MCMC

Dexter Barrows February 27, 2016

1 Intro

Markov Chain Monte Carlo (MCMC) is part of a general class of methods designed to sample from the posterior distribution of model parameters. It is an algorithm used when we wish to fit a model M that depends on some parameter (or more typically vector of parameters) θ to observed data D. MCMC works by constructing a Markov Chain whose stationary or equilibrium distribution is used to approximate the desired posterior distribution.

is equal to? (distinguish between theoretical properties -- stationary distribution converges to desired post dist -- and numerical sampling, which approximates ...)

2 Markov Chains

(Figure 1).

Consider a finite state machine with 3 states $S = \{x_1, x_2, x_3\}$, where the probability of transitioning from one particular state to another is shown as a transition graph in Figure 1) below.

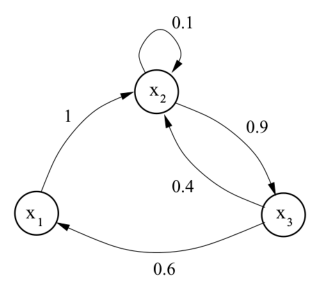


Figure 1: Finite state machine (Andrieu et al., 2003) talk about transition probabilities in the caption

The transition probabilities can be summarized as a matrix as

$$T = \begin{bmatrix} 0 & 1 & 0 \\ 0 & 0.1 & 0.9 \\ 0.6 & 0.4 & 0 \end{bmatrix}.$$

The probability vector $\mu(x^{(1)})$ for a state $x^{(1)}$ can be evolved using T by evaluating $\mu(x^{(1)})T$, then again by evaluating $\mu(x^{(1)})T^2$, and so on. If we take the limit as the number of transitions approaches infinity, we find

$$\lim_{t \to \infty} \mu(x^{(1)}) T^t = (27/122, 50/122, 45/122).$$

This indicates that no matter what we pick for the initial probability distribution $\mu(x^{(1)})$, the chain will always stabilize at the equilibrium distribution.

Note that this property holds when the chain satisfies the following conditions

- Irreducible Any state A can be reached from any other state B with non-zero probability
- Positive Recurrent The number of steps required for the chain to reach state A from state B must be finite
- Aperiodic The chain must be able to explore the parameter space without becoming trapped in a cycle

(maybe note that there are formal definitions for this in terms of linear algebra (Perron-Frobenius theorem), but we're not going there) Note that MCMC sampling generates a Markov chain $(\theta^{(1)}, \theta^{(2)}, ..., \theta^{(N)})$ that does indeed satisfy these conditions, and uses the chain's equilibrium distribution to approximate the posterior distribution of the parameter space.

see comments above

3 Likelihood

MCMC and similar methods hinge on the idea that the weight or support bestowed upon a particular set of parameters θ should be proportional to the probability of observing the data D given the model output using that set of parameters $M(\theta)$. In order to do this we need a way to evaluate whether or not $M(\theta)$ is a good fit for D; this is done by specifying a likelihood function $\mathcal{L}(\theta)$ such that

MCMC is usually defined in a Bayesian context, such that P(D|theta) (likelihood) naturally gives us P(theta|D) (once we specify a prior), via Bayes' rule. You're stating this in a sort of phenomenological way; normally it's done more formally. However, as I think I said early, Bayesian/frequentist approaches often get mixed up in the dynamical estimation literature (e.g. sometimes we do maximum a posteriori estimation rather than finding the mean; bit.ly/1LPBvRg gives a pointer to computing CIs by sampling from the "normalized likelihood"

$$\mathcal{L}(\theta) \propto P(D|\theta)$$
.

In standard Maximum Likelihood approaches, $\mathcal{L}(\theta)$ is searched to find a value of θ that maximizes $\mathcal{L}(\theta)$, then this θ is taken to be the most likely true value. Here our aim is to not just maximize the likelihood but to also explore the area around it.

in the standard approach one would typically compute *interval estimates* by profile likelihood. Perhaps be slightly more specific about point vs. interval estimates ...

4 Prior distribution

Another significant component of MCMC is the user-specified prior distribution for θ or distributions for the individual components of θ (Priors). Priors serve as a way for us to tell the MCMC algorithm what we think consist of good values for the parameters.

priors are *necessary* in the Bayesian framework, not just a convenient way to add info; if not specified they are still implicit ... Note that if very little is known about the parameters, or we are worried about biasing our estimate of the posterior, we can simply use a a wide uniform distribution. However, this handicaps the algorithm in two ways: convergence of the chain may become exceedingly slow, and more pressure is put on the likelihood function to be as good as possible – it will now be the only thing informing the algorithm of what constitutes a "good" set of parameters, and what should be considered poor.

maybe comment on the extreme of this case -- an improper prior (i.e. limit of Gaussian as std dev -> infty, or uniform as width -> infty) *can* lead to an improper posterior if data are weak enough. I don't normally think of weak priors as obstacles to convergence (if you're having that kind of problem with convergence then you probably have bigger inferential problems ...

5 Proposal distribution

As part of the MCMC algorithm, when we find a state in the parameter space that is accepted as part of the Markov chain construction process, we need a good way of generating a good next step to try. Unlike basic rejection sampling in which we would just randomly sample from our prior distribution, MCMC attempts to optimise our choices by choosing a step that is close enough to the last accepted step so as to stand a decent chance of also being accepted, but far enough away that it doesn't get "trapped" in a particular region of the parameter space.

This is done through the use of a proposal or candidate distribution. This will usually be a distribution centred around our last accepted step and with a dispersion potential narrower than that of out prior distribution.

The hoice of this distribution is theoretically not of the utmost importance, but in practice becomes important so as to not waste computer time.

6 Algorithm

Now that we have all the pieces necessary, we can discuss the details of the MCMC algorithm.

We will denote the previously discussed quantities as

- $p(\cdot)$ the prior distribution
- $q(\cdot|\cdot)$ the proposal distribution
- $\mathcal{L}(\cdot)$ the Likelihood function
- $\mathcal{U}(\cdot, \cdot)$ the uniform distribution

and the define the acceptance ratio, r, as

maybe refer this as the Metropolis-Hastings rule?

$$r = \frac{\mathcal{L}(\theta^*)p(\theta^*)q(\theta^*|\theta)}{\mathcal{L}(\theta)p(\theta)q(\theta|\theta^*)},$$

where θ^* is the proposed sample to draw from the posterior, and θ is the last accepted sample.

In the special case of the Metropolis Hastings variation of MCMC, the proposal distribution is symmetric, meaning $q(\theta^*|\theta) = q(\theta|\theta^*)$, and so the acceptance ratio simplifies to

$$r = \frac{\mathcal{L}(\theta^*)p(\theta^*)}{\mathcal{L}(\theta)p(\theta)}.$$

I believe this is just "Metropolis" (M-H refers to the version with the correction for asymmetric candidate distributions)

Thus, the MCMC algorithm is as follows.

Algorithm 1: Metropolis-Hastings MCMC

/* Samples from approximated posterior distribution

Output: Chain of samples $(\theta^{(1)}, \theta^{(2)}, ..., \theta^{(N)})$

```
/* Select a starting point
                                                                                                                               */
   Input : Initialize \theta^{(1)}
1 for i = 2 : N do
                                                                   pretty! I have to find out how to do this
        /* Sample
                                                                                                                               */
                                                                  formatting (listings package, or something
        \theta^* \sim q(\cdot|\theta^{(i-1)})
                                                                  else?)
        u \sim \mathcal{U}(0,1)
        /* Evaluate acceptance ratio
                                                                                                                               */
        r \leftarrow \frac{\mathcal{L}(\theta^*)p(\theta^*)}{2}
4
        /* Step acceptance criterion
        if u < \min\{1, r\} then
             \theta^{(i)} = \theta^*
6
```

In this way we are ensuring that steps that lead to better likelihood outcomes are likely to be accepted, but steps that do not will not be accepted as frequently. Note that these less "advantageous" moves will still occur but that this is by design – it ensures that as much of the parameter space as possible will be explored but more efficiently than using pure brute force.

*/

This is very much an optimization-centric approach; the more statistical approach is "this algorithm will lead to a stationary distribution equal to the posterior distribution" (I don't mind the optimization-centric approach, but you want to at least recognize the other perspective)

7 Burn-in

 $\theta^{(i)} = \theta^{(i-1)}$

One critical aspect of MCMC-based algorithms has yet to be discussed. The algorithm requires an initial starting point θ to be selected, but as the proposal distribution is supposed to restrict moves to an area close to the last accepted state, then the posterior distribution will be biased towards this starting point. This issue is avoided through the use of a Burn-in period.

Burning in a chain is the act of running the MCMC algorithm normally without saving first M samples. As we are seeking a chain of length N, the total computation will be equivalent to generating a chain of length M + N.

Thinning

8

really trivial but might want
\$n^{\textrm{th}}\$ or \$n\$th

Some models will require very long chains to get a good approximation of the posterior, which will consequently require a non-trivial amount of computer storage. One way to reduce the burden of storing so many samples is by thinning. This involves saving only every n^{th} step, which should still give a decent approximate of the posterior (since the chain has time to explore a large portion of the parameter space), but require less room to store.

9 Hamiltonian Monte Carlo

The Metropolis-Hastings algorithm has a primary drawback in that the parameter space may not be explored efficiently – a consequence of the rudimentary proposal mechanism. Instead, smarter moves can be proposed through the use of Hamiltonian dynamics, leading to a better exploration of the target distribution and a decrease in overall computational complexity.

perhaps? note adaptive/block M-H MCMC as an intermediate (although probably still dominated by HMC)

From physics, we will borrow the ideas of potential and kinetic energy. Here potential energy is analogous to the negative log likelihood of the parameter selection given the data, formally

$$U(\theta) = -log(\mathcal{L}(\theta)p(\theta)). \qquad \text{use log}$$
 (1)

Kinetic energy will serve as a way to "nudge" the parameters along a different moment for each component of θ . We introduce n auxiliary variables $r = (r_1, r_1, ..., r_n)$, where n is the number of components in θ . Note that the samples drawn for r are not of interest, they are only used to inform the evolution of the Hamiltonian dynamics of the system. We can now define the kinetic energy as

$$K(r) = \frac{1}{2}r^{T}M^{-1}r,$$
 (2)

where M is an $n \times n$ matrix. In practice M can simply be chosen as the identity matrix of size n, however it can also be used to account for correlation between components of θ .

probably don't need to warn you, but make sure that you're sufficiently paraphrasing technical discussions, and cite appropriately The Hamiltonian of the system is defined as

$$H(\theta, r) = U(\theta) + K(r), \tag{3}$$

w (lowercase)

Where the Hamiltonian dynamics of the combined system can be simulated using the following system of ODEs. colon?

$$\frac{d\theta}{dt} = M^{-1}r$$

$$\frac{dr}{dt} = -\nabla U(\theta)$$
 period?
$$(4)$$

.

It is tempting to try to integrate this system using the standard Euler evolution scheme, but in practice this leads to instability. Instead the "Leapfrog" scheme is used. This scheme is very similar to Euler scheme, except instead of using a fixed step size h for all evolutions, a step size of ε is used for most evolutions, with a half step size of $\varepsilon/2$ for evolutions of $\frac{dr}{dt}$ at the first step, and last step L. In this way the evolution steps "leapfrog" over each other while using future values from the other set of steps, leading to the scheme's name.

The end product of the Leapfrog steps are the new proposed parameters (θ^*, r^*) . These are either accepted or rejected using a mechanism similar to that of standard Metropolis-Hastings MCMC. Now, however, the acceptance ratio r is defined as

$$r = \exp\left[H(\theta, r) - H(\theta^*, r^*)\right],\tag{5}$$

where (θ, r) are the last values in the chain.

Algorithm 2: Hamiltonian MCMC /* Select a starting point */ Input : Initialize $\theta^{(1)}$ 1 for i = 2 : N do /* Resample moments */ for i = 1 : n do $\mathbf{2}$ $r(i) \leftarrow \mathcal{N}(0,1)$ 3 /* Leapfrog initialization */ $\theta_0 \leftarrow \theta^{(i-1)}$ 4 $r_0 \leftarrow r - \nabla U(\theta_0) \cdot \varepsilon / 2$ $\mathbf{5}$ /* Leapfrog intermediate steps */ for j = 1: L-1 do 6 $\begin{vmatrix} \theta_j \leftarrow \theta_{j-1} + M^{-1} r_{j-1} \cdot \varepsilon \\ r_j \leftarrow r_{j-1} - \nabla U(\theta_j) \cdot \varepsilon \end{vmatrix}$ 7 /* Leapfrog last steps */ $\theta^* \leftarrow \theta_{L-1} + M^{-1} r_{L-1} \cdot \varepsilon$ 9 $r^* \leftarrow \nabla U(\theta_L) \cdot \varepsilon/2 - r_{L-1}$ 10 /* Evaluate acceptance ratio */ $r = \exp\left[H(\theta^{(i-1)}, r) - H(\theta^*, r^*)\right]$ 11 /* Sample */ $u \sim \mathcal{U}(0,1)$ 12 /* Step acceptance criterion */ **if** $u < \min\{1, r\}$ **then** 13 $\theta^{(i)} = \theta^*$ 14 $_{ m else}$ 15 $\theta^{(i)} = \theta^{(i-1)}$ 16

Note that the parameters ε and L have to be tuned in order to maintain stability and maximize efficiency, a sometimes non-trivial process.

*/

but in Stan the No U-Turn Sampler kind of takes care of this \dots

/* Samples from approximated posterior distribution

Output: Chain of samples $(\theta^{(1)}, \theta^{(2)}, ..., \theta^{(N)})$

10 Fitting an SIR Model to Synthetic Epidemic Data

Here we will examine a test case in which Hamiltonian MCMC will be used to fit a Susceptible-Infected-Removed (SIR) epidemic model to mock infectious count data.

The synthetic data was produced by taking the solution to a basic SIR ODE model, sampling it at regular intervals, and perturbing those values by adding in observation noise. The SIR model used was

don't remember if you've defined SIR model already. Again, might be a good thing to get out of the way in the introduction

$$\frac{dS}{dt} = -\beta IS
\frac{dI}{dt} = \beta IS - rI
\frac{dR}{dt} = rI$$
(6)

where S is the number of individuals susceptible to infection, I is the number of infectious individuals, R is the number of recovered individuals, $\beta = R_0 r/N$ is the force of infection, R_0 is the number of secondary cases per infected individual, r is the recovery rate, and N is the population size.

this material could conceivably go in a technical appendix ...

The solution to this system was obtained using the ode() function from the deSolve package. The required derivative array function in the format required by ode() was specified as

```
SIR ← function(Time, State, Pars) {
    with(as.list(c(State, Pars)), {
             \leftarrow R0*r/N
                            # calculate Beta
         \texttt{BSI} \leftarrow \texttt{B*S*I}
                            # save product
         rI \leftarrow r*I
                            # save product
         dS = -BSI
                            # change in Susceptible people
         dI = BSI - rI
                            # change in Infected people
                            # change in Removed (recovered people)
         return(list(c(dS, dI, dR)))
    })
}
```

The true parameter values were set to $R_0 = 3.0, r = 0.1, N = 500$ by

```
pars \leftarrow c(R0 \leftarrow 3.0, # new infected people per infected person \leftarrow 0.1, # recovery rate N \leftarrow 500) # population size
```

The system was integrated over [0, 100] with infected counts drawn at each integer time step. These timings were set using

The initial conditions were set to 5 infectious individuals, 495 people susceptible to infection, and no one had yet recovered from infection and been removed. These were set using

```
y_{ini} \leftarrow c(S = 495, I = 5, R = 0) # initial conditions
```

The ode() function is called as

```
odeout \leftarrow ode(y_ini, times, SIR, pars)
```

where odeout is a $(T+1) \times 4$ matrix where the rows correspond to solutions at the given times (the first row is the initial condition), and the columns correspond to the solution times and S-I-R counts at those times.

The observation error was taken to be $\varepsilon_{obs} \sim \mathcal{N}(0, \sigma)$, where individual values were drawn for each synthetic data point.

These "true" values were perturbed to mimic observation error by

```
set.seed(1001) # set RNG seed for reproducibility
sigma ← 5 # observation error standard deviation
infec_counts_raw ← odeout[,3] + rnorm(101, 0, sigma)
infec_counts ← ifelse(infec_counts_raw < 0, 0, infec_counts)
```

or pmax(0, infec_counts_raw)

where the last two lines simply set negative observations (impossible) to 0.

Plotting the data using the ggplot2 package by

probably a little more detail than needed (certainly more than in the other chapter I just read) we obtain

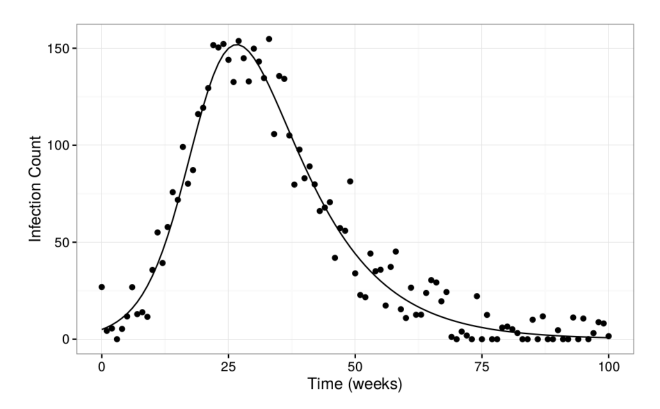


Figure 2: True SIR ODE solution infected counts, and with added observation noise

The Hamiltonian MCMC model fitting was done using Stan (http://mc-stan.org/), a program written in C++ that does Baysian statistical inference using Hamiltonian MCMC. Stan's R interface (http://mc-stan.org/interfaces/rstan.html) was used to ease implementation.

In order to use an Explicit Euler-like stepping method in the later Stan model (both for speed and for integration method homogeneity with other methods against which HMCMC was compared), the synthetic observation counts were treated as weekly observations in which the counts on the other six days of the week were unobserved. For computational and organizational simplicity, these vales were set to -1 (all valid observations are non-negative). This is done in R using

```
sPw ← 7  # steps per week
datlen ← (T-1)*7 + 1  # size of sparse data vector

data ← matrix(data = -1, nrow = T+1, ncol = sPw)
data[,1] ← infec_counts
standata ← as.vector(t(data))[1:datlen]
```

The data to be fed into the R Stan interface is packed as

```
sir_data ← list( T = datlen,  # simulation time
y = standata,  # infection count data
```

```
N = 500, # population size h = 1/sPw ) # step size per day
```

For efficiency we allow Stan to save compiled code to avoid recompilation, and allow multiple chains to be run simultaneously on separate CPU cores

```
rstan_options(auto_write = TRUE)
options(mc.cores = parallel::detectCores())
```

Now we call the Stan fitting function

```
stan_{options} \leftarrow list(
                              chains = 4,
                                              # number of chains
2
                                      = 2000, # iterations per chain
                              warmup = 1000, # warmup interations
4
                                      = 2 )
                                             # thinning number
    fit \leftarrow stan(file)
                          = "d_sirode_euler.stan",
                         = sir_data,
                 chains = stan_options$chains,
                         = stan_options$iter,
                 iter
                 warmup
                         = stan_options$warmup,
                 thin
                          = stan_options$thin )
```

which fits the model in the file d_sirode_euler.stan to the data passed in through sir_data. The options here specify that 10 chains will be run, each with a burn in period of 1000 steps, with 5000 steps to sample over, and only sampling every 10th step. Options are saved so they can be accessed later.

The Stan file contains three blocks that together specify the model. First, the data block specifies the information the model expects to be given. Here, this is

```
data {
          int
                   <lower=1>
                                Τ;
                                        // total integration steps
4
          real
                                y[T];
                                        // observed number of cases
          int
                                        // population size
                   <lower=1>
                                N;
          real
                                h;
                                        // step size
      }
```

where each of the data variables correspond to data passed in through the previously shown R code.

Next the parameters block specifies what Stan is expected to estimate. Here this is

```
parameters {

real <lower=0, upper=10> sigma; // observation error
real <lower=0, upper=10> R0; // R0
real <lower=0, upper=10> r; // recovery rate
```

```
real <lower=0, upper=500> y0[3]; // initial conditions
}
```

from which to draw new parameter values (?) Finally we have the model block. This crucial part of the code specifies the interaction between the parameters and the data. The core component of the model indicates we are fitting an approximation of an ODE model using Euler integration steps (one per day), with the initial conditions and SIR parameters unknown. Further, we can also specify the prior distributions to draw new parameter values from. The initial conditions are taken to be close to the initial data point, with adjustment for observation error, while the other parameters are assumed to be coming from log-normal distributions with relatively small means. Together, we have

```
model {
           real S[T];
4
           real I[T]:
           real R[T];
           S[1] <- y0[1];
8
          I[1] <- y0[2];
9
          R[1] <- y0[3];
          y[1] \sim normal(y0[2], sigma);
          for (t in 2:T) {
               S[t] \leftarrow S[t-1] + h*( - S[t-1]*I[t-1]*R0*r/N );
               I[t] \leftarrow I[t-1] + h*(S[t-1]*I[t-1]*R0*r/N - I[t-1]*r);
               R[t] \leftarrow R[t-1] + h*(I[t-1]*r);
               if (y[t] > 0) {
                   y[t] ~ normal( I[t], sigma );
               }
          }
          y0[1] ~ normal(N - y[1], sigma);
           y0[2] ~ normal(y[1], sigma);
           theta[1]
                          lognormal(1,1);
                          lognormal(1,1);
           theta[2]
                        ~ lognormal(1,1);
           sigma
      }
```

Examining the traceplot for the post-warmup chain data returned by the stan() function in the fit object, we see that the chains are mixing well and convergence has likely been reached. This is shown below in Figure 3

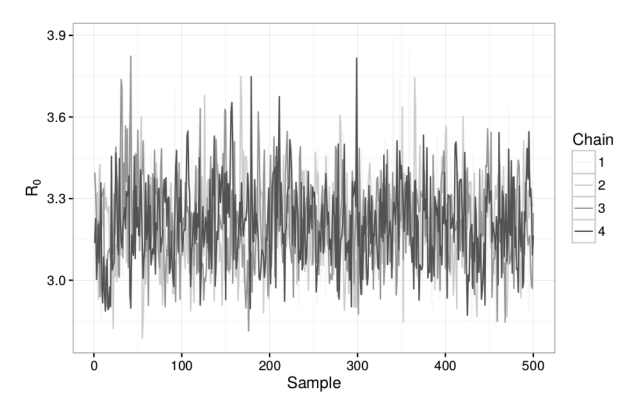


Figure 3: Traceplot of samples drawn for parameter R_0 , excluding warmup

Further, if we look at the chain data including the warmup samples in Figure 4, we can see why is is wise to discard these samples (note the scale).

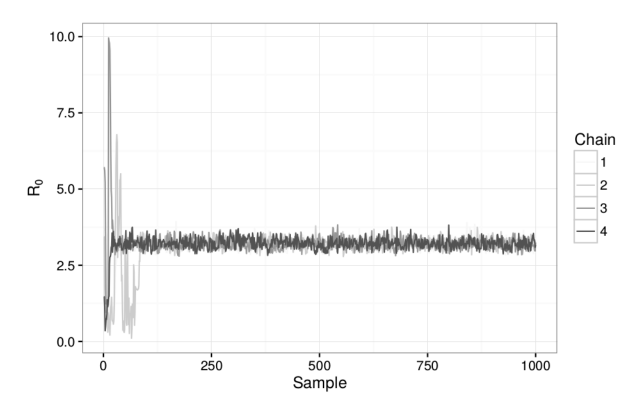


Figure 4: Traceplot of samples drawn for parameter R_0 , including warmup

Now if we look at the kernel density estimates for each of the model parameters and the initial number of cases, we see that while the estimates are not perfect, they are fairly decent. This is shown below in Figure 5.

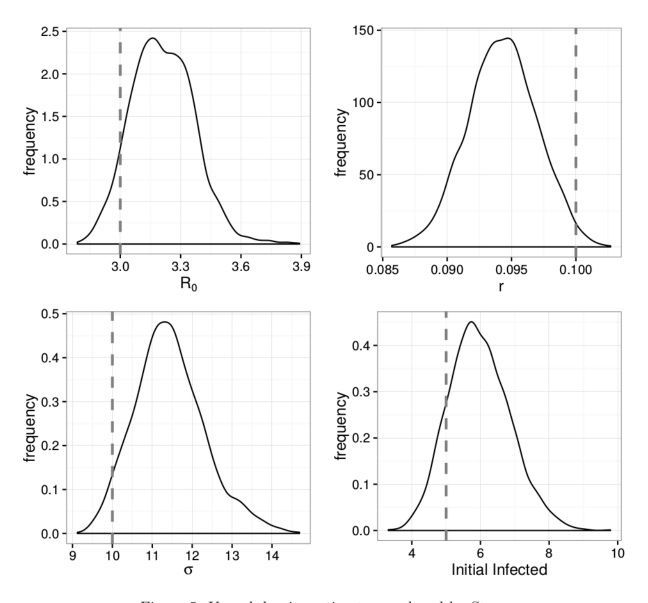


Figure 5: Kernel density estimates produced by Stan

Appendices

A Full R code

This code will run all the indicated analysis and produce all plots.

```
## Dexter Barrows
 2 ## McMaster University
 3 ## 2016
 5 library(deSolve)
 6 library(rstan)
 7 library(shinystan)
 8 library(ggplot2)
 9 library(RColorBrewer)
10 library(reshape2)
12 | SIR \leftarrow function(Time, State, Pars) {
        with(as.list(c(State, Pars)), {
                 \leftarrow R0*r/N
             BSI \leftarrow B*S*I
18
             \texttt{rI} \ \leftarrow \texttt{r} \! \star \! \texttt{I}
             dS = -BSI
             dI = BSI - rI
             dR = rI
             return(list(c(dS, dI, dR)))
        })
28 }
                                 # average number of new infected individuals per
   pars \leftarrow c(R0 \leftarrow 3.0,
       infectious person
                 r
                     \leftarrow 0.1,
                                 # recovery rate
                     \leftarrow 500)
                                 # population size
34 T ← 100
35 | y_ini \leftarrow c(S = 495, I = 5, R = 0)
36 times \leftarrow seq(0, T, by = 1)
38 odeout \leftarrow ode(y_ini, times, SIR, pars)
40 set.seed(1001)
41 sigma \leftarrow 10
42 infec_counts_raw \leftarrow odeout[,3] + rnorm(101, 0, sigma)
43 infec_counts \leftarrow ifelse(infec_counts_raw < 0, 0, infec_counts_raw)
```

```
44
  g \leftarrow qplot(0:T, odeout[,3], geom = "line", xlab = "Time (weeks)", ylab = "
      Infection Count") +
       geom_point(aes(y = infec_counts)) +
       theme_bw()
49 print(g)
50 ggsave(g, filename="dataplot.pdf", height=4, width=6.5)
52 sPw \leftarrow 7
53 datlen \leftarrow (T-1)*7 + 1
55 data \leftarrow matrix(data = -1, nrow = T+1, ncol = sPw)
56 \mid data[,1] \leftarrow infec\_counts
57 standata \leftarrow as.vector(t(data))[1:datlen]
59 | sir_data \leftarrow list(T = datlen,
                                   # simulation time
                      y = standata, # infection count data
                      N = 500.
                                    # population size
                      h = 1/sPw )
                                    # step size per day
64 rstan_options(auto_write = TRUE)
65 options(mc.cores = parallel::detectCores())
66 stan_options \leftarrow list(
                            chains = 4,
                                          # number of chains
                                   = 2000, # iterations per chain
                             iter
                             warmup = 1000, # warmup interations
                                    = 2) # thinning number
                             thin
                       = "d_sirode_euler.stan",
70 fit \leftarrow stan(file
                        = sir_data,
                data
                        = stan_options$chains,
                chains
                iter
                         = stan_options$iter,
                warmup = stan_options$warmup,
                         = stan_options$thin )
                thin
  exfit ← extract(fit, permuted = TRUE, inc_warmup = FALSE)
79 R0points \leftarrow exfit$R0
80 R0kernel \leftarrow qplot(R0points, geom = "density", xlab = expression(R[0]), ylab
      = "frequency") +
           geom_vline(aes(xintercept=R0), linetype="dashed", size=1, color="
               grey50") +
           theme_bw()
84 print (R0kernel)
  ggsave(R0kernel, filename="kernelR0.pdf", height=3, width=3.25)
86
87 rpoints \leftarrow exfit$r
88 rkernel \leftarrow qplot(rpoints, geom = "density", xlab = "r", ylab = "frequency")
            geom_vline(aes(xintercept=r), linetype="dashed", size=1, color="
               grev50") +
90
           theme_bw()
92 print(rkernel)
```

```
93 ggsave(rkernel, filename="kernelr.pdf", height=3, width=3.25)
95 sigmapoints \leftarrow exfit$sigma
96 sigmakernel ← qplot(sigmapoints, geom = "density", xlab = expression(sigma)
       , ylab = "frequency") +
            geom_vline(aes(xintercept=sigma), linetype="dashed", size=1, color=
                "grey50") +
            theme_bw()
100 print(sigmakernel)
101 ggsave(sigmakernel, filename="kernelsigma.pdf", height=3, width=3.25)
103 infecpoints \leftarrow exfit$y0[,2]
104 infeckernel \leftarrow qplot(infecpoints, geom = "density", xlab = "Initial Infected
       ", ylab = "frequency") +
            geom_vline(aes(xintercept=y_ini[['I']]), linetype="dashed", size=1,
                 color="grey50") +
106
            theme_bw()
108 print(infeckernel)
109 ggsave(infeckernel, filename="kernelinfec.pdf", height=3, width=3.25)
111 exfit 		 extract(fit, permuted = FALSE, inc_warmup = FALSE)
112 plotdata ← melt(exfit[,,"R0"])
113 tracefitR0 \leftarrow ggplot() +
                   geom_line(data = plotdata,
                             aes(x = iterations,
                             y = value,
                             color = factor(chains, labels = 1:stan_options$
                                 chains))) +
                   labs(x = "Sample", y = expression(R[0]), color = "Chain") +
                   scale_color_brewer(palette="Greys") +
                   theme_bw()
122 print(tracefitR0)
123 ggsave(tracefitR0, filename="traceplotR0.pdf", height=4, width=6.5)
|125| exfit \leftarrow extract(fit, permuted = FALSE, inc_warmup = TRUE)
126 plotdata \leftarrow melt(exfit[,,"R0"])
127 tracefitR0 \leftarrow ggplot() +
                   geom_line(data = plotdata,
                             aes(x = iterations,
                             y = value,
                             color = factor(chains, labels = 1:stan_options$
                                 chains))) +
                   labs(x = "Sample", y = expression(R[0]), color = "Chain") +
                   scale_color_brewer(palette="Greys") +
                   theme_bw()
136 print(tracefitR0)
137 ggsave(tracefitR0, filename="traceplotR0_inc.pdf", height=4, width=6.5)
139 sso \leftarrow as.shinystan(fit)
|140| sso \leftarrow launch_shinystan(sso)
```

B Full Stan code

Stan model code to be used with the preceding R code.

```
## Dexter Barrows
  ## McMaster University
  ## 2016
  data {
               <lower=1>
      int
                            Τ;
                                    // total integration steps
      real
                            y[T];
                                    // observed number of cases
      int
               <lower=1>
                            N;
                                    // population size
                                    // step size
      real
                            h;
12
  }
  parameters {
      real <lower=0, upper=10>
                                    R0;
      real <lower=0, upper=10>
                                             // recovery rate
                                    r;
      real <lower=0, upper=20>
                                    sigma; // observation error
      real <lower=0, upper=500>
                                    y0[3];
                                            // initial conditions
  }
23 model {
      real S[T];
      real I[T];
      real R[T];
      S[1] <- y0[1];
      I[1] <- y0[2];
      R[1] <- y0[3];
      y[1] ~ normal(y0[2], sigma);
      for (t in 2:T) {
           S[t] \leftarrow S[t-1] + h*( - S[t-1]*I[t-1]*R0*r/N );
38
           I[t] \leftarrow I[t-1] + h*(S[t-1]*I[t-1]*R0*r/N - I[t-1]*r);
           R[t] \leftarrow R[t-1] + h*(I[t-1]*r);
           if (y[t] > 0) {
               y[t] \sim normal(I[t], sigma);
           }
```

I'm a little torn by the step-by-step discussion in the body of this chapter. While it would be a really useful *pedagogical* tool, I'm not sure it belongs in the body of a master's thesis -- basically, your examiners all know these techniques and will trust that you're doing it right/sensibly. If you have really strong feelings about it, you could keep it, but otherwise I would suggest you cut most of it (sorry) and/or move it to the appendix.