

# Hierarchical Spatial Structure in Compartmental Epidemiological models

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## **Abstract**

Epidemiological modelling is essential to understand and predict the spread of infectious diseases and plan effective interventions. Disease spread is an inherently spatial process, particularly in diseases that spread through contact with body fluids, and spatially structured epidemiological models provide an opportunity to incorporate a wealth of geographical information into the modelling process. Here, we present a compartmental meta-population model encoding the hierarchical spatial structure of the Australian Statistical Geography Standard (ASGS). Relationships between spatial groupings are represented in a mixing matrix, which influences the spread of infection between patches (i.e. Patches in the same spatial grouping are more likely to spread disease to each other compared to patches outside this grouping). As an example, we model the Greater Melbourne Statistical Area (SA) with patches representing SA2, SA4, and SA4 level groupings, and stochastically simulate the course of an epidemic as a Continuous Time Markov Chain (CTMC). We compare the results of simulations with varying influence of the spatial relationships between patches (through the construction of the mixing matrix), different levels of spatial resolution (which level of the spatial hierarchy is represented by compartmental patches), and different infectious disease parameters (e.g.  $R_0$ ), as well as with the results of a spatial compartmental model incorporating an alternative source of geographic information (i.e. empirically derived mobility data). We conclude by speculating on the potential use of such geographically informed epidemiological models in the management of infectious diseases.

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This thesis is written using Quarto with renv (Ushey, 2022) to create a reproducible environment. All materials (including the data sets and source files) required to reproduce this document can be found at the Github repository github.com/SusanSu/thesis.

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## **Chapter 1**

## Introduction & Background

#### 1.1 Modelling Infectious Epidemics

"Epidemic models are used to inform decisions on disease prevention, surveillance, control and treatment and can be applied to new epidemics" (bjørnstad2020?)

#### 1.2 Compartment Models

"Compartmental models are the most frequently used type of epidemic model. In this class of models, individuals can be in a finite number of discrete states. Some of these states are simply labels that specify the various traits of individuals. Of these, some will be changing with time, such as age class, and others will be fixed, such as sex or species. Other states indicate the progress of an infection: for example, an individual can upon becoming infected, typically first enter a state of latency, then progress to a state of infectiousness, and then lose infected status to progress to a recovered/immune state. With each state one can associate the subpopulation of individuals who are in that particular state at the given time (e.g. a female in a latent state of infection). Often the same symbol is used as a label for a state and to denote the corresponding subpopulation size, either as a fraction or as a number (e.g. I or Y for individuals in an infectious state)" - (diekmann2010?)

#### 1.2.1 The SIR compartmental model

The simplest compartmental model of infectious disease spread is the SIR Compartmental Model, with three compartments: S - for individuals *susceptible* to the disease; I - for *infected* individuals; and R - for previously infected individuals who have *recovered* (or been otherwise *removed* from that compartment). With the simplifying assumption of a constant population size, *N*, i.e.

$$N = S + I + R \tag{1.1}$$

Individuals move between compartments in a fixed set of ways, they may either become infected (moving from  $S \rightarrow I$ ), or recover from infection ( $I \rightarrow R$ ).

At each time unit an infected individual can come into contact with, on average,  $k\frac{S}{N}$  Susceptible individuals.  $\pi$  is the probability of infecting somebody on coming in contact, so  $\beta=k\pi$  is the average rate at which an infected individual will infect a susceptible. Infected individuals recover at the constant rate,  $\gamma$ , with  $1/\gamma$  the mean recovery time.

Parameter	Interpretation		
β	Transmission rate		
$\gamma$	Recovery rate		

Thus, an SIR model can be represented by the schematic

### 1.3 SIR Dynamics

#### 1.3.1 ODE representation

Given the rate parameters defined above, the change in compartment composition over time can be described by the system of differential equations

$$\frac{dS}{dt} = -\beta I \frac{S}{N} 
\frac{dI}{dt} = \beta I \frac{S}{N} - \gamma I 
\frac{dR}{dt} = \gamma I$$
(1.2)

Importantly, if the total population size is known, given Equation 1.1, R = N - S + I and the entire system can be described by two of the equations in Equation 1.2.

#### **EXAMPLE**

#### **Limitations of Deterministic ODE**

While Equation 1.2 provides a neat solution for the expected behaviour of a epidemic, it fails to caputure the variability inherent in a complex process like disease spread.

#### 1.3.2 Stochastic Process (Markov Chain) Representation

A Stochastic Process is a collection of random variables,  $X_t$ .

#### Markov chains

A Markov chain is a sequence of random variables  $X_0, X_1, \ldots$  taking values in S with the property that

$$P(X_{n+1} = j \mid X_0 = x_0, \dots, X_{n-1} = x_{n-1}, X_n = i)$$

$$= P(X_{n+1} = j \mid X_n = i),$$
(1.3)

for all  $x_0, ..., x_{n-1}, i, j \in S$ , and  $n \ge 0$ . That is, the state at the next time step is determined only by the state at the current time step.

#### **Time Homogeneity**

While not a general property of Markov chains, all of those considered in this work will have the additional property of *time homogeneity*:

$$P(X_{n+1} = j \mid X_n = i) = P(X_1 = j \mid X_0 = i)$$
,

i.e. the probabilities in Equation 1.3 do not vary with t

#### **SIR DTMC**

An SIR Compartment model an be described by a Markov chain of two independent random variables S(t) and I(t) representing the number of susceptible or individuals infected at time

t respectively (as in the ODE case, a third RV R(t) denoting the number of recovered individuals is fully determined when the population size is known, and can be left out from this characterisation).

There are two potential events (tbl-SIR\_Events) resulting in a change of state i.e. from  $X_t$  to  $X_{t+\delta t}$ 

$$p_{(s+k,i+j),(s,i)}(\Delta t) = \begin{cases} \frac{\beta is}{N} \Delta t, & (k,j) = (-1,1) \\ \gamma i \Delta t, & (k,j) = (0,-1) \\ 1 - \left[ \frac{\beta is}{N} + \gamma i \right] \Delta t, & (k,j) = (0,0) \\ 0 & otherwise \end{cases}$$
(1.4)

**Transition Matrix** 

**SIR CTMC** 

# **Chapter 2**

# **Bibliography**

Ushey, K. (2022). renv: Project environments. https://CRAN.R-project.org/package=renv R package version 0.16.0

## **Appendix A**

## **Additional stuff**

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