Parameter estimation and optimization for biological mathematical models using Bayesian statistics

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PARAMETER ESTIMATION AND OPTIMIZATION FOR BIOLOGICAL MATHEMATICAL MODELS USING BAYESIAN STATISTICS

A Thesis

Submitted to the Graduate Faculty of the Louisiana State University and Agricultural and Mechanical College in partial fulfillment of the requirements for the degree of Masters in Applied Statistics in

The Department of Experimental Statistics

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I would like to thank Dr. Guo for encouraging me to stick with it, and Dr. Escobar for suggesting I continue in the initial probability class. I would also like to thank Dr. Li for serving on my committee.
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Abstract

In the field of biology, mathematical models are increasingly used to address biological questions and the large data sets generated in experimental studies. Mathematical models traditionally are simplified and structured to be analytically tractable, but computing power allows for more complex, larger models. Bayesian statistics lends itself naturally to address parameter estimation problems in these large models. Bayesian statistical inference is utilized in this thesis to obtain parameter estimates from a sparse data set on populations in the HIV epidemic. Current estimates of the HIV epidemic indicate a decrease in the incidence of the disease in the undiagnosed subpopulation over the past 10 years. A lack of access to care, however, has not been considered when modeling the population. Populations at high risk for contracting HIV are twice as likely to lack access to reliable medical care. In this thesis, we consider three contributors to the HIV population dynamics: susceptible pool exhaustion, lack of access to care, and usage of anti-retroviral therapy (ART) by diagnosed individuals. An extant problem in the mathematical study of this system is deriving parameter estimates due to a portion of the population being unobserved. We approach this problem by looking at the proportional change in the infected subpopulations. We obtain estimates for the proportional change of the infected subpopulations using hierarchical Bayesian statistics. The estimated proportional change is used to derive epidemic parameter estimates for a system of stochastic differential equations (SDEs). Model fit is quantified to determine the best parametric explanation for the observed dynamics in the infected subpopulations. Parameter estimates derived using these methods provide interpretability and recovery of the system. Simulations suggest that the undiagnosed population may be larger than currently estimated without significantly affecting the population dynamics.
Chapter 1. Bayesian Statistics and Mathematical Biology

1.1. Mathematical modeling in biology

Advances in technology allow scientists to collect and analyze massive data sets. Utilizing these data sets requires training in fields related to computer science, mathematics, and statistics. The field demands additional computational skills out of biologists, trending away from the traditional paradigm of primarily experimental work. These trends create a subset of modelers that are not quite traditional applied mathematicians, aren’t statisticians, and are mostly self-taught programmers. An existing challenge is providing this growing population with the myriad of techniques available outside the training methods available to them. The NSF’s Directorate for Biological Sciences (BIO) found that the unmet needs of its funded investigators centered around computational and mathematical training [2]. The NSF recently began a ’Big Ideas’ program to address problems that require long-term solutions, and not surprisingly this issue is in focus [1].

One area in particular where these resources have been lacking is Bayesian statistics. A solution to this problem has been developing computational tools that are easy to use [5, 6]. These tools perform Bayesian inference on SMBL (systems biology markup language) models. SMBL models are procedurally generated, converting labeled wire diagrams of biological systems into models that can be simulated.

One issue facing widespread usage of these kinds of tools is that to seek them out, you need to have some idea of the problems existing with the tools you currently use. Current parameter estimation methods get the job done, and modelers may not be motivated to look elsewhere, especially considering the hurdles to utilizing these methods - additional programming, more computationally intensive, and a lack of background knowledge. However, obtaining estimates of parameter distributions instead of point estimates, or capturing the variability of a system or stochasticity of a process motivate increasing the usage of
Bayesian statistical inference in mathematical biology.

1.1.1. Approaches to modeling biological systems

Mathematical biology as a field attempts to represent biological systems directly with equations, while managing to simplify these systems to allow for mathematical or numerical analysis. The observed data $D$ is a function of the response $Y$ over time $t$. Normally $Y$ consists of a set of $n$ observable and unobservable components, and the length of the time under observation $t$ is imposed based on financial or temporal constraints, or prior beliefs about the system behavior. Mathematical biologists try to construct $Y$ so that it directly represents the system in question. Two famous models that illustrate this is the Lotka-Volterra model of population growth and the Michaelis-Menten model of enzyme kinetics.

- Population modeling

The Lotka-Volterra model represents a system of predator and prey species. The objective of the model is to understand how predator and prey dynamics affect each other, and this objective is used to simplify the model. We consider some birth rate of the prey population $\alpha$, and some death rate $\beta$ as a result of the predator population. The natural death rate of the prey is ignored for simplification purposes, as we assume that $\alpha$ is much larger. Finally, we assume that the predator population can only grow at some rate $\delta \leq \beta$, if prey are available as their primary or sole food source, and the predator die at some rate $\epsilon$. This provides the following equations:

$$\frac{dx}{dt} = \alpha x - \beta xy$$

$$\frac{dy}{dt} = \delta xy - \epsilon y$$
This model contains many simplifications of a very complex, large system with variability, but it allows for biological inference on data of few observables. Most of the parameters are not observable, but we can roughly count the number of prey or predators at a given time to estimate the value of these rate constants; or predict the ability of a population to survive predation when assuming some parameter values.

- **Kinetics modeling**

  The Michaelis-Menten model of enzyme kinetics represents the following system:

  \[
  E + S \rightleftharpoons ES \rightarrow P + E
  \]

  This system consists of an enzyme E combining with a substrate molecule S. These molecules bind and separate at some rate \( k_{on} \) and \( k_{off} \). When they exist in the ES conformation, the product molecule P is formed at some rate \( k_{cat} \), and we assume that the enzyme separates and is again able to bind to a substrate. We also assume that all these rates are constants, and that we are looking at homogeneous populations of E, S, and P. In reality, some E, S, P will have different parameters describing their behavior, and these features are time and space dependent. A model based on this system is as follows.

  \[
  \begin{align*}
  \frac{dE}{dt} &= -k_{on}E \cdot S + k_{off}ES + k_{cat}ES \\
  \frac{dS}{dt} &= -k_{on}E \cdot S + k_{off}ES \\
  \frac{dES}{dt} &= k_{on}E \cdot S - k_{off}ES - k_{cat}ES \\
  \frac{dP}{dt} &= k_{cat}ES
  \end{align*}
  \]

  The major contribution of Michaelis-Menten was not this system of equations, but rather another simplification. They showed that for the majority of enzyme systems, it appeared you could ignore the change in the ES population. This is because in general the
amount of ES reaches some level pretty quickly, and if the number of S molecules present is much larger than the number of E molecules, the change is not appreciable. Setting $\frac{dES}{dt} = 0$, the following model can be obtained:

$$\nu = \frac{dP}{dt} = \frac{V_{max}S}{K_M + S}$$

The value $V_{max}$ represents the maximum velocity observed in the production of P, and $K_M$ refers to the amount of S required to achieve half of the maximum velocity.

- **Different approaches to handle parameter problems**

  In both of these models, we simplify the system in an attempt to obtain a manageable equation that contains the observables. In the Lotka-Volterra model, we minimize the number of unknown parameters since we can only get a rough estimate of the population of predator or prey. In the Michaelis-Menten model, we formulate our equation to be time-independent, containing two observables ($V_{max}$ and S) and one unknown parameter.

  Historically, clever simplifications and reductions have been applied to systems in order to make them tractable. Parameter estimation and identifiability in models with large numbers of parameters is challenging and computationally intensive. Biological networks can have disparate issues when it comes to available data. Some biological networks are very difficult to observe, such as large populations; some networks are not observable, such as what happens inside of a cell. These cases require special treatment to consider how to estimate the parameters, and whether we can estimate them at all. Other biological systems have large data sets associated with them. Bayesian statistical methods can be used to address many of these existing issues.
1.2. Bayesian statistics and parameter estimation

Bayes’ rule describes the probability of event A occurring given event B occurred.

\[ \Pr(A|B) = \frac{\Pr(B|A) \Pr(A)}{\Pr(B)} \]

The probability of A occurring given B occurred is equivalent to the probability of event B occurring given A is occurred, multiplied by the probability of event A independent of event B, divided by the probability of event B occurring independent of event A. In Bayesian statistics, we take \( \Pr(A) \) as the initial belief in the probability of A occurring. Parameter estimation for biological models also considers the current evidence and \textit{a priori} beliefs about the system when constructing equations and considering parameters.

1.2.1. Handling big data

It has become relatively common place to collect data on the effect of experimental conditions on the expression of the entire genome within a large population of cells. It is also possible to use fluorescent tagging of proteins or cellular components to track the behavior of a large population of cells visually. One example of a Bayesian approach to handling this type of data is the Naive Bayes classifier [10]. The classifier was used to determine the log odds of different cellular markers in determining if a cell was likely to die or to continue reproducing. The methodology allowed the researchers to avoid making assumptions about the connections within this biological network, instead using Bayesian statistics to determine the likely contributors to the cell's "decision" to reproduce or to die.

1.2.2. Parameter estimation for large biological networks

Traditional parameter estimation for differential equation models consists of analytically solving equations for some initial conditions or knowns. This is not feasible for larger nonlinear models. In this case various computational techniques are often used, such as
least-squares fitting or gradient descent. These methods involve setting initial parameter estimates and parameter bounds, ideally based on previous experimental data.

A problem with this approach is that we are often more confident in some parameters than in others. Bayesian parameter estimation techniques are an improvement on such existing techniques by allowing the incorporation of parameter uncertainty [12].

The goal of biological mathematical modeling is normally not just producing accurate simulations, but also generating new hypotheses. The proteins JAK2 and STAT5 form a feedback loop that prevents treatment of some breast cancers [11]. A mathematical model of this system has 29 components, 25 equations, and 113 parameters [9]. In total, 513 data points were available to analyze, obtained from 24 different experimental conditions, making it a challenging problem to obtain parameter estimates. The authors used Markov Chain Monte Carlo to obtain Bayesian inference on the parameters. The model then allowed them to identify new observables in the system, generating new hypotheses.

1.2.3. Studying stochastic systems

Many biological systems may be better represented as stochastic systems, from the biochemical level to the population level [7, 8]. Some non-Bayesian methods available to study these systems include various stochastic simulation algorithms, like the Gillespie algorithm. However, it is challenging to estimate parameters for stochastic models without Bayesian statistics [12].

Bayesian methods for stochastic parameter estimates include using the stochastic nature of the model to create a likelihood, to allow for Bayesian inference on these models. Approximate Bayesian computation (ABC) uses the difference between the data and stochastic simulations of the model to replace the likelihood. In the following chapter, we
use Bayesian statistics to estimate two parameters that summarize the available information on two HIV populations. Then we construct several stochastic differential equation models and numerically solve to obtain parameter estimates for these equations. We compare our stochastic simulations of these models against the data, and quantify the goodness-of-fit by calculating the likelihood of observing the data given the model.
Chapter 2. Estimating Epidemiological Parameters of a Stochastic Differential Model of HIV Dynamics Using a Hierarchical Bayesian Model

2.1. Introduction

The human immunodeficiency virus (HIV) progresses in three stages. The first stage lasts approximately 3 months and individuals in this stage are approximately 10 to 25 times more effective at transmitting the disease [29]. The chronic stage can last from 5-10 years without medication [38]. This is followed by acquired immunodeficiency syndrome (AIDS) and death shortly thereafter [29]. Individuals with HIV may go many years without diagnosis, during which time they may expose uninfected individuals to HIV. Efforts to improve the diagnosis rate include educational programs, as an individual’s perceived risk was shown to be highly correlated with the individual obtaining multiple HIV tests [19, 22, 24, 35]. Several studies have found a 50% reduction in risky behaviors after diagnosis, including safer sex practices, reduction in partner number, and medications that reduce viral load [27, 39]. Diagnosis events resulting in behavior modification are not thought to be sufficient to eradicate the epidemic [20, 27].

After diagnosis, infected individuals have the opportunity to take anti-retroviral therapy (ART) that reduces their viral load and retards the progression of the disease. The earlier that ART is received the higher the reduction in transmission events. ART therapies could eradicate the epidemic in a population with high prevalence of infection even without the additional effect of behavioral changes [20]. Mathematical models estimate that the HIV epidemic could be reduced to less than 1% of the population infected (elimination phase) with universal testing and by providing ART consistently to newly diagnosed individuals [36]. However, issues with adherence and resistance are well documented in the
literature [13–15, 25, 28, 32]. Patients tend to report their adherence as much higher than it actually is, but studies indicate that even low adherence may be sufficient for control of the epidemic [13, 39]. Transmission is rare for individuals on ART, even with relatively high plasma HIV concentrations [30, 33].

The largest barrier to eradication of the epidemic is lack of access to care, including diagnostic services and ART costs or prescriptions. A lack in access to care could create pockets of undiagnosed individuals while the overall trend appears to be a reduction in the size of the epidemic [16]. Various studies report between 50 - 96% of diagnosed individuals in the U.S. rely on public medical care to obtain their ART medications [18, 21, 32, 34, 39]. Access to care remains critical, but this has not been considered when modeling the dynamics of the epidemic [19, 22].

Estimates using CD4 levels of newly diagnosed individuals suggest that the undiagnosed population is decreasing between 2005 - 2013 [37, 38]. CD4 levels can be used to estimate the progression of HIV [40]. We consider three possible causes for this decrease including exhaustion of the susceptible population. The size of the susceptible population is not easy to estimate since it depends on behavior risk. High risk populations include individuals in poverty and men who have sex with men (MSM) [38]. This is particularly critical in the southern U.S., where individuals tend to be poor and lack access to medical care [16, 38]. As the at-risk population decreases the number of diagnoses will also decrease, which will cause the estimated number of undiagnosed individuals to decrease.

An additional possibility is that the reduction in number of diagnoses is due to individuals lacking access to care. HIV is over-represented in impoverished populations where access to diagnosis and treatment may be more difficult to obtain. In this case, the number of newly diagnosed individuals is not representative of the number of undiagnosed individ-
uals, and the estimates will be inaccurate. Finally, the usage of anti-retroviral therapies reduces the viral load and transmission potential of infected individuals.

The difficulty in studying this system mathematically lies in parameter estimates. A minimal model of this system requires at least three parameters: transmission of the disease, diagnosis of the disease, and death due to the disease. Since knowledge about the undiagnosed population is restricted to those who have been diagnosed, estimates of these parameters are generally forced to assume that this population is representative of the whole. Our motivation to model the system stochastically arises from heterogeneity due to reporting delays associated with population-level data [31]. Stochastic modeling will allow us to better understand both the effectiveness of our estimates and the quality of model fit.

In this work we use coupled statistical and mathematical methodology to study the relationships between the three hypothesized causes and their respective population dynamics. We use hierarchical Bayesian statistics to get estimates for the size of the infected populations and their proportional changes across the years. These estimates are used to calculate epidemiological parameters for a system of stochastic differential equations. Currently we are not aware of any similar methods in the literature. Such a problem is challenging as the proportional change across the populations is a hyperparameter controlling the yearly proportions, which each have their own statistical model. This results in a large model that must be studied numerically.

The resulting simulations give insight into the implications of the estimated undiagnosed population on epidemiological parameters. Our model suggests that the undiagnosed population may be larger than current estimates while recovering population dynamics. The best recovery occurs when the increase in the diagnosed population due to diagnosis
is greater than the decrease in the undiagnosed population. We hope this study will help inform future efforts to improve the situation of infected individuals and prevent future outbreaks.

2.2. Materials and methods

2.2.1. Bayesian statistics

A Markov model where $p_t$ centered at $qp_{t-1}$ is used to estimate the proportional change in the infected populations over time, where $p_t$ is the proportion in the current year and $q$ is the proportional change. These random variables are estimated using Bayesian statistics.

The sampling model is $x_t \sim Bin(n_t, p_t)$, where $n_t$ is population size in the current year. The random variable $q$ is taken as a hyperparameter for $p_t$. The random variables to be estimated for each infected subpopulation are $q$ and $p_t$, where $t = 2005, ..., 2013$. We estimate the random variables of undiagnosed and diagnosed subpopulations independently.

- Prior

The prior for the proportional change $q$ is a gamma distribution.

$$
\pi(q) \propto q^{a-1}e^{-\beta q}
$$

The parameters were chosen so that the prior distribution was centered at the arithmetic estimates of $q$ obtained from the CDC [38]. The arithmetic estimates were obtained by calculating:

$$
\frac{1}{n} \sum_{i=2}^{n} \frac{p_i}{p_{i-1}}
$$

The arithmetic estimate for the undiagnosed $q$ ($q_u$) is 0.979, and for the diagnosed $q$ ($q_d$) 1.025. The priors used were GAM(9.79,10) and GAM(10.25,10) so they were centered at 1.
The prior for the random variable $p_t$, the undiagnosed proportion, is a beta distribution centered at the previous proportion times $q$. The parameters of the beta distribution are $\alpha = 0.1n_{t-1} \times qp_{t-1}$, and $\beta = 0.1n_{t-1} - \alpha$.

$$\pi(p_t) \propto p_t^{\alpha-1}(1-p_t)^{\beta-1}$$

In the case where $t = 1$, the previous undiagnosed proportion is taken to be the expert opinion of 20%, and the diagnosed proportion to be $1 - p_0(\text{undiagnosed})$ [23]. The prior for the diagnosed population is formulated in the same way. Population sizes were considered in units of thousands.

- **Likelihood**

The likelihood is a binomial distribution, representing the chance of selecting an undiagnosed or diagnosed individual at random from the total infected population. For a given year $t$, the proportion of undiagnosed individuals depends on the total number of individuals:

$$\mathcal{L}(p_t|x_t, n_t) \propto p_t^{x_t}(1-p_t)^{n_t-x_t}$$

where $x_t$ is the total number of undiagnosed or diagnosed individuals, and $n_t$ is the total number of infected individuals. The likelihood across all the years is the product of each year’s likelihood.

$$\mathcal{L}(p_1, p_2, ..., p_9|x_1, x_2, ..., x_9, n_1, n_2, ..., n_9) \propto \Pi_{t=1}^{9} p_t^{x_t}(1-p_t)^{n_t-x_t}$$

The likelihood for the diagnosed population was formulated in the same way.
• **Posterior**

The joint posterior distribution is proportional to the priors multiplied by the likelihoods for all 9 years:

\[
f(p_1, p_2, ..., p_9, q) \propto \pi(q) \times \prod_{t=1}^{9} \pi(p_t|q, p_{t-1}) \times L(p_1|x_{t=1}) \times L(p_2|x_{t=2}) \times ... \times L(p_9|x_{t=9}) \\
\quad \propto q^{\alpha-1}e^{-\beta q} \times \prod_{t=1}^{9} \left( p_t^{\alpha-1}(1-p_t)^{\beta-1} \times p_t^{x_t}(1-p_t)^{n_t-x_t} \right)
\]

The posterior full conditional of \(p_t\) for \(t = 2005, ..., 2012\) is:

\[
f(p_t|q, p_{t-1}, p_{t+1}) \propto \prod_{t=1}^{9} \left( L(p_t|x_t) \times \pi(p_t|q, p_{t-1}) \times \pi(p_{t+1}|q, p_t) \right)
\]

The posterior full conditional of 2013, the 9th year, is:

\[
f(p_9|x_9, p_8, q) \propto L(p_9|x_{t=9}) \times \pi(p_9|q, p_8)
\]

The full conditional of \(q\) does not have a closed form. The forms of the diagnosed random variables are the same. Random variable estimates were obtained using Metropolis-Hastings nested within a Gibbs sampler over 100,000 iterations with R version 3.3.3 [41]. The proposal distribution was a truncated normal distribution, using package rmutil [42], centered at the previous value of the parameter. Proportions 1 through 9 were sampled consecutively, followed by hyperparameter \(q\). The trace plots converged quickly, and the first 2000 samples were removed. Code is provided in Appendices B and C.

### 2.2.2. Stochastic differential equations

The hyperparameter \(q\) was estimated to be 0.978 for the undiagnosed population and 1.036 for the diagnosed population. These were used as a boundary to solve for the epi-
demographical parameters in a simple stochastic differential model.

\[ dU = (q_u - 1)U dt + d\omega_t dt \]

\[ dD = (q_d - 1)D dt + d\omega_t dt \]

where \( U \) is the undiagnosed and \( D \) is the diagnosed populations, and \( d\omega_t \sim \text{Nor}(0, \sigma) \) is Brownian white noise with units \( \sqrt{t} \). The variance \( \sigma \) is chosen to be 10% of the size of the population.

The simplest model is constructed describing the dynamics of the infected subpopulations. The values of parameters transmission \( (\tau) \), diagnosis \( (\delta) \), and death \( (\epsilon) \) are calculated using the constraining \( q \):

\[ dU = [\tau(U + D) - \delta U - \epsilon] dt \tag{2.1} \]

\[ \cong (1 - q_u)U dt + d\omega_t dt \tag{2.2} \]

\[ = -0.022U dt + d\omega_t dt \tag{2.3} \]

\[ dD = (\delta U - \epsilon D) dt \tag{2.4} \]

\[ \cong (1 - q_d)D dt + d\omega_t dt \tag{2.5} \]

\[ = 0.036D dt + d\omega_t dt \tag{2.6} \]

We consider the parameters pseudo-steady state, and use the 2005 population sizes to estimate them. In addition, we assume that the general population are at steady state, and consider only the increased death rate due to infection \( \epsilon \) as 0.01 [38]. The diagnosis rate \( \delta \) is estimated by:

\[ \delta U = q_d D + \epsilon D = \frac{0.046D_{2005}}{U_{2005}} = 0.165 \]
and the transmission rate $\tau$ is estimated by:

$$
\tau(U + D) = (1 - q_u)U + \delta U + \epsilon U
$$

$$
\tau = \frac{0.153U_{2005}}{U_{2005} + D_{2005}} = 0.0334
$$

Due to the magnitude of the scale of this system we assume that all events will happen, and the source of the stochasticity is primarily reporting issues. Tau leap algorithm was used to preform the stochastic simulations. A time step of 1 year was selected, and the population at time $t+1$ is the numerical solution of the population at time $t$ and random noise from a $NOR(0,\sigma)$, where $\sigma$ is 10% of the population at time $t_0$ with units $\sqrt{t}$. The initial conditions for the infected populations were sampled from the posterior distributions obtained by the Bayesian estimates. Calculations were performed in Matlab [43] and code is available upon request.

The diagnosis rate was calculated using data from [37]. The susceptible population is estimated as twice the national average rate of self-identified MSM among adults. The mortality rate increase due to HIV was estimated using data from [38]. All calculations, including the effective parameter rates, are provided in Appendix A (in progress).

2.3. Results

2.3.1. Bayesian model

Bayesian estimates for the proportions of diagnosed or undiagnosed individuals was obtained concurrently with the estimated proportional change. The prior distribution was chosen to be a beta for the proportions and a gamma for the proportional changes. The likelihood function was a binomial, representing the chance of randomly selecting a diagnosed or undiagnosed individual from a pool of infected individuals. The posterior did not have a closed form. Due to the symmetry of the posterior samples we summarize them using their mean and variance. The posterior means of the proportions for both undiagnosed
and diagnosed estimates are very close to the original data (Fig. 2.1) [37]. The posterior mean of $q_u$ is 0.96, and $q_d$ is 1.02. This means that 96% of the undiagnosed population is preserved from year to year, or is dropping by about 4% per year. Similarly, the diagnosed population is increasing by 2% per year. Posterior means and variances are given in Table 2.1.

Table 2.1. Summary statistics of the posterior distribution. The parameter $p$ represents the estimated size of the proportion in that year. The hyperparameter $q$ represents the estimated proportional change of the population across all years.

<table>
<thead>
<tr>
<th></th>
<th>Undiagnosed</th>
<th>Diagnosed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameters</td>
<td>Mean</td>
<td>Variance</td>
</tr>
<tr>
<td>$p_{2005}$</td>
<td>0.78</td>
<td>0.0006</td>
</tr>
<tr>
<td>$p_{2006}$</td>
<td>0.79</td>
<td>0.0011</td>
</tr>
<tr>
<td>$p_{2007}$</td>
<td>0.80</td>
<td>0.0012</td>
</tr>
<tr>
<td>$p_{2008}$</td>
<td>0.81</td>
<td>0.0012</td>
</tr>
<tr>
<td>$p_{2009}$</td>
<td>0.82</td>
<td>0.0012</td>
</tr>
<tr>
<td>$p_{2010}$</td>
<td>0.82</td>
<td>0.0012</td>
</tr>
<tr>
<td>$p_{2011}$</td>
<td>0.83</td>
<td>0.0012</td>
</tr>
<tr>
<td>$p_{2012}$</td>
<td>0.83</td>
<td>0.0012</td>
</tr>
<tr>
<td>$p_{2013}$</td>
<td>0.84</td>
<td>0.0012</td>
</tr>
<tr>
<td>$q_d$</td>
<td>1.036</td>
<td>0.1046</td>
</tr>
</tbody>
</table>

2.3.2. Stochastic differential model

The Bayesian estimates of the proportional change in the diagnosed and undiagnosed population from 2005 to 2013 were used to determine the epidemiological parameters for a system of stochastic differential equations. The parameters transmission ($\tau$), diagnosis ($\delta$), and death ($\epsilon$) were calculated using the proportional changes in the respective population.

\[
dU = (q_u - 1)U dt + d\omega_t dt = \left[\tau(U + D) - \delta U - \epsilon U\right] dt \tag{2.7}
\]

\[
dD = (q_d - 1)D dt + d\omega_t dt = \left[\delta U - \epsilon D\right] dt \tag{2.8}
\]

where $U$ is the undiagnosed and $D$ is the diagnosed populations, and $d\omega_t \sim N(0, \sigma)$ is the noise term. These base estimates fit the data very well (Fig. 2.2).
Figure 2.1. Posterior information obtained from hierarchical Bayesian statistics. Bayesian estimates are shown as hollow squares with error bars showing standard deviations. Estimated proportion of diagnosed (pink) and undiagnosed (blue) populations recover the estimated proportions (circles). Data obtained from [37].

- **Exhaustion of susceptibles**

  In the case where the susceptible population is not much larger than the infected population, the transmission is dependent on the size of both populations. We estimate the susceptible population size as a fraction of the total infected population:

  \[ S = fT \]

  Then this is substituted into the model. The transmission term becomes

  \[ \tau TS \approx \tau fT^2 \]

  This gives an effective increase of \( f\tau \) in the transmission rate - see Table 2.2. This increase causes the simulations to fail to recover the diagnosed and undiagnosed popula-
Figure 2.2. Method Validation. The method was tested by simulating with the epidemiological parameters calculated using the Bayesian estimates of the proportional changes as constraints. The mean of 100 stochastic simulations (pink line) is compared with the data (circles). Proportions are relative to initial proportion.
tion dynamics, although the susceptible population does decrease significantly (Fig. 2.3). This result is intuitive since the infection rate is increased, but the diagnosis rate is not representative of this rate.

- **Lack of access to care**

  Lack of access to care may be conceptualized as pockets of undiagnosed individuals who are not being diagnosed. To capture this, we consider the diagnosis rate to be independent of the size of the undiagnosed population. The diagnosis rate is estimated as:

  \[
  \delta U = q_d D + \epsilon D = \frac{0.046D_{2005}}{U_{2005}} \cdot U_{2005} = 0.036
  \]

  The resulting equation for the undiagnosed subpopulation then becomes:

  \[
  dU = \left[ \tau(U + D) - \delta_0 - \epsilon U \right] dt
  \]

  where \( \delta_0 \) is 0.036. This large reduction in the diagnosis rate recovers the population dynamics well (Fig. 2.4).

- **ART usage**

  Since ART results in a viral load that has low chance of infecting a susceptible individual, we removed these individuals from the pool of infected individuals able to transmit the disease. Since 96% of diagnosed individuals reported taking anti-retroviral therapies in a previous study, the transmission term was modified as follows [39].

  \[
  \tau[U + (1 - 0.96)D]
  \]

  Variable or poor adherence on the part of diagnosed individuals is ignored due to the body of literature indicating that large benefit is gained from even poor adherence [13, 25].
Figure 2.3. Exhaustion of Susceptibles. Transmission of the disease is altered to reflect the impact of the size of the susceptible population. The mean of 100 stochastic simulations (pink line) is compared to the data (circles). Proportions are relative to initial proportion.
Figure 2.4. Lack of Access to Care. The effect of undiagnosed individuals lacking access to care affects the rate of diagnosis of the undiagnosed individuals. The diagnosis rate is held constant to reflect this scenario. The mean of 100 stochastic simulations (pink line) is compared with the data (circles). Proportions are relative to initial proportion.
This gives good recovery of both subpopulation dynamics and agrees best with both undiagnosed and diagnosed estimates (Fig. 2.5).

2.3.3. Multiple causes

Since it seems likely that most or all of these scenarios affect the infected population simultaneously, we analyze all their possible combinations (Appendix A). The parameters were altered as described in Table 2.2. To determine the best cause, we quantify the goodness of fit by determining the relative likelihood of observing the data given the mean and variance of the stochastic simulations. These probabilities are given in Fig. 2.6 with numerical details in the Appendix A table, as well as the average probability over the 9 years.

Table 2.2. Transmission and diagnosis rates are different under the different hypotheses. Average likelihood across both populations and all years (Fig. 2.6, Supplemental Table in Appendix A.

<table>
<thead>
<tr>
<th>Model</th>
<th>Transmission Rate</th>
<th>Diagnosis Rate</th>
<th>Likelihood</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base model</td>
<td>( \tau(U + D) )</td>
<td>( \delta U )</td>
<td>0.83</td>
</tr>
<tr>
<td>Exhaustion of Susceptibles (ES)</td>
<td>( \tau f(U + D)^2 )</td>
<td>( \delta U )</td>
<td>0.26</td>
</tr>
<tr>
<td>Lack of Access to Care (LAC)</td>
<td>( \tau(U + D) )</td>
<td>( \delta_0 )</td>
<td>0.88</td>
</tr>
<tr>
<td>Anti-retroviral Therapies (ART)</td>
<td>( \tau(U + 0.04D) )</td>
<td>( \delta U )</td>
<td>0.75</td>
</tr>
<tr>
<td>ES and LAC</td>
<td>( \tau f(U + D)^2 )</td>
<td>( \delta_0 )</td>
<td>0.54</td>
</tr>
<tr>
<td>ES and ART</td>
<td>( \tau f(U + 0.04D)^2 )</td>
<td>( \delta U )</td>
<td>0.52</td>
</tr>
<tr>
<td>LAC and ART</td>
<td>( \tau(U + 0.04D) )</td>
<td>( \delta_0 )</td>
<td>0.78</td>
</tr>
<tr>
<td>ES, LAC, and ART</td>
<td>( \tau f(U + 0.04D)^2 )</td>
<td>( \delta_0 )</td>
<td>0.58</td>
</tr>
</tbody>
</table>

Lack of access to care, ART usage, or their combined resulted in the best recovery of the data for both the undiagnosed and diagnosed populations. Under the ART scenario, the diagnosed population has been reduced by 96%, resulting in a dynamic reduction in the transmission rate. We originally estimate the transmission rate to be 3.3% of the total infected population. This is close to the literature estimate of around 4% [17, 23]. With the majority of the diagnosed population removed, the effective transmission rate is much lower. Under the LAC scenario, there is a constant diagnosis rate. This represents a yearly reduction in the undiagnosed population and increase in the diagnosed population of 3.6%.
Figure 2.5. Anti-retroviral Therapy Usage. To reflect the high levels of ART prescription and usage reported by diagnosed individuals this percentage is removed from the pool of diagnosed individuals able to transmit the disease. The mean of 100 stochastic simulations (pink line) is compared to the data (circles). Proportions are relative to initial proportion.
Figure 2.6. Model fit was quantified by calculating the relative likelihood of observing the data within the simulations. A higher likelihood is represented by a hotter color. From left to right: Base model, Exhaustion of Susceptibles (ES), Lack of Access to Care (LAC), Anti-Retroviral Therapy usage (ART), ES and LAC, ES and ART, LAC and ART, and ES, LAC, and ART. Details provided in Appendix A.
This means the diagnosed population grows faster than the undiagnosed population is reduced for a diagnosis event. Although lack of access to care in the undiagnosed population would mean the data are inaccurate, in our case the best fitting model is consistent for both subpopulations.

2.4. Discussion

We were able to obtain conservative estimates of the proportional changes in the diagnosed and undiagnosed HIV-infected populations using hierarchical Bayesian statistics. Our estimates suggest that the proportion of infected individuals who are undiagnosed is decreasing by approximately 2.2% each year from 2005 to 2013, while the proportion of diagnosed individuals is increasing by approximately 3.6%. We used the proportional change as constraints on a system of stochastic differential equations. This allowed us to estimate the transmission and diagnosis rates. We were able to recover reasonable parameter estimates and population dynamics using this methodology. To learn more about the cause of the decrease in the undiagnosed population, we considered some scenarios that would affect the epidemiological parameters: exhaustion of the susceptible population, lack of access to care, and reduction in viral load by anti-retroviral therapy.

We were able to recover the diagnosed population dynamics when we altered the parameters to reflect these scenarios with the exception of including exhaustion of susceptibles. Including the size of the susceptible population dramatically increased the transmission rate and caused the size of the infected populations to increase rapidly. In the other scenarios some interesting dynamics could be observed in the undiagnosed population. Lack of access to care was simulated by considering diagnosis rate a constant unaffected by the size of the undiagnosed population. This resulted in an improvement in the likelihood of observing the data (Fig. 2.6, Appendix A). Anti-retroviral therapy usage also improved the overall recovery, but this effect was weaker for the undiagnosed population dynamics. Although
the undiagnosed population size is dependent on the quality of the data available on the diagnosed population of that year, these results indicate that the scenarios that maximizes the probability of observing the diagnosed population also maximizes the probability of observing the diagnosed population estimates.

The observed results suggest that lack of access to care and ART usage contribute to the infected population dynamics. This is not unexpected. Many individuals with HIV are reported to lack access to care [16, 26]. In areas with high poverty rates the death rate of infected individuals is much higher than that of the general population [38, 44]. In 2017 the New York Times reported groups of untreated individuals in the deep south dying due to their lack of access to care [16]. The effect of simulating a lack of access to care suggest this to be a significant contributing factor to the infected population dynamics. Both models and studies have shown that providing ART to infected individuals in the early stages of HIV reduces transmission events and frequency of death due to AIDS [20, 28, 30, 32, 33, 36]. Even poor adherence may be enough to control or eradicate the epidemic and increase quality of life for infected individuals [14, 15, 25]. Greater effort must be made to ensure these populations have access to life-saving treatments.
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Appendix B. Diagnosed Population R Code

```r
library(MCMCpack)
library(pracma)
library(msm)
library(rmutil)

# DATA


total = c(896.9,923.2,951.5,979.7,1006.5,1032.6,1057.8,1082.1,1104.6)
diagnosed=c(701.3758,729.328,760.2485,789.6382,818.2845,845.6994,871.6272,898.143,923.4456)
x=diagnosed

plot(total)
plot(year,undiagnosed,xlab=Year,ylab='',xaxt='n',yaxt='n')
axis(1,at=year,labels=year)
axis(2,at=undiagnosed,labels=undiagnosed)
title(main=Estimated Undiagnosed HIV Population)
plot(x)
v=matrix(0,8)
for (i in 1:8){v[i]=diagnosed[i+1]/diagnosed[i]}
mean(v)
var(v)

p0=.8 #expert prior
n0=.25*total[1]

posteriorp=function(pt,pt1,pt.1,nt,nt1,nt.1,xt,xt1,xt.1,q,f){
  # this year t
  talpha=pt.1*nt.1*f*q
}
```

29
tbeta=(1–pt.1*q)*nt.1*f

## next year t+1

t1alpha=pt*nt*q*f
t1beta=(1–pt*q)*nt*f

# use beta to approximate binomial to keep numbers unrounded
a=dbeta(pt,talpha,tbeta)*dbeta(pt,xt+1,nt–xt+1)
b=dbeta(pt1,t1alpha,t1beta)*dbeta(pt1,xt1+1,nt1–xt1+1)
a*b

posterior_individ=function(pt,pt.1,xt,xt.1,nt,nt.1,q,f){
talpha=pt.1*nt.1*f*q; tbeta=(1–pt.1*q)*nt.1*f
dbeta(pt,xt+1,nt–xt+1)*dbeta(pt,talpha,tbeta)
}

posterior_q=function(pt,pt.1,xt,xt.1,nt,nt.1,q,f){
talpha=pt.1*nt.1*f*q; tbeta=(1–pt.1*q)*nt.1*f
dbeta(pt,talpha,tbeta)
}

##### GIBBS SAMPLER #####

sample=100000

gibb.sample=matrix(0,sample,10) #columns = parameters
tmu=matrix(.9,9,1) # initial values for proportion
q=1 # initial value for q

var_star = 0 # dummy variable for each iter

sd.rw=matrix(.05,11);accept=matrix(0,11) # acceptance matrix

sd.rw[10]=.52
f=.1 # to scale the strength of the prior

for (i in 1:sample ){
  set.seed(i) # for reproducibility
  # # # # # # # # # # #
  ## p1 : marginal posterior involves p0 and p1
```r
# # # # # # # # # # # # # # #
## p2 — p8 : marginal posteriors involve previous, current, and next p's
for (j in 2 : 8)
var_star <- rnorm(1, tmu[j], sd.rw[j], 0, 1)
nom = posteriorp(var_star, tmu[j+1], tmu[j-1], total[j], total[j+1], total[j-1], x[j], x[j+1], 0, q, f)
denom = posteriorp(tmu[j], tmu[j+1], tmu[j-1], total[j], total[j+1], total[j-1], x[j], x[j+1], 0, q, f)
if (is.nan(nom))
    nom=0;
if (nom == -Inf)
    nom = 0;
alpha <- min(1, nom/denom)
if (is.nan(alpha))
    alpha=0
r <- runif(1,0,1)
if (r<=alpha )
    accept[j] <- accept[j]+1
tmu[j] <- var_star # accept proposed value
# # # # # # # # # # # # # # #
```
## p9 : marginal posterior involves only p8 and p9

```r
var_star <- rtnorm(1, tmu[9], sd.rw[9], 0, 1)
nom = posterior_individ(var_star, tmu[8], x[9], x[8], total[9], total[8], q, f)
denom = posterior_individ(tmu[9], tmu[8], x[9], x[8], total[9], total[8], q, f)
if (is.nan(nom)) {
  nom = 0;
} if (nom == -Inf) {
  nom = 0;
}
alpha <- min(1, nom/denom)
if (is.nan(alpha)) {
  alpha = 0
}
r <- runif(1, 0, 1)
if (r <= alpha) {
  tmu[9] <- var_star
} gibb.sample[i, 9] = var_star
```

## q : marginal posterior involves all priors

```r
var_star <- rtnorm(1, q, sd.rw[10], 0)
nom = dgamma(var_star, 10.254, 10) # prior centered at one
denom = dgamma(q, 10.254, 10)
nom = nom * posterior_q(tmu[1], p0, x[1], 0, total[1], n0, var_star, f)
denom = denom * posterior_q(tmu[1], p0, x[1], 0, total[1], n0, q, f)
for (k in 2:9) {
  nom = nom * posterior_q(tmu[k], tmu[k-1], x[k], x[k-1], total[k], total[k-1], var_star, f)
  denom = denom * posterior_q(tmu[k], tmu[k-1], x[k], x[k-1], total[k], total[k-1], q, f)
}
if (is.nan(nom)) {
  nom = 0;
} if (nom == -Inf) {
  nom = 0;
}
alpha <- min(1, nom/denom)
if (is.nan(alpha)) {
  alpha = 0
}
r <- runif(1, 0, 1)
if (r <= alpha) {
  q <- var_star
} gibb.sample[i, 10] = q
```
gibb.sample[i,10]=var_star
}
accept/sample
burnin=2000
plot(gibb.sample[burnin:sample,1])
plot(gibb.sample[burnin:sample,2])
plot(gibb.sample[burnin:sample,3])
plot(gibb.sample[burnin:sample,4])
plot(gibb.sample[burnin:sample,5])
plot(gibb.sample[burnin:sample,6])
plot(gibb.sample[burnin:sample,7])
plot(gibb.sample[burnin:sample,8])
plot(gibb.sample[burnin:sample,9])
plot(gibb.sample[burnin:sample,10])
m1=mean(gibb.sample[burnin:sample,1])
m2=mean(gibb.sample[burnin:sample,2])
m3=mean(gibb.sample[burnin:sample,3])
m4=mean(gibb.sample[burnin:sample,4])
m5=mean(gibb.sample[burnin:sample,5])
m6=mean(gibb.sample[burnin:sample,6])
m7=mean(gibb.sample[burnin:sample,7])
m8=mean(gibb.sample[burnin:sample,8])
m9=mean(gibb.sample[burnin:sample,9])
m10=mean(gibb.sample[burnin:sample,10])
v1=var(gibb.sample[burnin:sample,1])
v2=var(gibb.sample[burnin:sample,2])
v3=var(gibb.sample[burnin:sample,3])
v4=var(gibb.sample[burnin:sample,4])
v5=var(gibb.sample[burnin:sample,5])
v6=var(gibb.sample[burnin:sample,6])
v7=var(gibb.sample[burnin:sample,7])
v8=var(gibb.sample[burnin:sample,8])
v9=var(gibb.sample[burnin:sample,9])
v10=var(gibb.sample[burnin:sample,10])
hist(gibb.sample[burnin:sample,10],ylim=c(0,1.75),prob=TRUE,xlab='',col=00a1c0,main=expression('Posterior'),cex=1)
hist(gibb.sample[burnin:sample,1],prob=TRUE,xlab='Posterior',main='Posterior Histogram of P(Undiagnosed), 2005')
hist(gibb.sample[burnin:sample,2],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2006')
hist(gibb.sample[burnin:sample,3],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2007')

hist(gibb.sample[burnin:sample,4],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2008')

hist(gibb.sample[burnin:sample,5],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2009')

hist(gibb.sample[burnin:sample,6],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2010')

hist(gibb.sample[burnin:sample,7],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2011')

hist(gibb.sample[burnin:sample,8],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2012')

hist(gibb.sample[burnin:sample,9],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2013')
Appendix C. Undiagnosed Population R Code

```r
# # # # # # # library(MCMCpack) library(pracma) library(msm) library(rmutil) # # # # # # DATA # # # # # # total = c(896.9,923.2,951.5,979.7,1006.5,1032.6,1057.8,1082.1,1104.6) undiagnosed = c(.218,.21,.201,.194,.187,.181,.176,.17,.164) x=undiagnosed*total plot(total) plot(year,undiagnosed,xlab=Year,ylab=,xaxt='n',yaxt='n') axis(1,at=year,labels=year) axis(2,at=undiagnosed,labels=undiagnosed) title(main=Estimated Undiagnosed HIV Population) plot(x) v=matrix(0,8) for (i in 1:8){v[i]=undiagnosed[i+1]/undiagnosed[i]} mean(v) var(v) # # # # # # # p0=.2 #expert prior — eqsual to proportion n0=.25*total[1] qs0=1 # assume no change: beta0_qs=.5; alpha0_qs=2; #uninformative qsprior=function(alpha,beta,q){ q^(alpha-1)*exp(-q/beta) } posteriorp=function(pt,pt1,pt.1,nt,nt1,nt.1,xt,xt1,xt.1,q,f){ ## this year t talpha=pt.1*nt.1*f*q tbeta=(1-pt.1*q)*nt.1*f ## next year t+1 t1alpha=pt*nt*q*f t1beta=(1-pt*q)*nt*f }```

a = dbeta(pt, talpha, tbeta) * dbeta(pt, xt + 1, nt - xt + 1)

b = dbeta(pt1, t1alpha, t1beta) * dbeta(pt1, xt1 + 1, nt1 - xt1 + 1)

a * b

posterior_individ = function(pt, pt.1, xt, xt.1, nt, nt.1, q, f){
  talpha = pt.1 * nt.1 * f * q; tbeta = (1 - pt.1 * q) * nt.1 * f
  dbeta(pt, xt + 1, nt - xt + 1) * dbeta(pt, talpha, tbeta)
}

posterior_q = function(pt, pt.1, xt, xt.1, nt, nt.1, q, f){
  talpha = pt.1 * nt.1 * f * q; tbeta = (1 - pt.1 * q) * nt.1 * f
  dbeta(pt, talpha, tbeta)
}

##### GIBBS SAMPLER ######

sample = 100000

gibb.sample = matrix(0, sample, 11)  # columns = parameters; check for convergence
tmu = matrix(.2, 9, 1)
q = .9
var_star = 0  # dummy variable for each iter

sd.rw = matrix(.05, 11); accept = matrix(0, 11)
sd.rw[10] = .3

f = .1
for (i in 1:sample) {
  set.seed(i)
  # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # #
  var_star <- rtnorm(1, tmu[1], sd.rw[1], 0, 1)
nom = posteriorp(var_star, tmu[2], p0, total[1], total[2], n0, x[1], x[2], 0, q, f)
denom = posteriorp(tmu[1], tmu[2], p0, total[1], total[2], n0, x[1], x[2], 0, q, f)
  if (is.nan(nom)){

nom=0;
}
if (nom == -Inf){
  nom = 0;
}
alpha <- min(1, nom/denom)
if (is.nan(alpha)){
  alpha=0
}

r <- runif(1,0,1)
if (r<=alpha ) {
  tmu[1] <- var_star
}
gibb.sample[i,1]=var_star

## 2 − 8

for (j in 2 : 8){
  var_star <- rtnorm(1,tmu[j],sd.rw[j],0,1)
  nom = posteriorp(var_star,tmu[j+1],tmu[j-1],total[j],total[j+1],total[j-1],x[j],x[j+1],0,q,f)
  denom = posteriorp(tmu[j],tmu[j+1],tmu[j-1],total[j],total[j+1],total[j-1],x[j],x[j+1],0,q,f)
  if (is.nan(nom)){
    nom=0;
  }
  if (nom == -Inf){
    nom = 0;
  }
  alpha <- min(1, nom/denom)
  if (is.nan(alpha)){
    alpha=0
  }
  r <- runif(1,0,1)
  if (r<=alpha ) {
    accept[j] <- accept[j]+1
    tmu[j] <- var_star
  }
gibb.sample[i,j]=var_star

}

## 9

var_star <- rtnorm(1,tmu[9],sd.rw[9],0,1)
nom = posterior_individ(var_star,tmu[8],x[9],x[8],total[9],total[8],q,f)
denom = posterior_individ(tmu[9], tmu[8], x[9], x[8], total[9], total[8], q, f)
if (is.nan(nom)){
    nom=0;
}
if (nom == -Inf){
    nom = 0;
}
alpha = min(1, nom/denom)
if (is.nan(alpha)){
    alpha=0
}
if (r <= alpha) {
    tmu[9] = var_star
    likelihood[i,9]=nom;
} else {
    gibb.sample[i,9]=var_star
    var_star = rtnorm(1,q,sd.rw[10],0)
    upper = q+sd.rw[10]; lower = q-sd.rw[10]
    if (lower<0){lower=0}
    var_star= runif(1,lower,upper)
    nom=dgamma(var_star,9.788,10)
    denom=dgamma(q,9.788,10)
    nom = nom*posterior_q(tmu[1], p0, x[1], 0, total[1], n0, var_star, f)
    denom = denom*posterior_q(tmu[1], p0, x[1], 0, total[1], n0, q, f)
    for (k in 2:9){
        nom=nom*posterior_q(tmu[k], tmu[k-1], x[k], x[k-1], total[k], total[k-1], var_star, f)
        denom=denom*posterior_q(tmu[k], tmu[k-1], x[k], x[k-1], total[k], total[k-1], q, f)
    }
if (is.nan(nom)){
    nom=0;
}
if (nom == -Inf){
    nom = 0;
}
alpha = min(1, nom/denom)
if (is.nan(alpha)){
    alpha=0
}
r = runif(1,0,1)
if (r <= alpha) {
q ← var_star
}

gibb.sample[i,10]=var_star
}

accept/sample
mean(gibb.sample[burnin:sample,10])

burnin=2000
plot(gibb.sample[burnin:sample,1])
plot(gibb.sample[burnin:sample,2])
plot(gibb.sample[burnin:sample,3])
plot(gibb.sample[burnin:sample,4])
plot(gibb.sample[burnin:sample,5])
plot(gibb.sample[burnin:sample,6])
plot(gibb.sample[burnin:sample,7])
plot(gibb.sample[burnin:sample,8])
plot(gibb.sample[burnin:sample,9])
plot(gibb.sample[burnin:sample,10])

00FF00 55FF00 AAFF00 FFFF00

hist(gibb.sample[burnin:sample,10],ylim=c(0,1.75),prob=TRUE,xlab='',col=00a1c0,main=expression('Posterior Histogram of q'[u]),cex=1)

curve(dgamma(x,9.79,10),xlim=c(0,10),lwd=6,col=b57786,add=TRUE,cex=.75)

hist(gibb.sample[burnin:sample,1],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2005')
hist(gibb.sample[burnin:sample,2],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2006')
hist(gibb.sample[burnin:sample,3],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2007')
hist(gibb.sample[burnin:sample,4],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2008')
hist(gibb.sample[burnin:sample,5],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2009')
hist(gibb.sample[burnin:sample,6],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2010')
hist(gibb.sample[burnin:sample,7],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2011')
hist(gibb.sample[burnin:sample,8],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2012')
hist(gibb.sample[burnin:sample,9],prob=TRUE,xlab='',main='Posterior Histogram of P(Undiagnosed), 2013')

m1=mean(gibb.sample[burnin:sample,1])
m2=mean(gibb.sample[burnin:sample,2])
m3 = mean(gibb.sample[burnin:sample,3])

m4 = mean(gibb.sample[burnin:sample,4])

m5 = mean(gibb.sample[burnin:sample,5])

m6 = mean(gibb.sample[burnin:sample,6])

m7 = mean(gibb.sample[burnin:sample,7])

m8 = mean(gibb.sample[burnin:sample,8])

m9 = mean(gibb.sample[burnin:sample,9])

m10 = mean(gibb.sample[burnin:sample,10])

means2 = c(m1, m2, m3, m4, m5, m6, m7, m8, m9)


plot(year, means2, xlab='', ylab='', ylim=c(.1,.25), type=p, pch=21, cex=2, col=red)

boxplot(gibb.sample[burnin:sample,1:9], xlab='', ylab='', xaxt='n')

lines(year, undiagnosed, type=p, pch=22, cex=2, col=blue)

axis(1, at=1:9, labels=year)

arrows(year, means2-stdevs, year, means2+stdevs, length=0.05, angle=90, code=3, col=red)

legend(2005,.155, c('Estimate', 'Observed'), pch = c(21,22), col=c('red', 'blue'))

title(main=Posterior Estimates of Undiagnosed Proportion)

plot(year, means2, xlab='', ylab='', ylim=c(.1,.25), type=p, pch=21, cex=2, col=red)

lines(year, undiagnosed, type=p, pch=22, cex=2, col=blue)

arrows(year, means2-stdevs, year, means2+stdevs, length=0.05, angle=90, code=3, col=red)

legend(2005,.155, c('Estimate', 'Observed'), pch = c(21,22), col=c('red', 'blue'))

title(main=Posterior Estimates of Undiagnosed Proportion)

v1 = var(gibb.sample[burnin:sample,1])

v2 = var(gibb.sample[burnin:sample,2])

v3 = var(gibb.sample[burnin:sample,3])

v4 = var(gibb.sample[burnin:sample,4])

v5 = var(gibb.sample[burnin:sample,5])

v6 = var(gibb.sample[burnin:sample,6])

v7 = var(gibb.sample[burnin:sample,7])

v8 = var(gibb.sample[burnin:sample,8])

v9 = var(gibb.sample[burnin:sample,9])

v10 = var(gibb.sample[burnin:sample,10])

plot(vars/n)

plot(c(v1,v2,v3,v4,v5,v6,v7,v8,v9))

vars = c(v1,v2,v3,v4,v5,v6,v7,v8,v9)

stdevs = sqrt(vars)

arrows(year, means2-stdevs, year, means2+stdevs, length=0.05, angle=90, code=3)
quantile((gibb.sample[burnin:sample,1]), c(.05,.95))
quantile((gibb.sample[burnin:sample,2]), c(.05,.95))
quantile((gibb.sample[burnin:sample,3]), c(.05,.95))
quantile((gibb.sample[burnin:sample,4]), c(.05,.95))
quantile((gibb.sample[burnin:sample,5]), c(.05,.95))
quantile((gibb.sample[burnin:sample,6]), c(.05,.95))
quantile((gibb.sample[burnin:sample,7]), c(.05,.95))
quantile((gibb.sample[burnin:sample,8]), c(.05,.95))
quantile((gibb.sample[burnin:sample,9]), c(.05,.95))
quantile((gibb.sample[burnin:sample,10]), c(.05,.95))
References


Vita

Renee Dale obtained a bachelor’s in biology and philosophy from LSU in 2013. After taking a class on mathematical modeling in her senior year, she entered grad school studying mathematical biology, eventually completing a master’s in 2015, and her PhD in 2019. During this time, she was inspired to study statistics after managing to pass Dr Escobar’s probability courses and learning about Bayesian statistics. Thanks to this, she was able to learn about data mining and regression, and how to do some cool stuff. Despite pandemic delays, she plans to receive her master’s in Statistics in May 2021.