Data augmentation in the general epidemic model

SISMID/July 18-20, 2022

Instructors: Vladimir Minin, Kari Auranen, Elizabeth Halloran



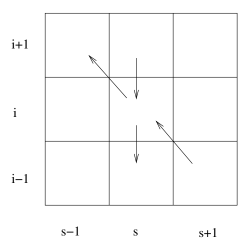
Outline

- The general epidemic model
 - A simple Susceptible–Infected–Removed (SIR) model of an outbreak of infection in a closed population
- Likelihood function for the infection and removal rates
 - Assuming complete data: both infection and removal times are observed
 - Under Gamma priors for the infection and removal rates, their full conditionals are also Gamma, so Gibbs updating steps can be used
- Incomplete data: only removal times are observed
 - Augment the unknown infection times to be able to apply the complete-data likelihood
 - Additional Metropolis-Hastings steps are required to sample the infection times

SIR model

- Consider a closed population of M individuals
- One introductory case (infective) introduces the infection into a population of initially susceptible individuals, starting an outbreak
- ▶ Once the outbreak has started, the hazard of infection for a still susceptible individual at time t depends on the number of infectives I(t) in the population: $(\beta/M)I(t)$
- If an individual becomes infected, the hazard of clearing infection (and stopping being infective) is γ , i.e., he/she remains infective for an exponentially distributed period of time. He/she then becomes *removed* and does not contribute to the outbreak any more
- ► There is no latency

Transitions in the state space



Complete data

- Assume one introductory case whose infection takes place at time t = 0 (this fixes the time origin)
- For M individuals followed from time 0 until the end of the outbreak at time T (after which time the number of infectives I(t) = 0), the *complete data* record all event times
- ▶ This is equivalent to observing the n-1 infection times and the n removal times, and the fact the M-n individuals escaped infection throughout the outbreak

$$\overbrace{\left\{0=\textit{i}_1<\textit{i}_2<...<\textit{i}_n\right\}}^{\text{infection times}} \text{ and } \overbrace{\left\{r_1<...<\textit{r}_{n-1}<\textit{r}_n=\textit{T}\right\}}^{\text{removal times}}$$

N.B. Here, the i_k and r_k do not correspond to the same individual (we will discuss this assumption later; see p. 19)

Counting the numbers of infectives and susceptibles

- Denote the ordered event times i_1, \ldots, i_n and r_1, \ldots, r_n jointly as $0 = u_1 < u_2 < \ldots < u_{2n} = T$
- ▶ Denote the indicators for time u_k being an infection or removal time by D_k and R_k , respectively
- ▶ Denote the number of infectives at time t by I(t)
 - I(t) is a piecewise constant (left-continuous) function, assuming values in the set $\{0, 1, ..., M\}$
 - ▶ I(t) jumps at times $u_2 < \ldots < u_{2n}$
- ▶ Denote the number of susceptibles at time t by S(t)
 - ▶ S(t) is a piecewise constant (left-continuous) function, jumping at times $i_2 < ... < i_n$
- Note that both I(t) and S(t) are fully determined by the complete data



Example S(t) 4 3 2 1 3 **I** *I(t)* 2 1 Time t i_4 $i_1 = 0$ i_3 i_2 $r_4 = T$ r_1 r_2 $u_1 = 0$ u_2 U_3 U_4 u_5 u_6 $u_8 = T$ U_7 $D_k = 0$ 0 0 1 0

0

0

 $R_k = 0$

The process of infections

- New infections occur as a non-homogeneous Poisson process with rate $\beta I(t)S(t)/M$
 - ▶ the rate is a piecewise constant (left-continuous) function
 - ▶ the rate jumps at times $u_2 < ... < u_{2n}$, with levels $\beta I(u_2)S(u_2)/M$, $\beta I(u_3)S(u_3)/M$, ..., $\beta I(u_{2n})S(u_{2n})/M$
- ▶ The probability density of infections occurring at i_1, \ldots, i_n can now be written by multiplying contributions from each of the 2n-1 subintervals:

The process of removals

- Removals occur as a non-homogeneous Poisson process with rate $\gamma I(t)$
 - ▶ the rate is a piecewise constant (left-continuous) function
 - he rate jumps at times $u_2 < \ldots < u_{2n}$, with levels $\gamma I(u_2), \gamma I(u_3), \ldots, \gamma I(u_{2n})$
- ▶ The probability density of removals occurring at $r_1, ..., r_n$ is thus

$$\prod_{k=2}^{2n} \left[(\gamma I(u_k))^{R_k} e^{-\gamma I(u_k)(u_k - u_{k-1})} \right]$$

$$\underbrace{-\gamma \sum_{k=2}^{2n} I(u_k)(u_k - u_{k-1})}_{\text{total time spent infective}}$$

$$= \prod_{k=2}^{2n} (\gamma I(u_k))^{R_k} \times e^{-\gamma I(u_k)(u_k - u_{k-1})}$$

Complete data likelihood

- The so called complete-data likelihood $L(\beta, \gamma; i, r)$ of parameters β and γ is based on the joint probability density $f(i, r|\beta, \gamma)$ of the infection and removal times
- ► The complete-data likelihood is obtained by putting together the expressions on pages 8 and 9:

$$\underbrace{L(\beta, \gamma; \boldsymbol{i}, \boldsymbol{r})}_{L(\beta, \gamma; \boldsymbol{i}, \boldsymbol{r})} = \prod_{k=2}^{2n} (\beta I(u_k) S(u_k))^{D_k} \prod_{k=2}^{2n} (\gamma I(u_k))^{R_k}$$

$$\times e^{-[(\beta/M)\sum_{k=2}^{2n}I(u_k)S(u_k)(u_k-u_{k-1})+\gamma\sum_{k=2}^{2n}I(u_k)(u_k-u_{k-1})]}$$

$$= \prod_{k=2}^{n} \{\beta I(i_{k}) S(i_{k})\} \prod_{k=1}^{n} \{\gamma I(r_{k})\}$$

$$-[(\beta/M)\sum_{k=2}^{2n}I(u_k)S(u_k)(u_k-u_{k-1})+\gamma\sum_{k=2}^{2n}I(u_k)](u_k-u_{k-1})]$$
× e

Simplifying the notation

Note that the total time of infectious pressure can be written simply as an integral, i.e.

$$\sum_{k=2}^{2n} I(u_k)S(u_k)(u_k - u_{k-1}) = \int_0^T I(u)S(u)du$$

Similarly the total time spent infective is

$$\sum_{k=2}^{2n} I(u_k)(u_k - u_{k-1}) = \int_0^T I(u) du$$

▶ The complete-data likelihood can thus be written as

$$\prod_{k=2}^{n} \{\beta I(i_k) S(i_k)\} \prod_{k=1}^{n} \{\gamma I(r_k)\}$$

$$\times \exp\left(-\int_{0}^{T} [(\beta/M) I(u) S(u) + \gamma I(u)] du\right)$$

Computation of the integral terms

▶ In practice, the integral terms can be calculated as follows:

 $\overbrace{\int_{0}^{T} I(u) du}^{\text{total time spent infective}} = \sum_{k=1}^{n} (r_k - i_k)$

total time of infectious pressure

$$\int_{0}^{T} I(u)S(u)du = \sum_{k=1}^{n} \sum_{j=1}^{M} (\min(r_{k}, i_{j}) - \min(i_{k}, i_{j}))$$

where $i_j = \infty$ for j > n, i.e., for those never infected

► These expressions are invariant to choice of which r_k corresponds to which i_k

Poisson likelihood and Gamma priors

- ▶ The complete-data likelihood of the two parameters, β and γ , is often called a Poisson likelihood
- In particular, Gamma distributions can be used as conjugate priors for β and γ
- It follows that the full conditional distributions of β and γ are also Gamma and can be updated by Gibbs steps (see pages 14–16)

Gamma prior distributions

 \blacktriangleright The two rate parameters β and γ are given independent Gamma priors

$$f(\beta) \propto \beta^{
u_{eta}-1} \exp(-\lambda_{eta} eta)$$

 $f(\gamma) \propto \gamma^{
u_{\gamma}-1} \exp(-\lambda_{\gamma} \gamma)$

- Note that these choices pf the prior the full conditional distributions of both β and γ are gamma distributions (the next two pages)
- ▶ In practice, this means that updating β and γ within an MCMC algorithm can be implemented as Gibbs steps

The full conditional of β

Parameter β can be updated through a Gibbs step because the full conditional of β is a Gamma distribution:

$$f(\beta|\mathbf{i},\mathbf{r},\gamma) \propto f(\beta,\gamma,\mathbf{i},\mathbf{r}) \propto f(\mathbf{i},\mathbf{r}|\beta,\gamma)f(\beta)$$

$$\propto \beta^{n-1} \exp\left(-(\beta/M) \int_0^T I(u)S(u)du\right) \beta^{\nu_\beta-1} \exp(-\lambda_\beta\beta)$$

$$= \beta^{n+\nu_\beta-2} \exp\left(-[(1/M) \int_0^T I(u)S(u)du + \lambda_\beta]\beta\right)$$

The full conditional distribution of β is thus the following Gamma distribution:

$$eta|(oldsymbol{i},oldsymbol{r},\gamma)\sim\Gamma\left(n+
u_eta-1,(1/M)\int_0^TI(u)S(u)du+\lambda_eta
ight)$$

The full conditional of γ

Also the full conditional of parameter γ is a Gamma distribution:

$$f(\gamma|\mathbf{i},\mathbf{r},\beta) \propto f(\beta,\gamma,\mathbf{i},\mathbf{r}) \propto f(\mathbf{i},\mathbf{r}|\beta,\gamma)f(\gamma)$$

$$\propto \gamma^n \exp\left(-\gamma \int_0^T I(u)du\right) \gamma^{\nu_{\gamma}-1} \exp(-\lambda_{\gamma}\gamma)$$

$$= \gamma^{n+\nu_{\gamma}-1} \exp\left(-\left[\int_0^T I(u)du + \lambda_{\gamma}\right]\gamma\right)$$

lacktriangle The full conditional of γ thus is

$$\gamma | (\boldsymbol{i}, \boldsymbol{r}, \beta) \sim \Gamma \left(n + \nu_{\gamma}, \int_{0}^{T} I(u) du + \lambda_{\gamma} \right)$$

Incomplete data

- Assume now that only the removal times $\mathbf{r} = (r_1, \dots, r_n)$ have been observed
- Augment the set of unknowns (β and γ) with infection times $\mathbf{i} = (i_2, \dots, i_n)$
- The aim is to do statistical inference about rates β and γ (and times i), based on their posterior distribution $f(\beta, \gamma, i|r)$
- ► The posterior distribution is proportional to the joint distribution of all model quantities:

$$f(\beta, \gamma, \boldsymbol{i} | \boldsymbol{r}) \propto f(\beta, \gamma, \boldsymbol{i}, \boldsymbol{r}) = \overbrace{f(\boldsymbol{i}, \boldsymbol{r} | \beta, \gamma)}^{\text{complete data likelihood}} \overbrace{f(\beta, \gamma, \boldsymbol{i}, \boldsymbol{r})}^{\text{prior}},$$

Updating infection times

- lacktriangle The full conditional distributions of eta and γ are as above
- ► The unknown infection times *t* require a Metropolis–Hastings step, including explicit evaluations of the Poisson likelihood
- ▶ If the current iterate of i_k is $i_k^{(j)}$, a new value \tilde{i}_k is first proposed (e.g.) from a uniform distribution on [0, T]
- lacktriangle The proposal is then accepted, i.e., $i_k^{(j+1)}:= ilde{i}$, with probability

$$\min\{1, \frac{f(\tilde{\pmb{i}}, \pmb{r}|\beta, \gamma)}{f(\pmb{i}, \pmb{r}|\beta, \gamma)}\}$$

▶ Here \tilde{i} is i except for the kth entry which is \tilde{i}_k (instead of $i_k^{(j)}$)

Augmenting individual histories

- ➤ The likelihood above was constructed for the aggregate processes, i.e., counting the total numbers of susceptibles and infectives
- ▶ In such a case, the corresponding augmentation model must not consider individuals
 - ▶ In particular, times i₂,..., i_n must not be tied to particular removal times, i.e., individual event histories must not be reconstructed
- ▶ However, if one considers individual event histories as pairs of times (i_k, r_k) for individuals k = 1, ..., M, the appropriate complete data likelihood is

$$\gamma^n \prod_{k=2}^n \{\beta I(i_k)\} \exp\left(-\int_0^T (\gamma I(u) + (\beta/M)I(u)S(u))du\right)$$

▶ N.B. The likelihood construction in the computer lab is based on the above *individual-based* approach



Example: a smallpox outbreak

- ► The Abakaliki smallpox outbreak
 - A village of M = 120 inhabitants
 - One introductory case
 - ▶ 29 subsequent cases; this means that n = 1 + 29 = 30
- We will assume that the index case started being infectious on day 0 and that she/he entered the village starting the outbreak at the same day
- ► The observed data are the 30 removal times (in days) with respect to the time origin:

 \blacktriangleright The problem: to estimate rates β and γ from these data; see the computer lab data



A useful reference

- ► The computer lab analysis of the Abakaliki data is not realistic as it omits
 - the relevant stages of infection (latent time from exposure to infectiousness, possibly varying infectiousness during fever and the subsequent rash/symptoms)
 - the fact that isolation of cases at their symptomatic stage was only implemented at some point during the outbreak
 - community structure (compounds in the village and the larger community)
 - two more cases that occurred outside the particular group of faith with the 30 cases considered here
- ➤ For example, we assumed for simplicity that removal occurred at the time of symptoms although in reality removal only occurred at recovery/death *or* isolation, and only after some delay since symptom onset
- ► A proper analysis is given by Stockdale et al. (2017)

References

- [1] O'Neill Ph. and Roberts G. Bayesian inference for partially observed stochastic processes. Journal of the Royal Statistical Society, Series A, 1999; 162: 121–129.
- [2] O'Neill Ph. A tutorial introduction to Bayesian inference for stochastic epidemic models using Markov chain Monte Carlo methods. Mathematical Biosciences 2002; 180:103-114.
- [3] Becker N. Analysis of infectious disease data. Chapman and Hall, New York 1989.
- [4] Andersen et al. Statistical models based on counting processes. Springer Verlag, New York, 1993.
- [5] Stockdale J, Kypraios Th., O'Neill Ph. Modelling and Bayesian analysis of the Abakaliki smallpox data. Epidemics 2017; 19:13–23.