**Supplementary Materials – Under Ascertainment (UE in another file)**

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# Simple Parameter Estimation with SIR:

## Introduction:

*SIR model is the simplest way to model a pandemic, hence, within a constant rate of infection and the absence of governmental control over the outbreak in the initial stage, this method is useful to see the underascertainment rate.*

* + N = I + S + R
  + Goal: Estimate beta, gamma (infection rate and recovery rate, respectively)
  + As I have introduced before, the Bayesian Inference helps us to determine the proportion of population who are highly immune to the current pandemic, hence, Exposed Population Susceptible Population N (population excluded those who are immune). At the first day of pandemic (N=S):
* + - , we will find a non-linear function for computing the m:

∴ **I(t) ~ I0** (since I grows exponentially)

#*with t is time interval and I0 is the number of infection in day* 1

|  |  |
| --- | --- |
| ∴ **ln(I) = mt + ln(I0)** | **I.1** |

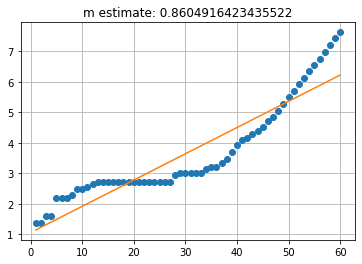
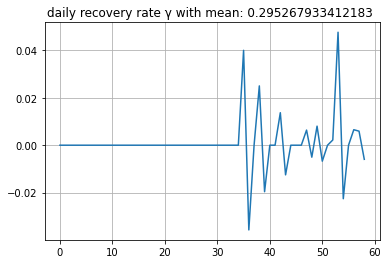
# From equation [I.1], we can easily estimate the value of m using log plot by substituting the value of In from official data then using least square method to fit the line.

**Note:** Different time intervals might produce different results of m, the higher m means the more uncontrolled pandemic was at the time t interval.

* + Then we can estimate the value of recovery rate by using 2 ways: using the number of days in infection period certified by health authorities or we can using [iii] equation to reckon the rate
    - R(t) =
    - *#I got a little concern in this part, if we account for the incubation period, it should also be: with is the length of incubation period. Let’s just stick with R(t+1) in this paper because my final outputs do not show much difference*

## Application on AUSSIE dataset:

* The scope of this research is the first 60 days of pandemic (because it is the initial stage of pandemic where number of infected = as defined by (Diekmann, 2011))

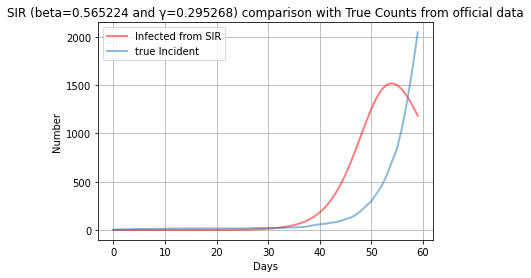


**Figure I‑1: Estimations of m and (from notebook –** [**ADD LINK HERE**](https://github.com/QUT-Trusted-Networks-Lab/n10648771-stephen/blob/main/Baseline%20Models.ipynb)**)**

Figure 1 displays that the value of m and gamma is 0.8 and 0.29 respectively. We have final table of parameter estimation for the first method

|  |  |  |
| --- | --- | --- |
| β=0.565224 | γ=0.295268 | R0 = β/ γ = 1.91427 |

* Compare with the number of true infection counts, we have:



**Figure I‑2 Compare number of infected**

## Problem of this method?

The fact that SIR is always the one to be chosen as the base model for any infectious disease simulation was due to its simplicity. However, the problem of this method is that it is a heavily data-driven technique that would be bias based on the inputs. This means that SIR only reflects and predicts based on what data we have, ignoring the reporting errors might occur during the process. Let’s look back at the equation I.1: **ln(I) = mt + ln(I0).** When we build our model based on this equation, we somehow assume that the number of cases in the first day (I0) are fully reported (which definitely is not true). Therefore, a better approach that includes anonymous estimations should be introduced.

# Model Pandemic using contact network and transmission risk in a structured population:

## Introduction

The second method is more specific and based on the result of some other peer-reviewed literature upon the rate of contact made by an individual and the risk of infectious transmission. At the effort of attaining the most reliable rate of infection which is calculated by equation (II.1), I have to assume some :

|  |  |
| --- | --- |
| **β = D \* c** | **II.1** |

The rate of infection is extremely important when analyzing an infectious disease, many models are built on the foundation of this parameter such as ODE’s models, Reed-Frost, Branching Process... In this section, I would like to use 3 approaches which are ODE’s models for structured population and spatial data, the incident model and branching process from R0

### Calculating the average number of contacts c in a structured population:

#### Data Collection Method:

From the survey conducted by European Commission, 7,290 participants recorded characteristics of 97,904 contacts with different individuals during one day, including age, sex, location, duration, frequency, and occurrence of physical contact. Respondents are demanded to fill in Table II-1 and its dataset is attached in the APPENDIX A.

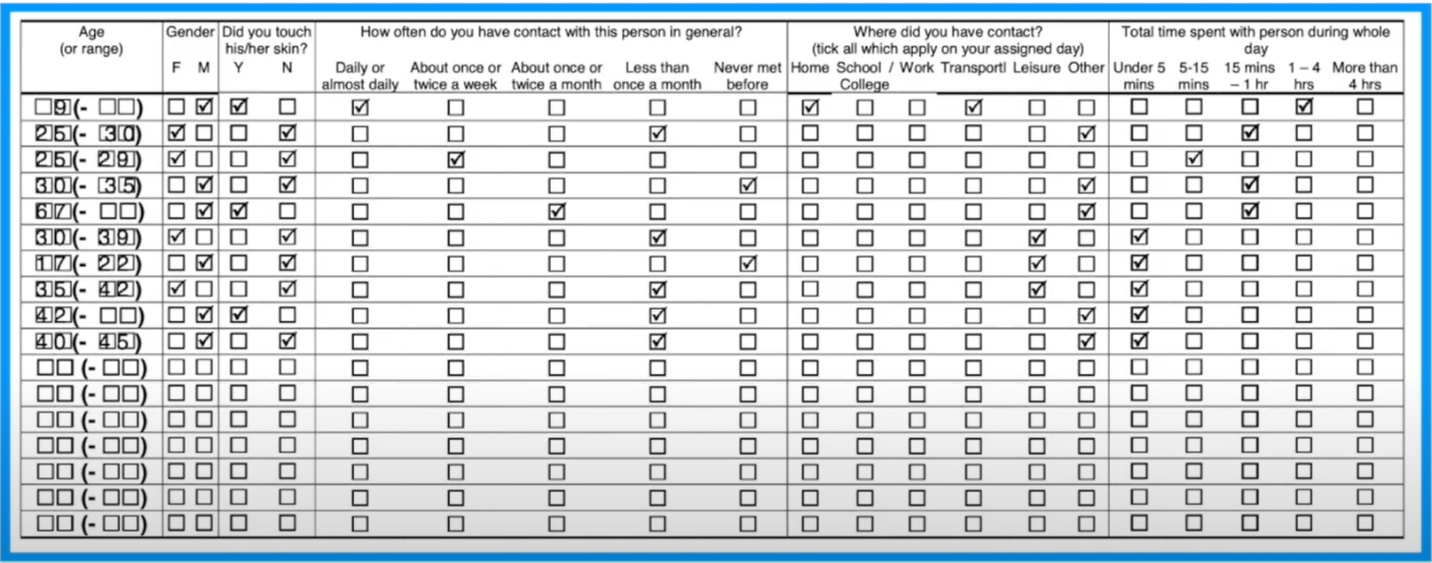


Table II‑1: Survey form recording the pattern of contact made by an individual (Source: POSMOD)

Some prominent insights of the dataset are illustrated in Table II-2. It can be instantly acknowledged that the average number of contacts are 19.6 for the whole population (aged from 0-90). However, it would be biased if we ignored assumed that all age ranges have the similar rate of contacts (in reality, infants and senior citizens might not attend public avenues as frequently as young generations since they do not work or travel). Hence, reckoning the true contact rate corresponding to respective age groups is important to identify which groups are the most infective and what measures should be imposed on them.

# The implication for this method is that if we can apply the same survey with the form given above to Australian population, we can trace their movements and interactions which can helps authorities to control the pandemic.

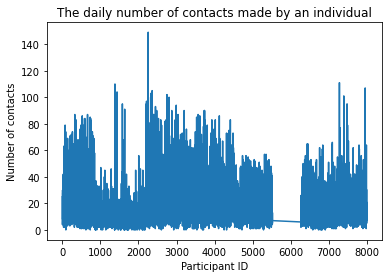
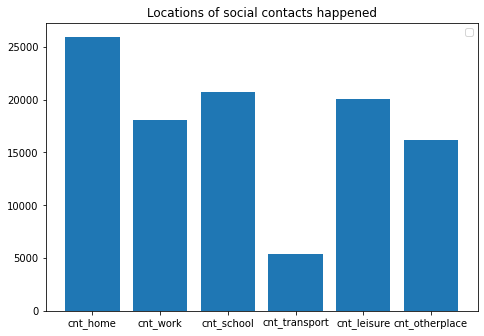


Figure II‑1: Contact pattern

#### Method:

A contacts matrix is used to account for the total number of contacts. Let Cij be the number of contacts of people in age group with age group j; Pi be The population of participants reporting their contact incidents; Ni be the total population in group i (including both participants and reported people).

The number of people in group j that an average person from age group i contact with is calculated as:

The total number of contacts made by an age group to other groups is:

Age groups of participants

Age groups of people that participants contact with

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  | 0-10 (P1) | 10-20 (P2) | ... | 80+ (Pn) |
| 0-10 | T11 | T12 | ... | . |
| 10-20 | T21 | T22 | ... | . |
| ... | ... | .. | ... | . |
| 80+ | . | . | . | . |

The limitation is that: Logically, T21  should be= T12 , however, the data from EC is just a sample of the whole population. Therefore, inconsistency in computation can be easily seen where Tij ≠ Tji. To smoothen the inconsistency, I transpose matrix A and take out the mean of 2 matrices to compute the new matrix B with a more consistent data. Hence:

#### Application:

Applying the above method to the dataset from EC (POSMOD), we retrieve the following matrice:

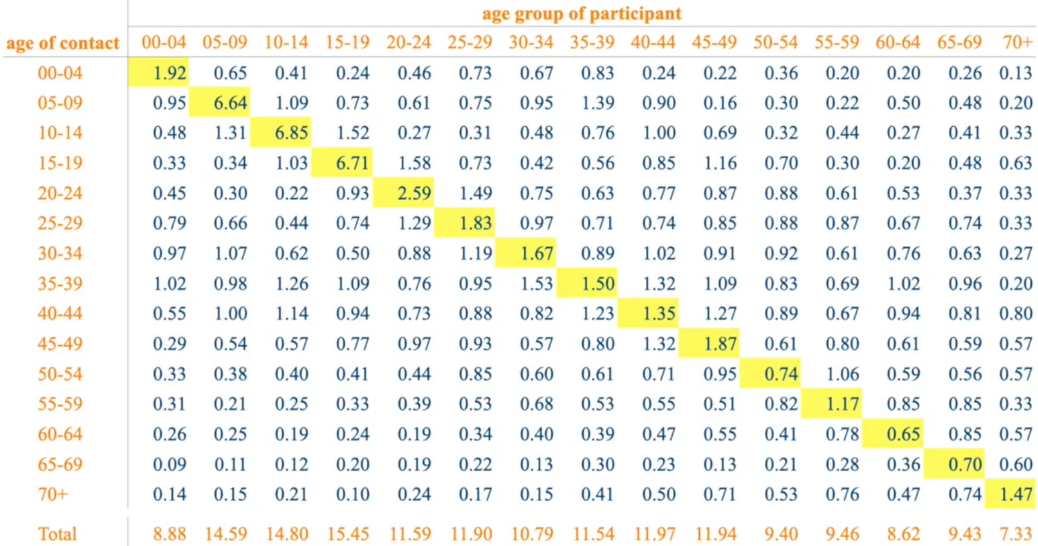


Table II‑2 Matrix of contact numbers

We can inferred from the above table that infants and old people have the lowest number of contacts while children at the primary school age have a higher chance of exposing to an infected individual.

### Estimating the risk of transmission risk by an infective individual

Transmission risk requires a strong understanding in biomedical domain to compute the desired outputs. There are several methods that are verified to be effective such as the equation of transmission risk proposed by John Hopskin Uni:

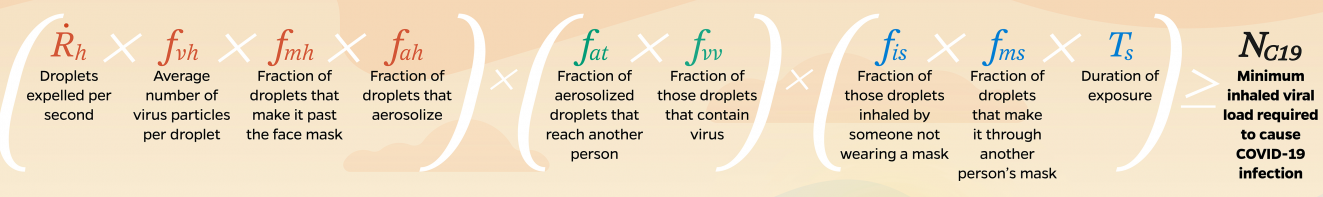


Figure II‑2 Transmission Risk Estimation Equation (source: John Hopskin, 2020)

**However,** this method is hard to apply on a larger scale as only medical specialists are capable of measuring such variables.

A simpler approach of capturing infection risks via the aerosol emission was introduced by a group of chemistry experts from Max Planck Institute that can help individuals to check the risk for themselves (see more: [link](https://www.mpic.de/4747361/risk-calculator?en) ). ***Despite of its drawbacks that many assumptions were made to perform calculations, but the result from it can be referred on a theoretical level.***

From the article, some inferences can be extracted as the infection risk of a single viral RNA is calculated as :

With x = 0.5 represents for the number of susceptible objects to be infected by taking a mean influx of RNA (Dose\_mean). The rate of getting infected by a person exposed to an infective in a close proximity is:

*R*(%) = [1 - ]× 100,

With s is the people involved in the environment – in our case, s equals to the number of contacts made by an individual, Depisode is the number of viral RNA copies inhaled by a person. It would be hard to reckon the value for such Dosemean and Depisode since I possess no relevant knowledge to bio-medical field. Therefore, it would be better to take the value from the literation which is 91.9% that a person can be infected by an infective agent if he/she shares a close contact for a duration of 2 hours. That infection rate decreases when the distance of interactions enlarges. However, upon the dataset proposed by POSMOD, the reported contacts are physical ones with varying frequencies, hence, the space of interaction can be equivalent to be in proximity with an infective host. Therefore, the transmission rate is in the range of 60% to 91.9%.

## Methods

### Incident Model:

The incident model measures the growth of infective cases in the initial phase of the pandemic. New case at time t result from contacts with individuals that were infected before t and become infectious at t (we call incubation period ). The incident i(t) is the number of new cases arising per time t will be executed in the equation:

With D = 76% (mean of the probability range in section A.2) and C is the average contact number. We can easily compute the growth rate of incident in the age-structured population, however, I will simplify the work by summing up all the cases to compare with the true counts from official data later on. Set incubation period from 4-6, we have the following model:

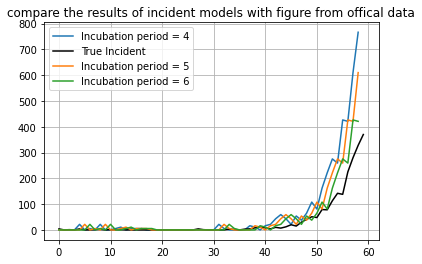


Figure II‑3 Comparisons with Incident Model

A quick analysis from the figure II-3 is that it is comprehensible that the lower incubation duration leads to the higher rate of infected cases.

### ODE SIR application for age-structured population:

The infection rate equation (II.2) will be transformed into:

***βij = D \****

cij: Contact made by an individual from group age i to group age j

The recovery rate varies upon the age groups (as we may know that older generations have lower immune system than the younger). The new variant SEIR would be as following:

***### stuck with estimation of gamma for different age group***

# Mixed model retrieve the fully ascertained number of infected:

## Introduction

All models above can help us to identify the underascertainment to some extent by simply subtracting the new outputs of each model to the original dataset. However, the bias can be clear since some of my assumptions (transmission risks, first day of infection,...) might not be correct. A mixed model of ODE with incidence function can be an option to validate the findings. I came up with a SII model which is illustrated as in the figure III-1 with S is the population of susceptible, Irep is the population of the infected who are reported and Iunrep is the population of the infective individuals who are not reported [12].

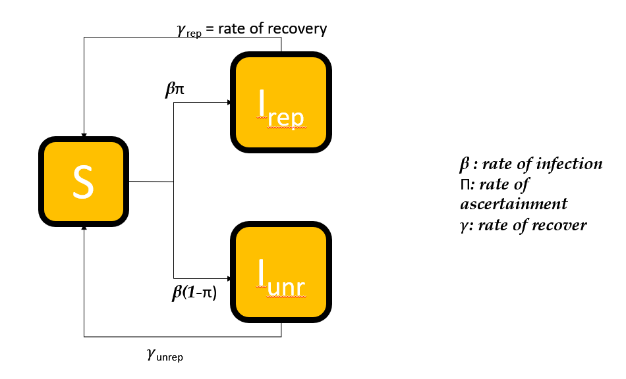


Figure III‑1 SII model for the initial phase

It is hard to estimate the value of parameter π within official available data online or some panel reports from surveillance reports. Panel data for estimating the underreporting rate is unreliable since it is a passive method that only attracts the responses from voluntary people. A marginal error of sampling from the survey data will occur and might influence the result of the study substantially. A prominent example of this issue was the presidential debate in USA between Roosevelt and Landon. Among 1.2 million people participating in an opinion poll conducted by some intelligence agencies, 43% of them voted for Roosevelt. However, at the end of the election day, President Roosevelt claimed the victory within 62% of votes. This was because they did not account for the sample error that would occur in such passive surveys (error here is that they only asked for a certain type of people about their voting opinions, many others in population might have different point of view from people they interviewed). Similarly, for infectious disease issues, the number of cases captured by official records might not reflect fully the current situation of the outbreak since:

* There might be some people who are infected but refused to visit testing clinics because they are afraid of being quarantined or the financial cost it might arise.
* There might be some people who are infective but they do not experience any symptoms of the disease (we call this the infective host or patient “0” – a term for a virus carrier who appears with perfect medical condition but can infect other people)

Therefore, parameter π can be understood as the sampling error of an observational study which should be estimated to produce the most exact number of incidents. Since the passive survey methods are ineffective and unreliable, I propose 2 active approach to resolve the issue in the Method section.

After getting the rate of overreporting(?)/underreporting, we will substitute the value of π for the following compartment model below. SII actually is an improved version of SIS (SIS can also be called SI), which is used for the type of infectious disease that possesses no certified cures (meaning that all recovered individuals will have a high chance of second attack from the pandemic

Our ODE now will be as following:

According to a peer-reviewed publication by Nishiura, H., [12], he has worked with the Travel & Tourism department of Japan to conduct a survey of under-ascertainment. In his report, 592 of 655 people from the survey showed COVID19-related symptoms and 8 of them are infected but seemed asymptomatic #review later

## Methods:

### **Method 1: Estimating the sampling error by detecting the disease as soon as possible, then integrating with SII/SIR model to retrieve the number of true counts (or even branching processes model can be used here): I0**

#### Sampling Technique:

Let D(t) denote the set of events that the disease is detected from a sample of size s. The probability of detecting disease, P(D(t) > 0), can be modeled as a binomial random variable. Since we are planning to deploy the SII technique, the proportion of the infecteds at time t is given by I(t)/S(t) (if we use SEIR then the rate would be I(t)/E(t)).

In a sample of size s, P(D(t)=0) is the complement of not detecting any infected individual at time t, and calculated as following:

At the initial stage, the ratio of I/S would be relatively minimal (<0), therefore:

We also consider the probability of detecting the disease for the first time in the kth sample when sampling occurs regularly at discrete time intervals. A model of detecting the disease in the current sampling period, but not before, is a geometric distribution with time-dependent detection probabilities (geometric distribution is to measure the chance of getting infected cases after having P(D(t)=0) of not detecting any infective individuals). Hence:

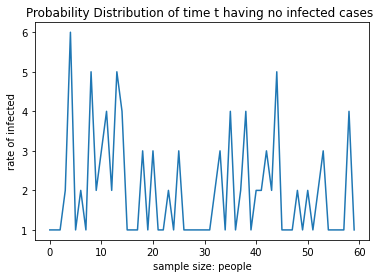
P(k) = (1-p)k-1p = (

with π is the probability of positive detection in a single pooled sample at time t, we have:

P(k) = (

* Next step: Simulate data with a pooled sample within a random π variable and constraint (a cost function display correlations between the related factors - that impact on the decision of a person reporting his/her condition or not - like quarantine, financial cost... – and the underreporting rate ) .
* To MLE the cost function, we would use Karush-Kuhn-Tucker (KKT) conditions

.... under process

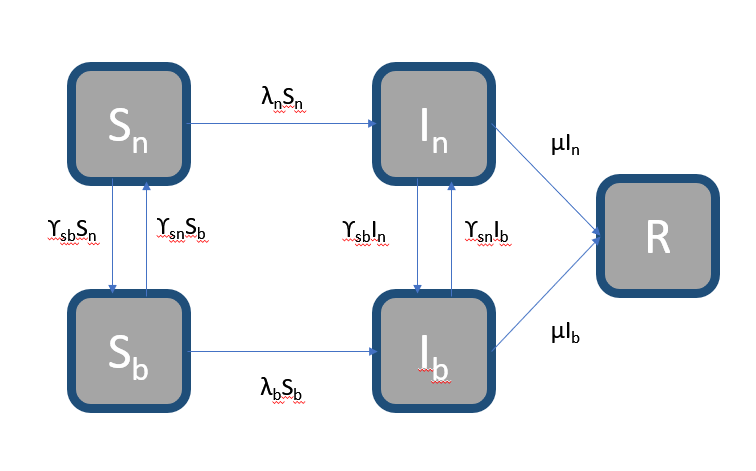


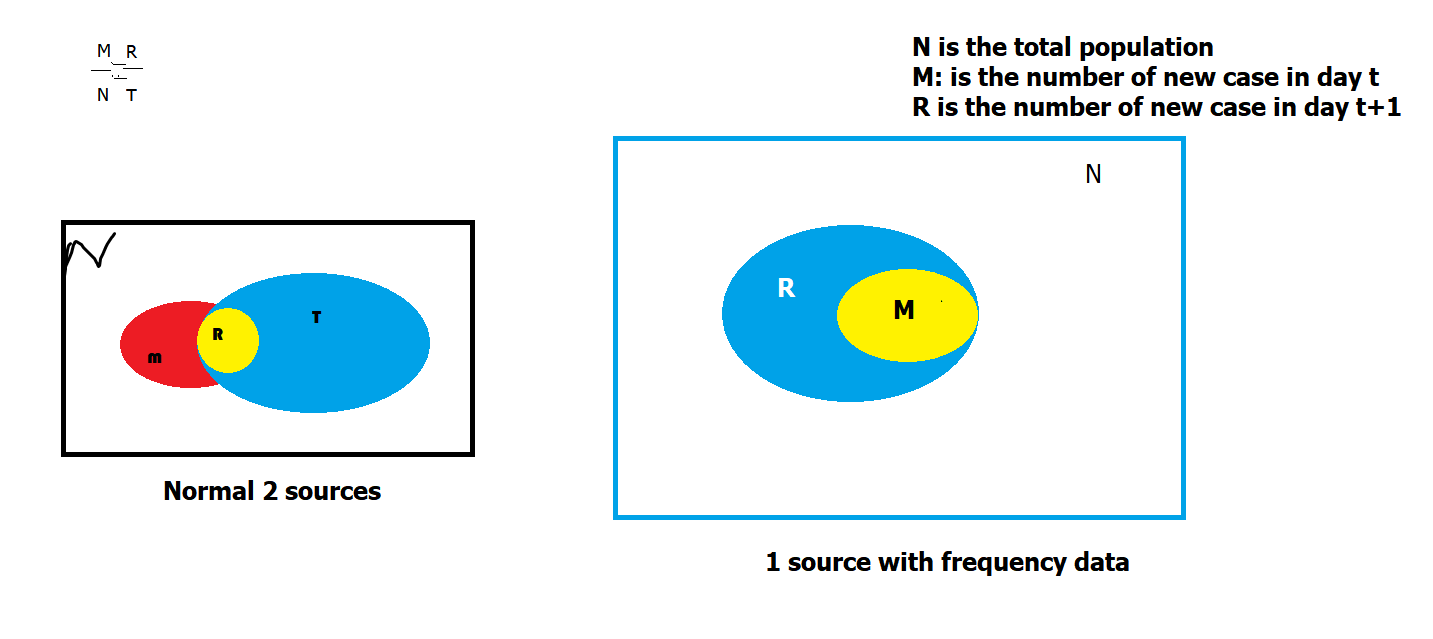
# just a visualization of what I am trying to do here.

### **Method 2: Estimating π by using likelihood based methodologies**

### In detail, we will perform an estimation of R0 then using the equilibrium equation of initial phase of the outbreak to make estimation for transmission rate and recovery rate. Finally, using SII model to estimate the whole under-ascertainment problems.

# ODE for next stage of epidemic: Social Distancing – Within results of under-ascertainment rate from (III)





Let H denote the number of hidden case in Australia, inversing the equation [] results in the following equation:

Replace M: ΔN(t-1), R: ΔN(t) – ΔD(t) :

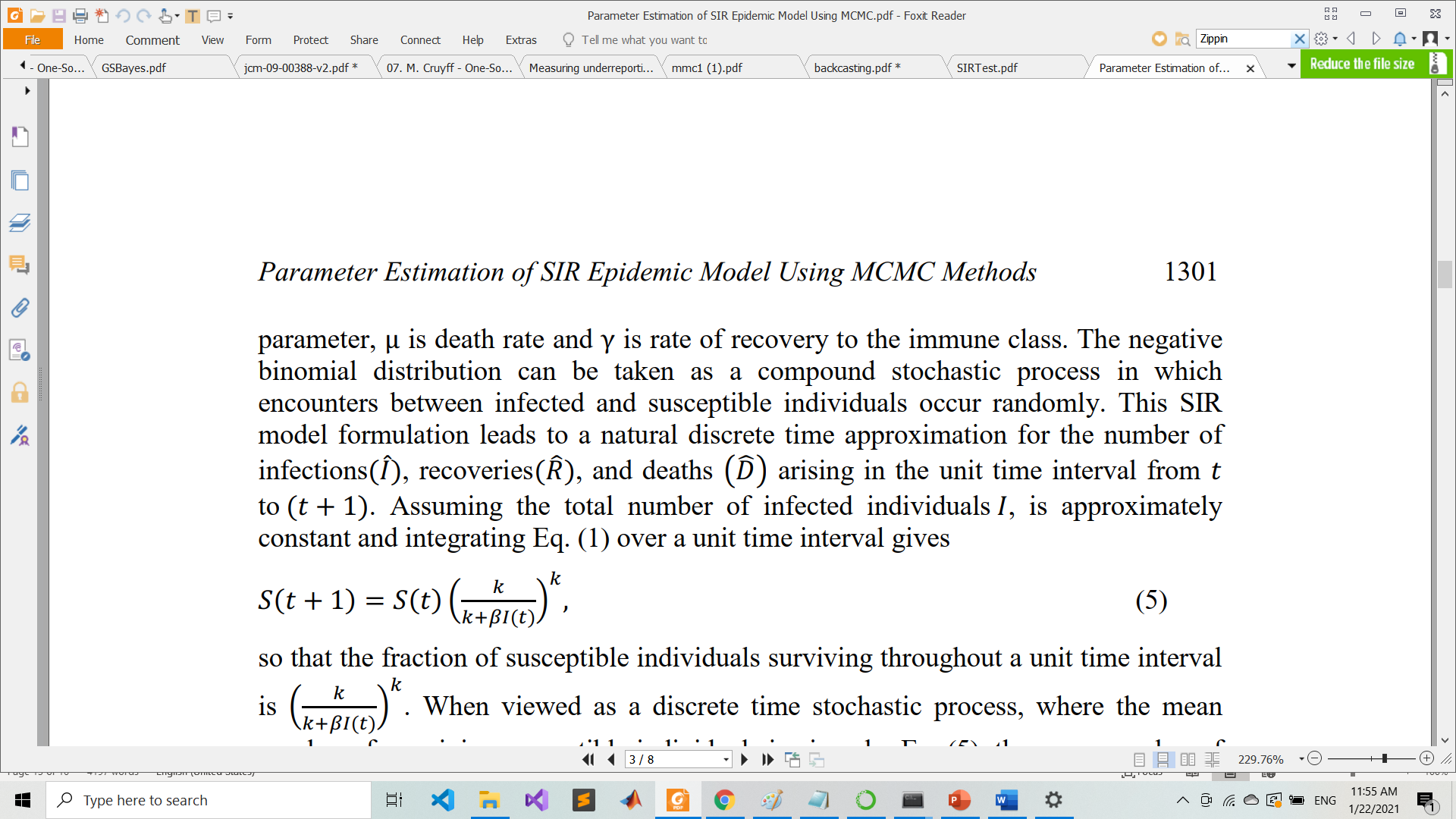
A picture containing diagram

Description automatically generated

# Monte Carlo Markov Chain (gibb sampling for μ and Metropolis-Hasting for beta)

According to Allawah et al, transmission rate is the most challenging factor to be estimated stochastically. Many bottom-up analysis are proposed to extract the most exact value for this parameter. Since most statistical methods are heavily dependent on the deterministic characteristics of the data, MCMC can be an escape from the novelty that can provide the range of possible values for the target variables.

With Parameter Estimation of SIR Epidemic Model Using MCMC Methods β is transmission rate, k is over dispersion parameter, μ is death rate. The **negative binomial distribution** can be taken as a compound stochastic process in which encounters between infected and susceptible individuals occur randomly (Markov Chain)

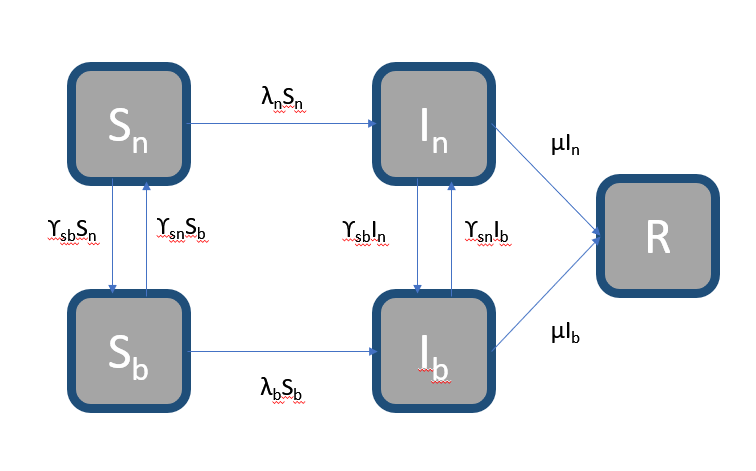


If we assume S(t) = s and I(t) = i, we may sensibly take the new infections Î at time(t + 1) to follow

P(𝐼̂|𝑠, 𝑖)~𝐵𝑖𝑛(𝑠, 𝑝𝑖(𝑖, 𝛽, 𝑘)), where 𝑝𝑖(𝑖, 𝛽, 𝑘) = 1 - ()

I used Markov Chain Monte Carlo (MCMC) [9, 10] to find the posterior distributions of 𝛽, 𝑘, 𝛾

∏𝑇 𝑡=1 𝐵𝑖𝑛(𝑖̂𝑡|𝑆(𝑡 - 1), 𝑝𝑖(𝐼(𝑡 - 1), 𝛽, 𝑘)) × ∑𝑇 𝑡=1 𝐵𝑖𝑛(𝑟̂𝑡|𝑝𝑟) × ∏𝑇 𝑡=1 𝐵𝑖𝑛(𝑑̂𝑡|𝐼(𝑡) - 𝑟̂𝑡, 𝑝𝑑)



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