as a function of temperature and mosquito density at fixed rainfall (and in the absence of seasonality). B) Predicted doubling times for *P. falciparum* and *P. vivax* with and without seasonality in temperature (where *m* = 40 and all other parameters as per (A)).]
systems is beyond the scope of this chapter. However, approximate preliminary insight may be obtained by substituting (10), with parameter values given in Table 4, into (19) for parameters that depend on temperature and the results are shown in Figure 4B for *P. falciparum* and *P. vivax*. In the absence of seasonality, there is little difference between *P. falciparum* and *P. vivax*, with the latter spreading marginally quicker and potentially resulting in a greater outbreak threat at higher altitudes given the lower critical temperature for transmission (around 14.5°C and 16°C for *P. vivax* and *P. falciparum* respectively). Figures 4A,B combined also highlight that malaria invasion is considerably more sensitive to vector density than the strain of *Plasmodium* parasite. The crude inclusion of moderate seasonality in temperature, around three times greater than seasonality observed across Tanzania, is found to have little effect on the doubling times, although a more thorough analysis is required to fully understand the effects of variability in environmental variables on malarial invasion.

**Implications for $R_0$ and Mapping Risk**

It is readily shown from (1) that successful establishment and invasion of malaria into a previously unaffected population will lead to endemicity when $R_0 > 1$. Given the dependence of the vector population dynamics on temperature and rainfall, together with the functional forms contributing to $R_0$ in Table 2, an analytical expression for $R_0$ as a function of mean temperature and rainfall may be derived by substituting into (6). For the purpose of this analysis, we assume that the vector population is in equilibrium, so that $M(R,T)/b(R,T)/u(T)$ and there is no seasonality in rainfall nor temperature (so $R_0 = T; = 0$). For brevity, we do not write down the full expression, but $R_0$ as a function of these two variables is plotted in Figure 5.

Parameter uncertainty in the vector population model means that while precise quantitative conclusions should not be drawn from Figure 5, the qualitative dependence of $R_0$, capturing both the population risk of malaria becoming endemic and individual infection risk, on temperature and rainfall remains robust to these uncertainties. As with the invasion dynamics, we observe a clear window for optimum malaria transmission around 32-33°C, corresponding to where the balance between mosquitoes surviving long enough for completion of the sporogonic cycle and the rapid decline in mosquito survival at high temperatures is optimised. The approximate quadratic dependence of $R_0$ on temperature also offers insight into the question of how changes in local transmission risk may shift due to increases in temperature. Consider an endemic region currently at mean temperature $T$ experiencing an increase in temperature by an amount $\Delta T$. If

![Figure 5. The dependence of $R_0$ on temperature and rainfall.](image-url)
$T_{\text{max}}$ represents the temperature at which the $R_0(T)$ profile peaks, it is clear that if $T + \Delta T < T_{\text{max}}$, malaria prevalence will get significantly worse in that region, while if $T > T_{\text{max}}$, global warming will lead to a decline in mosquito survival and a reduction in transmission.

For the case of Tanzania, Figure 6 plots the predicted mean temperature distribution for April (when rainfall peaks) under the A2a emission scenario$^6$ (data from WorldClim and where the calculation of mean temperature assumes uniformly distributed temperatures between monthly maxima and minima). Analysis of the expression quantifying the relationship between $R_0$ and temperature, and substitution of parameter values in Table 3, shows that $T_{\text{max}}$ is around 32.9°C for $P. falciparum$ and $P. vivax$. Temperature data presented in Figure 6 shows that the maximum mean temperature is predicted to be around 32.6°C for Tanzania and thus, given the monotonic relationship between $R_0$ and prevalence from (5) and the fact that we are in the regime $T + \Delta T < T_{\text{max}}$, we predict malaria prevalence to increase across Tanzania around the peak rainy season. The severity of the increase, however, will be regionally dependent and this highlights the need for implementation of control and mitigation strategies at a local level. Improvements in parameterisation of the model will lead to improved ability to make quantitative predictions about spatiotemporal dynamics and control, as well as addressing how changes in rainfall distribution, as well as temperature, may lead to changes in transmission risk. The impact of excessive rainfall, for instance, on vector abundance and transmission is only qualitatively captured here. Such an analysis also highlights the usefulness of assessing control strategies within a dynamic framework. Questions such as the relative magnitude and timing of interventions, as well as the impact of parameter uncertainty, variability and heterogeneity, be they human, mosquito or parasite, may also be considered, along with evaluating such questions within the context of limited resources. The direct impact of seasonality in vector dynamics on $R_0$ has been considered from a theoretical perspective elsewhere and we postpone a more thorough analysis of the dependence of control on variability in environmental conditions to future work.

Figure 6. Predicted mean temperature distribution across Tanzania in April under the A2a emission scenario (data from WorldClim).
Conclusion

The complex transmission dynamics of malaria are strongly influenced by environmental conditions, with temperature, rainfall, humidity, wind speed and altitude, among others, shown to affect incidence. However, despite the opinion that malaria may represent one of the most sensitive VBDs to climate change over the next 100 years, most previous work on interactions between climate and malaria has been largely based on the use of statistical modelling approaches with little corresponding work carried out on using mathematical models to better understand the dynamical effects of this influence on disease transmission dynamics. Such process-based models may not only represent powerful tools for understanding how future trends and variability in key climate variables may affect global changes in malaria risk via their capacity to deal with the likely nonlinear and complex feedbacks in climate-pathogen relationships, but when combined with the ability to simulate ‘what-if’ scenarios, they may also represent valuable tools for policy-makers and future public health planning.1,7,11

In this chapter, we have shown the crucial need for the development of dynamic modelling frameworks to better understand the dependence of malaria transmission on changes in temperature and rainfall. The most general and probably most realistic modelling framework one could develop here would be a stochastic transmission model incorporating key environmental variables that are continuously changing. While we expect the field to undoubtedly progress towards this end, it is invaluable at the early stages of our understanding to construct simpler climate-linked transmission models that permit an understanding of how global warming may affect the burden of disease. Thus, using a relatively simple deterministic transmission model, we have been able to demonstrate the importance of accounting for the dependence of mosquito abundance on temperature and rainfall, which by entering the dynamical model through the nonlinear transmission term, we show can strongly influence the establishment probability of malaria in previously disease-free regions. The potential for mosquito extinction with large seasonality in temperature highlights a second key result, namely that it is important not only to examine the effects of future average trends in climate variables, but also the variability about these trends.

In reality, this variability may represent genuine fluctuations of climate variables or uncertainty from climate model predictions or human behaviour (corresponding to a range of possible emission scenarios). Indeed, it is clear from this chapter that seasonality and variability in climate on longer timescales can have profound effects on establishment, invasion and persistence and this remains a priority area for future research. Changes in patterns of malaria incidence due to changes in environmental conditions on longer timescales, e.g., due to the effects of El Niño-Southern Oscillation (ENSO), have been considered in the literature44 47 and this continues to merit further research. Similarly, it is clear that when these studies are combined with corresponding concepts from infection system dynamics, such as the dependence of $R_0$ on temperature and rainfall and the identification of an optimal transmission window, such integrated analyses will prove crucial to improving understanding about how long-term global climate change will affect local environmental conditions and, in turn, whether a region is likely to experience a worsening or improvement in prevalence as global warming becomes more severe.

The results also highlight that considerable work, experimental, theoretical, modelling and policy-based, still remains to be satisfactorily carried out and modelled in an integrated fashion if we are to more realistically capture the impact of climate change on disease transmission. Although certain aspects of transmission dynamics are physiological (and therefore deterministic) drivers of transmission, it is clear that heterogeneities across the human, mosquito and parasite populations introduce considerable uncertainty into the system. It should also be borne in mind that a more realistic modelling approach should take spatial heterogeneities into account and thus realistic transmission models need to be spatial if they are to better predict spatiotemporal disease persistence and spread. A key challenge in incorporating environmental variables into malaria models is also selecting the appropriate scale at which to model, not only spatially, but over the most appropriate timescale and hierarchical level.48 This is driven not only by the resolution of available climate data (either from remote sensing or output from climate models), but also from the knowledge that modelling at too fine a scale may translate poorly into global observables, while oversimplifying
local heterogeneities may neglect key environmental or biological processes influencing observations. Another challenge is how best to integrate ecological drivers with sociological processes of disease transmission in vulnerable communities, an area only now beginning to be examined.\textsuperscript{1,2,23,25,26,43}

The quality of incidence data, as well as the reliability and depth of experimental knowledge on the effects of environmental variables on the transmission cycle, also requires considerable research and this remains an important priority. The impact of excessive rainfall, for instance, on vector abundance and transmission is only qualitatively captured here. Mathematical models can thus act as a useful guide for data collection by identifying areas where improvements in data quality may lead to substantial improvements in their explanatory power.

Finally, our results also underscore the usefulness of assessing control strategies within a dynamic framework. In particular, we suggest that only such frameworks will allow fuller exploration of key questions such as the relative magnitude and timing of interventions, as well as the impact of parameter uncertainty, variability and heterogeneity, be they human, mosquito or parasite, along with reliable evaluation of these questions within the context of limited resources. The dynamical modelling approach described here may thus provide a useful framework not only for obtaining a better understanding of the integrated impact of climate change on disease transmission dynamics, but also to serve as a tool for policy-makers developing mitigation and intervention strategies that may be used to tackle the potentially tough challenges that lie ahead.

\textbf{Acknowledgements}

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\textbf{References}

Modelling the Transmission of *Trypanosoma cruzi*: The Need for an Integrated Genetic Epidemiological and Population Genomics Approach

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Abstract

This chapter describes what should be an integrated approach to the genetic epidemiology and population genomics of Chagas disease. Many studies have been conducted on the genetic diversity of *Trypanosoma cruzi* and the various triatomine bug species able to transmit Chagas disease. Far less research has analyzed the role played by the host's genetic variability on the transmission and severity of the disease. An integrated genetic epidemiology/population genomics approach would analyze these three components of the transmission chain together as well as their possible interactions (co-evolution phenomena). This is facilitated by the recent impressive progress in mega biotechnologies and by the fact that Chagas disease is an ideal model for experimental evolution approaches.

Introduction

For several years, I have pleaded for an integrated approach to the epidemiology of infectious diseases. Most authors focus on only one component of the transmission chain of a given disease: either the host or the pathogenic agent, or (in the case of vector-borne diseases such as Chagas disease) the vector. It is clear that these three actors play a role in the same performance as part of a single biological phenomenon: the coevolution between the host, the pathogen and the vector. An integrated approach to this global phenomenon is therefore sorely needed. Scientists have a natural tendency to specialize, even to overspecialize. For example, my experience tells me that people working on the population genetics of African trypanosomes are poorly informed about similar studies conducted on *T. cruzi*. This is all the more distressing since comparative approaches are extremely informative, delineating the general laws that govern the evolution of organisms, while underscoring the specificities of each case.

When transmission and severity of infectious diseases are concerned, it is very hard to know whether different genotypes of the pathogen are able to cause different clinical forms without knowing what the role of the host's genetic diversity could also be. An integrated approach consists in analyzing both phenomena jointly and in including studies on the vector when researching vector-borne diseases. Even more specifically, it aims at dissecting the possible interactions between the three (co-evolution phenomena). The international congresses MEEGID (Molecular Epidemiology and Evolutionary Genetics of Infectious Diseases) and the journal *Infection, Genetics*...
Modelling the Transmission of Trypanosoma cruzi and Evolution (http://www.elsevier.com/locate/meegid) both aim to broaden the scope of this research by encouraging and practicing such an integrated approach. Interestingly, Chagas disease is an exemplary case for this approach.5 I will explain below why this is more than ever true today. I will start by describing today’s situation for each of the three actors: the pathogen, the vector(s) and the host(s).

Trypanosoma cruzi, World Champion of Pathogens for Population Genetics

The scientific community that works on Chagas is a tiny one. In fact, almost all of us know each other personally. It is like a big family, in which the contribution of the Latin American element has been and still is, prominent. Considering the small size of this group, T. cruzi would not be expected to be the pathogenic agent whose genetic diversity is among the best known, possibly more than the heavy weights Escherichia coli and Candida albicans. This fact leads me to advocate setting up T. cruzi as one of the landmark models of modern biology, together with the legendary E. coli, Drosophila melanogaster, Caenorhabditis elegans and Mus musculus. A historical view of how this unexpected situation came about is useful.

The Isoenzyme Saga

In the late 1960s and the 1970s, isoenzyme markers became immensely popular.6 Isoenzymes are electrophoretic variants of the same enzyme that reveal the sequence diversity of the genes that code for them (Fig. 1). For the first time, they made it possible to directly unravel the genetic diversity of organisms: population genetics stopped being a speculative affair and entered the enchanted world of direct observation. Thousands of papers based on what had quickly become the gold standard have been published, covering virtually the entire living reign. This made it possible to firmly establish the mendelian inheritance (codominant markers) of isoenzymes. Fortunately, bacteriologists and parasitologists did not miss the train and took advantage of the interesting properties of isoenzymes to clarify the subspecific variability and population structure of their pet bugs.7,8 This has been especially true for T. cruzi, in particular through the pioneering studies of Miles and collaborators9 and studies based on a population genetics approach by our group.10 It is worth noting that all the results on the population structure and evolutionary pattern of T. cruzi based on isoenzymes have been fully confirmed by more fashionable molecular methods (see below).

Figure 1. An isoenzyme gel for the genetic locus Glucose phosphate isomerase (Gpi) showing the genetic polymorphism of different genetic subdivisions (discrete typing units; see text) of T. cruzi (experiment and photograph by Jenny Telleria, IRD Montpellier, France).
The Molecular Biology Wave
In the 1980s, “the Maniatis”11 became the laboratory bible and young researchers dangerously started spotting their fingers with ethidium bromide, fascinated by the orange fluorescent banding patterns visible on gels under UV light. In a flash, isoenzymers became nerdish. In the Chagas community, the first group to step on stage was Morel’s group,12 who showed off the very esthetic schizodeme profiles of strains (RFLP patterns of kinetoplast DNA). Many other molecular techniques were later added to the display of strain typers, including miniexon gene polymorphism,13 microsatellites14 and random primed polymorphic amplified DNA or RAPD,15 among others. All these studies revealed a striking pattern: they showed a constantly converging picture of \( T. cruzi \) subspecific variability, also congruent with the picture drawn by the nerdy isoenzymes. This was all grist for the population geneticists’ mill (see below).

The Sequencing, Genomic and Postgenomic Era
We are still in the midst of this revolution. \( T. cruzi \) was not left by the wayside in this new wave. Its genome is now fully sequenced.16 Several other strains are currently being sequenced as well (El Sayed, personal communication). Microarrays and proteomic analyses are on the way. The challenge is now to channel and filter the coming flood of data so that it remains informative. Fortunately, in the case of \( T. cruzi \), theory came before the data flood and we have a robust population model available that provides a relevant framework for all studies investigating the genetic, genomic and phenotypic diversity of this parasite. The story is briefly expounded below.

Is \( T. cruzi \) a “Good” Species?
This is the first question to raise from an epidemiological point of view, especially where molecular epidemiology is concerned. Defining a good species refers to the definition of a species itself. Here is not the place to rekindle the debate.17 Briefly, species are generally defined as: (i) a mating community (the biological species concept) or (ii) a clade (a monophyletic line with only one ancestor; the phylogenetic species concept) or (iii) a set of organisms that share remarkable phenotypic traits (the phenotypic species concept). Undoubtedly, \( T. cruzi \) meets the criteria for (ii) and (iii). All phylogenetic studies have brought all \( T. cruzi \) strains into a unique clade that is distinguishable from closely related taxa (\( T. cruzi \) marenkellei, a close cousin of \( T. cruzi \) that parasitizes bats, is the best example of such an outgroup). Moreover, all \( T. cruzi \) strains share a set of specific phenotypic characters (morphological aspects, vectorial transmission by triatomine bugs, potential host range extended to all mammals, but restricted to them, geographical distribution limited to the new world, potential pathogenicity). Consequently, from the point of view of the phylogenetic and phenotypic concepts, \( T. cruzi \) is a good species. The fact that \( T. cruzi \) is a unique clade makes it possible to design many molecular markers that will be specifically shared by all strains of the taxon (in the cladistic jargon, synapomorphic characteristics; see also the concepts of DTU and tags below).

The Population Structure of \( T. cruzi \): Sex or No Sex?
In the 1980s, \( T. cruzi \) found itself enrolled in the noisy clonality/sexuality debate that roused bacteriologists and parasitologists.18 There is now consensus that this parasite is somewhat of a paradigm of the preponderant clonal evolution model.19,20 This means that its multi-locus genotypes copy themselves like genetic photocopies and are extremely stable in space and time, even at an evolutionary scale. Two important features of the model, often neglected by scientists who read only the abstracts of the papers, are that: (i) sex (in the broad sense: any kind of genetic exchange between two different cells) is not supposed to be totally absent, but only rare and not frequent enough to break the prevalent pattern of clonality; (ii) clonality is taken here in its genetic meaning and refers to all situations where descendant multi-locus genotypes are virtually identical to parental genotypes, whatever the actual mating system. It could be mitotic clonality, or parthenogenesis, or an extreme selfing situation, or extreme homogamy.3

Scarcity or absence of genetic recombination has been established in \( T. cruzi \) by evidencing an extreme linkage disequilibrium (LD; nonrandom association of genotypes occurring at separated
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loci, as shown in Figure 2. A striking manifestation of LD is that phylogenetic trees established from different genetic markers are very similar (Fig. 3). The contrary arises in those pathogens where recombination is abundant, such as the bacterium *Helicobacter pylori*.

The model has considerable implications in terms of applied research: (i) the stability of multi-locus genotypes makes them ideal targets for molecular epidemiology (strain typing and tracking); (ii) since genotypes are genetically separated from each other, their evolutionary fate is to accumulate more and more divergent mutations, including for those genes that govern medically relevant properties (pathogenicity, drug resistance). Clones that are genetically similar should tend to be also phenotypically similar and vice versa.

There is also a consensus on the number of genetic subdivisions that are observable within *T. cruzi*: there are two main clusters, which have been named by a group of anonymous experts *T. cruzi* I and *T. cruzi* II (TC I and II). TC II is itself subdivided into five lesser clusters (TC IIa-e). There has been a debate on the actual evolutionary nature of these clusters. The presence of

Figure 2. Four electrophoretic experiments, two isoenzyme gels with two different genetic loci (top), two random primed amplified polymorphic DNA (RAPD) surveying two different genomic regions (bottom). The same *T. cruzi* strains are surveyed on the four gels; however, RAPD experiments have two additional strains. On the four gels, only two main genotypes (DTU 1 and DTU 2; see text) are observed; lines 1-7 and 8-12 for isoenzymes, lines 1-9 and 10-14 for RAPD. M lines on RAPD gels are molecular weight ladders. Genotypes DTU 1 are constantly linked together and the same is observed for genotypes DTU 2. Crossed genotypes, which would be the result of genetic recombination (for example gel A line 1/gel B line 8 + gel C line 1 + gel D line 10), have never been observed on more than 600 *T. cruzi* strains characterized by our team to date. This kind of strong association between genotypes occurring at different loci is by definition a linkage disequilibrium (see text) and is a manifestation of preponderant clonal evolution in *T. cruzi*. Copyright 1993, Proc Natl Acad Sci USA. Tibayrenc M et al. 90:1335-39.
Figure 3. Two phylogenetic trees depicting the evolutionary relationships among *T. cruzi* genotypes: isoenzymes (left) and RAPD (right). The strong similarity between the two trees is an extreme manifestation of linkage disequilibrium (see Fig. 2). Copyright 2000, Int J parasitol. Brisse S et al. 30:35-44.
some genetic recombination and hybrid genotypes prevents one from calling them clades. We have proposed the operational term of discrete typing units (DTU):\(^1\) sets of strains that are genetically more similar to each other than to any other strains and that share common molecular or serological markers (tags). This terminology is widely accepted in the Chagas research community.\(^4\) We will henceforth refer to DTU I, IIa, IIb, IIc, IId and IIe. Figure 1 shows that isoenzyme electrophoresis shows drastically distinct patterns between \textit{T. cruzi} DTUs.

As already noted, the clonal model does not state that sex (recombination) is absent, but only that it is rare and is not a mandatory mechanism of reproduction, as it is in humans and fruitflies. Several natural \textit{T. cruzi} genotypes appear to have a hybrid origin, subsequently stabilized by clonal propagation.\(^23\)-\(^26\) Moreover, the potential for genetic recombination in \textit{T. cruzi} has been fully confirmed by experiments.\(^27\)

The story therefore can be summarized as follows: \textit{T. cruzi} undergoes predominant clonal evolution, which permits it to stabilize favorable genotypic combinations. Occasionally it mates, which makes it possible to rapidly generate new genotypes subsequently stabilized by clonal propagation. It is a typical case of reticulate evolution, a pattern also observed in many plant species.

The important facts to be kept in mind for the topic of this chapter: (i) the above-mentioned picture of subspecific variability could be refined; however, it is improbable that further studies will upset this picture. (ii) \textit{T. cruzi} DTUs are robust units of analysis corroborated by many different studies, easy to specifically identify with appropriate tags. They actually behave like distinct taxa. They constitute convenient units of analysis for epidemiological tracking, applied research (vaccine and drug design) and experimental evolution studies.

However, a wide gap in our knowledge on \textit{T. cruzi}'s genetic variability persists: the biological and epidemiological differences among \textit{T. cruzi} clonal genotypes and DTUs remain imperfectly known. A relation between \textit{T. cruzi} isoenzyme genotypes (zymodemes) and clinical forms of the disease has been long suspected,\(^28\) but never fully confirmed. Long-term experimental surveys in our laboratory have shown significant statistical associations between the genetic distances recorded among \textit{T. cruzi} genotypes on one hand and biomedical differences on the other hand\(^29\)-\(^34\) (see also refs. 35-36). The biomedical properties that have been surveyed include growth speed in acellular and cell cultures, transmissibility through vectors, pathogenicity in mice and in vitro in vivo susceptibility to antichagasic drugs (série). Only one study investigating in vitro drug sensitivity showed no statistical association.\(^37\) There is clearly “something” there when the working hypothesis of a link between \textit{T. cruzi} genotypic and biological diversity is tested. However, the picture is far from black and white and the statistical sets are limited for the moment. An interesting hypothesis states that different \textit{T. cruzi} genotypes have different organ tropisms.\(^38\) However, no firm conclusions have been reached on this point either. This long-term debate therefore remains partly unanswered.

**The Second Actor: The Vector**

It would be more accurate to say the vectors, for there are many of them. Triatomine bugs are true bugs (heteropterous). They make up a subfamily (Triatominae) within the family reduviidae, which are basically predator bugs. The triatominae turned out to be obligatory blood feeders, adults of both genders and larvae of all stages.

From a population genetics and evolutionary point of view, the harvest is not as plentiful as for \textit{T. cruzi}. However, many studies have been conducted on triatomines, which makes them a rather well-known group.

It is now strongly suspected that the adaptative trait of strict hematophagy occurred several times in the evolutionary history of reduviidae. The triatominae are thought to be a polyphyletic group (Schofield, personal communication). Three main genera are recorded in the triatomines: \textit{Rhodnius},

\[ \text{a. In an expert meeting held in Buzios (Brazil), August 2009, in which the author participated, these DTUs have been validated. However, they have been renumbered I, IV, II, III, V, and VI, respectively.} \]
Triatoma and Panstrongylus. It is being debated whether each of these genera considered alone is monophyletic. Within these genera, many different species are able to transmit T. cruzi.\textsuperscript{39}

Triatomines exhibit various interesting phenomena of partial speciation and sibling speciation, which makes them informative models for scientists interested in speciation processes. These situations have been conveniently explored by population genetics approaches.

Since the first pioneering isoenzyme studies of the 1980s,\textsuperscript{40} many population genetics and phylogenetics analyses have been conducted on various triatomine species with markers ranging from microsatellites\textsuperscript{41,42} to RAPD,\textsuperscript{43,44} mitochondrial genes\textsuperscript{45} and gene sequences.\textsuperscript{46-48} Interestingly, this rather classical molecular display has been completed by the complementary tools of cytogenetics\textsuperscript{49} and morphometric analysis.\textsuperscript{42,50,51}

The world of triatomine bugs is vast, with an extreme taxonomic, phylogenetic and ecological diversity, making this group a gold mine for population genetic and evolutionary studies.

Within the theme of the present chapter, an immense field of knowledge remains to be explored, with several unanswered key questions: (i) have all species of triatomines the same vectorial capacity (the answer is probably no); (ii) do species of the same genus tend to have comparable vectorial capacities; (iii) within the same species, do different populations have the same vectorial capacity; (iv) are different strains, genetic clones and DTUs of T. cruzi equally transmitted by triatomine bugs; (v) more generally, what complex phenomena of co-evolution and co-adaptation exist between the vector and the pathogen.

The Host

Again, it is preferable to say the hosts, since all mammalian species are potentially able to be contaminated by T. cruzi. Triatomine bugs feed on birds as well; however, these animals are resistant to Chagas disease. The reservoir of the disease is therefore virtually unlimited, since wild mammals as well as domestic animals are hosts. Chagas disease is a typical zoonosis. The transmission is enhanced by the fact that humans and domestic animals often live close together. For example, in Bolivia, farmers often have pet guinea pigs in their kitchen.

As for vector species, this wide range of mammalian host species provides abundant matter for co-evolution analyses, either in field surveys or in experimental studies. Dogs with their many breeds have been a choice model for experimental studies on T. cruzi pathogenicity.\textsuperscript{52-55} Of course mice remain the easiest model to handle in experimental Chagas disease.\textsuperscript{33,56,57} However, we still lack an overall picture of differential Chagas pathogenicity among different mammalian species and different populations and breeds of the same species. Moreover, the animal models do not clearly identify the candidate genes\textsuperscript{58} that could be involved in the susceptibility to Chagas disease.

As for the human species, it is quite unexpected that our knowledge of the impact of human genetic diversity on the severity and clinical diversity of Chagas disease is poorly known. By comparison, in this field, we know much more about malaria,\textsuperscript{59} tuberculosis,\textsuperscript{60} leprosy,\textsuperscript{61} schistosomiasis\textsuperscript{62} and leishmaniosis.\textsuperscript{63} This is all the more astonishing since Chagas disease should constitute a very favorable case to study this problem, for two reasons:

i. As for leprosy, the clinical phenotypes of the disease are quite clearly defined. It starts with an acute phase that can be discreet or on the contrary shows a severe septicemic syndrome. Approximately 10% of patients die at this stage. Those who survive enter the undetermined phase with no symptoms. Parasites hide in the cells. Unfortunately, after at few years, roughly 30% of patients begin showing symptomatic Chagas disease. The majority of them suffer from a cardiac form that leads to severe cardiac insufficiency. Digestive forms account for 3% of patent Chagas disease cases. They have the form of megacolon or megaesophagus. Some patients have a cardiac and digestive form. Some patients have no clinical symptoms, but have electrocardiogram abnormalities. Others have symptoms but negative serological tests.\textsuperscript{64} Lastly, there are great differences in the way that antichagasic drugs act on different patients.\textsuperscript{36} There is therefore a wide span of well-defined clinical forms to analyze in linkage studies.

ii. Chagas disease strikes a range of genetically diverse human populations, including different ethnic groups (mainly Amerindians, Causasians, Africans and people of mixed ancestry)
that could exhibit different susceptibilities to the disease. For example, Aymara Indians in Bolivia and Peru have lived only on the Altiplano for several thousand years. They have therefore long been protected from Chagas disease, since triatomine bugs do not survive at such altitudes. Their immunity against the disease could be lesser than that of other Amerindian populations who have coexisted with triatomine bugs for a long time.

A debate of importance for studying genetic susceptibility to Chagas disease is to know whether chronic Chagas disease is actually an autoimmune disease\(^{56}\) or if the parasite is still present in the host's cells and causes a chronic inflammatory response. Both PCR experiments and the classical xenodiagnosis strongly suggest that the second hypothesis holds true.

The meager results obtained on human genetic susceptibility to Chagas diseases and its various clinical forms are now summarized. A familial component in the cardiac form of chronic Chagas disease has been suggested from a Brazilian study.\(^{66}\) These results clashed with the results obtained from an Argentinian survey.\(^{67}\) HLA polymorphism associated with Chagas disease has been analyzed in many studies. Some of them reported a total lack of association,\(^{68}\) while on the contrary other research observed associations.\(^{69}\) Some serological parameters seem to exhibit a notable heritability, as evidenced by extended pedigrees. In Brazil, the heritability of Chagas seropositivity would be no less than 0.556.\(^{70}\) More specifically, the heritability of the IgA and IgG levels would be 0.33.\(^{71}\) I have myself observed that many subjects in Bolivia seem to be quite resistant to Chagas disease contamination. In some areas where 100% of the thousands of triatomine bugs I have collected harbored *Trypanosoma cruzi*, only 50% of the children who lived in those areas were seropositive. Nevertheless, all these children had been bitten hundreds or thousands of times in their lifetime.

To make a long story short, our knowledge on the role of human genetic diversity on Chagas transmission, severity and clinical polymorphism is extremely patchy and contradictory. This is the weak spot in the array of knowledge needed to design an integrated genetic epidemiology approach of Chagas disease.

**The Future**

Chagas disease could become a paradigm academic case for an integrated genetic epidemiology approach to transmissible diseases.\(^{5}\) However, we still are far from this goal. Our knowledge on the parasite's genetic diversity is fairly advanced. Thanks to phylogenetic/population genetic/morphometric studies, the complexity of the world of triatomine bugs is being deciphered little by little. Although the role of the host's genetics remains poorly known, it is only a matter of applying the necessary effort. A black hole remains in our knowledge on the interactions and reciprocal impact between these three components. Actually, this is the case for all transmissible diseases. Research is strongly compartmentalized and parasitologists, entomologists and human/mammal geneticists rarely interact.

In the case of Chagas disease, the gaps in our knowledge could be filled using two approaches:

**Field Studies**

New powerful technologies (high-throughput sequencing, genome-wide scanning, microarrays, real-time PCR, high-resolution morphometric analysis) should make it possible to considerably improve our knowledge on the parasite, the vector and the host.

When humans are considered, as it is the case for any disease, studies on Chagas disease could greatly benefit from the megaprojects presently running on human genetic diversity, mainly the HapMap project (http://www.hapmap.org/). Genomics is presently making exponential progress and the few scientists working on human genetic susceptibility to Chagas disease should climb aboard this high-speed train. As emphasized above, in Chagas genetic epidemiology, the human side is the weak link in the chain. Of course studies dealing with the parasite and the vector would also greatly benefit from the impressive progress reached in genomics, proteomics and bioinformatics analysis.
Although it is important that specialists working on the parasite, the vector and the host continue developing their specific fields, our knowledge on these actors sorely needs them to interact more, including in field studies. It is crucial that epidemiologists, mammalogists, entomologists, human geneticists and clinicians include parasite isolation and characterization in their programs, so that it is better known, for example, which *T. cruzi* genotypes are more frequently harbored by given populations of given triatomine bug species, or more frequently isolated from given clinical forms of the disease. Such goals should be attainable by setting up multidisciplinary research consortia. Networking organizations would be an efficient means to attract valuable sources of funding. The long-term goal is to build an integrated population genomic approach,72 joining together genomic and postgenomic studies investigating the pathogen, the vector and the host at the population level. Obviously, such an ambitious approach can be accomplished by extended collaborations between many teams of diversified expertise only.

**Experimental Evolution**

Integrated genetic epidemiology in the field is hard and involves heavy research protocols. Experiments on the evolution of Chagas disease are easier to design, since Chagas disease is an extremely favorable model for this kind of approach5 and a complete Chagas transmission cycle is easy to maintain in experiments. The parasite is easy (although harmful) to culture, either in acellular or in cell (Vero cells) cultures. Many triatomine bug species are easy to rear in the laboratory. Various mice breeds can be infected by *T. cruzi*. Other laboratory animals such as dogs can also be used. A complete transmission cycle can even be maintained without laboratory animals by using artificial feeding devices for triatomine bugs. It is therefore quite feasible to survey the interactions between the parasite’s, the host’s and the vector’s genetic variability by varying only one parameter at a time. A promising avenue of research is to explore the interactions between mixtures of *T. cruzi* genotypes, a situation that is frequent in natural cycles, in vectors as well as in Chagas disease patients. Our working hypothesis is that there is some sort of cooperation between different clonal genotypes that infect a single host (Ann Rev gen), so that the whole is more than the sum of the parts. This seems to have been verified in a number of preliminary experiments involving such mixtures of clonal genotypes.34

**Conclusion**

Chagas disease potentially constitutes a paradigm model for the integrated genetic epidemiology and integrated population genomic approaches. However, the Chagas scientific community, although talented, is tiny, a handicap in reaching this goal. It is therefore indispensable to attract other scientists to the enterprise. This is why it is crucial to sell *T. cruzi* and Chagas disease as a reference model for basic biology and evolution, as it is the case for *Escherichia coli*, *Candida albicans*, *Caenorhabditis elegans*, *Mus musculus* and *Drosophila melanogaster*.

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