FUN WITH FORECASTING USING STOCHASTIC NONLINEAR DYNAMICS

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Abstract

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Dedication

To Mom and Dad

Acknowledgements

Soooooo many people

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Chapter 1

Introduction

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Chapter 2

Hamiltonian MCMC

2.1 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.

2.2 Markov Chains

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 [2.1].

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

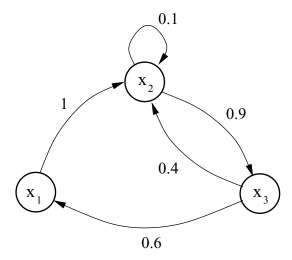


Figure 2.1: Finite state machine. (Andrieu et al., 2003)

 $\mu(x^{(1)})T$, 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

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.

2.3 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

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

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.

2.4 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.

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.

2.5 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.

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

2.6 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

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

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

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

Thus, the MCMC algorithm shown in Algorithm [1].

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.

Algorithm 1: Metropolis-Hastings MCMC /* Select a starting point */ **Input**: Initialize $\theta^{(1)}$ 1 for i = 2 : N do /* Sample */ $\theta^* \sim q(\cdot|\theta^{(i-1)})$ 2 $u \sim \mathcal{U}(0,1)$ /* Evaluate acceptance ratio */ $\underline{\mathcal{L}(\theta^*)p(\theta^*)}$ /* Step acceptance criterion */ if $u < \min\{1, r\}$ then $\mathbf{5}$ $\theta^{(i)} = \theta^*$ 6 else 7 $\theta^{(i)} = \theta^{(i-1)}$ 8 /* Samples from approximated posterior distribution */ Output: Chain of samples $(\theta^{(1)}, \theta^{(2)}, ..., \theta^{(N)})$

2.7 Burn-in

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.

2.8 Thinning

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.

2.9 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.

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

$$U(\theta) = -\log(\mathcal{L}(\theta)p(\theta)). \tag{2.1}$$

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

$$K(r) = \frac{1}{2}r^{T}M^{-1}r,$$
(2.2)

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 .

The Hamiltonian of the system is defined as

$$H(\theta, r) = U(\theta) + K(r), \tag{2.3}$$

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

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

$$\frac{dr}{dt} = -\nabla U(\theta)$$
(2.4)

.

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

$$r = \exp\left[H(\theta, r) - H(\theta^*, r^*)\right],\tag{2.5}$$

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

Together, we have Algorithm [2].

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

2.10 Fitting

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

$$\frac{dS}{dt} = -\beta IS$$

$$\frac{dI}{dt} = \beta IS - rI$$

$$\frac{dR}{dt} = rI$$
(2.6)

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 $r_0 \leftarrow r - \nabla U(\theta_0) \cdot \varepsilon/2$ 5 /* Leapfrog intermediate steps */ for j = 1 : L - 1 do 6 $\begin{bmatrix} \ddot{\theta_j} \leftarrow \theta_{j-1} + M^{-1}r_{j-1} \cdot \varepsilon \\ r_j \leftarrow r_{j-1} - \nabla U(\theta_j) \cdot \varepsilon \end{bmatrix}$ 7 8 /* 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 $\star/$ $r = \exp\left[H(\theta^{(i-1)}, r) - H(\theta^*, r^*)\right]$ 11 /* Sample */ $u \sim \mathcal{U}(0,1)$ 12/* Step acceptance criterion */ if $u < \min\{1, r\}$ then **13** $\theta^{(i)} = \theta^*$ 14 else 15 $\theta^{(i)} = \theta^{(i-1)}$ 16 /* Samples from approximated posterior distribution */ Output: Chain of samples $(\theta^{(1)}, \theta^{(2)}, ..., \theta^{(N)})$

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.

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
         BSI \leftarrow B*S*I
                          # save product
         rI \leftarrow r*I
                          # save product
         dS = -BSI
                          # change in Susceptible people
         dI = BSI - rI
                          # change in Infected people
         dR = rI
                          # 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

```
T \leftarrow 100  # total integration time times \leftarrow seq(0, T, by = 1)  # times to draw solution values
```

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)
```

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

Plotting the data using the ggplot2 package by

we obtain

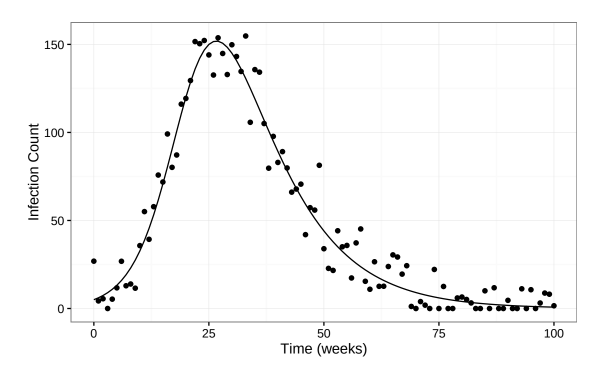


Figure 2.2: True SIR ODE solution infected counts, and with added observation noise

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

```
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
                                 = 2000, # iterations per chain
                          iter
                          warmup = 1000, # warmup interations
                                 = 2 )
                                        # thinning number
                     = "d_sirode_euler.stan",
fit \leftarrow stan(file)
             data
                     = sir_data,
             chains
                     = stan_options$chains,
                     = stan_options$iter,
             iter
                     = stan_options$warmup,
             warmup
                     = stan_options$thin )
             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> T; // total integration steps
real y[T]; // observed number of cases
int <lower=1> N; // population size
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 {

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
}
```

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];
    real I[T];
    real R[T];
    S[1] \leftarrow y0[1];
    I[1] \leftarrow y0[2];
    R[1] \leftarrow 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);
                    lognormal(1,1);
    theta[1]
                    lognormal(1,1);
    theta[2]
    sigma
                   lognormal(1,1);
```

Examining the traceplot for the post-warmup chain data returned by the stan()

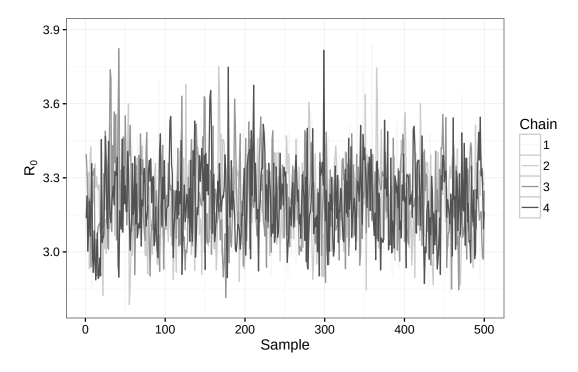


Figure 2.3: Traceplot of samples drawn for parameter R_0 , excluding warmup

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 [2.3].

Further, if we look at the chain data including the warmup samples in Figure [2.4], we can see why is is wise to discard these samples (note the scale).

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 [2.5].

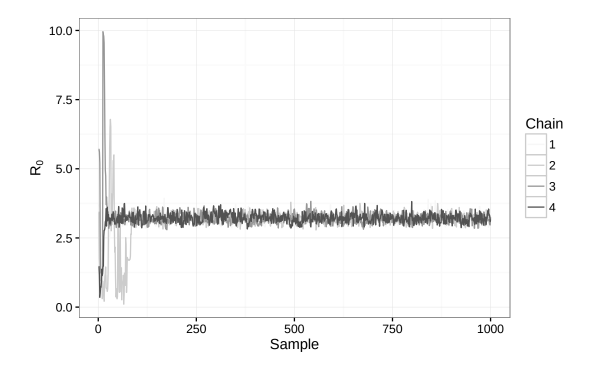


Figure 2.4: Traceplot of samples drawn for parameter R_0 , including warmup.

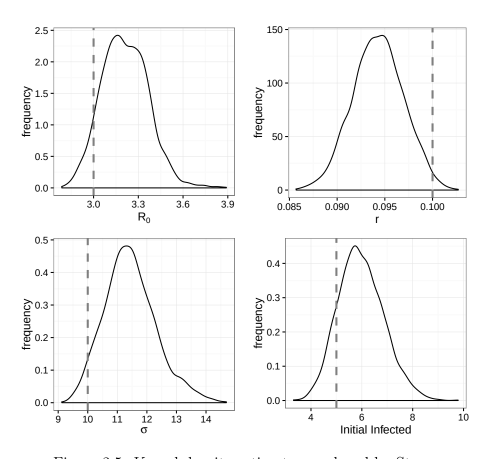


Figure 2.5: Kernel density estimates produced by Stan

Chapter 3

Iterated Filtering

3.1 Intro

Particle filters are similar to MCMC-based methods in that they attempt to draw samples from an approximation of the posterior distribution of model parameters θ given observed data D. Instead of constructing a Markov chain and approximating its stationary distribution, a cohort of "particles" are used to move through the data in an on-line (sequential) fashion with the cohort being culled of poorly-performing particles at each iteration via importance sampling. If the culled particles are not replenished, this will be a Sequential Importance Sampling (SIS) particle filter. If the culled particles are replenished from surviving particles, in a sense setting up a process not dissimilar from Darwinian selection, then this will be a Sequential Importance Resampling (SIR) particle filter.

3.2 Formulation

Particle filters, also called Sequential Monte-Carlo (SMC) or bootstrap filters, feature similar core functionality as the venerable Kalman Filter. As the algorithm moves through the data (sequence of observations), a prediction-update cycle is used to simulate the evolution of the model M with different particular parameter selections, track how closely these predictions approximate the new observed value, and update the current cohort appropriately.

Two separate functions are used to simulate the evolution and observation processes. The "true" state evolution is specified by

$$X_{t+1} \sim f_1(X_t, \theta), \tag{3.1}$$

And the observation process by

$$Y_t \sim f_2(X_t, \theta). \tag{3.2}$$

Note that components of θ can contribute to both functions, but a typical formulation is to have some components contribute to $f_1(\cdot, \theta)$ and others to $f_2(\cdot, \theta)$.

The prediction part of the cycle utilises $f_1(\cdot, \theta)$ to update each particle's current state estimate to the next time step, while $f_2(\cdot, \theta)$ is used to evaluate a weighting w for each particle which will be used to determine how closely that particle is estimating the true underlying state of the system. Note that $f_2(\cdot, \theta)$ could be thought of as a probability of observing a piece of data y_t given the particle's current state estimate and parameter set, $P(y_t|X_t,\theta)$. Then, the new cohort of particles is drawn from the old cohort proportional to the weights. This process is repeated until the set of observations D is exhausted.

3.3 Algorithm

Now we will formalize the particle filter.

We will denote each particle $p^{(j)}$ as the j^{th} particle consisting of a state estimate at time t, $X_t^{(j)}$, a parameter set $\theta^{(j)}$, and a weight $w^{(j)}$. Note that the state estimates will evolve with the system as the cohort traverses the data.

The algorithm for a Sequential Importance Resampling particle is shown in Algorithm 3.

3.4 Particle Collapse

Not uncommonly, a situation may arise in which a single particle is assigned a normalized weight very close to 1 and all the other particles are assigned weights very close to 0. When this occurs, the next generation of the cohort will overwhelmingly consist of descendants of the heavily-weighted particle, termed particle collapse or degeneracy.

Algorithm 3: SIR particle filter

```
/* Select a starting point
  Input: Observations D = y_1, y_2, ..., y_T, initial particle distribution P_0 of size
              J
  /* Setup
                                                                                           */
1 Initialize particle cohort by sampling (p^{(1)}, p^{(2)}, ..., p^{(J)}) from P_0
2 for t = 1 : T do
      /* Evolve
                                                                                           */
      for j = 1:J do
3
       /* Weight
                                                                                           */
      for j = 1:J do
5
       w^{(j)} \leftarrow P(y_t | X_t^{(j)}, \theta^{(j)}) = f_2(X_t^{(j)}, \theta^{(j)})
      /* Normalize
                                                                                           */
      for j = 1:J do
7
       /* Resample
                                                                                           */
      p^{(1:J)} \leftarrow \text{sample}(p^{(1:J)}, \text{prob} = w, \text{replace} = true)
  /* Samples from approximated posterior distribution
                                                                                           */
  Output: Cohort of posterior samples (\theta^{(1)}, \theta^{(2)}, ..., \theta^{(J)})
```

Since the basic SIR particle filter does not perturb either the particle system states or system parameter values, the cohort will quickly consist solely of identical particles, effectively halting further exploration of the parameter space as new data is introduced.

A similar situation occurs when a small number of particles (but not necessarily a single particle) split almost all of the normalized weight between them, then jointly dominate the resampling process for the remainder of the iterations. This again halts the exploration of the parameter space with new data.

In either case, the hallmark feature used to detect collapse is the same – at some point the cohort will consist of particles with very similar or identical parameter sets which will consequently result in their assigned weights being extremely close together.

Mathematically, we are interested in the number of effective particles, N_{eff} , which represents the number of particles that are acceptably dissimilar. This is estimated by evaluating

$$N_{eff} = \frac{1}{\sum_{1}^{J} (w^{(j)})^2}.$$
(3.3)

This can be used to diagnose not only when collapse has occurred, but can also indicate when it is near.

3.5 Iterated Filtering and Data Cloning

A particle filter hinges on the idea that as it progresses through the data set D, its estimate of the posterior carried in the cohort of particles approaches maximum likelihood. However, this convergence may not be fast enough so that the estimate it produces is of quality before the data runs out. One way around this problem is to "clone" the data and make multiple passes through it as if it were a continuation of the original time series. Note that the system state contained in each particle will have to be reset with each pass.

Rigorous proofs have been developed (references to Ionides et. al. work) that show that by treating the parameters as stochastic processes instead of fixed values, the multiple passes through the data will indeed force convergence of the process mean toward maximum likelihood, and the process variance toward 0.

3.6 IF2

The successor to Iterated Filtering 1 (reference), Iterated Filtering 2 is simpler, faster, and demonstrated better convergence toward maximum likelihood (reference). The core concept involves a two-pronged approach. First, Data cloning is used to allow more time for the parameter stochastic process means to converge to maximum likelihood, and frequent cooled perturbation of the particle parameters allow better exploration of the parameter space while still allowing convergence to good point estimates.

It is worth noting that IF2 is not designed to estimate the full posterior distribution, but in practice can be used to do so within reason. Further, IF2 thwarts the problem of particle collapse by keeping at least some perturbation in the system at all times. It is important to note that while true particle collapse will not occur, there is still risk of a pseudo-collapse in which all particles will be extremely close to one another so as to be virtually indistinguishable. However this will only occur with the use of overly-aggressive cooling strategies or by specifying an excessive number of passes through the data.

An important new quantity is the particle perturbation density denoted $h(\theta|,\sigma)$. Typically this is multi-normal with σ being a vector of variances proportional to the expected values of θ . In practice the proportionality can be derived from current means or specified ahead of time. Further, these intensities must decrease over time. This can be done via exponential or geometric cooling, a decreasing step function, a combination of these, or though some other similar scheme.

The algorithm for IF2 can be seen in Algorithm 4.

3.7 Fitting

Here we will examine a test case in which IF2 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

Algorithm 4: IF2

```
/* Select a starting point
    Input: Observations D = y_1, y_2, ..., y_T, initial particle distribution P_0 of size
                  J, decreasing sequence of perturbation intensity vectors
                  \sigma_1, \sigma_2, ..., \sigma_M
    /* Setup
                                                                                                                */
 1 Initialize particle cohort by sampling (p^{(1)}, p^{(2)}, ..., p^{(J)}) from P_0
    /* Particle seeding distribution
                                                                                                                */
 \mathbf{2} \ \Theta \leftarrow P_0
 3 for m = 1 : M do
         /* Pass perturbation
                                                                                                                */
         for j = 1:J do
 4
          p^{(j)} \sim h(\Theta^{(j)}, \sigma_m)
         for t = 1 : T do
 6
              for j = 1:J do
 7
                   /* Iteration perturbation
                                                                                                               */
                  p^{(j)} \sim h(p^{(j)}, \sigma_m)
 8
                  /* Evolve X_t^{(j)} \leftarrow f_1(X_{t-1}^{(j)}, \theta^{(j)})
 9
                  /* Weight w^{(j)} \leftarrow P(y_t | X_t^{(j)}, \theta^{(j)}) = f_2(X_t^{(j)}, \theta^{(j)})
                                                                                                               */
10
              /* Normalize
                                                                                                               */
              for j = 1:J do
11
               w^{(j)} \leftarrow w^{(j)} / \sum_{1}^{J} w^{(j)}
12
              /* Resample
                                                                                                               */
             p^{(1:J)} \leftarrow \text{sample}(p^{(1:J)}, \text{prob} = w, \text{replace} = true)
13
         /* Collect particles for next pass
                                                                                                                */
         for j = 1 : J \text{ do}
14
            \Theta^{(j)} \leftarrow p^{(j)}
15
    /* Samples from approximated posterior distribution
                                                                                                                */
    Output: Cohort of posterior samples (\theta^{(1)}, \theta^{(2)}, ..., \theta^{(J)})
```

$$\frac{dS}{dt} = -\beta IS
\frac{dI}{dt} = \beta IS - rI
\frac{dR}{dt} = rI$$
(3.4)

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.

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
        BSI \leftarrow B*S*I
                          # save product
         rI \leftarrow r*I
                          # save product
         dS = -BSI
                          # change in Susceptible people
         dI = BSI - rI
                          # change in Infected people
         dR = rI
                          # 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 = 3.0, # new infected people per infected person r = 0.1, # recovery rate N = 500) # population size
```

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

The ode() function is called as

```
odeout ← ode(y = true_init_cond, times = 0:(T-1), func = SIR, parms = true_pars)
```

where **odeout** is a $T \times 4$ 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 ← 10 # observation error standard deviation
infec_counts_raw ← odeout[,3] + rnorm(101, 0, sigma)
infec_counts ← ifelse(infec_counts_raw < 0, 0, infec_counts)
```

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

Plotting the data using the ggplot2 package by

we obtain Figure [7.1].

The IF2 algorithm was implemented in C++ for speed, and integrated into the R workflow using the Rcpp package. The C++ code is compiled using

```
sourceCpp(paste(getwd(),"if2.cpp",sep="/"))
```

Then run and packed into a data frame using

```
paramdata ← data.frame(if2(infec_counts[1:Tlim], Tlim, N))
colnames(paramdata) ← c("R0", "r", "sigma", "Sinit", "Iinit", "
Rinit")
```

The final kernel estimates for four of the key parameters are shown in Figure [3.2].

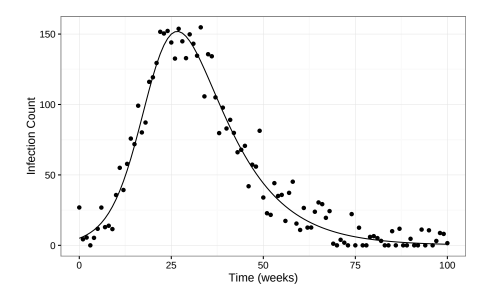


Figure 3.1: True SIR ODE solution infected counts, and with added observation noise. $\,$

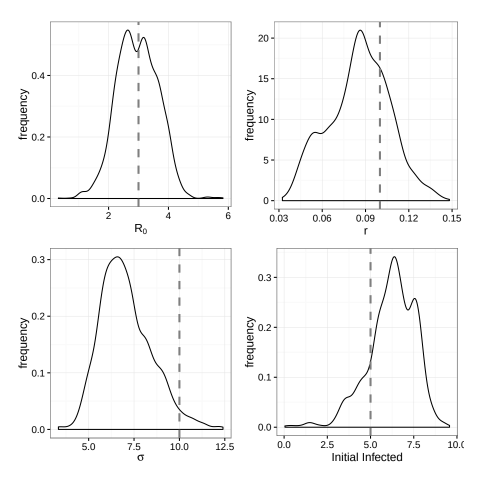


Figure 3.2: Kernel estimates for four essential system parameters. True values are indicated by solid vertical lines, sample means by dashed lines.

Chapter 4

Parameter Fitting

4.1 Fitting Setup

Now that we have established which methods we wish to evaluate the efficacy of for epidemic forecasting, it is prudent to see how they perform when fitting parameters for a known epidemic model. We have already seen how they perform when fitting parameters for a model with a deterministic evolution process and observation noise, but a more realistic model will have both process and observation noise.

To form such a model, we will take a deterministic SIR ODE model given by

$$\frac{dS}{dt} = -\beta SI$$

$$\frac{dI}{dt} = \beta SI - \gamma I$$

$$\frac{dR}{dt} = \gamma I,$$
(4.1)

and add process noise by allowing β to embark on a geometric random walk given by

$$\beta_{t+1} = \exp\left(\log(\beta_t) + \eta(\log(\bar{\beta}) - \log(\beta_t)) + \epsilon_t\right). \tag{4.2}$$

We will take ϵ_t to be normally distributed with standard deviation ρ^2 such that $\epsilon_t \sim \mathcal{N}(0, \rho^2)$. The geometric attraction term constrains the random walk, the force of which is $\eta \in [0, 1]$. If we take $\eta = 0$ then the walk will be unconstrained; if we let $\eta = 1$ then all values of β_t will be independent from the previous value (and consequently all other values in the sequence).

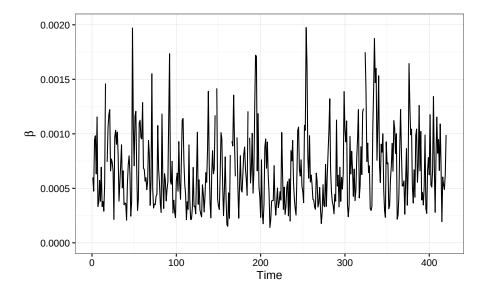


Figure 4.1: Simulated geometric autoregressive process show in Equation [4.2].

When $\eta \in (0,1)$, we have an autoregressive process of order 1 on the logarithmic scale of the form

$$X_{t+1} = c + \rho X_t + \epsilon_t, \tag{4.3}$$

where ϵ_t is normally distributed noise with mean 0 and standard deviation σ_E . This process has a theoretical expected mean of $\mu = c/(1-\rho)$ and variance $\sigma = \sigma_E^2/(1-\rho^2)$. If we choose $\eta = 0.5$, the resulting log-normal distribution has a mean of 6.80×10^{-4} and standard deviation of 4.46×10^{-4} .

Simulating the process in Equation [4.2] with $\eta=0.5$ gives us the plot in Figure [4.1] below.

We can obtain the corresponding density plot of the values in Figure [4.1], shown in Figure [4.2].

We see a density plot similar in shape to the desired density, and the geometric random walk displays dependence on previous values. Further the mean of this distribution was calculated to be 6.92×10^{-4} and standard the deviation to be 3.99×10^{-4} , which are very close to the theoretical values.

If we take the full stochastic SIR system and evolve it using an Euler stepping scheme with a step size of h = 1/7, for 1 step per day, we obtain the plot in Figure [4.3].

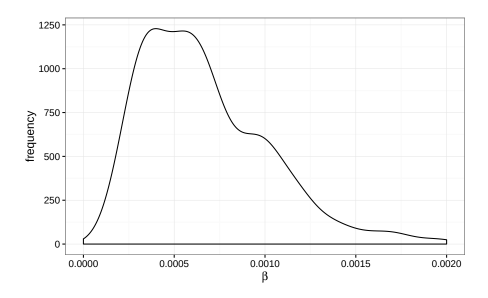


Figure 4.2: Density plot of values shown if Figure [4.1].

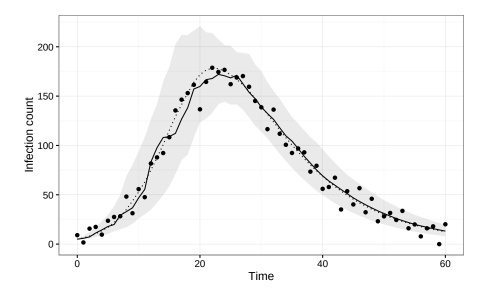


Figure 4.3: Stochastic SIR model simulated using an explicit Euler stepping scheme. The solid line is a single random trajectory, the dots show the data points obtained by adding in observation error defined as $\epsilon_{obvs} = \mathcal{N}(0, 10)$, and the grey ribbon is centre 95th quantile from 100 random trajectories.

4.2 Calibrating Samples

In order to compare HMCMC and IF2 we need to set up a fair and theoretically justified way to select the number of samples to draw for the HMCMC iterations and the number of particles to use for IF2. We assume that we are working with a problem that has an unknown real solution, so we use the Monte Carlo Standard Error (MCSE).

Suppose we are using a Monte-Carlo based method to obtain an estimate $\hat{\mu}_n$ for a quantity μ , where n is the number of samples. Then the Law of Large Numbers says that $\hat{\mu}_n \to \mu$ as $n \to \infty$. Further, the Central Limit Theorem says that the error $\hat{\mu}_n - \mu$ should shrink with number of samples such that $\sqrt{n}(\hat{\mu}_n - \mu) \to \mathcal{N}(0, \sigma^2)$ as $n \to \infty$, where σ^2 is the variance of the samples drawn.

We of course do not know μ , but the above allows us to obtain an estimate $\hat{\sigma}_n$ for σ given a number of samples n as

$$\hat{\sigma}_n = \sqrt{\frac{1}{n} \sum_{i=1}^n (X_i - \hat{\mu})},\tag{4.4}$$

which is known as the Monte Carlo Standard Error.

We can modify this formula to account for multiple variables by replacing the single variance measure sum by

$$\Theta^* V(\Theta^*)^T \tag{4.5}$$

where Θ^* is a row vector containing the reciprocals of the means of the parameters of interest, and V is the variance-covariance matrix with respect to the same parameters. This in effect scales the variances with respect to their magnitudes and accounts for covariation between parameters in one fell swoop. We also divide by the number of parameters, yielding

$$\hat{\sigma}_n = \sqrt{\frac{1}{n} \frac{1}{P} \Theta^* V(\Theta^*)^T} \tag{4.6}$$

where P is the number of particles.

The goal here is to then pick the number of HMCMC samples and IF2 particles to yield similar MCSE values. To do this we picked a combination of parameters for RStan that yielded decent results when applied to the stochastic SIR model specified above, calculated the resulting mean MCSE across several model fits, and isolated

TT3 (C) (C

the expected number of IF2 particles needed to obtain the same value. This was used as a starting value to "titrate" the IF2 iterations to the same point.

The resulting values were 1000 HMCMC warm-up iterations with 1000 samples drawn post-warm-up, and 2500 IF2 particles sent through 50 passes, each method giving an approximate MCSE of 0.0065.

4.3 IF2 Fitting

Now we will use an implementation of the IF2 algorithm to attempt to fit the stochastic SIR model to the previous data. The goal here is just parameter inference, but since IF2 works by applying a series on particle filters we essentially get the average system state estimates for a very small additional computational cost. Hence, we will will also look at that estimated behaviour in addition the the parameter estimates.

The code used here is a mix of R and C++ implemented using RCpp. The fitting was undertaken using 2500 particles with 50 IF2 passes and a cooling schedule given by a reduction in particle spread determined by 0.975^p, where p is the pass number starting with 0.

The MLE parameter estimates, taken to be the mean of the particle swarm values after the final pass, are shown in the table in Figure [4.4], along with the true values and the relative error.

TDO

		IF2		HMCMC	
Name	True	Fit	Error	Fit	Error
R_0	3.0	3.27	9.08×10^{-2}	3.12	1.05×10^{-1}
r	10^{-1}	1.04×10^{-1}	3.61×10^{-2}	9.99×10^{-2}	-7.56×10^{-4}
Initial Infected	5	7.90	5.80×10^{-1}	6.64	3.28×10^{-1}
σ	10	8.84	-1.15×10^{-1}	8.5	-1.50×10^{-1}
η	5×10^{-1}	5.87×10^{-1}	1.73×10^{-1}	4.57×10^{-1}	-8.27×10^{-2}
$arepsilon_{err}$	5×10^{-1}	1.63×10^{-1}	-6.73×10^{-1}	1.60×10^{-1}	-6.80×10^{-1}

Figure 4.4: Fitting errors.

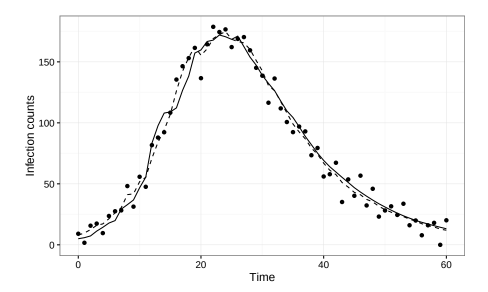


Figure 4.5: True system trajectory (solid line), observed data (dots), and IF2 estimated real state (dashed line).

From last IF2 particle filtering iteration, the mean state values from the particle swarm at each time step are shown with the true underlying state and data in the plot in Figure [4.5].

4.4 IF2 Convergence

Since IF2 is an iterative algorithm where each pass through he data is expected to push the parameter estimates towards the MLE, we can see the evolution of these estimates as a function of the pass number. Plots showing evolution of the mean estimates are shown if Figure [4.6] for the six most critical parameters.

Similarly, we can look at the evolution of the standard deviations of the parameter estimates from the particle swarm as a function of the pass number, shown in Figure [4.7] below.

As expected there is a downward trend in all plots, with a very strong trend in all but two of them.

4.5 IF2 Densities

Of diagnostic importance are the densities of the parameter estimates given by the final parameter swarm. These are shown below if Figure [4.8].

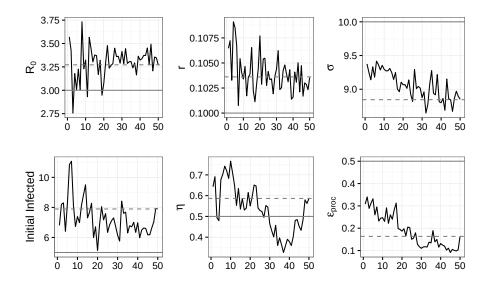


Figure 4.6: The horizontal axis shows the IF2 pass number. The solid black lines show the evolution of the ML estimates, the solid grey lines show the true value, and the dashed grey lines show the mean parameter estimates from the particle swarm after the final pass.

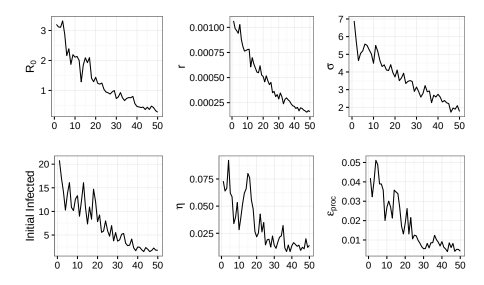


Figure 4.7: The horizontal axis shows the IF2 pass number and the solid black lines show the evolution of the standard deviations of the particle swarm values.

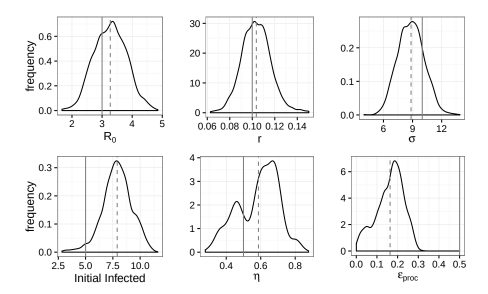


Figure 4.8: As before, the solid grey lines show the true parameter values and the dashed grey lines show the density means.

It is worth noting that the IF2 parameters chosen were in part chosen so as to not artificially narrow these densities; a more aggressive cooling schedule and/or an increased number of passes would have resulted in much narrower densities, and indeed have the potential to collapse them to point estimates.

4.6 HMCMC Fitting

We can use the Hamiltonian Monte Carlo algorithm implemented in the 'Rstan' package to fit the stochastic SIR model as above. This was done with a single HMC chain of 2000 iterations with 1000 of those being warm-up iterations.

The MLE parameter estimates, taken to be the means of the samples in the chain, were shown in the table in Figure [4.4] along with the true values and relative error.

4.7 HMCMC Densities

The parameter estimation densities from the Stan HMCMC fitting are shown in Figure [4.9] below.

the densities shown here represent a "true" MLE density estimate in that they represent HMC's attempt to directly sample from the parameter space according to

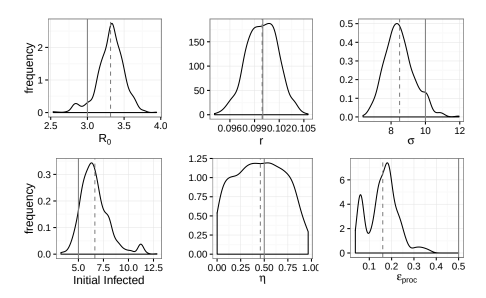


Figure 4.9: As before, the solid grey lines show the true parameter values and the dashed grey lines show the density means.

the likelihood surface, unlike IF2 which is in theory only trying to get a ML point estimate. Hence, these densities are potentially more robust than those produced by the IF2 implementation.

4.8 HMCMC and Bootstrapping

Unlike particle particle-filtering-based approaches, HMC does not produce state estimates as a by-product of parameter fitting, but we can use information about the stochastic nodes related to the noise in the β geometric random walk to reconstruct state estimates. The results of 100 bootstrap trajectories is shown in Figure [4.10].

4.9 Multi-trajectory Parameter Estimation

Here we fit the stochastic SIR model to 200 random independent trajectories using each method and examine the density of the point estimates produced.

The densities by and large display similar coverage, with the IF2 densities for r and ε_{proc} showing slightly wider coverage than the HMCMC densities for the same parameters.

The running times for each algorithm are summarized in Figure [7.4] below.

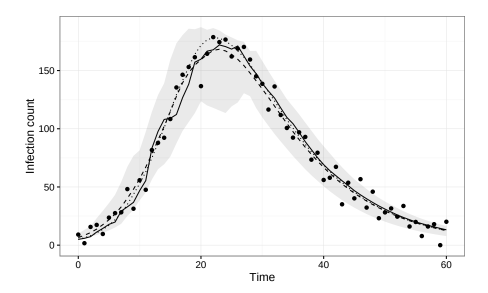


Figure 4.10: Result from 100 HMCMC bootstrap trajectories. The solid line shows the true states, the dots show the data, the dotted line shows the average system behaviour, the dashed line shows the bootstrap mean, and the grey ribbon shows the centre 95th quantile of the bootstrap trajectories.

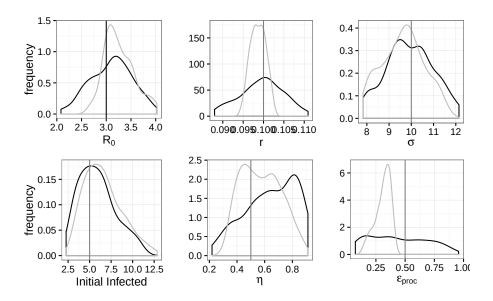


Figure 4.11: IF2 point estimate densities are shown in black and HMCMC point estimate densities are shown in grey. The vertical black lines show the true parameter values.

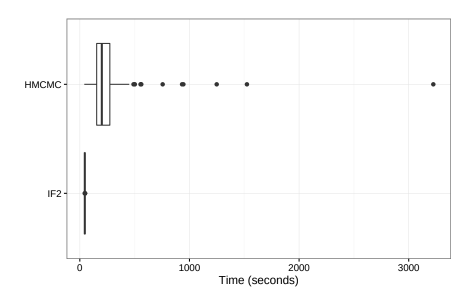


Figure 4.12: Fitting times for IF2 and HMCMC, in seconds. The centre box in each plot shows the centre 50th quantile, with the bold centre line showing the median.

The average running times were approximately 45.5 seconds and 257.4 seconds for IF2 and HMCMC respectively, representing a 5.7x speedup for IF2 over HMCMC. While IF2 may be able to fit the model to data faster than HMCMC, we are obtaining less information; this will become important in the next section. Further, the results in Figure [7.4] show that while the running time for IF2 is relatively fixed, the times for HMCMC are anything but, showing a wide spread of potential times.

Chapter 5

Forecasting Frameworks

5.1 Data Setup

This section will focus on taking the stochastic SIR model from the previous section, truncating the synthetic data output from realizations of that model, and seeing how well IF2 and HMCMC can reconstruct out-of-sample forecasts.

An example of a simulated system with truncated data can be seen in Figure [7.1].

In essence we want to be able to give either IF2 of HMCMC only the data points and have it reconstruct the entirety of the true system states.

5.2 IF2

For IF2, we will take advantage of the fact that the particle filter will produce state estimates for every datum in the time series given to it, as well as producing parameter maximum likelihood point estimates. Both of these sources of information will be used to produce forecasts by parametric bootstrapping using the final parameter estimates from the particle swarm after the last IF2 pass, then using the newly generated parameter sets along with the system state point estimates from the first fitting to simulate the systems forward into he future.

We will truncate the data at half the original time series length (to T = 30), and fit the model as previously described.

First, we can see the state estimates for each time point produced by the last IF2 pass in Figure [5.2].

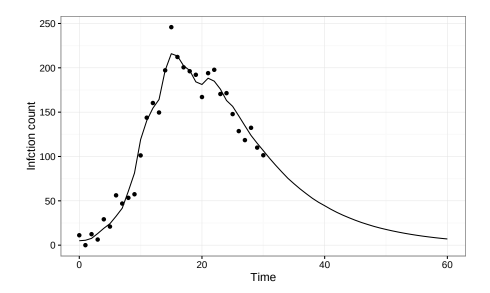


Figure 5.1: Infection count data truncated at T=30. The solid line shows the true underlying system states, and the dots show those states with added observation noise. Parameters used were $R_0 = 3.0$, r = 0.1, $\eta = .05$, $\sigma_{proc} = 0.5$, and additive observation noise was drawn from $\mathcal{N}(0, 10)$.

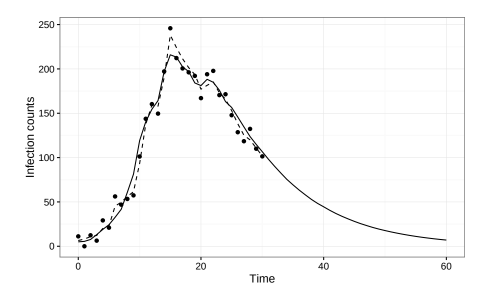


Figure 5.2: Infection count data truncated at T=30 from Figure [7.1] above. The dashed line shows IF2's attempt to reconstruct the true underlying state from the observed data points.

Recall that IF2 is not trying to generate parameter estimation densities, but rather produce a point estimate. Since we wish to determine the approximate distribution of each of the parameters in addition to the point estimate, we must turn to another method, parametric bootstrapping.

5.2.1 Parametric Bootstrapping

The goal of the parametric bootstrap is use an initial density sample θ^* to generate further samples $\theta_1, \theta_2, ..., \theta_M$. It works by using θ to generate artificial data sets $D_1, D_2, ..., D_M$ to which we can refit our model of interest and generate new parameter sets.

[I'm still trying to dig up a good paper that talks about applicability to dynamical systems, there will be a paragraph here about it.]

An algorithm for parametric bootstrapping using IF2 and our stochastic SIR model is shown in Algorithm [5].

5.2.2 IF2 Forecasts

Using the parameter sets $\theta_1, \theta_2, ..., \theta_M$ and the point estimate of the state provided by the initial IF2 fit, we can use a normal bootstrap to produce estimates of the future state. A plot showing a projection of the data from the previous plots can be seen in Figure [5.3].

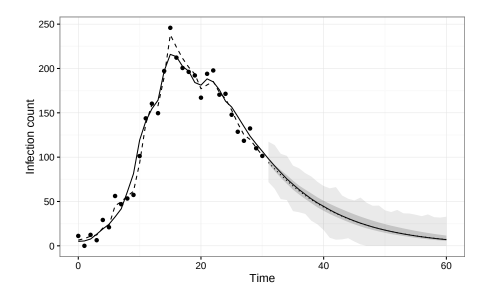


Figure 5.3: Forecast produced by the IF2 / parametric bootstrapping framework. The dotted line shows the mean estimate of the forecasts, the dark grey ribbon shows the centre 95th quantile of the true state estimates, and the lighter grey ribbon shows the centre 95th quantile of the true state estimates with added observation noise drawn from $\mathcal{N}(0, \sigma)$.

We can define a metric to gauge forecast effectiveness by calculating the SSE and dividing that value by the number of values predicted to get the average squared error per point. For the data in Figure [5.3] the value was $\overline{SSE} = 1.67$.

5.3 HMCMC

For HMCMC we can use a simpler bootstrapping approach. We do not get state estimates directly from the RStan fitting due to the way we implemented the model, but we can construct them using the process noise latent variables. Once we've done this we can forward simulate the system from the state estimate into the future.

As before we fit the stochastic SIR model to the partial data, but now perform bootstrapping as described above, and obtain the plot in Figure [5.4].

And as before we can evaluate the averaged SSE of the forecast for the data shown, giving $\overline{SSE} = 20.27$.

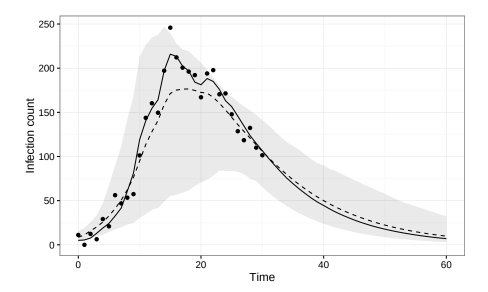


Figure 5.4: Forecast produced by the HMCMC / bootstrapping framework with M=200 trajectories. The dotted line shows the mean estimate of the forecasts, and the grey ribbon shows the centre 95th quantile.

5.4 Truncation vs. Error

Of course the above mini-comparison only shows one truncation value for one trajectory. Really, we need to know how each method performs on average given different trajectories and truncation amounts. In effect we wish to "starve" each method of data and see how poor the estimates become with each successive data point loss.

Using each method, we can fit the stochastic SIR model to successively smaller time series to see the effect of truncation on forecast averaged SSE. This was performed with 10 new trajectories drawn for each of the desired lengths. The results are shown in Figure [5.5].

IF2 and HMCMC perform very closely, with IF2 maintaining a small advantage up to a truncation of about 25-30 data points.

Since the parametric bootstrapping approach used by IF2 requires a significant number of additional fits, its computational cost is significantly higher than the simpler bootstrapping approach used by the HMCMC framework, about 35.5x as expensive. However the now much longer running time can somewhat alleviated by parallelizing the parametric bootstrapping process; as each of the parametric bootstrap fittings in entirely independent, this can be done without a great deal of additional effort. The code used here has this capability, but it was not utilised in

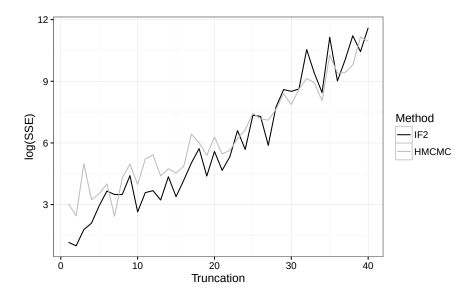


Figure 5.5: Error growth as a function of data truncation amount. Both methods used 200 bootstrap trajectories. Note that the y-axis shows the natural log of the averaged SSE, not the total SSE.

the comparison so as to accurately represent total computational cost, not potential running time.

Chapter 6

S-map and SIRS

6.1 S-maps

A family of forecasting methods that shy away from the mechanistic model-based approaches outlined in the previous sections have been developed by Sugihara (references) over the last several decades. As these methods do not include a mechanistic model in their forecasting process, they also do not attempt to perform parameter inference. Instead they attempt to reconstruct the underlying dynamical process as a weighted linear model from a time series.

One such method, the sequential locally weighted global linear maps (S-map), builds a global linear map model and uses it to produce forecasts directly. Despite relying on a linear mapping, the S-map does not assume the time series on which it is operating is the product of linear system dynamics, and in fact was developed to accommodate non-linear dynamics.

The S-map works by first constructing a time series embedding of length E, known as the library and denoted $\{\mathbf{x_i}\}$. Consider a time series of length T denoted $x_1, x_2, ..., x_T$. Each element in the time series with indices in the range E, E+1, ..., T will have a corresponding entry in the library such that a given element x_t will correspond to a library vector of the form $\mathbf{x_i} = (x_t, x_{t-1}, ..., x_{t-E+1})$. Next, given a forecast length E (representing E time steps into the future), each library vector $\mathbf{x_i}$ is assigned a prediction from the time series $y_i = x_{t+L}$, where x_t is the first entry in $\mathbf{x_i}$. Finally, a forecast \hat{y}_t for specified predictor vector $\mathbf{x_t}$ (usually from the library itself), is generated using an exponentially weighted function of the library $\{\mathbf{x_i}\}$, predictions $\{y_i\}$, and predictor vector $\mathbf{x_t}$.

This function is defined as follows:

First construct a matrix A and vector b defined as

$$A(i,j) = w(||\mathbf{x_i} - \mathbf{x_t}||)\mathbf{x_i}(j)$$

$$b(i) = w(||\mathbf{x_i} - \mathbf{x_t}||)y_i$$
(6.1)

where i ranges over 1 to the length of the library, and j ranges over [0, E]. It should be noted that in the above equations and the ones that follow, $x_t(0) = 1$ to account for the linear term in the map.

The weighting function w is defined as

$$w(d) = \exp\left(\frac{-\theta d}{\bar{d}}\right),\tag{6.2}$$

where d is the euclidean distance between the predictor vector and library vectors in Equation (6.1) and \bar{d} is the average of these distances. We can then see that θ serves as a way to specify the appropriate level of penalization applied to poorly-matching library vectors – if θ is 0 all weights are the same (no penalization), and increasing θ increases the level of penalization.

Now we solve the system Ac = b to obtain the linear weightings used in to generate the forecast according to

$$\hat{y}_t = \sum_{j=0}^E c_t(j) \mathbf{x_t}(j). \tag{6.3}$$

In this way we have produced a forecast value for a single time. This process can be repeated for a sequence of times T+1, T+2, ... to project a time series into the future.

6.2 S-map Algorithm

The above description can be summarized in Algorithm [6].

Algorithm 6: S-map /* Select a starting point */ **Input**: Time series $x_1, x_2, ..., x_T$, embedding dimension E, distance penalization θ , forecast length L, predictor vector $\mathbf{x_t}$ /* Construct library $\{x_i\}$ */ 1 for i = E : T do $\mathbf{z} \mid \mathbf{x_i} = (x_i, x_{i-1}, ..., x_{i-E-1})$ /* Construct mapping from library vectors to predictions */ 3 for $i = 1 : (T_E + 1)$ do for j = 1 : E do 6 for $i = 1 : (T_E + 1)$ do 7 $b(i) = w(||\mathbf{x_i} - \mathbf{x_t}||)y_i$ /* Use SVD to solve the mapping system, Ac = b */ s SVD(Ac = b)/* Compute forecast */ 9 $\hat{y_t} = \sum_{j=0}^{E} c_t(j) \mathbf{x_t}(j)$ /* Forecasted value in time series Output: Forecast $\hat{y_t}$

6.3 SIRS Model

In an epidemic or infectious disease context, the S-map algorithm will only really work on time series that appear cyclic. While there is nothing mechanically that prevents it from operating on a time series that do not appear cyclic, S-mapping requires a long time series in order to build a quality library. Without one the forecasting process would produce unreliable data.

With that in mind, the only fair way to compare the efficacy of s-mapping to IF2 or Hamiltonian MCMC is to generate data from a SIRS model with a seasonal component, and have all methods operate on the resulting time series.

The basic skeleton of the SIRS model is similar to the stochastic SIR model described previously. The deterministic ODE component of the model is as follows.

$$\frac{dS}{dt} = -\Gamma(t)\beta SI + \eta R$$

$$\frac{dI}{dt} = \Gamma(t)\beta SI - \gamma I$$

$$\frac{dR}{dt} = \gamma I - \eta R,$$
(6.4)

There are two new features here. We have a re-susceptibility rate η through which people become able to be reinfected, and a seasonality factor Γ defined as

$$\Gamma(t) = \exp\left(2\cos\left(\frac{2\pi}{365}t\right) - 2\right). \tag{6.5}$$

This function oscillates between 1 and e^{-4} (close to 0) and is meant to represent transmission damping during the off-season, for example summer for influenza. Further, it displays flatter troughs and sharper peaks to exaggerate its effect in peak season.

As before, β is allowed to walk restricted by a geometric mean, described by

$$\beta_{t+1} = \exp\left(\log(\beta_t) + \eta(\log(\bar{\beta}) - \log(\beta_t)) + \epsilon_t\right). \tag{6.6}$$

When simulated for the equivalent of 5 years (260 weeks), and adding noise drawn from $\mathcal{N}(0,\sigma)$ we obtain Figure [7.1].

We can see how the S-map can reconstruct the next cycle in the time series in Figure [6.2].

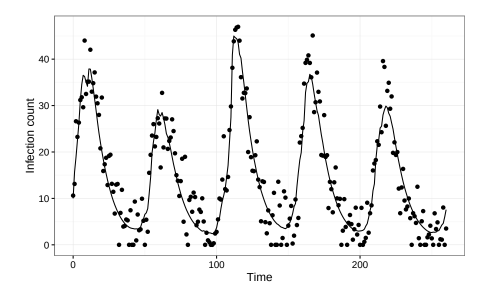


Figure 6.1: Five cycles generated by the SIRS function. The solid line the true number of cases, dots show case counts with added observation noise. The Parameter values were R0 = 3.0, $\gamma = 0.1$, $\eta = 1$, $\sigma = 5$, and 10 initial cases.

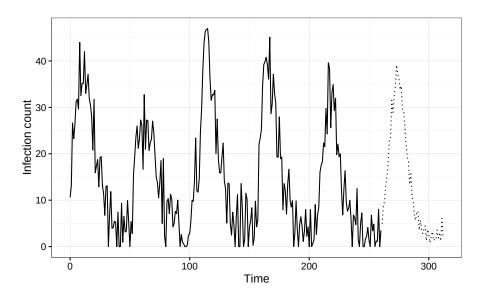


Figure 6.2: S-map applied to the data from the previous figure. The solid line shows the infection counts with observation noise form the previous plot, and the dotted line is the S-map forecast. Parameters chosen were E=14 and $\theta=3$.

The parameters used in the S-map algorithm to obtain the forecast used in Figure [6.2] were obtained using a grid search of potential parameters outlined in (Sugihara

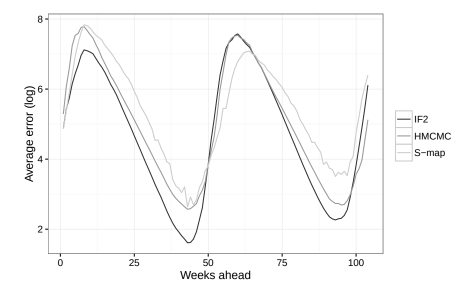


Figure 6.3: Error as a function of forecast length.

ref). The script is included in the appendices.

6.4 SIRS Model Forecasting

Naturally we wish to compare the efficacy of this comparatively simple technique against the more complex and more computationally taxing frameworks we have established to perform forecasting using IF2 and HMCMC.

To do this we generated a series of artificial time series of length 260 meant to represent 5 years of weekly incidence counts and used each method to forecast up to 2 years into the future. Our goal here was to determine how forecast error changed with forecast length.

The results of the simulation are shown in Figure [7.3].

Interestingly, all methods produce roughly the same result, which is to say the spike in each outbreak cycle are difficult to accurately predict. IF2 produces better results than either HMCMC and the S-map for the majority of forecast lengths, with the S-map producing the poorest results with the exception of the second rise in infection rates in which it outperforms the other methods.

While the S-map may not provide the same fidelity or forecast as IF2 or HMCMC, it shines when it comes to running time. Figure [7.4] shows the running times over 20 simulations.

It is clear from the above figure that the S-map running times are minute compared

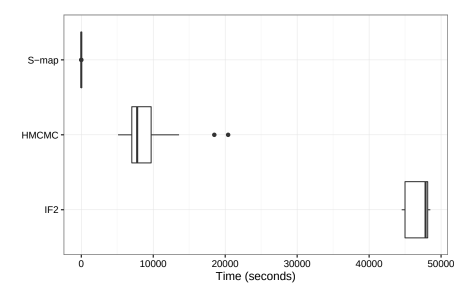


Figure 6.4: Runtimes for producing SIRS forecasts. The box shows the middle 50th quantile, the bold line is the median, and the dots are outliers.

to the other methods, but to emphasize the degree: The average running time for the S-map is about 1.49×10^{-1} seconds, for IF2 it is about 4.70×10^4 , and for HMCMC it is about 9.20×10^3 . This is a speed-up of over 316,000x compared to IF2 and over 61,800x compared to HMCMC.

Chapter 7

Spatial Epidemics

7.1 Spatial SIR

Spatial epidemic models provide a way to capture not just the temporal trend in an epidemic, but to also integrate spatial data and infer how the infection is spreading in both space and time. One such model we can use is a dynamic spatiotemporal SIR model.

We wish to construct a model build upon the stochastic SIR compartment model described previously but one that consists of several connected spatial locations, each with its own set of compartments. Consider a set of locations numbered i = 1, ..., N, where N is the number of locations. Further, let N_i be the number of neighbours location i has. The model is then

$$\frac{dS_i}{dt} = -\left(1 - \phi \frac{N_i}{N_i + 1}\right) \beta_i S_i I_i - \left(\frac{\phi}{N_i + 1}\right) S_i \sum_{j=0}^{N_i} \beta_j I_j$$

$$\frac{dI_i}{dt} = \left(1 - \phi \frac{N_i}{N_i + 1}\right) \beta_i S_i I_i + \left(\frac{\phi}{N_i + 1}\right) S_i \sum_{j=0}^{N_i} \beta_j I_j - \gamma I$$

$$\frac{dR_i}{dt} = \gamma I,$$
(7.1)

Neighbours for a particular location are numbered $j = 1, ..., N_i$. We have a new parameter, $\phi \in [0, 1]$, which is the degree of connectivity. If we let $\phi = 0$ we have total spatial isolation, and the dynamics reduce to the basic SIR model. If we let $\phi = 1$ then each of the neighbouring locations will have weight equivalent to the parent location.

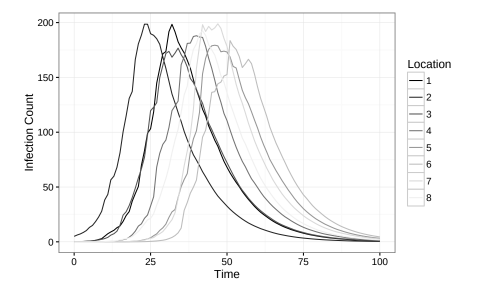


Figure 7.1: Evolution of a spatial epidemic in a ring topology. The outbreak was started with 5 cases in Location 2. Parameters were $R_0 = 3.0$, $\gamma = 0.1$, $\eta = 0.5$, $\sigma_{err} = 0.5$, and $\phi = 0.5$.

As before we let β embark on a geometric random walk defined as

$$\beta_{i,t+1} = \exp\left(\log(\beta_{i,t}) + \eta(\log(\bar{\beta}) - \log(\beta_{i,t})) + \epsilon_t\right). \tag{7.2}$$

Note that as β is a state variable, each location has its own stochastic process driving the evolution of its β state.

If we imagine a circular topology in which each of 10 locations is connected to exactly two neighbours (i.e. location 1 is connected to locations N and 2, location 2 is connected to locations 1 and 3, etc.), and we start each location with completely susceptible populations except for a handful of infected individuals in one of the locations, we obtain a plot of the outbreak progression in Figure [7.1].

If we add noise to the data from Figure [7.1], we obtain Figure [7.2], below.

7.2 Dewdrop Regression

Dewdrop regression (references) aims to overcome the primary disadvantage suffered by methods such as the S-map or its cousin Simplex Projection: the requirement of long time series from which to build a library. Suggested by Sugihara's group in 2008, Dewdrop Regression works by stitching together shorter, related, time series, in order to give the S-map or similar methods enough data to operate on. The

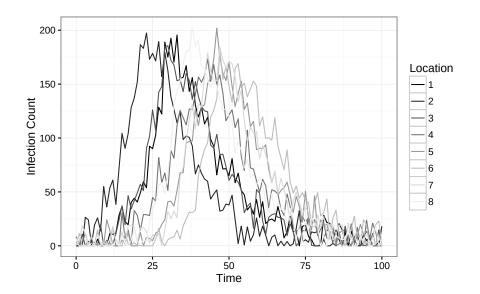


Figure 7.2: Evolution of a spatial epidemic as in Figure [7.1], with added observation noise drawn from $\mathcal{N}(0, 10)$.

underlying idea is that as long as the underlying dynamics of the time series display similar behaviour (such as potentially collapsing to the same attractor), they can be treated as part of the same overarching system.

It is not enough to simply concatenate the shorter time series together – several procedures must be carried out and a few caveats observed. First, as the individual time series can be or drastically differing scales and breadths, they all must be rescaled to unit mean and variance. Then the library is constructed as before with an embedding dimension E, but any library vectors that span any of the seams joining the time series are discarded. Further, and predictions stemming from a library vector must stay within the time series from which they originated. In this way we are allowing the "shadow" of of the underlying dynamics of the separate time series to infer the forecasts for segments of other time series. Once the library has been constructed, S-mapping can be carried out as previously specified.

This procedure is especially well-suited to a the spatial model we are using. While the dynamics are stochastic, they still display very similar means and variances. This means the rescaling process in Dewdrop Regression is not necessary and can be skipped. Further, the overall variation between the epidemic curves in each location is on the smaller side, meaning the S-map will have a high-quality library from which to build forecasts.

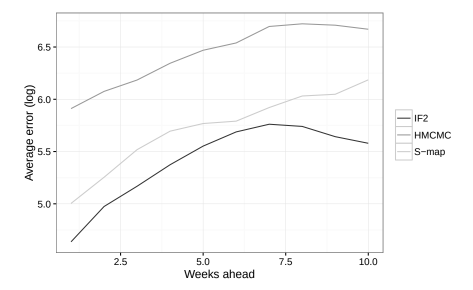


Figure 7.3: Average SSE (log scale) across each location and all trials as a function of the number of weeks ahead in the forecast.

7.3 Spatial Model Forecasting

In order to compare the forecasting efficacy of Dewdrop Regression with S-mapping against IF2 and HMCMC, we generated 20 independent spatial data sets up to time T=50 weeks in each of L=10 locations and forecasted 10 weeks into the future. Forecasts were compared to that of the true model evolution, and the average SSE for each week ahead in the forecast were computed. The number of bootstrapping trajectories used by IF2 and HMCMC was reduced from 200 to 50 to curtail running times.

The results are shown in Figure [7.3].

The results show a clear delineation in forecast fidelity between methods. IF2 maintains an advantage regardless of how long the forecast produced. Interestingly, Dewdrop Regression with S-mapping performs almost as well as IF2, and outperforms HMCMC. HMCMC lags behind both methods by a healthy margin.

If we examine the runtimes for each forecast framework, we obtain the data in Figure [7.4].

As before, the S-map with Dewdrop Regression runs faster than the other two methods with a huge margin. It is again hard to see exactly how large the margin is from the figure due to the scale, but we can examine the average values: the average running time for S-mapping with Dewdrop Regression was about 249 seconds, whereas the average times for IF2 and HMCMC were about 2.90×10^4 and

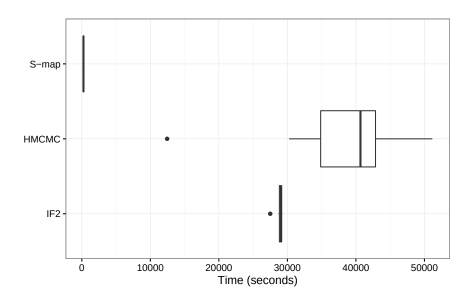


Figure 7.4: Runtimes for producing spatial SIR forecasts. The box shows the middle 50th quantile, the bold line is the median, and the dots are outliers.

 $3.88\times10^4,$ respectively. This is a speed-up of just over 116x over IF2 and 156x over HMCMC.

Considering how well S-mapping performed with regards to forecast error, it shows a significant advantage over HMCMC in particular – it outperforms it in both forecast error and running times.

Appendix A

Hamiltonian MCMC

A.1 Full R code

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

```
1 ## 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)), {
                 ← R0*r/N
             \texttt{BSI} \leftarrow \texttt{B} {\star} \texttt{S} {\star} \texttt{I}
             rI \ \leftarrow r*I
             dS = -BSI
             dI = BSI - rI
             dR = rI
             return(list(c(dS, dI, dR)))
        })
28 }
```

```
30 pars \leftarrow c(R0 \leftarrow 3.0, # average number of new infected individuals
       per infectious person
               r \leftarrow 0.1, # recovery rate
                   \leftarrow 500)
                             # population size
34 \mid T \leftarrow 100
35 | y_i = 5, R = 0
36 times \leftarrow seq(0, T, by = 1)
38 | \text{odeout} \leftarrow \text{ode}(y_{\text{ini}}, \text{times}, \text{SIR}, \text{pars})
40 set.seed(1001)
41 | sigma \leftarrow 10
42 infec_counts_raw \leftarrow odeout[,3] + rnorm(T+1, 0, sigma)
43 infec_counts \leftarrow ifelse(infec_counts_raw < 0, 0, infec_counts_raw)
45 \mid g \leftarrow qplot(0:T, odeout[,3], geom = "line", xlab = "Time (weeks)",
      ylab = "Infection Count") +
       geom_point(aes(y = infec_counts)) +
       theme_bw()
48
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 data[,1] ← infec_counts
57 standata \leftarrow as.vector(t(data))[1:datlen]
59 sir_data ← 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())
                            chains = 4,  # number of chains
66 stan_options \leftarrow list(
                              iter = 2000, # iterations per chain
                              warmup = 1000, # warmup interations
                                    = 2) # thinning number
                              thin
70 fit \leftarrow stan(file
                        = "d_sirode_euler.stan",
                data
                        = sir_data,
                chains = stan_options$chains,
                iter
                         = stan_options$iter,
                warmup = stan_options$warmup,
                thin
                         = stan_options$thin )
```

```
77 exfit ← extract(fit, permuted = TRUE, inc_warmup = FALSE)
79 R0points \leftarrow exfit$R0
80 R0kernel \leftarrow qplot(R0points, geom = "density", xlab = expression(R[0])
       , ylab = "frequency") +
            geom_vline(aes(xintercept=R0), linetype="dashed", size=1,
                color="grey50") +
            theme_bw()
84 print(R0kernel)
85 ggsave(R0kernel, filename="kernelR0.pdf", height=3, width=3.25)
87
   rpoints \leftarrow exfit$r
88 rkernel \leftarrow qplot(rpoints, geom = "density", xlab = "r", ylab = "
       frequency") +
            geom_vline(aes(xintercept=r), linetype="dashed", size=1,
                color="grey50") +
            theme_bw()
92 print(rkernel)
93 ggsave(rkernel, filename="kernelr.pdf", height=3, width=3.25)
95 sigmapoints \leftarrow exfit$sigma
96 sigmakernel \leftarrow qplot(sigmapoints, geom = "density", xlab = expression
       (sigma), ylab = "frequency") +
            geom_vline(aes(xintercept=sigma), linetype="dashed", size=1,
                 color="grey50") +
            theme_bw()
100 print(sigmakernel)
101 ggsave(sigmakernel, filename="kernelsigma.pdf", height=3, width
       =3.25)
|103| infecpoints \leftarrow exfit$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") +
            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| \operatorname{sso} \leftarrow \operatorname{as.shinystan}(\operatorname{fit})
|140| sso \leftarrow launch_shinystan(sso)
```

A.2 Full Stan code

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

```
## Dexter Barrows
2 ## McMaster University
3 ## 2016
  data {
      int
               <lower=1>
                                   // total integration steps
                           Τ;
                                    // observed number of cases
      real
                           y[T];
      int
               <lower=1>
                           N;
                                    // population size
      real
                                    // step size
                           h;
12 }
13
```

```
14 parameters {
       real <lower=0, upper=10>
                                     R0;
                                            // 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
21 }
23 model {
       real S[T];
       real I[T];
       real R[T];
       S[1] \leftarrow y0[1];
       I[1] \leftarrow y0[2];
       R[1] \leftarrow 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 );
           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);
48
       R0
               \sim lognormal(1,1);
               ~ lognormal(1,1);
       r
              ~ lognormal(1,1);
       sigma
54 }
```

Appendix B

Iterated Filtering

B.1 Full R code

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

```
Author: Dexter Barrows
 2 ## Github: dbarrows.github.io
 4 library (deSolve)
 5 library(ggplot2)
6 library(reshape2)
 7 library(gridExtra)
   library(Rcpp)
10 \mid SIR \leftarrow function(Time, State, Pars) {
        with(as.list(c(State, Pars)), {
                 ← R0*r/N
             \texttt{BSI} \leftarrow \texttt{B} {\star} \texttt{S} {\star} \texttt{I}
             rI \leftarrow r*I
             dS = -BSI
             dI = BSI - rI
             dR = rI
21
             return(list(c(dS, dI, dR)))
        })
26 }
28 T
          \leftarrow 100
```

```
29 N
        ← 500
30 \mid \text{sigma} \leftarrow 10
31 | i_i | fec \leftarrow 5
33 ## Generate true trajecory and synthetic data
34 ##
36 true_init_cond \leftarrow c(S = N - i_infec,
                         I = i_infec,
                         R = 0)
40 true_pars \leftarrow c(R0 = 3.0,
                   r = 0.1,
                   N = 500.0
43
44 odeout ← ode(true_init_cond, 0:T, SIR, true_pars)
45 trueTraj \leftarrow odeout[,3]
47 set.seed(1001)
49 infec_counts_raw ← odeout[,3] + rnorm(T+1, 0, sigma)
50 infec_counts \leftarrow ifelse(infec_counts_raw < 0, 0, infec_counts_raw)
52 \mid g \leftarrow qplot(0:T, odeout[,3], geom = "line", xlab = "Time (weeks)",
      ylab = "Infection Count") +
       geom_point(aes(y = infec_counts)) +
       theme_bw()
56 print(g)
57 ggsave(g, filename="dataplot.pdf", height=4, width=6.5)
59 ## Rcpp stuff
60 ##
62 sourceCpp(paste(getwd(), "d_if2.cpp", sep="/"))
64 paramdata \leftarrow data.frame(if2(infec_counts, T+1, N))
65 colnames(paramdata) \leftarrow c("R0", "r", "sigma", "Sinit", "Iinit", "Rinit"
67 ## Parameter density kernels
68 ##
70 R0points \leftarrow paramdata$R0
71 R0kernel \leftarrow qplot(R0points, geom = "density", xlab = expression(R[0])
      , ylab = "frequency") +
72
            geom_vline(aes(xintercept=true_pars[["R0"]]), linetype="
               dashed", size=1, color="grey50") +
            theme_bw()
```

```
75 print(R0kernel)
76 ggsave(R0kernel, filename="kernelR0.pdf", height=3, width=3.25)
78 rpoints \leftarrow paramdata$r
79 rkernel \leftarrow qplot(rpoints, geom = "density", xlab = "r", ylab = "
       frequency") +
            geom_vline(aes(xintercept=true_pars[["r"]]), linetype="
               dashed", size=1, color="grey50") +
            theme_bw()
82
83 print(rkernel)
84 ggsave(rkernel, filename="kernelr.pdf", height=3, width=3.25)
86 sigmapoints \leftarrow paramdata\$sigma
87 sigmakernel \leftarrow qplot(sigmapoints, geom = "density", xlab = expression
       (sigma), ylab = "frequency") +
            geom_vline(aes(xintercept=sigma), linetype="dashed", size=1,
                color="grey50") +
89
            theme_bw()
91 print(sigmakernel)
92 ggsave(sigmakernel, filename="kernelsigma.pdf", height=3, width
       =3.25)
94 infecpoints ← paramdata$Iinit
95 infeckernel \leftarrow qplot(infecpoints, geom = "density", xlab = "Initial
       Infected", ylab = "frequency") +
            geom_vline(aes(xintercept=true_init_cond[['I']]), linetype="
               dashed", size=1, color="grey50") +
            theme_bw()
99 print(infeckernel)
100 ggsave(infeckernel, filename="kernelinfec.pdf", height=3, width
       =3.25)
102 # show grid
103 grid.arrange(R0kernel, rkernel, sigmakernel, infeckernel, ncol = 2,
      nrow = 2)
105 \text{ pdf}("if2kernels.pdf", height = 6.5, width = 6.5)
106|grid.arrange(R0kernel, rkernel, sigmakernel, infeckernel, ncol = 2,
       nrow = 2)
107 dev.off()
108 #ggsave(filename="if2kernels.pdf", g2, height=6.5, width=6.5)
```

B.2 Full C++ code

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

```
Author: Dexter Barrows
      Github: dbarrows.github.io
      */
6 #include <stdio.h>
7 #include <math.h>
8 #include <sys/time.h>
9 #include <time.h>
10 #include <stdlib.h>
11 #include <string>
12 #include <cmath>
13 #include <cstdlib>
14 #include <fstream>
16 //#include "rand.h"
17 //#include "timer.h"
18
19 #define Treal
                   100
                               // time to simulate over
20 #define R0true 3.0
                               // infectiousness
21 #define rtrue
                   0.1
                               // recovery rate
22 #define Nreal
                   500.0
                               // population size
23 #define merr
                   10.0
                               // expected measurement error
                               // Initial infected individuals
24 #define I0
                   5.0
26 #include <Rcpp.h>
  using namespace Rcpp;
30 struct Particle {
      double R0;
      double r;
      double sigma;
      double S;
      double I;
      double R;
      double Sinit;
      double Iinit;
      double Rinit;
40 };
42 struct ParticleInfo {
                           double R0sd;
      double R0mean;
                           double rsd;
      double rmean;
      double sigmamean;
                           double sigmasd;
      double Sinitmean;
                           double Sinitsd;
```

```
double Iinitmean;
                          double Iinitsd;
      double Rinitmean;
                          double Rinitsd;
49 };
52 int timeval_subtract (double *result, struct timeval *x, struct
     timeval *y);
53 int check_double(double x, double y);
54 void exp_euler_SIR(double h, double t0, double tn, int N, Particle *
      particle);
55 void copyParticle(Particle * dst, Particle * src);
56 void perturbParticles(Particle * particles, int N, int NP, int
      passnum, double coolrate);
57 bool isCollapsed(Particle * particles, int NP);
58 void particleDiagnostics(ParticleInfo * partInfo, Particle *
      particles, int NP);
59 NumericMatrix if2(NumericVector * data, int T, int N);
60 double randu();
61 double randn();
63 // [[Rcpp::export]]
64 NumericMatrix if2(NumericVector data, int T, int N) {
      int
              NP
                          = 2500;
      int
              nPasses
                          = 50;
      double coolrate
                          = 0.975;
      int
              i\_infec
                           = I0;
      NumericMatrix paramdata(NP, 6);
      srand(time(NULL)); // Seed PRNG with system time
      double w[NP];
                          // particle weights
      Particle particles[NP];  // particle estimates for current
      Particle particles_old[NP]; // intermediate particle states for
          resampling
      printf("Initializing particle states\n");
81
82
      // initialize particle parameter states (seeding)
84
      for (int n = 0; n < NP; n++) {
86
          double R0can, rcan, sigmacan, Iinitcan;
87
          do {
              R0can = R0true + R0true*randn();
          } while (R0can < 0);</pre>
```

```
particles[n].R0 = R0can;
          do {
               rcan = rtrue + rtrue*randn();
           } while (rcan < 0);
96
          particles[n].r = rcan;
          do {
               sigmacan = merr + merr*randn();
          } while (sigmacan < 0);</pre>
          particles[n].sigma = sigmacan;
          do {
               Iinitcan = i_infec + i_infec*randn();
          } while (Iinitcan < 0 || N < Iinitcan);</pre>
          particles[n].Sinit = N - Iinitcan;
          particles[n].Iinit = Iinitcan;
          particles[n].Rinit = 0.0;
      }
      // START PASSES THROUGH DATA
      printf("Starting filter\n");
      printf("----\n");
      printf("Pass\n");
      for (int pass = 0; pass < nPasses; pass++) {</pre>
          printf("...%d / %d\n", pass, nPasses);
          perturbParticles(particles, N, NP, pass, coolrate);
          // initialize particle system states
          for (int n = 0; n < NP; n++) {
               particles[n].S = particles[n].Sinit;
               particles[n].I = particles[n].Iinit;
               particles[n].R = particles[n].Rinit;
          }
          // between-pass perturbations
          for (int t = 1; t < T; t++) {
               // between-iteration perturbations
               perturbParticles(particles, N, NP, pass, coolrate);
```

```
// generate individual predictions and weight
               for (int n = 0; n < NP; n++) {
                   exp_euler_SIR(1.0/10.0, 0.0, 1.0, N, &particles[n]);
                   double merr_par = particles[n].sigma;
                   double y_diff = data[t] - particles[n].I;
                  w[n] = 1.0/(merr_par*sqrt(2.0*M_PI)) * exp( - y_diff
                      *y_diff / (2.0*merr_par*merr_par) );
               }
               // cumulative sum
               for (int n = 1; n < NP; n++) {
                  w[n] += w[n-1];
               }
              // save particle states to resample from
               for (int n = 0; n < NP; n++){
                   copyParticle(&particles_old[n], &particles[n]);
               }
               // resampling
               for (int n = 0; n < NP; n++) {
                   double w_r = randu() * w[NP-1];
                   int i = 0;
                   while (w_r > w[i]) {
                      i++;
                   }
                  // i is now the index to copy state from
                  copyParticle(&particles[n], &particles_old[i]);
               }
           }
       }
181
       ParticleInfo pInfo;
       particleDiagnostics(&pInfo, particles, NP);
       printf("Parameter results (mean | sd)\n");
       printf("----\n");
185
                       %f %f\n", pInfo.R0mean, pInfo.R0sd);
       printf("R0
       printf("r
187
                       %f %f\n", pInfo.rmean, pInfo.rsd);
       printf("sigma %f %f\n", pInfo.sigmamean, pInfo.sigmasd);
       printf("S_init %f %f\n", pInfo.Sinitmean, pInfo.Sinitsd);
```

```
190
       printf("I_init
                         %f %f\n", pInfo.Iinitmean, pInfo.Iinitsd);
       printf("R_init
                         %f %f\n", pInfo.Rinitmean, pInfo.Rinitsd);
       printf("\n");
       // Get particle results to pass back to R
       for (int n = 0; n < NP; n++) {
           paramdata(n, 0) = particles[n].R0;
           paramdata(n, 1) = particles[n].r;
           paramdata(n, 2) = particles[n].sigma;
           paramdata(n, 3) = particles[n].Sinit;
           paramdata(n, 4) = particles[n].Iinit;
           paramdata(n, 5) = particles[n].Rinit;
       }
       return paramdata;
212 }
215 /*
       Use the Explicit Euler integration scheme to integrate SIR model
       forward in time
       double h - time step size
       double t0 - start time
       double tn - stop time
218
       double * y - current system state; a three-component vector
          representing [S I R], susceptible-infected-recovered
222 void exp_euler_SIR(double h, double t0, double tn, int N, Particle *
       particle) {
       int num_steps = floor( (tn-t0) / h );
       double S = particle->S;
       double I = particle->I;
       double R = particle->R;
       double R0
                   = particle->R0;
       double r
                   = particle->r;
       double B
                   = R0 * r / N;
       for(int i = 0; i < num\_steps; i++) {
           // get derivatives
           double dS = -B*S*I;
236
```

```
double dI = B*S*I - r*I;
           double dR = r*I;
           // step forward by h
           S += h*dS;
           I += h*dI;
           R += h*dR;
       }
       particle -> S = S;
       particle ->I = I;
       particle ->R = R;
248
249 }
252 /* Particle pertubation function to be run between iterations and
       passes
255 void perturbParticles(Particle * particles, int N, int NP, int
       passnum, double coolrate) {
       double coolcoef = pow(coolrate, passnum);
       double spreadR0
                            = coolcoef * R0true / 10.0;
       double spreadr
                            = coolcoef * rtrue / 10.0;
       double spreadsigma = coolcoef * merr
                                                 / 10.0;
       double spreadIinit = coolcoef * I0
                                                 / 10.0;
       double R0can, rcan, sigmacan, Iinitcan;
       for (int n = 0; n < NP; n++) {
           do {
                R0can = particles[n].R0 + spreadR0*randn();
270
            } while (R0can < 0);</pre>
           particles[n].R0 = R0can;
272
273
           do {
                rcan = particles[n].r + spreadr*randn();
           } while (rcan < 0);</pre>
           particles[n].r = rcan;
           do {
                sigmacan = particles[n].sigma + spreadsigma*randn();
           } while (sigmacan < 0);</pre>
           particles[n].sigma = sigmacan;
           do {
                Iinitcan = particles[n].Iinit + spreadIinit*randn();
```

```
} while (Iinitcan < 0 || Iinitcan > 500);
           particles[n].Iinit = Iinitcan;
           particles[n].Sinit = N - Iinitcan;
       }
290
291 }
294 /*
       Convinience function for particle resampling process
296
297 void copyParticle(Particle * dst, Particle * src) {
       dst->R0
                   = src -> R0;
300
       dst->r
                   = src -> r;
       dst->sigma = src->sigma;
       dst->S
                   = src->S;
       dst->I
                   = src -> I;
       dst->R
                   = src -> R;
       dst->Sinit = src->Sinit;
       dst->Iinit = src->Iinit;
       dst->Rinit = src->Rinit;
309 }
312 /*
       Checks to see if particles are collapsed
       This is done by checking if the standard deviations between the
313
          particles' parameter
       values are significantly close to one another. Spread threshold
          may need to be tuned.
317 bool isCollapsed(Particle * particles, int NP) {
       bool retVal;
       double R0mean = 0, rmean = 0, sigmamean = 0, Sinitmean = 0,
           Iinitmean = 0, Rinitmean = 0;
       // means
       for (int n = 0; n < NP; n++) {
           R0mean
                        += particles[n].R0;
           rmean
                        += particles[n].r;
                        += particles[n].sigma;
           sigmamean
                       += particles[n].Sinit;
           Sinitmean
           Iinitmean
                        += particles[n].Iinit;
```

```
Rinitmean += particles[n].Rinit;
       }
       R0mean
                   /= NP;
       rmean
                   /= NP;
       sigmamean
                   /= NP;
                   /= NP;
       Sinitmean
       Iinitmean
                   /= NP;
       Rinitmean
                   /= NP;
       double R0sd = 0, rsd = 0, sigmasd = 0, Sinitsd = 0, Iinitsd =
          0, Rinitsd = 0;
       for (int n = 0; n < NP; n++) {
           R0sd
                   += ( particles[n].R0 - R0mean ) * ( particles[n].R0
               - R0mean );
                   += ( particles[n].r - rmean ) * ( particles[n].r -
           rsd
               rmean );
           sigmasd += ( particles[n].sigma - sigmamean ) * ( particles[
              n].sigma - sigmamean );
           Sinitsd += ( particles[n].Sinit - Sinitmean ) * ( particles[
               n].Sinit - Sinitmean );
           Iinitsd += ( particles[n].Iinit - Iinitmean ) * ( particles[
               n]. Iinit - Iinitmean );
           Rinitsd += ( particles[n].Rinit - Rinitmean ) * ( particles[
              n].Rinit - Rinitmean );
       }
       R0sd
                   /= NP;
       rsd
                   /= NP;
                   /= NP;
       sigmasd
                   /= NP;
       Sinitsd
       Iinitsd
                   /= NP;
       Rinitsd
                   /= NP;
       if (R0sd + rsd + sigmasd) < 1e-5)
           retVal = true;
       else
           retVal = false;
       return retVal;
369
370 }
372 void particleDiagnostics(ParticleInfo * partInfo, Particle *
      particles, int NP) {
373
```

```
double
                    = 0.0,
        R0mean
        rmean
                    = 0.0,
        sigmamean
                    = 0.0,
                    = 0.0,
        Sinitmean
        Iinitmean
                    = 0.0,
        Rinitmean
                    = 0.0;
// means
for (int n = 0; n < NP; n++) {
    R0mean
                += particles[n].R0;
    rmean
                += particles[n].r;
                += particles[n].sigma;
    sigmamean
    Sinitmean
                += particles[n].Sinit;
    Iinitmean
                += particles[n]. Iinit;
    Rinitmean
                += particles[n].Rinit;
}
R0mean
            /= NP;
rmean
            /= NP;
            /= NP;
sigmamean
Sinitmean
            /= NP;
            /= NP;
Iinitmean
Rinitmean
            /= NP;
// standard deviations
double R0sd
                = 0.0,
        rsd
                = 0.0.
        sigmasd = 0.0,
        Sinitsd = 0.0,
        Iinitsd = 0.0,
        Rinitsd = 0.0;
for (int n = 0; n < NP; n++) {
            += ( particles[n].R0 - R0mean ) * ( particles[n].R0
       - R0mean );
            += ( particles[n].r - rmean ) * ( particles[n].r -
       rmean );
    sigmasd += ( particles[n].sigma - sigmamean ) * ( particles[
       n].sigma - sigmamean );
    Sinitsd += ( particles[n].Sinit - Sinitmean ) * ( particles[
       n].Sinit - Sinitmean );
    Iinitsd += ( particles[n].Iinit - Iinitmean ) * ( particles[
       n]. Iinit - Iinitmean );
    Rinitsd += ( particles[n].Rinit - Rinitmean ) * ( particles[
       n].Rinit - Rinitmean );
```

```
418
       }
       R0sd
                   /= NP;
       rsd
                   /= NP;
       sigmasd
                   /= NP;
       Sinitsd
                   /= NP;
                   /= NP;
       Iinitsd
       Rinitsd
                   /= NP;
       partInfo->R0mean
                           = R0mean;
       partInfo->R0sd
                           = R0sd;
       partInfo->sigmamean = sigmamean;
       partInfo->sigmasd = sigmasd;
       partInfo->rmean
                           = rmean;
       partInfo->rsd
                           = rsd;
       partInfo->Sinitmean = Sinitmean;
       partInfo->Sinitsd = Sinitsd;
       partInfo->Iinitmean = Iinitmean;
       partInfo->Iinitsd = Iinitsd;
       partInfo->Rinitmean = Rinitmean;
       partInfo->Rinitsd = Rinitsd;
441 }
443 double randu() {
       return (double) rand() / (double) RAND_MAX;
447 }
450 /* Return a normally distributed random number with mean 0 and
       standard deviation 1
       Uses the polar form of the Box-Muller transformation
       From http://www.design.caltech.edu/erik/Misc/Gaussian.html
       */
454 double randn() {
       double x1, x2, w, y1;
       do {
           x1 = 2.0 * randu() - 1.0;
           x2 = 2.0 * randu() - 1.0;
           w = x1 * x1 + x2 * x2;
       } while ( w >= 1.0 );
       w = sqrt((-2.0 * log(w)) / w);
       y1 = x1 * w;
```

```
467 return y1;
468
469 }
```

Appendix C

Parameter Fitting

Appendix D

Forecasting Frameworks

D.1 IF2 Parametric Bootstrapping Function

The parametric bootstrapping machinery used to produce forecasts.

```
1 # Dexter Barrows
2 #
3 # IF2 parametric bootstrapping function
5 library(foreach)
6 library(parallel)
7 library(doParallel)
  library(Rcpp)
10| if2_paraboot \leftarrow function(if2data_parent, T, Tlim, steps, N, nTrials,
      if2file, if2_s_file, stoc_sir_file, NP, nPasses, coolrate) {
    source(stoc_sir_file)
    if (nTrials < 2)
      ntrials \leftarrow 2
    # unpack if2 first fit data
18
    # ...parameters
    paramdata_parent ← data.frame( if2data_parent$paramdata )
    names(paramdata_parent) \leftarrow c("R0", "r", "sigma", "eta", "berr", "
        Sinit", "Iinit", "Rinit")
21
    parmeans_parent ← colMeans(paramdata_parent)
    names(parmeans_parent) \leftarrow c("R0", "r", "sigma", "eta", "berr", "
        Sinit", "Iinit", "Rinit")
    # ...states
    statedata_parent ← data.frame( if2data_parent$statedata )
    names(statedata_parent) ← c("S","I","R","B")
```

```
statemeans_parent ← colMeans(statedata_parent)
27
    names(statemeans_parent) ← c("S","I","R","B")
    ## use parametric bootstrapping to generate forcasts
     trajectories \leftarrow foreach(i = 1:nTrials, .combine = rbind, .packages
         = "Rcpp") %dopar% {
       source(stoc_sir_file)
      ## draw new data
       pars ← with( as.list(parmeans_parent),
                      c(R0 = R0,
                      r = r,
                      N = N,
                      eta = eta,
                      berr = berr) )
       init_cond ← with( as.list(parmeans_parent),
                           c(S = Sinit,
                             I = Iinit,
                             R = Rinit))
       # generate trajectory
       sdeout \leftarrow StocSIR(init\_cond, pars, Tlim + 1, steps)
       colnames(sdeout) ← c('S','I','R','B')
       # add noise
       counts_raw ← sdeout[,'I'] + rnorm(dim(sdeout)[1], 0, parmeans_
          parent[['sigma']])
                    ← ifelse(counts_raw < 0, 0, counts_raw)</p>
         counts
         ## refit using new data
         ##
         rm(if2) # because stupid things get done in packages
         sourceCpp(if2file)
         if2time \leftarrow system.time( if2data \leftarrow if2(counts, Tlim+1, N, NP,
            nPasses, coolrate) )
         paramdata \leftarrow data.frame( if2data\$paramdata )
       names(paramdata) \leftarrow c("R0", "r", "sigma", "eta", "berr", "Sinit",
           "Iinit", "Rinit")
       parmeans ← colMeans(paramdata)
       names(parmeans) ← c("R0", "r", "sigma", "eta", "berr", "Sinit",
          "Iinit", "Rinit")
```

```
## generate the rest of the trajectory
       # pack new parameter estimates
       pars ← with( as.list(parmeans),
                      c(R0 = R0,
                      r = r,
                      N = N,
                      eta = eta,
                      berr = berr) )
81
       init\_cond \leftarrow c(S = statemeans\_parent[['S']],
82
                         I = statemeans_parent[['I']],
                         R = statemeans_parent[['R']])
84
       # generate remaining trajectory part
       sdeout_future \( \text{StocSIR(init_cond, pars, T-Tlim, steps)} \)
87
       colnames(sdeout_future) ← c('S','I','R','B')
       return ( c( counts = unname(sdeout_future[,'I']),
                    parmeans,
                    time = if2time[['user.self']]) )
    }
96
     return(trajectories)
98 }
```

D.2 RStan Forward Simulator

The code used to reconstruct the state estimates, then project the trajectory forward past data.

```
StocSIRstan ← function(y, pars, T, steps, berrvec, bveclim) {

out ← matrix(NA, nrow = (T+1), ncol = 4)

R0 ← pars[['R0']]
r ← pars[['r']]
N ← pars[['N']]
eta ← pars[['eta']]
berr ← pars[['berr']]

S ← y[['S']]
I ← y[['I']]
R ← y[['I']]
```

```
14
      B0 \leftarrow R0 * r / N
16
      B \leftarrow B0
18
      out[1,] \leftarrow c(S,I,R,B)
      h \leftarrow 1 \text{ / steps}
       for ( i in 1:(T*steps) ) {
             if (i <= bveclim) {</pre>
                B \leftarrow \exp(\log(B) + eta*(\log(B0) - \log(B)) + berrvec[i])
             } else {
                   B \leftarrow \exp(\log(B) + eta*(\log(B0) - \log(B)) + rnorm(1, 0,
                        berr))
28
             }
          \texttt{BSI} \leftarrow \texttt{B} {\star} \texttt{S} {\star} \texttt{I}
          rI \leftarrow r*I
          \mathsf{dS} \leftarrow \mathsf{-BSI}
          dI \leftarrow BSI - rI
          \text{dR} \leftarrow \text{rI}
          S \leftarrow S + h*dS #newInf
          I \leftarrow I + h*dI #newInf - h*dR
          R \leftarrow R + h*dR #h*dR
41
          if (i %% steps == 0)
             out[i/steps+1,] \leftarrow c(S,I,R,B)
      }
45
       return(out)
47
48 }
```

Appendix E

S-map and SIRS

E.1 SIRS R Function Code

R code to simulate the outlines SIRS function.

```
StocSIRS \leftarrow function(y, pars, T, steps) {
      out \leftarrow matrix(NA, nrow = (T+1), ncol = 4)
      R0 ← pars[['R0']]
      r \leftarrow pars[['r']]
      N \leftarrow pars[['N']]
      eta ← pars[['eta']]
      berr ← pars[['berr']]
         re ← pars[['re']]
      S \leftarrow y[['S']]
      I \leftarrow y[['I']]
      R \leftarrow y[['R']]
      B0 \leftarrow R0 * r / N
      B \leftarrow B0
      out[1,] \leftarrow c(S,I,R,B)
21
      h \leftarrow 1 / steps
      for ( i in 1:(T*steps) ) {
24
               \#Bfac \leftarrow 1/2 - \cos((2*pi/365)*i)/2
               \mathsf{Bfac} \leftarrow \mathsf{exp}(2*\mathsf{cos}((2*\mathsf{pi}/365)*\mathsf{i}) - 2)
28
         B \leftarrow \exp(\log(B) + eta*(\log(B0) - \log(B)) + rnorm(1, 0, berr))
```

```
BSI \leftarrow Bfac*B*S*I
        \texttt{rI} \leftarrow \texttt{r} {\star} \texttt{I}
             reR \leftarrow re*R
        dS \leftarrow -BSI + reR
        \texttt{dI} \leftarrow \texttt{BSI} - \texttt{rI}
        dR \leftarrow rI - reR
        S \leftarrow S + h*dS #newInf
        I \leftarrow I + h*dI #newInf - h*dR
        R \leftarrow R + h*dR #h*dR
        if (i %% steps == 0)
           out[i/steps+1,] \leftarrow c(S,I,R,B)
      }
      colnames(out) ← c("S","I","R","B")
      return(out)
50 }
52 ### suggested parameters
53 #
54 # T ← 200
55 # i_infec \leftarrow 10
56 # steps \leftarrow 7
57 # N ← 500
58 # sigma \leftarrow 5
59 #
60 # pars \leftarrow c(R0 = 3.0, # new infected people per infected person
          r = 0.1, # recovery rate
61 #
            N = 500, # population size
eta = 0.5, # geometric random walk
62 #
63 #
             berr = 0.5, # Beta geometric walk noise
64 #
65 #
            re = 1) # resuceptibility rate
```

E.2 SMAP Code

This code implements an SMAP function on a user-provided time series.

```
library(pracma)
smap ← function(data, E, theta, stepsAhead) {
    # construct library
```

```
6
       tseries ← as.vector(data)
       liblen \leftarrow length(tseries) - E + 1 - stepsAhead
               ← matrix(NA, liblen, E)
       for (i in 1:E) {
           lib[,i] \leftarrow tseries[(E-i+1):(liblen+E-i)]
       }
       # predict from the last index
       tslen \leftarrow length(tseries)
       predictee \leftarrow rev(t(as.matrix(tseries[(tslen-E+1):tslen])))
       predictions ← numeric(stepsAhead)
       #allPredictees ← matrix(NA, stepsAhead, E)
21
       # for each prediction index (number of steps ahead)
       for(i in 1:stepsAhead) {
24
           # set up weight calculation
           predmat ← repmat(predictee, liblen, 1)
           distances ← sqrt( rowSums( abs(lib - predmat)^2 ) )
27
           meanDist ← mean(distances)
           # calculate weights
           weights ← exp( - (theta * distances) / meanDist )
           # construct A, B
           preds ← tseries[(E+i):(liblen+E+i-1)]
           A \leftarrow cbind(rep(1.0, liblen), lib) * repmat(as.matrix(
               weights), 1, E+1)
           B ← as.matrix(preds * weights)
           # solve system for C
           Asvd \leftarrow svd(A)
           C \leftarrow Asvd$v %*% diag(1/Asvd$d) %*% t(Asvd$u) %*% B
           # get prediction
           predsum \leftarrow sum(C * c(1, predictee))
           # save
           predictions[i] \leftarrow predsum
           # next predictee
           #predictee ← c( predsum, predictee[-E] )
```

```
#allPredictees[i,] ← predictee

#allPredictees[i,] ← predictee

return(predictions)

return(predictions)
```

E.3 SMAP Parameter Optimization Code

This code determines the optimal parameter values to be used by the S-map algorithm.

```
library(deSolve)
2 library(ggplot2)
3 library (RColorBrewer)
4 library(pracma)
6 set.seed(1010)
8 ## external files
10 | stoc_sirs_file \leftarrow paste(getwd(), "../sir-functions", "StocSIRS.r",
      sep = "/")
11 smap_file
                 ← paste(getwd(), "smap.r", sep = "/")
12 source(stoc_sirs_file)
13 source(smap_file)
17 ## parameters
18 ##
19 T
         \leftarrow 6 * 52
20 Tlim \leftarrow T - 52
21 | i_i = 10
22 steps \leftarrow 7
23 N
        ← 500
24 \mid \text{sigma} \quad \leftarrow 5
26 true_pars \leftarrow c( R0 = 3.0, # new infected people per infected
      person
                  r = 0.1, # recovery rate
              N = 500, # population size
              eta = 0.5, # geometric random walk
              berr = 0.5, # Beta geometric walk noise
                  re = 1) # resuceptibility rate
```

```
33 true_init_cond \leftarrow c(S = N - i_infec,
                        I = i_infec,
                        R = 0)
37 ## trial parameter values to check.options
38 ##
39 Elist \leftarrow 1:20
40 thetalist \leftarrow 10*exp(-(seq(0,9.5,0.5)))
41 | \text{nTrials} \leftarrow 100
43 ssemat \leftarrow matrix(NA, 20, 20)
45 for (i in 1:length(Elist)) {
  for (j in 1:length(thetalist)) {
       ssemean \leftarrow 0
      for (k in 1:nTrials) {
         E \leftarrow Elist[i]
         theta ← thetalist[j]
         ## get true trajectory
         sdeout ← StocSIRS(true_init_cond, true_pars, T, steps)
         ## perturb to get data
         ##
         infec_counts_raw \( \text{sdeout[1:(Tlim+1),'I'] + rnorm(Tlim+1,0,} \)
         raw)
         predictions ← smap(infec_counts, E, theta, 52)
         err ← sdeout[(Tlim+2):dim(sdeout)[1],'I'] - predictions
         sse \leftarrow sum(err^2)
         ssemean ← ssemean + (sse / nTrials)
      }
73
       ssemat[i,j] \leftarrow ssemean
    }
  }
78
79 quartz()
80 image(-ssemat)
```

```
81 quartz()
82 filled.contour(-ssemat)
83
84 #print(ssemat)
85 \text{ #cms} \leftarrow \text{colMeans(ssemat)}
86 \text{ #rms} \leftarrow \text{rowMeans(ssemat)}
87
88 #Emin ← Elist[which.min(rms)]
89 |#thetamin \leftarrow thetalist[which.min(cms)]
90 #print(Emin)
91 #print(thetamin)
93 mininds \leftarrow which(ssemat==min(ssemat), arr.ind=TRUE)
95 Emin ← Elist[mininds[,'row']]
96 thetamin \leftarrow thetalist[mininds[,'col']]
98 print(Emin)
99 print(thetamin)
```

E.4 RStan SIRS Code

This code implements a periodic SIRS model in Rstan.

```
data {
                                   // total integration steps
      int
               <lower=1>
                                   // observed number of cases
      real
                           y[T];
                                   // population size
      int
               <lower=1>
                           N;
      real
                           h;
                                   // step size
7
8
  }
10 parameters {
      real <lower=0, upper=10>
                                       R0;
                                                // R0
      real <lower=0, upper=10>
                                                // recovery rate
                                       r;
      real <lower=0, upper=10>
                                                // resusceptibility rate
                                       re;
                                                // observation error
      real <lower=0, upper=20>
                                       sigma;
      real <lower=0, upper=30>
                                       Iinit;
                                                 // initial infected
      real <lower=0, upper=1>
                                       eta;
                                                // geometric walk
          attraction strength
      real <lower=0, upper=1>
                                       berr;
                                                // beta walk noise
      real <lower=-1.5, upper=1.5>
                                       Bnoise[T]; // Beta vector
20
21 }
22
```

```
23 //transformed parameters {
24 //
          real B0 \leftarrow R0 \star r / N;
25 //}
27 model {
        real S[T];
        real I[T];
       real R[T];
        real B[T];
        real B0;
        real pi;
       real Bfac;
       pi \leftarrow 3.1415926535;
       B0 \leftarrow R0 * r / N;
       B[1] \leftarrow B0;
       S[1] \leftarrow N - Iinit;
       I[1] \leftarrow Iinit;
       R[1] \leftarrow 0.0;
       for (t in 2:T) {
            Bnoise[t] ~ normal(0,berr);
            Bfac \leftarrow \exp(2*\cos((2*pi/365)*t) - 2);
            B[t] \leftarrow \exp(\log(B0) + eta * (\log(B[t-1]) - \log(B0)) +
                 Bnoise[t] );
            S[t] \leftarrow S[t-1] + h*( - Bfac*B[t]*S[t-1]*I[t-1] + re*R[t-1] );
            I[t] \leftarrow I[t-1] + h*( Bfac*B[t]*S[t-1]*I[t-1] - I[t-1]*r );
            R[t] \leftarrow R[t-1] + h*( I[t-1]*r - re*R[t-1] );
            if (y[t] > 0) {
                 y[t] ^ normal( I[t], sigma );
            }
       }
                 ~ lognormal(1,1);
       R0
                 ^{\sim} lognormal(1,1);
       r
                 ^{\sim} lognormal(1,1);
        sigma
                 ~ lognormal(1,1);
        re
                ~ normal(y[1], sigma);
       Iinit
70 }
```

E.5 IF2 SIRS Code

This code implements a periodic SIRS model using IF2 in C++.

```
Author: Dexter Barrows
      Github: dbarrows.github.io
      */
6 #include <stdio.h>
7 #include <math.h>
8 #include <sys/time.h>
9 #include <time.h>
10 #include <stdlib.h>
11 #include <string>
12 #include <cmath>
13 #include <cstdlib>
14 #include <fstream>
16 //#include "rand.h"
17 //#include "timer.h"
18
19 #define Treal
                       100
                                   // time to simulate over
20 #define R0true
                       3.0
                                   // infectiousness
21 #define rtrue
                       0.1
                                   // recovery rate
22 #define retrue
                       0.05
                                   // resusceptibility rate
23 #define Nreal
                       500.0
                                   // population size
24 #define etatrue
                       0.5
                                   // real drift attraction strength
25 #define berrtrue
                       0.5
                                   // real beta drift noise
26 #define merr
                       5.0
                                   // expected measurement error
27 #define I0
                                   // Initial infected individuals
                       5.0
                       0.5
29 #define PSC
                                   // scale factor for more sensitive
     parameters
31 #include <Rcpp.h>
32 using namespace Rcpp;
34 struct State {
      double S;
      double I;
      double R;
38 };
40 struct Particle {
      double R0;
      double r;
      double re;
      double sigma;
45
      double eta;
```

```
double berr;
      double B;
      double S;
      double I;
      double R;
      double Sinit;
      double Iinit;
      double Rinit;
54 };
56 struct ParticleInfo {
                           double R0sd;
      double R0mean;
      double rmean;
                           double rsd;
      double remean;
                           double resd:
      double sigmamean;
                           double sigmasd;
      double etamean;
                           double etasd;
      double berrmean;
                           double berrsd;
      double Sinitmean; double Sinitsd;
      double Iinitmean; double Iinitsd;
      double Rinitmean;
                         double Rinitsd;
66 };
69 int timeval_subtract (double *result, struct timeval *x, struct
      timeval *y);
70 int check_double(double x, double y);
71 void exp_euler_SIRS(double h, double t0, double tn, int N, Particle
      * particle);
72 void copyParticle(Particle * dst, Particle * src);
73 void perturbParticles(Particle * particles, int N, int NP, int
      passnum, double coolrate);
74 void particleDiagnostics(ParticleInfo * partInfo, Particle *
      particles, int NP);
75 void getStateMeans(State * state, Particle* particles, int NP);
76 NumericMatrix if2(NumericVector * data, int T, int N);
77 double randu();
78 double randn();
80 // [[Rcpp::export]]
81 Rcpp::List if2_sirs(NumericVector data, int T, int N, int NP, int
      nPasses, double coolrate) {
82
      int npar = 9;
84
      NumericMatrix paramdata(NP, npar);
86
      NumericMatrix means(nPasses, npar);
      NumericMatrix sds(nPasses, npar);
87
      NumericMatrix statemeans(T, 3);
      NumericMatrix statedata(NP, 4);
90
```

```
srand(time(NULL));
                        // Seed PRNG with system time
double w[NP];
                    // particle weights
Particle particles[NP];  // particle estimates for current
Particle particles_old[NP]; // intermediate particle states for
   resampling
printf("Initializing particle states\n");
// initialize particle parameter states (seeding)
for (int n = 0; n < NP; n++) {
    double R0can, rcan, recan, sigmacan, Iinitcan, etacan,
       berrcan;
    do {
        R0can = R0true + R0true*randn();
    \} while (R0can < 0);
    particles[n].R0 = R0can;
    do {
        rcan = rtrue + rtrue*randn();
    } while (rcan < 0);</pre>
    particles[n].r = rcan;
    do {
        recan = retrue + retrue*randn();
    } while (recan < 0);</pre>
    particles[n].re = recan;
    particles[n].B = (double) R0can * rcan / N;
    do {
        sigmacan = merr + merr*randn();
    } while (sigmacan < 0);</pre>
    particles[n].sigma = sigmacan;
    do {
        etacan = etatrue + PSC*etatrue*randn();
    } while (etacan < 0 || etacan > 1);
    particles[n].eta = etacan;
    do {
        berrcan = berrtrue + PSC*berrtrue*randn();
    } while (berrcan < 0);</pre>
    particles[n].berr = berrcan;
    do {
```

```
138
               Iinitcan = I0 + I0*randn();
           } while (Iinitcan < 0 || N < Iinitcan);</pre>
           particles[n].Sinit = N - Iinitcan;
           particles[n].Iinit = Iinitcan;
           particles[n].Rinit = 0.0;
       }
       // START PASSES THROUGH DATA
       printf("Starting filter\n");
       printf("----\n");
       printf("Pass\n");
       for (int pass = 0; pass < nPasses; pass++) {</pre>
           printf("...%d / %d\n", pass, nPasses);
           // reset particle system evolution states
           for (int n = 0; n < NP; n++) {
               particles[n].S = particles[n].Sinit;
               particles[n].I = particles[n].Iinit;
               particles[n].R = particles[n].Rinit;
               particles[n].B = (double) particles[n].R0 * particles[n
                   ].r / N;
           }
           if (pass == (nPasses-1)) {
               State sMeans;
               getStateMeans(&sMeans, particles, NP);
               statemeans(0,0) = sMeans.S;
               statemeans(0,1) = sMeans.I;
               statemeans(0,2) = sMeans.R;
           }
           for (int t = 1; t < T; t++) {
               // generate individual predictions and weight
               for (int n = 0; n < NP; n++) {
                   exp_euler_SIRS(1.0/7.0, (double) t-1, (double) t, N,
                        &particles[n]);
181
                   double merr_par = particles[n].sigma;
                   double y_diff = data[t] - particles[n].I;
                   w[n] = 1.0/(merr_par*sqrt(2.0*M_PI)) * exp( - y_diff|
```

```
*y_diff / (2.0*merr_par*merr_par) );
               }
               // cumulative sum
               for (int n = 1; n < NP; n++) {
                   w[n] += w[n-1];
               }
               // save particle states to resample from
               for (int n = 0; n < NP; n++){
                    copyParticle(&particles_old[n], &particles[n]);
               }
               // resampling
200
               for (int n = 0; n < NP; n++) {
                    double w_r = randu() * w[NP-1];
                    int i = 0;
                    while (w_r > w[i]) {
                        i++;
                    }
                    // i is now the index to copy state from
                    copyParticle(&particles[n], &particles_old[i]);
               }
               // between-iteration perturbations, not after last time
                   step
               if (t < (T-1))
                    perturbParticles(particles, N, NP, pass, coolrate);
                if (pass == (nPasses -1)) {
                    State sMeans;
                    getStateMeans(&sMeans, particles, NP);
                    statemeans(t,0) = sMeans.S;
                    statemeans(t,1) = sMeans.I;
                    statemeans(t,2) = sMeans.R;
               }
           }
           ParticleInfo pInfo;
           particleDiagnostics(&pInfo, particles, NP);
           means(pass, 0) = pInfo.R0mean;
           means(pass, 1) = pInfo.rmean;
           means(pass, 2) = pInfo.remean;
           means(pass, 3) = pInfo.sigmamean;
```

```
means(pass, 4) = pInfo.etamean;
           means(pass, 5) = pInfo.berrmean;
           means(pass, 6) = pInfo.Sinitmean;
           means(pass, 7) = pInfo.Iinitmean;
           means(pass, 8) = pInfo.Rinitmean;
           sds(pass, 0) = pInfo.R0sd;
           sds(pass, 1) = pInfo.rsd;
           sds(pass, 2) = pInfo.resd;
           sds(pass, 3) = pInfo.sigmasd;
           sds(pass, 4) = pInfo.etasd;
           sds(pass, 5) = pInfo.berrsd;
           sds(pass, 6) = pInfo.Sinitsd;
           sds(pass, 7) = pInfo.Iinitsd;
           sds(pass, 8) = pInfo.Rinitsd;
           // between-pass perturbations, not after last pass
           if (pass < (nPasses + 1))
               perturbParticles(particles, N, NP, pass, coolrate);
       }
       ParticleInfo pInfo;
       particleDiagnostics(&pInfo, particles, NP);
       printf("Parameter results (mean | sd)\n");
       printf("----\n");
                         %f %f\n", pInfo.R0mean, pInfo.R0sd);
       printf("R0
                         %f %f\n", pInfo.rmean, pInfo.rsd);
       printf("r
       printf("re
                         %f %f\n", pInfo.remean, pInfo.resd);
       printf("sigma
                         %f %f\n", pInfo.sigmamean, pInfo.sigmasd);
       printf("eta
                         %f %f\n", pInfo.etamean, pInfo.etasd);
       printf("berr
                      %f %f\n", pInfo.berrmean, pInfo.berrsd);
       printf("S\_init ~\%f ~\%f\n", pInfo.Sinitmean, pInfo.Sinitsd);\\
       printf("I_init
                       %f %f\n", pInfo.Iinitmean, pInfo.Iinitsd);
       printf("R_init
                        %f %f\n", pInfo.Rinitmean, pInfo.Rinitsd);
       printf("\n");
272
       // Get particle results to pass back to R
       for (int n = 0; n < NP; n++) {
           paramdata(n, 0) = particles[n].R0;
           paramdata(n, 1) = particles[n].r;
           paramdata(n, 2) = particles[n].re;
           paramdata(n, 3) = particles[n].sigma;
           paramdata(n, 4) = particles[n].eta;
```

```
paramdata(n, 5) = particles[n].berr;
           paramdata(n, 6) = particles[n].Sinit;
           paramdata(n, 7) = particles[n].Iinit;
           paramdata(n, 8) = particles[n].Rinit;
       }
       for (int n = 0; n < NP; n++) {
           statedata(n, 0) = particles[n].S;
           statedata(n, 1) = particles[n].I;
           statedata(n, 2) = particles[n].R;
           statedata(n, 3) = particles[n].B;
       }
       return Rcpp::List::create( Rcpp::Named("paramdata") = paramdata
                                   Rcpp::Named("means") = means,
                                   Rcpp::Named("statemeans") =
                                       statemeans,
                                   Rcpp::Named("statedata") = statedata
                                   Rcpp::Named("sds") = sds);
308 }
311 /*
       Use the Explicit Euler integration scheme to integrate SIR model
       forward in time
       double h - time step size
       double t0 - start time
       double tn - stop time
       double * y - current system state; a three-component vector
          representing [S I R], susceptible-infected-recovered
318 void exp_euler_SIRS(double h, double t0, double tn, int N, Particle
      * particle) {
       int num_steps = floor( (tn-t0) / h );
       double S = particle->S;
       double I = particle->I;
       double R = particle->R;
       double R0
                   = particle->R0;
                 = particle->r;
       double r
```

```
328
       double re
                   = particle->re;
       double B0 = R0 * r / N;
       double eta = particle->eta;
       double berr = particle->berr;
       double B = particle->B;
       for(int i = 0; i < num\_steps; i++) {
           //double Bfac = 0.5 - 0.95*cos( (2.0*M_PI/365)*(t0*num_steps
              +i) )/2.0;
           double Bfac = exp(2*cos((2*M_PI/365)*(t0*num_steps+i)) - 2);
           B = \exp(\log(B) + eta*(\log(B0) - \log(B)) + berr*randn());
           double BSI = Bfac*B*S*I;
           double rI = r*I;
           double reR = re*R;
           // get derivatives
           double dS = - BSI + reR;
           double dI = BSI - rI;
           double dR = rI - reR;
           // step forward by h
           S += h*dS;
           I += h*dI;
           R += h*dR;
       }
       particle->S = S;
       particle ->I = I;
       particle -> R = R;
       particle ->B = B;
362 }
365 /* Particle pertubation function to be run between iterations and
      passes
368 void perturbParticles(Particle * particles, int N, int NP, int
       passnum, double coolrate) {
       //double coolcoef = exp( - (double) passnum / coolrate );
       double coolcoef = pow(coolrate, passnum);
       double spreadR0 = coolcoef * R0true / 10.0;
```

```
double spreadr
                           = coolcoef * rtrue / 10.0;
       double spreadre
                           = coolcoef * retrue / 10.0;
       double spreadsigma = coolcoef * merr / 10.0;
       double spreadIinit = coolcoef * I0 / 10.0;
       double spreadeta = coolcoef * etatrue / 10.0;
       double spreadberr = coolcoef * berrtrue / 10.0;
       double R0can, rcan, recan, sigmacan, Iinitcan, etacan, berrcan;
       for (int n = 0; n < NP; n++) {
           do {
               R0can = particles[n].R0 + spreadR0*randn();
           \} while (R0can < 0);
           particles[n].R0 = R0can;
           do {
               rcan = particles[n].r + spreadr*randn();
           } while (rcan < 0);
           particles[n].r = rcan;
           do {
               recan = particles[n].re + spreadre*randn();
           } while (recan < 0);</pre>
           particles[n].re = recan;
           do {
               sigmacan = particles[n].sigma + spreadsigma*randn();
           } while (sigmacan < 0);</pre>
           particles[n].sigma = sigmacan;
           do {
               etacan = particles[n].eta + PSC*spreadeta*randn();
           \} while (etacan < 0 || etacan > 1);
           particles[n].eta = etacan;
           do {
               berrcan = particles[n].berr + PSC*spreadberr*randn();
           } while (berrcan < 0);</pre>
           particles[n].berr = berrcan;
           do {
               Iinitcan = particles[n].Iinit + spreadIinit*randn();
           } while (Iinitcan < 0 || Iinitcan > 500);
           particles[n].Iinit = Iinitcan;
           particles[n].Sinit = N - Iinitcan;
       }
424
```

```
425 }
428 /*
       Convinience function for particle resampling process
431 void copyParticle(Particle * dst, Particle * src) {
       dst->R0
                   = src -> R0;
       dst->r
                   = src ->r;
       dst->re
                  = src->re;
436
       dst->sigma = src->sigma;
       dst->eta
                  = src->eta;
       dst->berr = src->berr;
       dst->B
                  = src->B;
       dst->S
                   = src->S;
       dst->I
                   = src->I;
       dst->R
                  = src->R;
       dst->Sinit = src->Sinit;
       dst->Iinit = src->Iinit;
       dst->Rinit = src->Rinit;
447 }
449 void particleDiagnostics(ParticleInfo * partInfo, Particle *
      particles, int NP) {
       double
               R0mean
                           = 0.0.
               rmean
                           = 0.0,
               remean
                           = 0.0,
               sigmamean
                           = 0.0,
               etamean
                           = 0.0.
               berrmean
                          = 0.0,
               Sinitmean = 0.0,
                           = 0.0,
               Iinitmean
               Rinitmean
                           = 0.0;
       // means
       for (int n = 0; n < NP; n++) {
           R0mean
                       += particles[n].R0;
           rmean
                       += particles[n].r;
                       += particles[n].re;
           remean
           etamean
                       += particles[n].eta,
                       += particles[n].berr,
           berrmean
                       += particles[n].sigma;
           sigmamean
                       += particles[n].Sinit;
           Sinitmean
           Iinitmean
                       += particles[n].Iinit;
473
           Rinitmean
                       += particles[n].Rinit;
```

```
}
                   /= NP;
       R0mean
       rmean
                   /= NP;
                   /= NP;
       remean
                   /= NP;
       sigmamean
                   /= NP;
       etamean
       berrmean
                   /= NP;
       Sinitmean
                   /= NP;
       Iinitmean
                   /= NP;
       Rinitmean
                   /= NP;
       // standard deviations
       double R0sd
                       = 0.0,
               rsd
                       = 0.0,
               resd
                       = 0.0,
               sigmasd = 0.0,
               etasd
                       = 0.0,
               berrsd = 0.0,
               Sinitsd = 0.0,
               Iinitsd = 0.0,
               Rinitsd = 0.0;
       for (int n = 0; n < NP; n++) {
           R0sd
                   += ( particles[n].R0 - R0mean ) * ( particles[n].R0
               - R0mean );
                   += ( particles[n].r - rmean ) * ( particles[n].r -
               rmean );
           resd += ( particles[n].re - rmean ) * ( particles[n].re -
               rmean );
           sigmasd += ( particles[n].sigma - sigmamean ) * ( particles[
              n].sigma - sigmamean );
           etasd += ( particles[n].eta - etamean ) * ( particles[n].
               eta - etamean );
           berrsd += ( particles[n].berr - berrmean ) * ( particles[n
               ].berr - berrmean );
           Sinitsd += ( particles[n].Sinit - Sinitmean ) * ( particles[
               n].Sinit - Sinitmean );
           Iinitsd += ( particles[n].Iinit - Iinitmean ) * ( particles[
              n].Iinit - Iinitmean );
           Rinitsd += ( particles[n].Rinit - Rinitmean ) * ( particles[
               n].Rinit - Rinitmean );
       }
513
       R0sd
                   /= NP;
514
                   /= NP;
       rsd
```

```
515
                   /= NP;
       resd
       sigmasd
                   /= NP;
       etasd
                   /= NP;
518
       berrsd
                   /= NP;
       Sinitsd
                   /= NP;
       Iinitsd
                   /= NP;
       Rinitsd
                   /= NP;
       partInfo->R0mean
                           = R0mean;
       partInfo->R0sd
                          = R0sd;
       partInfo->rmean
                         = rmean;
                         = rsd;
       partInfo->rsd
                         = remean;
       partInfo->remean
       partInfo->resd = resd;
       partInfo->sigmamean = sigmamean;
       partInfo->sigmasd = sigmasd;
       partInfo->etamean = etamean;
       partInfo->etasd = etasd;
       partInfo->berrmean = berrmean;
       partInfo->berrsd = berrsd;
       partInfo->Sinitmean = Sinitmean;
       partInfo->Sinitsd = Sinitsd;
       partInfo->Iinitmean = Iinitmean;
       partInfo->Iinitsd = Iinitsd;
       partInfo->Rinitmean = Rinitmean;
       partInfo->Rinitsd = Rinitsd;
542 }
544 double randu() {
       return (double) rand() / (double) RAND_MAX;
548 }
550 void getStateMeans(State * state, Particle* particles, int NP) {
       double Smean = 0, Imean = 0, Rmean = 0;
       for (int n = 0; n < NP; n++) {
           Smean += particles[n].S;
           Imean += particles[n].I;
           Rmean += particles[n].R;
       }
560
       state->S = (double) Smean / NP;
       state->I = (double) Imean / NP;
       state->R = (double) Rmean / NP;
564 }
```

```
565
567 /\star Return a normally distributed random number with mean 0 and
      standard deviation 1
       Uses the polar form of the Box-Muller transformation
       From http://www.design.caltech.edu/erik/Misc/Gaussian.html
       */
571 double randn() {
       double x1, x2, w, y1;
       do {
           x1 = 2.0 * randu() - 1.0;
           x2 = 2.0 * randu() - 1.0;
           w = x1 * x1 + x2 * x2;
       } while ( w >= 1.0 );
       w = sqrt((-2.0 * log(w)) / w);
581
       y1 = x1 * w;
       return y1;
586 }
```

Appendix F

Spatial Epidemics

F.1 Spatial SIR R Function Code

R code to simulate the outlined Spatial SIR function.

```
## ymat: Contains the initial conditions where:
2 # - rows are locations
          - columns are S, I, R
4 ## pars: Contains the parameters: global values for R0, r, N, eta,
      berr
5 ## T:
             The stop time. Since 0 in included, there should be T+1
      time steps in the simulation
6 ## neinum: Number of neighbors for each location, in order
7 | ## neibmat: Contains lists of neighbors for each location

    rows are parent locations (nodes)

           - columns are locations each parent is attached to (edges)
10 StocSSIR \leftarrow function(ymat, pars, T, steps, neinum, neibmat) {
    ## number of locations
       nloc \leftarrow dim(ymat)[1]
      ## storage
      ## dims are locations, (S,I,R,B), times
       # output array
      out \leftarrow array(NA, c(nloc, 4, T+1), dimnames = list(NULL, c("S","I"
           ,"R","B"), NULL))
      # temp storage
      BSI \leftarrow numeric(nloc)
21
       rI \leftarrow numeric(nloc)
      ## extract parameters
      R0 \leftarrow pars[['R0']]
       r \leftarrow pars[['r']]
```

```
N \leftarrow pars[['N']]
27
         eta ← pars[['eta']]
         berr ← pars[['berr']]
         phi ← pars[['phi']]
         B0 \leftarrow rep(R0*r/N, nloc)
         ## state vectors
         S \leftarrow ymat[,'S']
         I \leftarrow ymat[,'I']
         R \leftarrow ymat[,'R']
         B \leftarrow B0
         ## assign starting to output matrix
         out[,,1] \leftarrow cbind(ymat, B0)
         h \leftarrow 1 \text{ / steps}
43
         for ( i in 1:(T*steps) ) {
               B \leftarrow \exp(\log(B) + eta*(\log(B0) - \log(B)) + rnorm(nloc, 0,
                    berr) )
               for (loc in 1:nloc) {
                  n \, \leftarrow \, neinum[loc]
                  sphi \leftarrow 1 - phi*(n/(n+1))
                  ophi \leftarrow phi/(n+1)
                  nBIsum ← B[neibmat[loc,1:n]] %*% I[neibmat[loc,1:n]]
                  BSI[loc] \leftarrow S[loc]*( sphi*B[loc]*I[loc] + ophi*nBIsum )
               }
               #if(i == 1)
               # print(BSI)
59
               \texttt{rI} \leftarrow \texttt{r*I}
               \mathsf{dS} \leftarrow \mathsf{-BSI}
               \texttt{dI} \leftarrow \texttt{BSI} \ \texttt{-} \ \texttt{rI}
               dR \leftarrow rI
               S \leftarrow S + h*dS
               I \leftarrow I + h*dI
               R \leftarrow R + h*dR
               if (i %% steps == 0) {
                     out[,,i/steps+1] \leftarrow cbind(S,I,R,B)
71
               }
         }
74
```

```
\#out[,,2] \leftarrow cbind(S,I,R,B)
    return(out)
79 }
80
81 ### Suggested parameters
82 #
83 # T
            ← 60
84 \# i\_infec \leftarrow 5
85 # steps \leftarrow 7
86 # N
           ← 500
           ← 10
87 # sigma
88 #
89 # pars \leftarrow c(R0 = 3.0,
                            # new infected people per infected person
            r = 0.1,
                            # recovery rate
90 #
               N = 500,
91 #
                             # population size
92 #
               eta = 0.5,
                             # geometric random walk
               berr = 0.5) # Beta geometric walk noise
93 #
```

F.2 RStan Spatial SIR Code

This code implements a Spatial SIR model in Rstan.

```
data {
                                   // total integration steps
      int
               <lower=1>
                                   // number of locations
      int
               <lower=1>
                           nloc;
      real
                           y[nloc, T]; // observed number of cases
      int
               <lower=1>
                           N;
                                   // population size
                                   // step size
      real
               <lower=0>
                           neinum[nloc];
                                                 // number of neighbors
      int
          each location has
                           neibmat[nloc, nloc]; // neighbor list for
      int
          each location
11 }
13 parameters {
      real <lower=0, upper=10>
                                        R0;
                                                // R0
      real <lower=0, upper=10>
                                                // recovery rate
                                        r;
                                               // observation error
      real <lower=0, upper=20>
                                        sigma;
                                                        // initial
      real <lower=0, upper=30>
                                        Iinit[nloc];
          infected for each location
      real <lower=0, upper=1>
                                        eta;
                                                // geometric walk
          attraction strength
```

```
real <lower=0, upper=1>
                                            berr;
                                                     // beta walk noise
21
       real <lower=-1.5, upper=1.5>
                                            Bnoise[nloc,T];
                                                                // Beta vector
                                                     // interconnectivity
       real <lower=0, upper=1>
                                            phi;
           strength
24 }
26 model {
       real S[nloc, T];
       real I[nloc, T];
       real R[nloc, T];
       real B[nloc, T];
       real B0;
       real BSI[nloc, T];
       real rI[nloc, T];
       int n;
       real sphi;
       real ophi;
       real nBIsum;
       B0 \leftarrow R0 * r / N;
       for (loc in 1:nloc) {
            S[loc, 1] \leftarrow N - Iinit[loc];
            I[loc, 1] ← Iinit[loc];
            R[loc, 1] \leftarrow 0.0;
            B[loc, 1] \leftarrow B0;
       }
       for (t in 2:T) {
            for (loc in 1:nloc) {
                Bnoise[loc, t] ~ normal(0,berr);
                B[loc, t] \leftarrow exp(log(B[loc, t-1]) + eta * (log(B0) -
                    log(B[loc, t-1]) ) + Bnoise[loc, t]);
                n \leftarrow neinum[loc];
                sphi \leftarrow 1.0 - phi*(n/(n+1.0));
                ophi \leftarrow phi/(n+1.0);
                nBIsum \leftarrow 0.0;
                for (j in 1:n)
                     nBIsum \leftarrow nBIsum + B[neibmat[loc, j], t-1] * I[
                         neibmat[loc, j], t-1];
                BSI[loc, t] \leftarrow S[loc, t-1]*( sphi*B[loc, t-1]*I[loc, t-1]
                     + ophi*nBIsum );
                rI[loc, t] \leftarrow r*I[loc, t-1];
```

```
S[loc, t] \leftarrow S[loc, t-1] + h*( - BSI[loc, t] );
                I[loc, t] \leftarrow I[loc, t-1] + h*(BSI[loc, t] - rI[loc, t])
                R[loc, t] \leftarrow R[loc, t-1] + h*(rI[loc, t]);
                if (y[loc, t] > 0) {
                    y[loc, t] ~ normal( I[loc, t], sigma );
           }
       }
                 lognormal(1,1);
       R0
                ~ lognormal(1,1);
                ~ lognormal(1,1);
       sigma
81
       for (loc in 1:nloc) {
82
           Iinit[loc] ~ normal(y[loc, 1], sigma);
       }
84
85 }
```

F.3 IF2 Spatial SIR Code

This code implements a Spatial SIR model using IF2 in C++.

```
Author: Dexter Barrows
      Github: dbarrows.github.io
  #include <stdio.h>
  #include <math.h>
8 #include <sys/time.h>
  #include <time.h>
10 #include <stdlib.h>
11 #include <string>
12 #include <cmath>
13 #include <cstdlib>
14 #include <fstream>
16 //#include "rand.h"
17 //#include "timer.h"
19 #define Treal
                       100
                                   // time to simulate over
20 #define R0true
                       3.0
                                   // infectiousness
21 #define rtrue
                       0.1
                                   // recovery rate
```

```
22 #define Nreal
                       500.0
                                   // population size
23 #define etatrue
                       0.5
                                   // real drift attraction strength
24 #define berrtrue
                       0.5
                                   // real beta drift noise
25 #define phitrue
                       0.5
                                   // real connectivity strength
26 #define merr
                       10.0
                                   // expected measurement error
27 #define I0
                       5.0
                                   // Initial infected individuals
29 #define PSC
                                   // perturbation scale factor for
                       0.5
     more sensitive parameters
31 #include <Rcpp.h>
32 using namespace Rcpp;
34 struct Particle {
      double R0;
      double r;
      double sigma;
      double eta;
      double berr;
      double phi;
      double * S;
      double * I;
      double * R;
      double * B;
      double * Iinit;
46 };
49 int timeval_subtract (double *result, struct timeval *x, struct
      timeval *y);
50 int check_double(double x, double y);
51 void initializeParticles(Particle ** particles, int NP, int nloc,
      int N);
52 void exp_euler_SSIR(double h, double t0, double tn, int N, Particle
      * particle,
                       NumericVector neinum, NumericMatrix neibmat, int
                           nloc);
54 void copyParticle(Particle * dst, Particle * src, int nloc);
55 void perturbParticles(Particle * particles, int N, int NP, int nloc,
       int passnum, double coolrate);
56 double randu();
57 double randn();
59 // [[Rcpp::export]]
60 Rcpp::List if2_spa(NumericMatrix data, int T, int N, int NP, int
      nPasses, double coolrate, NumericVector neinum, NumericMatrix
      neibmat, int nloc) {
      NumericMatrix paramdata(NP, 6);  // for R0, r, sigma, eta,
          berr, phi
```

```
NumericMatrix initInfec(nloc, NP); // for Iinit
       NumericMatrix infecmeans(nloc, T); // mean infection counts for
           each location
       NumericMatrix finalstate(nloc, 4); // SIRB means for each
          location
       srand(time(NULL));
                              // Seed PRNG with system time
       double w[NP];
                              // particle weights
       // initialize particles
       printf("Initializing particle states\n");
       Particle * particles = NULL; // particle estimates for
          current step
       Particle * particles_old = NULL; // intermediate particle
          states for resampling
       initializeParticles(&particles, NP, nloc, N);
       initializeParticles(&particles_old, NP, nloc, N);
       // copy particle test
       copyParticle(&particles[0], &particles_old[0], nloc);
81
       // perturb particle test
       perturbParticles(particles, N, NP, nloc, 1, coolrate);
84
       // evolution test
86
       // reset particle system evolution states
       for (int n = 0; n < NP; n++) {
87
           for (int loc = 0; loc < nloc; loc++) {
               particles[n].S[loc] = N - particles[n].Iinit[loc];
               particles[n].I[loc] = particles[n].Iinit[loc];
               particles[n].R[loc] = 0.0;
               particles[n].B[loc] = (double) particles[n].R0 *
                  particles[n].r / N;
       printf("Before S:%f | I:%f | R:%f\n", particles[0].S[0],
          particles[0].I[0], particles[0].R[0]);
       exp_euler_SSIR(1.0/7.0, 0.0, 1.0, N, &particles[0], neinum,
          neibmat, nloc);
       printf("After S:%f | I:%f | R:%f\n", particles[0].S[0],
          particles[0].I[0], particles[0].R[0]);
       */
       // START PASSES THROUGH DATA
100
       printf("Starting filter\n");
       printf("----\n");
       printf("Pass\n");
```

```
for (int pass = 0; pass < nPasses; pass++) {
           printf("...%d / %d\n", pass, nPasses);
           // reset particle system evolution states
           for (int n = 0; n < NP; n++) {
               for (int loc = 0; loc < nloc; loc++) {
                   particles[n].S[loc] = N - particles[n].Iinit[loc];
                   particles[n].I[loc] = particles[n].Iinit[loc];
                   particles[n].R[loc] = 0.0;
                   particles[n].B[loc] = (double) particles[n].R0 *
                       particles[n].r / N;
               }
119
           }
           if (pass == (nPasses-1)) {
               double means[nloc];
               for (int loc = 0; loc < nloc; loc++) {
                   means[loc] = 0.0;
                   for (int n = 0; n < NP; n++) {
                       means[loc] += particles[n].I[loc] / NP;
                   infecmeans(loc, 0) = means[loc];
               }
           }
           for (int t = 1; t < T; t++) {
               // generate individual predictions and weight
               for (int n = 0; n < NP; n++) {
                   exp_euler_SSIR(1.0/7.0, 0.0, 1.0, N, &particles[n],
                       neinum, neibmat, nloc);
                   double merr_par = particles[n].sigma;
                   w[n] = 1.0;
                   for (int loc = 0; loc < nloc; loc++) {
                        double y_diff = data(loc, t) - particles[n].I[
                           loc];
                       w[n] *= 1.0/(merr_par*sqrt(2.0*M_PI)) * exp(-
                           y_diff*y_diff / (2.0*merr_par*merr_par) );
                   }
               }
               // cumulative sum
               for (int n = 1; n < NP; n++) {
```

```
w[n] += w[n-1];
        }
        // save particle states to resample from
        for (int n = 0; n < NP; n++){
            copyParticle(&particles_old[n], &particles[n], nloc)
        }
        // resampling
        for (int n = 0; n < NP; n++) {
            double w_r = randu() * w[NP-1];
            int i = 0;
            while (w_r > w[i]) {
                i++;
            }
            // i is now the index to copy state from
            copyParticle(&particles[n], &particles_old[i], nloc)
        }
        // between-iteration perturbations, not after last time
        if (t < (T-1))
            perturbParticles(particles, N, NP, nloc, pass,
               coolrate);
        if (pass == (nPasses-1)) {
            double means[nloc];
            for (int loc = 0; loc < nloc; loc++) {
                means[loc] = 0.0;
                for (int n = 0; n < NP; n++) {
                    means[loc] += particles[n].I[loc] / NP;
                infecmeans(loc, t) = means[loc];
            }
        }
    }
    // between-pass perturbations, not after last pass
    if (pass < (nPasses + 1))
        perturbParticles(particles, N, NP, nloc, pass, coolrate)
           ;
}
```

```
196
       // pack parameter data (minus initial conditions)
       for (int n = 0; n < NP; n++) {
           paramdata(n, 0) = particles[n].R0;
           paramdata(n, 1) = particles[n].r;
           paramdata(n, 2) = particles[n].sigma;
           paramdata(n, 3) = particles[n].eta;
           paramdata(n, 4) = particles[n].berr;
           paramdata(n, 5) = particles[n].phi;
       }
       // Pack initial condition data
       for (int n = 0; n < NP; n++) {
           for (int loc = 0; loc < nloc; loc++) {
               initInfec(loc, n) = particles[n].Iinit[loc];
           }
       }
       // Pack final state means data
       double Smeans[nloc], Imeans[nloc], Rmeans[nloc];
       for (int loc = 0; loc < nloc; loc++) {
           Smeans[loc] = 0.0;
           Imeans[loc] = 0.0;
218
           Rmeans[loc] = 0.0;
           Bmeans[loc] = 0.0;
           for (int n = 0; n < NP; n++) {
               Smeans[loc] += particles[n].S[loc] / NP;
               Imeans[loc] += particles[n].I[loc] / NP;
               Rmeans[loc] += particles[n].R[loc] / NP;
               Bmeans[loc] += particles[n].B[loc] / NP;
           }
           finalstate(loc, 0) = Smeans[loc];
           finalstate(loc, 1) = Imeans[loc];
           finalstate(loc, 2) = Rmeans[loc];
           finalstate(loc, 3) = Bmeans[loc];
       }
       return Rcpp::List::create( Rcpp::Named("paramdata") = paramdata
                                   Rcpp::Named("initInfec") = initInfec
                                   Rcpp::Named("infecmeans") =
                                       infecmeans,
                                   Rcpp::Named("finalstate") =
                                       finalstate);
240 }
```

```
242
243 /*
       Use the Explicit Euler integration scheme to integrate SIR model
       forward in time
       double h - time step size
                   - start time
       double t0
       double tn - stop time
       double * y - current system state; a three-component vector
           representing [S I R], susceptible-infected-recovered
250 void exp_euler_SSIR(double h, double t0, double tn, int N, Particle
      * particle,
                        NumericVector neinum, NumericMatrix neibmat, int
                            nloc) {
       int num_steps = floor( (tn-t0) / h );
       double * S = particle->S;
       double * I = particle->I;
       double * R = particle->R;
       double * B = particle->B;
       // create last state vectors
       double S_last[nloc];
       double I_last[nloc];
       double R_last[nloc];
       double B_last[nloc];
       double R0
                   = particle->R0;
       double r
                  = particle->r;
268
       double B0 = R0 * r / N;
       double eta = particle->eta;
       double berr = particle->berr;
       double phi = particle->phi;
272
273
       //printf("sphi \t\ ophi \t\ BSI \t\ rI \t\ dS \t\ dI \t\
           t \mid dR \mid t \mid S \mid t \mid I \mid t \mid R \mid n'';
275
       for(int t = 0; t < num\_steps; t++) {
           for (int loc = 0; loc < nloc; loc++) {
               S_last[loc] = S[loc];
               I_last[loc] = I[loc];
               R_last[loc] = R[loc];
               B_last[loc] = B[loc];
           }
           for (int loc = 0; loc < nloc; loc++) {
               B[loc] = exp(log(B_last[loc]) + eta*(log(B0) - log(
```

```
B_last[loc])) + berr*randn() );
               int n = neinum[loc];
               double sphi = 1.0 - phi*((double) n/(n+1.0));
               double ophi = phi/(n+1.0);
               double nBIsum = 0.0;
               for (int j = 0; j < n; j++)
                   nBIsum += B_last[(int) neibmat(loc, j) - 1] * I_last
                      [(int) neibmat(loc, j) - 1];
               double BSI = S_last[loc]*( sphi*B_last[loc]*I_last[loc]
                  + ophi*nBIsum );
               double rI = r*I_last[loc];
              // get derivatives
              double dS = -BSI;
              double dI = BSI - rI;
              double dR = rI;
              // step forward by h
              S[loc] += h*dS;
              I[loc] += h*dI;
              R[loc] += h*dR;
              //if (loc == 1)
               |%f\t|\n", sphi, ophi, BSI, rI, dS, dI, dR, S[1], I
                  [1], R[1]);
           }
       }
       /*particle->S = S;
       particle ->I = I;
       particle -> R = R;
       particle -> B = B; */
321 }
323 /*
      Initializes particles
325 void initializeParticles(Particle ** particles, int NP, int nloc,
      int N) {
       // allocate space for doubles
       *particles = (Particle*) malloc (NP*sizeof(Particle));
330
       // allocate space for arays inside particles
```

```
for (int n = 0; n < NP; n++) {
            (*particles)[n].S = (double*) malloc(nloc*sizeof(double));
            (*particles)[n].I = (double*) malloc(nloc*sizeof(double));
            (*particles)[n].R = (double*) malloc(nloc*sizeof(double));
            (*particles)[n].B = (double*) malloc(nloc*sizeof(double));
            (*particles)[n].Iinit = (double*) malloc(nloc*sizeof(double)
               );
       }
       // initialize all all parameters
       for (int n = 0; n < NP; n++) {
            double R0can, rcan, sigmacan, Iinitcan, etacan, berrcan,
               phican;
           do {
                R0can = R0true + R0true*randn();
            } while (R0can < 0);</pre>
            (*particles)[n].R0 = R0can;
           do {
                rcan = rtrue + rtrue*randn();
            } while (rcan < 0);
            (*particles)[n].r = rcan;
            for (int loc = 0; loc < nloc; loc++)
                (*particles)[n].B[loc] = (double) R0can * rcan / N;
           do {
                sigmacan = merr + merr*randn();
            } while (sigmacan < 0);</pre>
            (*particles)[n].sigma = sigmacan;
           do {
                etacan = etatrue + PSC*etatrue*randn();
            } while (etacan < 0 || etacan > 1);
            (*particles)[n].eta = etacan;
           do {
                berrcan = berrtrue + PSC*berrtrue*randn();
            } while (berrcan < 0);</pre>
            (*particles)[n].berr = berrcan;
           do {
                phican = phitrue + PSC*phitrue*randn();
            } while (phican <= 0 \mid \mid phican >= 1);
            (*particles)[n].phi = phican;
            for (int loc = 0; loc < nloc; loc++) {
378
                do {
```

```
Iinitcan = I0 + I0*randn();
                } while (Iinitcan < 0 || N < Iinitcan);</pre>
                (*particles)[n]. Iinit[loc] = Iinitcan;
           }
       }
386 }
388 /* Particle pertubation function to be run between iterations and
      passes
       */
391 void perturbParticles(Particle * particles, int N, int NP, int nloc,
       int passnum, double coolrate) {
       //double coolcoef = exp( - (double) passnum / coolrate );
       double coolcoef = pow(coolrate, passnum);
       double spreadR0
                            = coolcoef * R0true / 10.0;
       double spreadr
                           = coolcoef * rtrue / 10.0;
       double spreadsigma = coolcoef * merr / 10.0;
       double spreadIinit = coolcoef * I0 / 10.0;
       double spreadeta
                           = coolcoef * etatrue / 10.0;
       double spreadberr = coolcoef * berrtrue / 10.0;
       double spreadphi
                          = coolcoef * phitrue / 10.0;
       double R0can, rcan, sigmacan, Iinitcan, etacan, berrcan, phican;
       for (int n = 0; n < NP; n++) {
           do {
               R0can = particles[n].R0 + spreadR0*randn();
           } while (R0can < 0);</pre>
           particles[n].R0 = R0can;
           do {
               rcan = particles[n].r + spreadr*randn();
           } while (rcan < 0);</pre>
           particles[n].r = rcan;
           do {
                sigmacan = particles[n].sigma + spreadsigma*randn();
           } while (sigmacan < 0);</pre>
           particles[n].sigma = sigmacan;
           do {
               etacan = particles[n].eta + PSC*spreadeta*randn();
           \} while (etacan < 0 || etacan > 1);
           particles[n].eta = etacan;
```

```
do {
               berrcan = particles[n].berr + PSC*spreadberr*randn();
           } while (berrcan < 0);</pre>
           particles[n].berr = berrcan;
           do {
               phican = particles[n].phi + PSC*spreadphi*randn();
           } while (phican \leq 0 || phican \geq 1);
           particles[n].phi = phican;
           for (int loc = 0; loc < nloc; loc++) {
               do {
                    Iinitcan = particles[n].Iinit[loc] + spreadIinit*
                       randn();
                } while (Iinitcan < 0 || Iinitcan > 500);
                particles[n].Iinit[loc] = Iinitcan;
           }
       }
446 }
448 /*
       Convinience function for particle resampling process
450 void copyParticle(Particle * dst, Particle * src, int nloc) {
       dst->R0
                   = src -> R0;
       dst->r
                   = src ->r;
       dst->sigma = src->sigma;
                  = src->eta;
       dst->eta
       dst->berr
                   = src->berr;
       dst->phi
                  = src->phi;
       for (int n = 0; n < nloc; n++) {
           dst->S[n]
                           = src->S[n];
           dst->I[n]
                           = src->I[n];
           dst->R[n]
                           = src -> R[n];
           dst->B[n]
                          = src->B[n];
           dst->Iinit[n] = src->Iinit[n];
       }
467 }
471 double randu() {
       return (double) rand() / (double) RAND_MAX;
475 }
```

```
476
   /*
478 void getStateMeans(State * state, Particle* particles, int NP) {
       double Smean = 0, Imean = 0, Rmean = 0;
       for (int n = 0; n < NP; n++) {
           Smean += particles[n].S;
           Imean += particles[n].I;
           Rmean += particles[n].R;
       state->S = (double) Smean / NP;
       state->I = (double) Imean / NP;
       state->R = (double) Rmean / NP;
492 }
493 */
495 /* Return a normally distributed random number with mean 0 and
       standard deviation 1
496
       Uses the polar form of the Box-Muller transformation
       From http://www.design.caltech.edu/erik/Misc/Gaussian.html
499 double randn() {
       double x1, x2, w, y1;
       do {
           x1 = 2.0 * randu() - 1.0;
           x2 = 2.0 * randu() - 1.0;
           w = x1 * x1 + x2 * x2;
       } while ( w >= 1.0 );
       w = sqrt((-2.0 * log(w)) / w);
       y1 = x1 * w;
       return y1;
514 }
```