

# Computational Environments and Toolchains

## Topic 04 : Implementing Models

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### Lecture 03 : SIR Models

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#### Outline

- Types of epidemic models: Component/Stochastic/Agent-based models
- SIR and related models
- Parameters dependence

# Outline

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2.2. Python Implementation	15
3. SIS	26
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4.1. Parameter Dependence	34

# Epidemic Modeling

Epidemic models describe and predict the spread of infectious diseases within a population.

## Compartamental Models

Divide the population into distinct “compartments” representing different stages of the disease. Common compartments include Susceptible, Infectious, Recovered, and sometimes Exposed or Dead.

## Stochastic models

Incorporate randomness, recognising that disease transmission is not purely deterministic and that random events (e.g., super-spreader events) can affect the course of an epidemic.

Examples: Agent-based models, Monte Carlo simulations, Markov chains.

## Network Models

Network models represent individuals as nodes and interactions as edges in a network, simulating disease spread through social or spatial contacts.

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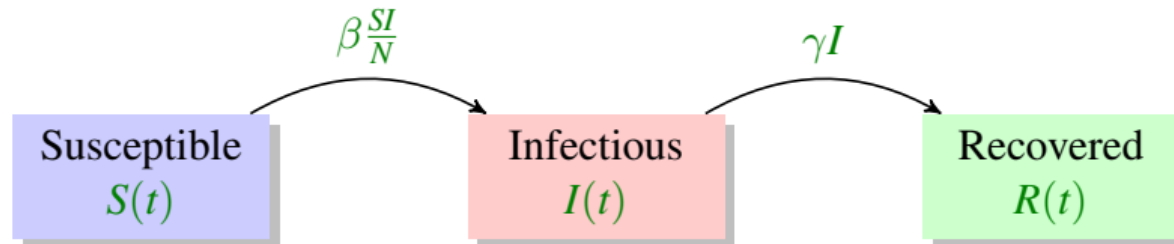
# SIR Model

The population of  $N$  individuals is divided into three compartments:

$S$  individuals susceptible to be infected;

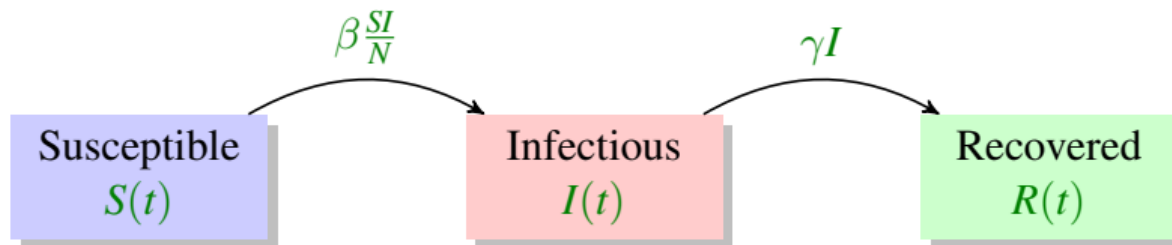
$I$  individuals infected;

$R$  individuals recovered from the disease (and now have acquired immunity to it).



- Individuals in  $S$  can become, rate  $\beta$ , infected after positive contact with an  $I$  individual. The number of contacts is  $SI/N$ .
- They develop immunity to the disease, at a  $\gamma$  cure rate, so they leave  $I$  compartment.

# SIR Model Equations



- $\beta > 0$ , the rate of contraction of the disease (transmission parameter).
- $\gamma > 0$ , mean recovery rate

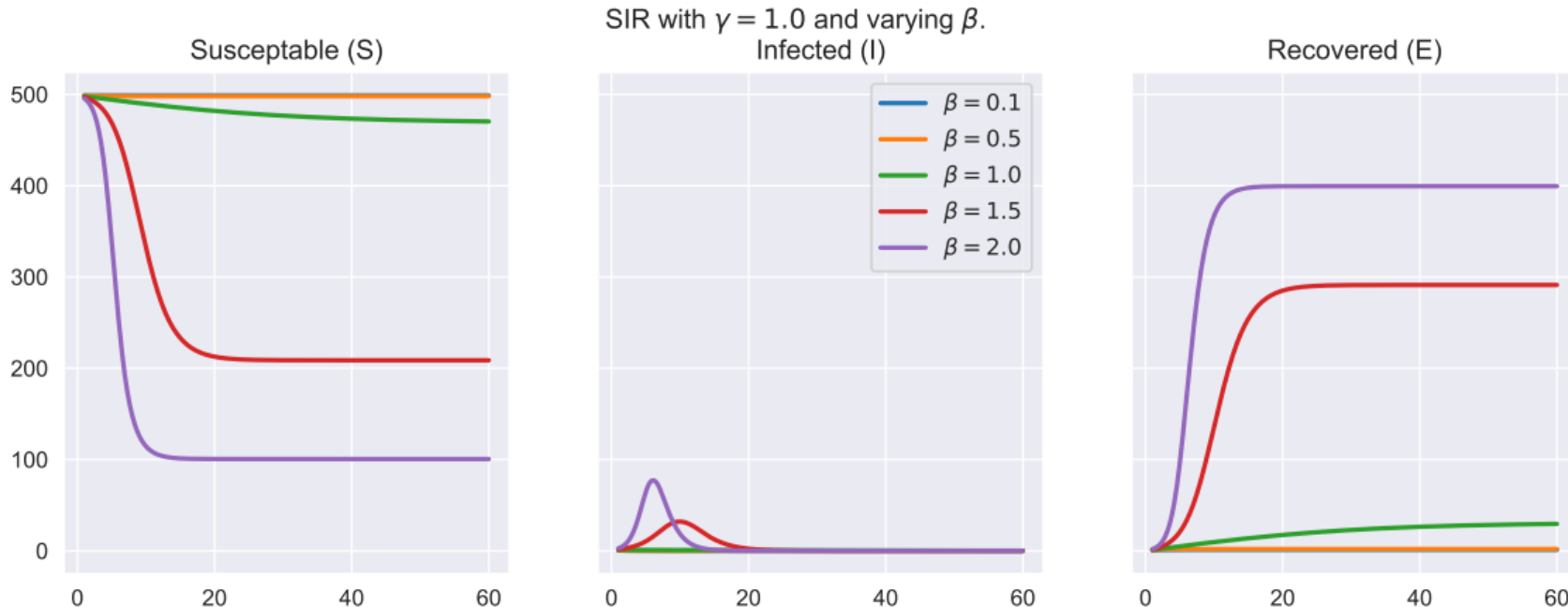
$$\left. \begin{array}{rcl} \frac{dS}{dt} & = & -\beta SI/N \\ \frac{dI}{dt} & = & \beta SI/N - \gamma I \\ \frac{dR}{dt} & = & \gamma I \end{array} \right\} \implies \begin{array}{l} N = S + I + R \\ \frac{dN}{dt} = 0 \end{array}$$

Assume disease spread is fast time scale so can ignore natural births/deaths/etc.

# Transmission rate, $\beta$

$\beta > 0$ : the rate of contraction of the disease (transmission parameter).

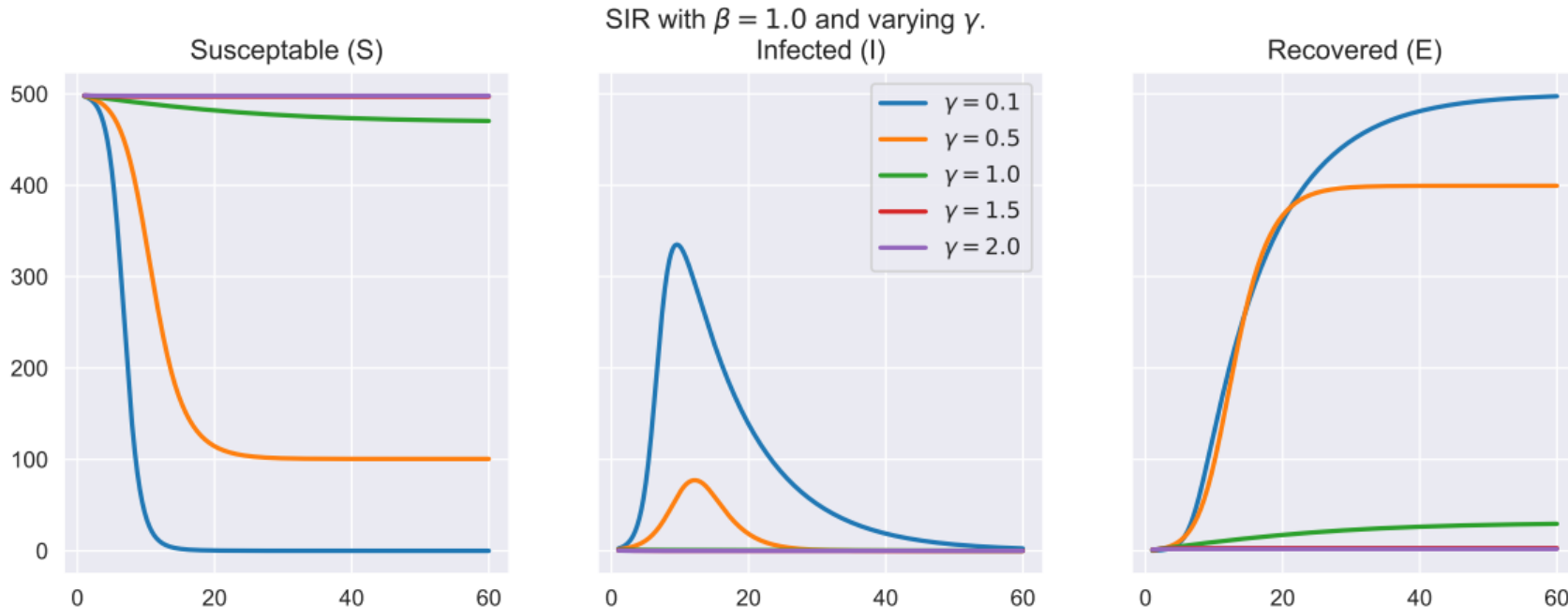
- Greater  $\beta$  reflects a more contagious infection  $\Rightarrow$  (in the limit) lower  $S$  and higher  $R$ .



# Recovery rate, $\gamma$

$\gamma > 0$ : mean recovery rate

- Greater  $\gamma$  reflects shorter infectious period  $\Rightarrow$  (in the limit) higher  $S$  and lower  $R$ .



# Basic Reproduction Number, $R_0$

$R_0$  is a critical concept in epidemiology that helps to quantify how contagious or transmissible an infectious disease is within a susceptible population. In the context of the SIR model,

$$R_0 = \frac{\beta}{\gamma}$$

is defined as the average number of secondary infections generated by a single infectious individual in a *fully susceptible population*.

The value of  $R_0$  indicates how the disease will spread in the early stages of an outbreak:

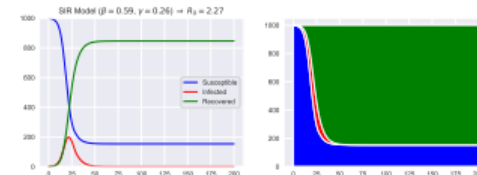
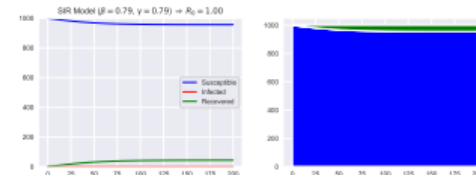
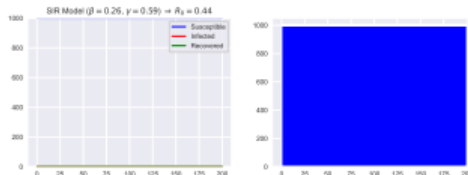
$$R_0 \begin{cases} < 1 & \text{Each infected individual infects fewer than one person on average, meaning the disease will eventually die out.} \\ = 1 & \text{The disease will be endemic, maintaining a steady state without growing or dying out.} \\ > 1 & \text{Each infected person, on average, transmits the infection to more than one other person, leading to an epidemic (the infection will spread).} \end{cases}$$

# Basic Reproduction Number, $R_0$

$$R_0 < 1$$

$$R_0 = 1$$

$$R_0 > 1$$



infects fewer than one person on average, meaning the disease will eventually die out.

disease will be endemic, maintaining a steady state without growing or dying out.

infects more than one other person, leading to an epidemic (the infection will spread).

# Basic Reproduction Number, $R_0$

## Factors Influencing $R_0$

- **Infectious period:**

Longer infectious periods (i.e., lower  $\gamma$ ) increase  $R_0$ .

- **Transmission rate:**

Higher contact rates or more effective transmission (i.e., higher  $\beta$ ) increase  $R_0$ .

## Limitations of $R_0$

- It assumes a fully susceptible population, which is rarely the case after an outbreak has begun.
- $R_0$  can vary across different environments or populations. (Think urban vs. rural.)
- Interventions like quarantine or social distancing are not reflected in the  $R_0$  value.

# Herd Immunity Threshold, $H$

The basic reproduction number also helps determine the **herd immunity threshold**:

$$H = 1 - \frac{1}{R_0}$$

This is the proportion of the population that must be immune (either through recovery or vaccination) to prevent the disease from spreading further.

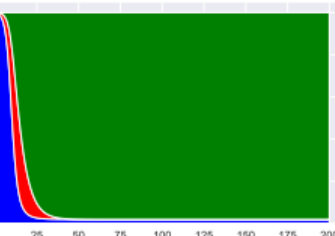
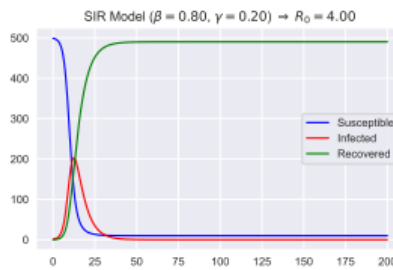
Higher  $R_0$  values require a larger immune proportion in the population to achieve herd immunity.

	$R_0$	$H$	
Measles	12–18	92%–94%	(a very high threshold, which is why high vaccination coverage is critical)
Mumps	10–12	90%–92%	
COVID-19	2–3	50%–67%	(initial estimate, current is 2.9–9.5)
Flu	$\approx 1.3$	$\approx 23\%$	
Ebola	1.3–2.0	23%–50%	

Example  $\beta = 0.8, \gamma = 0.2 \implies R_0 = 4, H = 0.75$

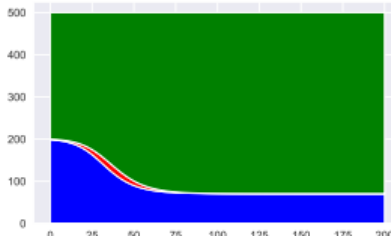
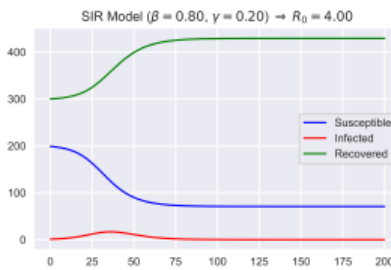
$H = 0.75\%$  so in a population of 500, this corresponds to 375 immune.

0% immune



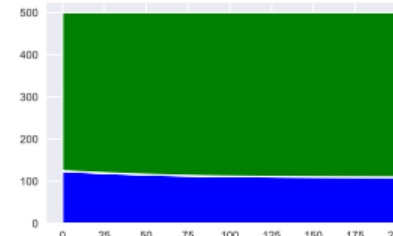
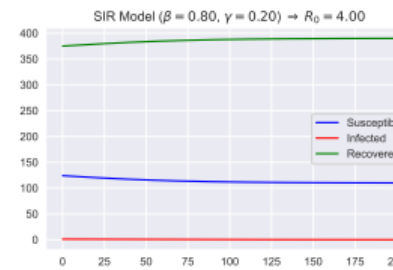
At outbreak there is no immune individuals and  $R_0 > 1$ , so disease spreads unchecked.

60% immune



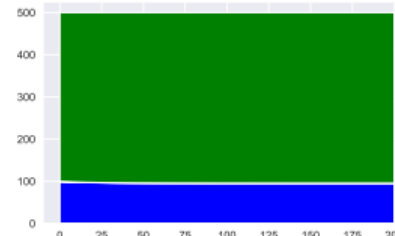
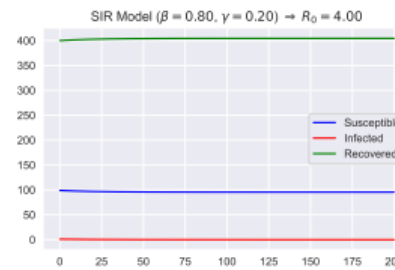
At outbreak, population has (300 individuals) 60% immune, just below threshold. Disease spread is more limited

75% immune



At outbreak, population has (375 individuals) 75% immune, at threshold. Disease outbreak does not occur (even with  $R_0 > 1$ )

80% immune



At outbreak, population has (400 individuals) 80% immune, above threshold. Disease outbreak does not occur (even with  $R_0 > 1$ )

# Achieving Herd Immunity Through Vaccination

Vaccination is the safest and most effective way to achieve herd immunity, especially for diseases with high  $R_0$ . The proportion of the population that needs to be vaccinated to achieve herd immunity is:

$$V = \frac{H}{E} = \frac{1 - \frac{1}{R_0}}{E}$$

where

$V$  is the vaccination coverage needed for herd immunity,

$E$  is the **vaccine efficacy** (the percentage of vaccinated people who actually develop immunity).

## Example

For example, if a disease has  $R_0 = 3$  and a vaccine efficacy of 90%, then:

$$V = \frac{1 - \frac{1}{3}}{0.9} \approx 74\%$$

So approximately 74% of the population would need to be vaccinated to stop the disease from spreading.

# Python Implementation — Setup

- Load our standard scientific python modules `numpy`, `matplotlib`, and `seaborn`.
- From `scipy` import `integrate` for solving differential equations and `optimise` for curve fitting/parameter estimation.
- Load `pandas` to read CSV data files.
- Local `ipywidgets` to control generation of results based on varying parameter values.

1

```
import numpy as np
import matplotlib.pyplot as plt
import seaborn as sns
sns.set_style('darkgrid')

from scipy import integrate, optimize

import pandas as pd          # to deal with CSV files
import ipywidgets as ipw      # interactive plots
```

Setup

# Python Implementation — Define ODE System

```
2 def rhs_sir(t, state, beta, gamma):  
3     S, I, R = state  
4  
5     N = S + I + R  
6     dS = - beta * S * I / N  
7     dI = + beta * S * I / N - gamma * I  
8     dR = gamma * I  
9  
10  
11  
12     beta = 0.4  
13     gamma = 0.1  
14  
15     rhs_sir(0, [1_000, 1, 0], beta, gamma)
```

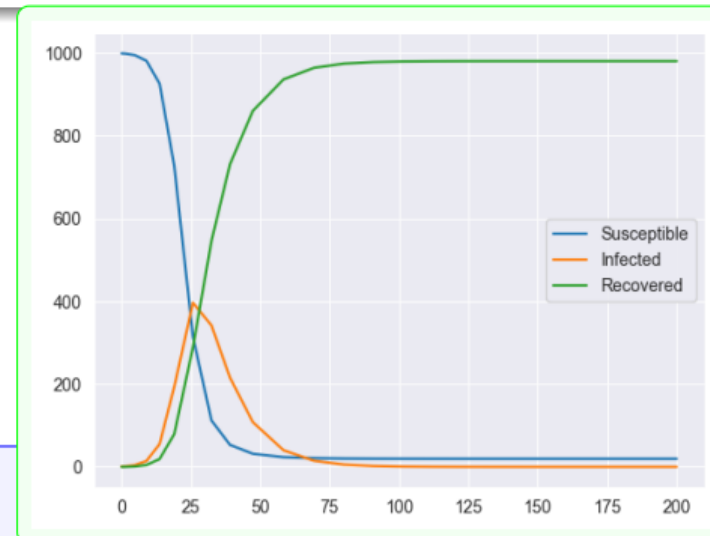
$$\begin{aligned}\frac{dS}{dt} &= -\beta SI/N \\ \frac{dI}{dt} &= \beta SI/N - \gamma I \\ \frac{dR}{dt} &= \gamma I\end{aligned}$$

RHS

# Python Implementation — Integrate

- We will use `solve_ivp` instead of the older `odeint` to integrate an ODE.
- Unlike `odeint`, we only specify the independent variable span (min, max), and `solve_ivp` picks enough points to get solution to sufficient accuracy.
- But visually we might want more points.

```
3  
1 T_MAX = 200  
2 N = 1_000  
3 I0 = 1  
4 ic = [N-I0, I0, 0]  
5  
6  
7 sol = integrate.solve_ivp(rhs_sir, (0,T_MAX), ic, args=(beta, gamma))  
8 t = sol.t  
9 S, I, R = sol.y
```



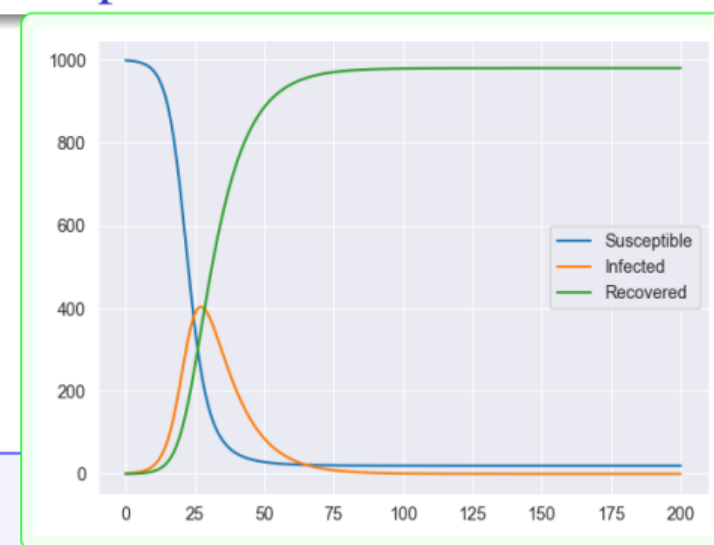
# Python Implementation — Integrate with Interpolation

- To get smoother plots, we use optional term `t_eval` to specify at which points to generate the state.
- This parameter will also be useful when comparing solutions from multiple calls to `solve_ivp`.
- Use `np.linspace`

```

1 T_MAX = 200
2 N = 1_000
3 I0 = 1
4 ic = [N-I0, I0, 0]
5
6 t_eval = np.linspace(0, T_MAX, 1000)
7 sol = integrate.solve_ivp(rhs_sir, (0,T_MAX), ic, args=(beta, gamma), t_eval=t_eval)
8 t = sol.t
9 S, I, R = sol.y

```



# Python Implementation — Visualisation

```
5
1 def update_sir(beta=0.2, gamma=0.1):
2
3     sol = integrate.solve_ivp(rhs_sir, (0,T_MAX), ic, args=(beta, gamma), t_eval=t_eval)
4     t, (S, I, R) = sol.t, sol.y
5     N = sum(ic)
6
7     fig = plt.figure(2, figsize=(12,4))
8     axs = fig.subplots(1, 2, sharex=0)
9
10    colors=['blue', 'red', 'green']
11    axs[0].plot(t, S, colors[0], label = "Susceptible")
12    axs[0].plot(t, I, colors[1], label = 'Infected')
13    axs[0].plot(t, R, colors[2], label = 'Recovered')
14    axs[0].legend()
15
16    axs[1].stackplot(t, S, I, R, colors=colors)
17    R_0 = np.nan if gamma==0 else beta/gamma
18    axs[0].set_title(r"SIR Model ($\beta=% .2f$, $\gamma=% .2f$) $\Rightarrow$ $R_0=% .2f$"
19                      % (beta,gamma,R_0))
20    plt.show()
```

Implement a function to draw  
the required plots

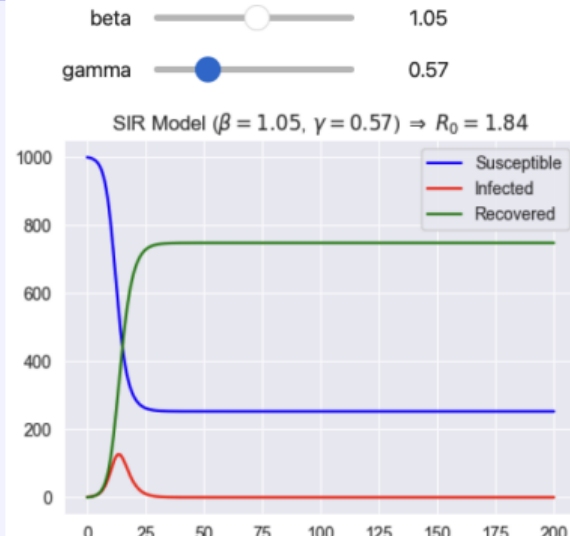
# Python Implementation — Interactive

We can control values of parameters  $\beta$  and  $\gamma$  using sliders ...

```

6 N = 1_000
1 I0 = 1
2 ic = [N-I0, I0, 0]
3 t_eval = np.linspace(0, T_MAX, 1000)
4
5
6 interactive_plot = ipw.interactive(update_sir,
7     beta=ipw.FloatSlider(0.0, min=0.0, max=2.0, step=0.01),
8     gamma=ipw.FloatSlider(0.0, min=0.0, max=2.0, step=0.01),
9 )
10 output = interactive_plot.children[-1]
11 output.layout.height = '350px'
12 interactive_plot

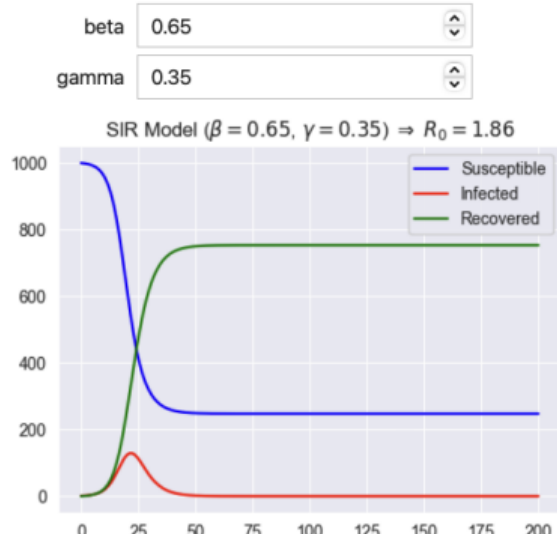
```



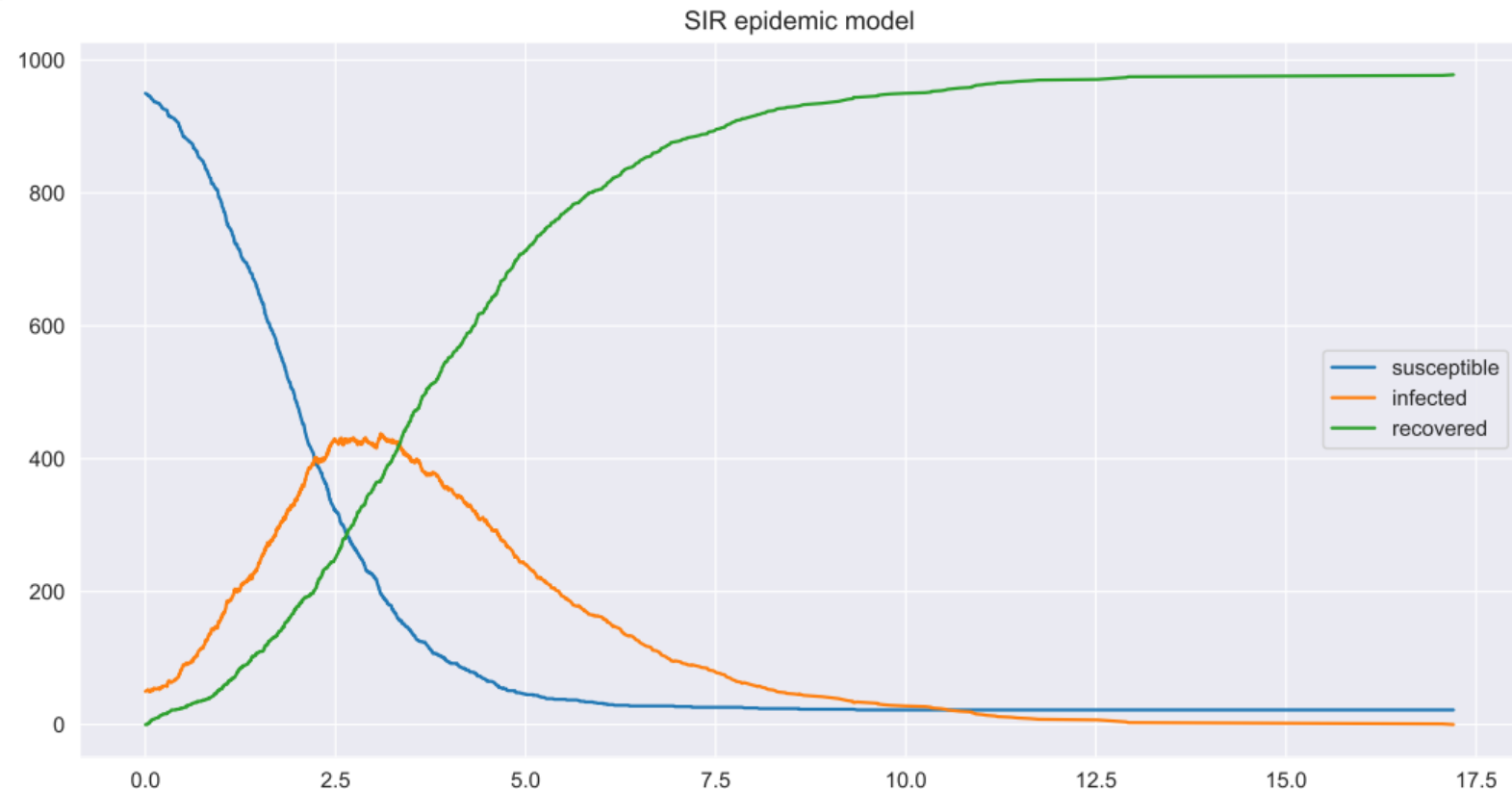
# Python Implementation — Interactive

Colab can be too slow so switch to `FloatText` widgets ...

```
7 N = 1_000
1 I0 = 1
3 ic = [N-I0, I0, 0]
4 t_eval = np.linspace(0, T_MAX, 1000)
5
6 interactive_plot = ipw.interactive(update_sir,
    beta=ipw.FloatText(value=0.0, step=0.05),
    gamma=ipw.FloatText(value=0.0, step=0.05),
)
10 output = interactive_plot.children[-1]
11 output.layout.height = '350px'
12 interactive_plot
```



# Python Implementation — Parameter Fitting



# Python Implementation — Parameter Fitting

Load Data ...

```
8 df = pd.read_csv("data.csv")
1 df.head()
```

Extract *t* and *y* values...

```
9 df = pd.read_csv("data.csv")
1
2
3 t_obs = df.t.values
4
5 y_obs = df[ ['S', 'I', 'R'] ].values
6 y_obs
```

```
array([[950, 50,  0],
       [949, 51,  0],
       [948, 52,  0],
       ...,
       [ 22,  2, 976],
       [ 22,  1, 977],
       [ 22,  0, 978]])
```

	<b>t</b>	<b>S</b>	<b>I</b>	<b>R</b>
<b>0</b>	0.000000	950	50	0
<b>1</b>	0.018729	949	51	0
<b>2</b>	0.024190	948	52	0
<b>3</b>	0.030419	948	51	1
<b>4</b>	0.031453	947	52	1

# Python Implementation — Parameter Fitting

- Use `optimize.minimize` instead of `optimise.curve_fit` as have more control over objective function.
- We want to minimise, the sum of squares of differences

$$\sum_k \left[ y_{\text{model},k} - y_{\text{obs},k} \right]^2$$

10

```
ic = y_obs[0]
T_MAX = t_obs[-1]

def f(x):
    beta, gamma = x

    sol = integrate.solve_ivp(rhs_sir, (0,T_MAX), ic, args=(beta, gamma), t_eval=t_obs)
    return np.linalg.norm(sol.y - y_obs.T)

sol = optimize.minimize(f, [0.5, 0.5], bounds=((0,3),(0,3)) )
sol
```

message: CONVERGENCE: REL\_REDUCTION\_OF\_F\_<=\_FACTR\*EPSMCH  
 success: **True**  
 status: 0  
 fun: 754.0829736360038  
 x: [ 1.923e+00 4.758e-01]  
 nit: 9  
 jac: [-2.246e-01 -1.756e-01]  
 nfev: 45  
 njev: 15  
 hess\_inv: <2x2 LbfgsInvHessProduct with dtype=float64>

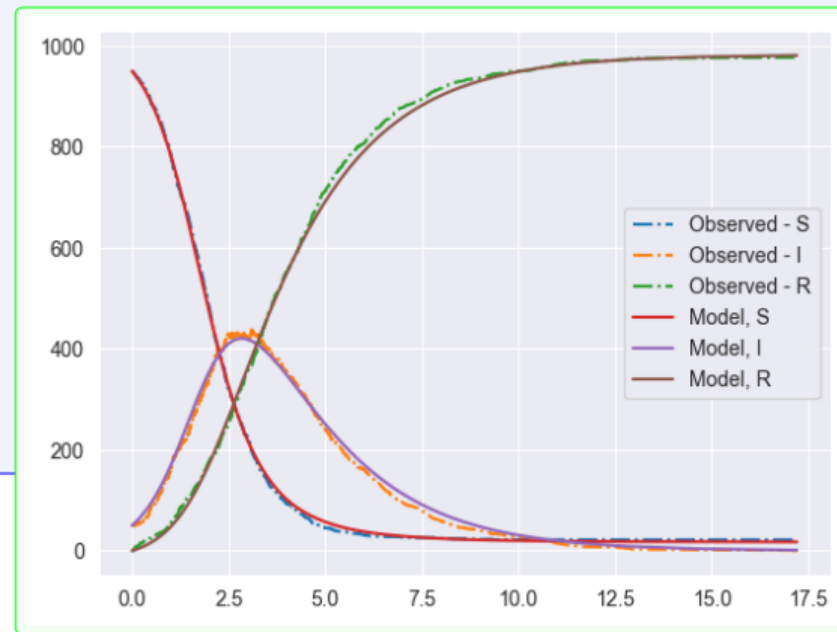
# Python Implementation — Parameter Fitting

11

```
sol_ivp = integrate.solve_ivp(rhs_sir, (0,t_obs[-1]), ic, args=sol.x, t_eval=t_obs)
t = sol_ivp.t

plt.plot(t, df.S, '_.', label='Observed - S')
plt.plot(t, df.I, '_.', label='Observed - I')
plt.plot(t, df.R, '_.', label='Observed - R')

plt.plot(t, sol_ivp.y[0], label='Model, S')
plt.plot(t, sol_ivp.y[1], label='Model, I')
plt.plot(t, sol_ivp.y[2], label='Model, R')
plt.legend()
plt.show()
```



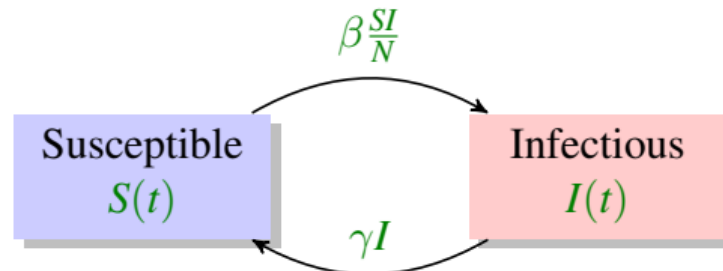
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# SIS Model

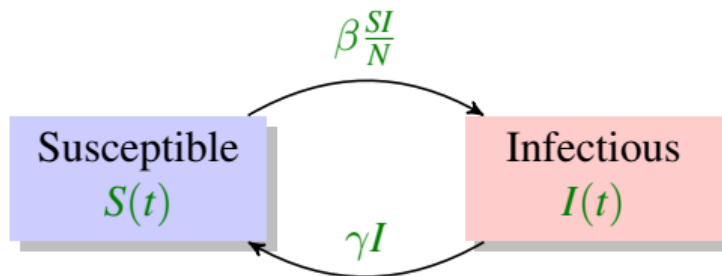
The SIS model is a compartmental model in epidemiology similar to the SIR model, but it's designed to describe diseases where individuals do not acquire long-term immunity after infection, such as the common cold and the flu.

In the SIS model, individuals move between the Susceptible (S) and Infected (I) states, without transitioning to a recovered state. After recovery, individuals return to the susceptible pool, capable of being reinfected.



- Susceptible individuals can become, rate  $\beta$ , infected after positive contact with an  $I$  individual. The number of contacts is  $SI/N$ .
- Infected individuals recover, at rate  $\gamma$ , and can be reinfected.

# SIS Model Equations



- $\beta > 0$ , the rate of contraction of the disease (transmission parameter).
- $\gamma > 0$ , mean recovery rate

$$\left. \begin{array}{rcl} \frac{dS}{dt} & = & -\beta SI/N + \gamma I \\ \frac{dI}{dt} & = & \beta SI/N - \gamma I \end{array} \right\} \implies \begin{array}{l} N = S + I \\ \frac{dN}{dt} = 0 \end{array}$$

Assume disease spread is fast time scale so can ignore natural births/deaths/etc.

# Basic Reproduction Number, $R_0$

The basic reproduction number,  $R_0$ , in the SIS model is calculated similarly to the SIR model:

$$R_0 = \frac{\beta}{\gamma}$$

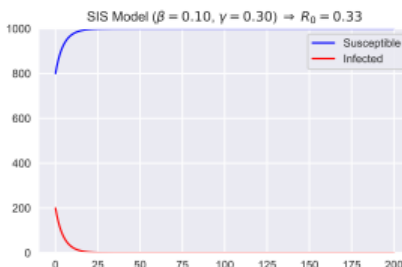
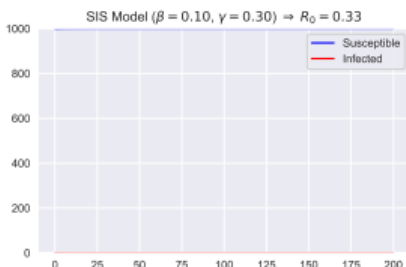
The value of  $R_0$  determines how the disease will spread:

$$R_0 \left\{ \begin{array}{ll} \leq 1 & \text{The infection dies out over time, and eventually, there will be no infectious individuals in the population.} \\ & \lim_{t \rightarrow \infty} I(t) = 0 \\ > 1 & \text{The infection will spread and reach an endemic equilibrium, where a constant proportion of the population remains infected.} \\ & \lim_{t \rightarrow \infty} I(t) = \left(1 - \frac{1}{R_0}\right) N \end{array} \right.$$

# Basic Reproduction Number, $R_0$

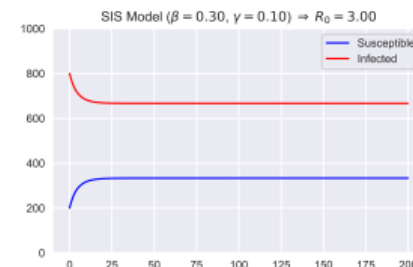
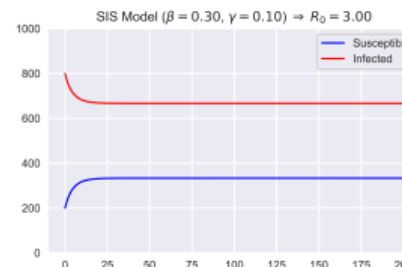
$$\longleftrightarrow R_0 \leq 1 \longleftrightarrow$$

$$I_0 = 1$$



$$\longleftrightarrow R_0 > 1 \longleftrightarrow$$

$$I_0 = 1$$



infects fewer than one person on average, meaning the disease will eventually die out.

the disease will eventually die out, even if initial pocket of infected

Number of infected trends towards limit of  $(1 - 1/R_0) N$ .

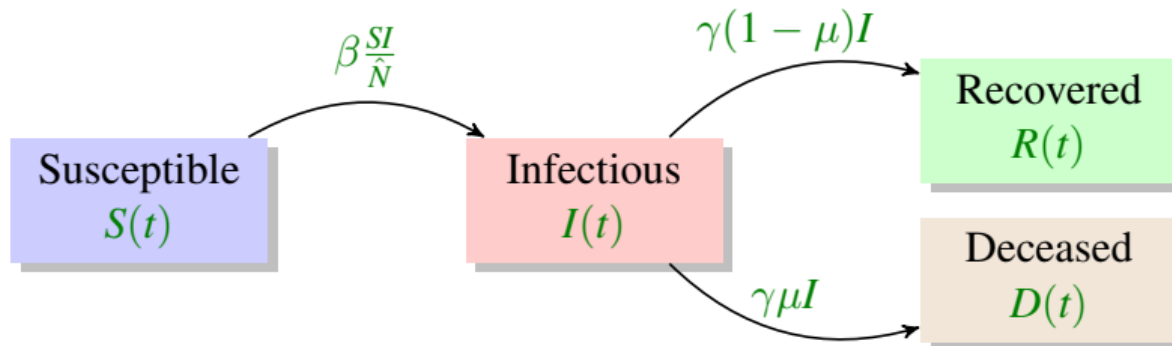
Number of infected trends towards limit of  $(1 - 1/R_0) N$ .

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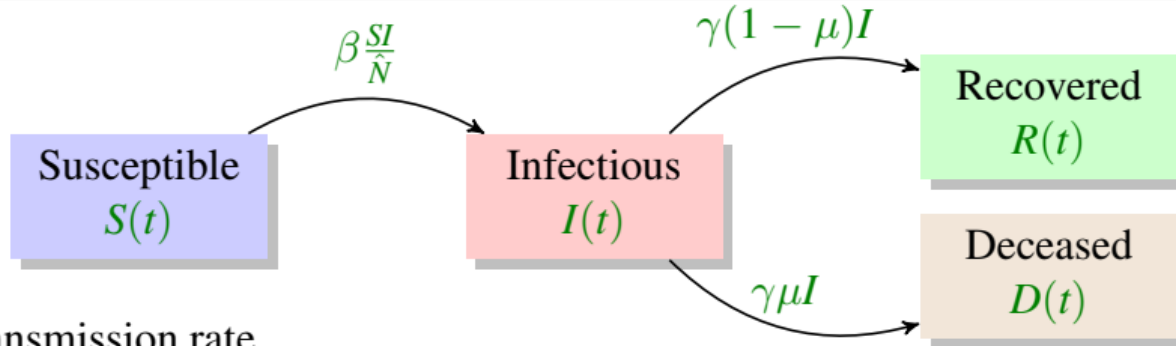
# SIRD Model

The SIRD model is an extension of the basic SIR model that incorporates an additional Death (D) compartment to represent individuals who die from the disease. This modification makes the SIRD model more realistic for diseases with significant mortality rates, such as COVID-19, Ebola, and other severe infections.



- Susceptible individuals can become, rate  $\beta$ , infected after positive contact with an  $I$  individual. The number of contacts is  $SI/\hat{N}$ , where  $\hat{N} = S + I + R$ , i.e. no necrophilism.
- Infected individuals can recover and develop immunity to the disease, at rate  $\gamma$  and move to Recovered, or die at a rate  $\mu$  and move to Deceased.

# SIRD Model Equations



- $\beta > 0$ , transmission rate.
- $\gamma(1 - \mu) > 0$ , mean recovery rate
- $\gamma\mu > 0$ , mean mortality rate

$$\left. \begin{array}{rcl} \frac{dS}{dt} & = & -\beta SI/\hat{N} \\ \frac{dI}{dt} & = & \beta SI/\hat{N} - \gamma I \\ \frac{dR}{dt} & = & \gamma(1 - \mu)I \\ \frac{dD}{dt} & = & \gamma\mu I \end{array} \right\} \quad \begin{aligned} N &= S + I + R + D \\ \frac{dN}{dt} &= 0 \\ \hat{N} &= S + I + R \end{aligned}$$

Assume disease spread is fast time scale so can ignore natural births/deaths/etc.

# Basic Reproduction Number, $R_0$

The basic reproduction number,  $R_0$ , in the SIRD model is:

$$R_0 = \frac{\beta}{\gamma}$$

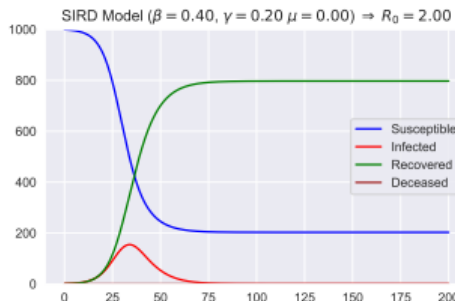
The value of  $R_0$  indicates how the disease will spread in the early stages of an outbreak:

$$R_0 \begin{cases} < 1 & \text{Each infected individual infects fewer than one person on average, meaning the disease will eventually die out.} \\ \geq 1 & \text{Each infected person, on average, transmits the infection to more than one other person, leading to an epidemic (the infection will spread).} \end{cases}$$

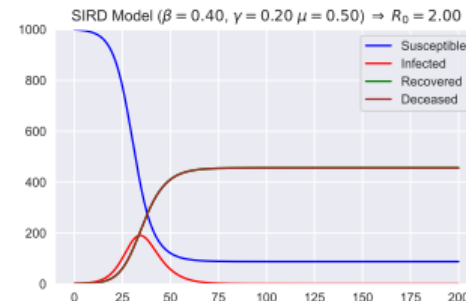
# Effect of varying $\mu$ .

$$\longleftrightarrow \beta = 0.4, \gamma = 0.2 \implies R_0 = 2 \longleftrightarrow$$

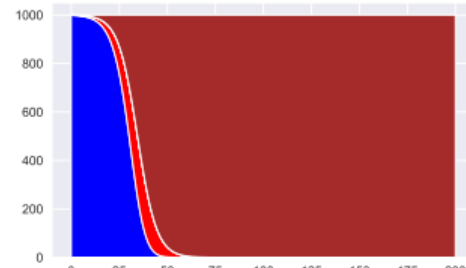
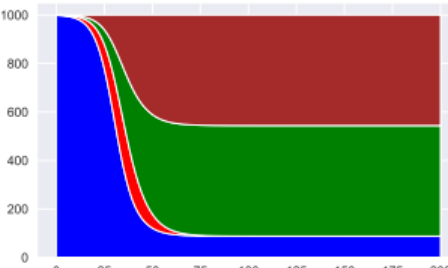
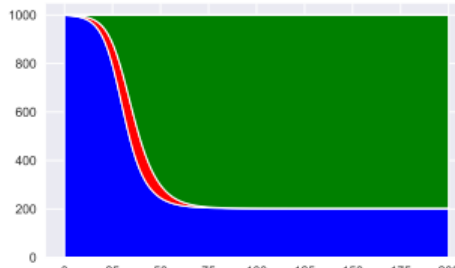
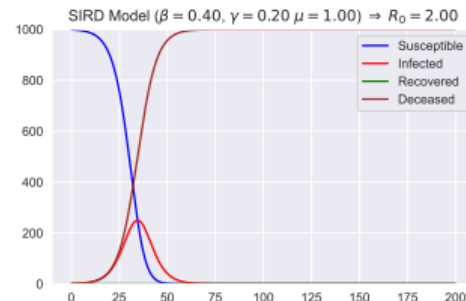
$\mu = 0.0$



$\mu = 0.5$

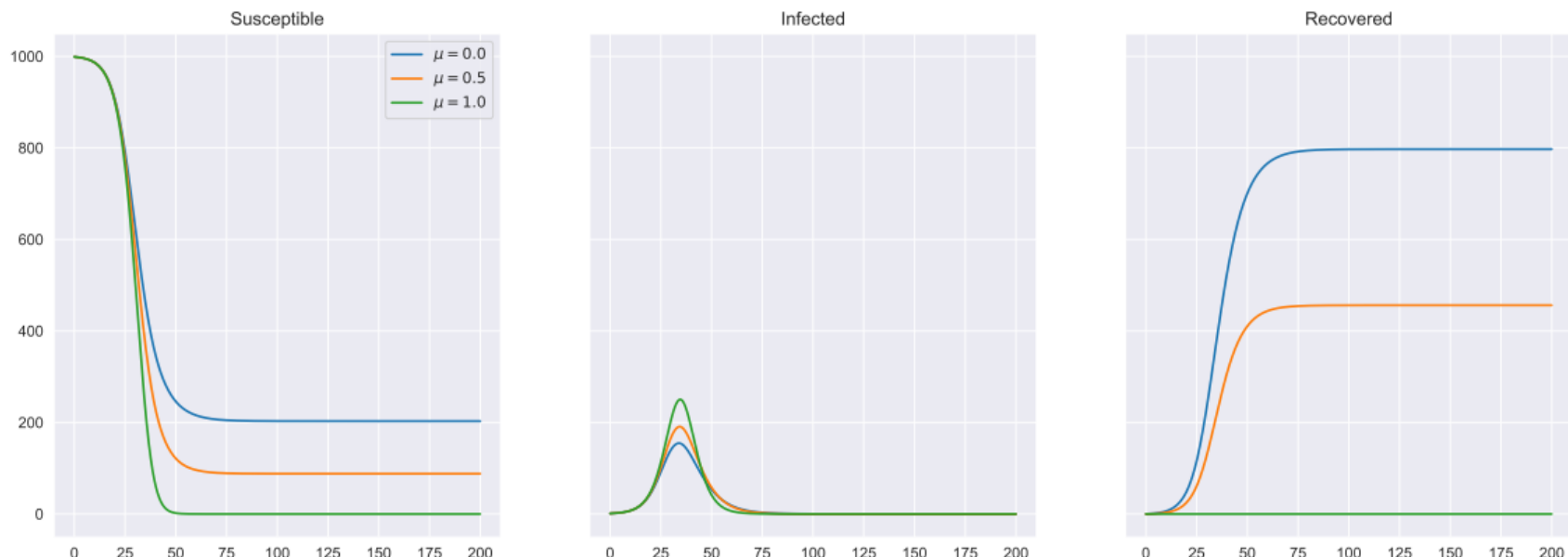


$\mu = 1.0$



# Effect of varying $\mu$ .

Effect of mortality rate on S, I, and R ( $\beta = 0.4$   $\gamma = 0.2$ ).



S and R behave as expected under varying  $\mu$ , but why does the infection spike higher and then decay faster as  $\mu$  increases?

# 1971 Influenza Outbreaks in Tristan da Cunha

