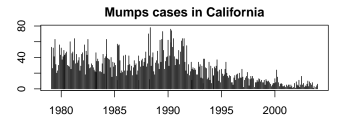
Stochastic Models

John M. Drake & Pejman Rohani

Epidemiological data are noisy

Two types of noise:

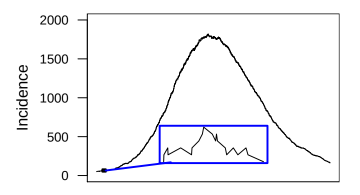
- Observation error: the data are probabilistically related to the true state of the system
- Process noise: the system progresses probabilistically
 - Environmental noise: some parameter is a random variable
 - Demographic noise: individual-level chance events



Noise is addressed using stochastic models



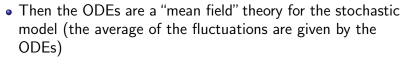
The SIR model is a continuum approximation



The SIR model (e.g., $dY/dt = \beta XY/N - \gamma Y$) implies that changes in the states X, Y, and Z are continuous. But, in reality individuals are either susceptible, infected, or recovered so that X, Y, and Z are integer-valued and changes in the system state occur as discrete steps. The differential equation is an idealization.

Demographic stochasticity

- What we seek is a stochastic model for which the system of ODEs is an appropriate idealization
- There are an infinite number of such mode but the simplest one is a continuous e, discrete ace Markov Chain with propensities given by the various terms in the differential equations



 This model may also be interpreted as an event-driven model with state transition probabilities

Analytic solution

"Master Equation"

$$\frac{Q}{dP_k/dt} = \sum_{l} A_{kl} P_l \tag{1}$$

where A is a matrix of transition propensities

This approach is only tractable for very simple models (e.g. *SI* and *SIS* epidemics)

Simulation approach

Exact simulation is straightforward via Gillespie's Direct method:

- Initialize
- Iteration of a two step process
 - Determine time of the next event
 - Determine change of state at the next event time
- Summarize



Step 1: time to next event

Given system state (S_{t}, S_{t}) be the sum of all the propensities for all changes of state and (S_{t}, S_{t}) be the probability that no event occurs in subsequent time interval S_{t} for system state N.

By the Markov assumption

$$G_N(s + \delta s) = Pr \{ \text{no event in}(t, t + \delta s) \}$$

= $Pr \{ \text{no event in}(t, t + s) \}$
= $G_N(s) \times \{1 - R(N) \times \delta s \}$ r {no event in}(t + s, t + s + \delta s) }

Step 1: time to next event

After rearranging

$$\frac{G_N(s+\delta s)-G_N(s)}{\delta s}=-R(N)\times G_N(s)$$

Letting $\delta s o 0$

$$\frac{dG_N}{ds} = -R(N) \times G_N(s)$$

With solution

$$G_N(s) = e^{-R(N)s}$$

Thus, the probability the next event occurs in (t, t + s) is

$$F_N(s) = 1 - e^{-R(N)s}$$







Step 1: time to next event

Given event time distribution F_N , an exponentially distributed random event time S can be obtained from a uniform random random variate U_1 by setting



$$U_1 = F_N(s) = 1 - e^{-R(N)S}$$

and solving to obtain

$$S = -\log(R(N))$$

Step 2: change of state



Let the propensities of event types $E_1, E_2, E_3, ...$ be denoted $R_1, R_2, R_3, ...$ with total rate $R_{sum} = R(N) = \sum_i R_i$. In the long run, events of each type should occur with relative frequency $R_i/R(N)$. We can randomly draw event classes with these frequencies by simulating a second uniform random variate U_2 and assigning event class E_i if

$$R_{sum}^{-1} \sum_{i=1}^{p-1} R_i < U_2 \le R_{sum}^{-1} \sum_{i=1}^{p} R_i.$$

Gillespie's direct method

- Label all possible events $E_1, E_2, E_3, ...$
- 2 Initialize t = 0 and state N
- Update step
 - Calculate propensities $R_1, R_2, R_3, ...$
 - 2 Calculate $R_{sum} = R(N) = \sum_{i} R_{i}$
 - **9** Generate U_1 and transform to obtain S
 - **9** Generate U_2 and determine event type E_i
 - **5** Update state based on E_i
 - **6** Update time t = t + S
- Go to step (3)

Example with SIR model

• Events:

- E_1 : Birth of susceptible individual $(X \to X + 1)$
- E_2 : Infection $(X \rightarrow X 1, Y \rightarrow Y + 1)$
- E_3 : Recovery $(Y \rightarrow Y 1, Z \rightarrow Z + 1)$
- E_4 : Death of susceptible individual $(X \to X 1)$
- E_5 : Death of infected individual ($Y \rightarrow Y 1$)
- ullet E_6 : Death of recovered individual (Z o Z 1)

Propensities

- R_1 : $\mu(X + Y + Z)$
- R₂: βXY/N
- R₃: γ Y
- R₄: μX
- R₅: μy
- R₆: μZ



We create a function SIR.onestep to perform calculations of each update step

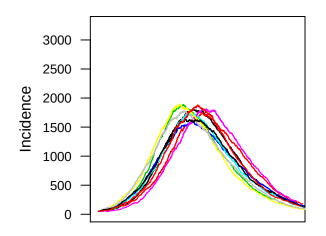
```
> SIR.onestep <- function (x. params) { #function to calculate one step of stochastic SIR
    X <- x [2]
                                           #local variable for susceptibles
   Y <- x[3]
                                           #local variable for infecteds
    Z \leftarrow x \lceil 4 \rceil
                                           #local variable for recovereds
   N \leftarrow X+Y+Z
                                           #total population size (subject to demographic change)
   with(
                                           #use with as in deterministic model to simplify code
         as.list(params),
           total.rate <- mu*N+beta*X*Y/N+mu*X+mu*Y+gamma*Y+mu*Z #calculate ``total rate''
           tau <- rexp(n=1,rate=total.rate)
                                                                   #inter-event time
           new.xyz <- c(X,Y,Z) #initialize a local variable at previous state variable values
           U \leftarrow runif(1)
                                #uniform random deviate
           new.xyz<-c(X,Y,Z-1) #death of recovered id `default''
           if (U<=(mu*N+beta*X*Y/N+mu*X+gamma*Y+mu*Y)/total.rate) new.xyz<-c(X,Y-1,Z) #death of infected
           if (U<=(mu*N+beta*X*Y/N+mu*X+gamma*Y)/total.rate) new.xyz<-c(X,Y-1,Z+1) #recovery of infected
           if (U<=(mu*N+beta*X*Y/N+mu*X)/total.rate) new.xyz<-c(X-1,Y,Z)
                                                                                #death of a susceptible
           if (U \le (mu \cdot N + beta \cdot X \cdot Y / N) / total.rate) new.xyz< -c(X-1, Y+1, Z)
                                                                                #transmission event
           if (U<=(mu*N/total.rate)) new.xyz<-c(X+1, Y, Z)
                                                                                #birth of susceptible
           c(tau,new.xyz) #store result
         }
         )
```

Now we write a function SIR.model that iteratively calls SIR.onestep to simulate an epidemic

```
> SIR.model <- function (x, params, nstep) {  #function to simulate stochastic SIR
+ output <- array(dim=c(nstep+1,4))  #set up array to store results
+ colnames(output) <- c("time","X","Y","Z") #name variables
+ output[1,] <- x  #first record of output is initial condition
+ for (k in 1:nstep) {  #iterate for nstep steps
+ output[k+1,] <- x <- SIR.onestep(x,params)
+ }
+ output  #return output
+ }</pre>
```

Finally, we write a code that calls SIR.model to simulate epidemics

```
> set.seed(38499583)
                                    #set seed
> nsims <- 10
                                    #number of simulations
> pop.size <- 10000
                                      #total population size
> YO <- 50
                                     #initial number infected
> X0 <- round(0.98*pop.size)
                                    #initial number suscepitlble (~98% of population)
> nstep <- 16000
                                     #number of events to simulate
> xstart <- c(time=0, X=X0, Y=Y0, Z=pop.size-X0-Y0) #initial conditions
> params <- list(mu=0.00001,beta=60,gamma=365/13) #parameters
> data <- vector(mode='list'.length=nsims) #initialize list to store the output
> for (k in 1:nsims) {
                                  #simulate nsims times
+ data[[k]] <- as.data.frame(SIR.model(xstart,params,nstep))</pre>
+ data[[k]]$cum.time <- cumsum(data[[k]]$time)
+ }
> max.time<-data[[1]]$cum.time[max(which(data[[1]]$Y>0))] #maximum time in first simulation
> max.y<-1.8*max(data[[1]]$Y) #find max infected in run 1 and increase by 80% for plo
> plot(Y~cum.time,data=data[[1]],xlab='Time',ylab='Incidence',col=1,xlim=c(0,max.time),ylim=
> hox()
> axis(2, cex.axis=0.8, las=2)
> for (k in 1:nsims) {
                                    #add multiple epidemics to plot
+ lines(Y~cum.time.data=data[[k]].col=k.tvpe='l')
+ }
```



Some stochastic phenomena

J-U transition in final outbreak size

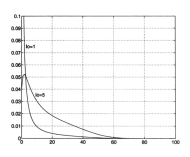


Figure 1. Size distribution of the general epidemic (N = 100, $R_0 = 0.9$).

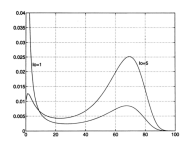
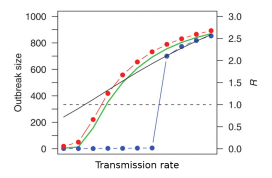


Figure 3. Size distribution of the general epidemic $(N = 100, R_0 = 1.5)$.

J-U transition illustrated by Nasell (1995) in *Epidemic models: their structure and relation to data*

Some stochastic phenomena

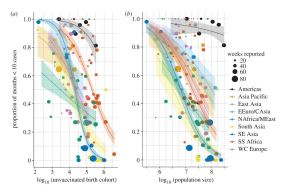
Difference between "likely" outcome (median: red points) and "worst case scenario" (95th percentile: blue points) compared with deterministic approximation (green line) and R_0 (black line)



Park et al. 2009. Science 326:726-728

Some stochastic phenomena

Critical community size



Ferrari et al. 2013. Philosophical Transactions of the Royal Society B 368:20120141

Summary

- Transmission is obscured by three sources of noise: observation error, environmental variability, and intrinsic demographic noise
- Gillespie's direct method is a straightforward way to study the effects of demographic stochasticity in small populations
- Demographic noise is especially important in systems where $R_0 \approx 1$