

# Model Documentation - Simple Network Sim

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## 1 Overview

Simple network sim (sns) is network-based simulation model built for COVID-19 and written in Python.

Each node in an edge-weighted network contains an age-structured population and fractional people within that population are recorded as in various disease states. We assume that the temporal scale of the simulation is such that individuals do not progress between age classes, nor are any new individuals introduced. These nodes might be used to model e.g. health boards, counties, or datazones.

After seeding disease at specified nodes, sns simulates progression of an outbreak due to both within- and between-node infectious contacts. Infection and disease progression can be run either deterministically (resulting in fractional people in each disease state) or stochastically (resulting in integer people in each nodes and disease state). The end result of the model is a timeseries of the number of people in each node, compartment and age.

## 2 Inputs and Notation

Data, parameters, and the compartmental model to be used are all passed to sns as inputs and accessed via the SCRC Data Pipeline API.

- Disease-specific
  - compartmental model
  - infectious classes
  - transition rates
  - infection probability of a contact
- Network characteristics
  - weighted edges

- Within-node information
  - age-structured population at nodes
  - mixing between age classes at nodes
- Time varying-modifiers
  - movement modifiers
  - contact rate modifiers

## 2.1 Notation

Related both to the inputs and the internal state of the model at each timestep, we define some notation for convenience.

- $w(u, v)$  is the number of expected contacts from node  $u$  to node  $v$ , supplied as the weighted edges input file
- $p$  the infection probability of a contact between an infectious person and a susceptible person (unique value for all age groups, nodes and times)
- $x_t$  is the (optional) movement modifier at time  $t$ , supplied as part of an input file
- $y_t$  is the (optional) contact modifier at time  $t$ , supplied as part of an input file
- $A_{u,k}$  is age class  $k$  at node  $u$ . Where it is clear that we are talking about individuals in all nodes we may abuse notation and use  $A_k$ , and when talking about individuals in a node of all ages, we use  $A_u$ .
- $X_t(A_{u,k})$  where  $X$  is a compartment in the compartmental model is the number of individuals in compartment  $X$  in age class  $A_k$  in node  $u$  at time  $t$
- $N_t(A_{u,k})$  is the total number of individuals in age class  $A_k$  in node  $u$  at time  $t$
- For convenience, we also define  $\mathcal{I}_t(A_{u,k})$  as the number of individuals in *any* infectious compartment in age class  $A_k$  in node  $u$  at time  $t$
- Similarly for convenience, we define as  $p_{t,\mathcal{I}}(A_{u,k}) = \frac{\mathcal{I}_t(A_{u,k})}{N_t(A_{u,k})}$  the proportion of age class  $A_k$  in node  $u$  at time  $t$  that is infectious, and as  $p_{t,S}(A_{u,k}) = \frac{S_t(A_{u,k})}{N_t(A_{u,k})}$  the proportion of age class  $A_k$  in node  $u$  at time  $t$  that is susceptible
- $C$  is a matrix describing contact between age classes, where  $C_{i,j}$  is the expected number of contacts that an individual in age class  $A_i$  has with an individual in age class  $A_j$ .

### 3 Algorithm of simulation

Simple network sim uses a notionally daily timestep, and forward-simulates by performing local and between-region infection followed by disease progression updating based on rates supplied as part of the compartmental model. The simulation can run deterministically using point estimates of rates and infection, or stochastically by drawing individuals to infect or progress from the population.

#### 3.1 Central infection mechanism: infectious contacts

The central mechanism of infection uses the notion of a number of infectious contacts that are distributed amongst the population in a node, structured by age class. A single infectious contact allocated to a node is a contact that is between a person in an infectious class either within the node or from another node and a person who is susceptible within the node. In addition to age-structured within-node infection, we take the total number of all of the infectious contacts exerted from other nodes, and then using that total number distribute the infectious contacts to a number of target individuals within the node using either a fixed combinatorial expression or in the stochastic case a hypergeometric distribution to draw (with replacement) the number of susceptible people who are subject to those infectious contacts, subject to a specified probability of transmission given a single infectious contact. This takes account of the possibility of a single susceptible individual in a node being the recipient of multiple infectious contacts. All infectious classes are considered to have the same infectivity.

##### 3.1.1 Calculating the number of within-node infectious contacts

Within node, the number of infectious contacts targeting susceptible individuals in age class  $A_i$  at time  $t$  is a function of the proportion of infectious individuals in each age class within the node and the expected number of contacts between an individual in  $A_i$  and individuals in each of the source age classes at time  $t - 1$ .

Within node  $u$ , for a single source age class  $A_j$  where  $C_{i,j}$  is the number of contacts that an individual in  $A_i$  expects to have with individuals in  $A_j$ ,  $y_{t-1}$  is the time-varying contact modifier (defaulting to 1.0 if none is supplied), then the number of within-node infectious contacts from  $A_j$  to  $A_i$  is defined as follows:

##### Deterministic mode

The number of infectious contacts is given by:

$$y_{t-1} C_{i,j} p_{t-1,\mathcal{I}}(A_{u,j}) p_{t-1,S}(A_{u,i})$$

##### Stochastic mode

For each infected person in  $A_j$ , we randomly sample from a Poisson distribution,  $\mathcal{P}(y_{t-1} C_{i,j})$

to generate how many contacts this person had with people in  $A_i$ . The average of the Poisson distribution will be the same value as the one in the deterministic model. Then for each infectious person, and it's random number of contacts *contacts*, we randomly sample from a hypergeometric distribution (the hypergeometric distribution is used instead of binomial because we assume the contacts are with unique people, hence the random sampling must be done without replacement), with parameters  $Hypergeometric(M = N_t(A_{u,i}), n = S_t(A_{u,i}), N = \text{contacts})$ , i.e. we sample *contact* times in a the total population of node  $u$ , with a "success" being a susceptible person. These infectious contacts for all people are then summed.

Remarks:

- This is a compound probability system, and it allows retaining individual level heterogeneity even with group level parameters. They are often used to allow more variability in systems where simple distributions do not allow for enough variability (in this case, running the hypergeometric with the same *contacts* for every individual)
- The modelling of super spreaders could be investigated by replacing the Poisson distribution by a heavier tailed one (Negative Binomial, Zeta, etc)
- Replacing all values by distribution averages converges to the result of the deterministic version

### 3.1.2 Calculating the number of between-node infectious contacts

For between-node infectious contact calculations, we use an input file of expected directional contacts, an optional time-varying movement modifier, and the disease state at the previous time of the nodes involved. For a particular pair of nodes  $u, v$  at time  $t$ , let  $w(u, v)$  be the number of expected directional contacts,  $x_t$  the appropriate time's movement modifier,  $p_{v,\mathcal{I}}$  the proportion of individuals in  $u$  in an infectious class, the expected number of infectious contacts from  $u$  targeting  $v$  is defined as following. At this stage, all age groups are mixed in together.

#### Deterministic mode

The number of infectious contacts is given by:

$$x_{t-1}p_{t-1,\mathcal{I}}(A_u)p_{t-1,S}(A_v)w(u, v)$$

#### Stochastic mode

The stochastic version attempts at randomly generating those commutes that are generated from infectious people in  $u$ , to susceptible people in  $v$ . Therefore, we first sample the number of commutes originating from infectious people in  $u$ , using a Binomial distribution,

$$\text{commutes from infectious} \sim \text{Bin}(p = p_{t-1,\mathcal{I}}(A_u), N = x_{t-1} * w(u, v))$$

Then given a number of such commutes, we select those that targeted susceptible people in  $v$ .

*commutes from infectious to susceptible  $\sim \text{Bin}(p = p_{t-1,S}(A_v), N = \text{commutes from infectious})$*

### Assignment to age groups

We then sum over all nodes that have directed contact toward  $v$ , and then distribute those infectious contacts amongst the population within  $v$ . In the deterministic mode we assign the contacts using the ratios of people in each age group to total number of people. In the stochastic mode we use a multinomial distribution to randomly assign the contacts into age groups, with the probabilities being the ratios of people in each age group.

## 3.2 Disease progression

Progression through disease states from exposed onward as well as the compartmental model in use is specified by an input file of compartmental transition rates. We have encoded two compartmental models (as in Figure 1) for use with simple network sim: both are essentially SEIR models, but with additional compartments for severely ill (assumed hospitalised) and asymptomatic infectious individuals. In a deterministic run of the model, at each time step the (fractional) number of people flowing from a state  $s_i$  to a state  $s_j$  given a rate of  $r_{i,j}$  is simply calculated as  $|s_i|r_{i,j}$ , where  $|s_i|$  is the number of people in  $s_i$ . In a stochastic run, the integer number of people flowing from  $s_i$  to  $s_j$  given rate  $r_{i,j}$  is a sample from a multinomial distribution with parameters  $|s_i|, r_{i,j}$ .

## 3.3 Infection Seeding

Infection seeding is currently from an input file which specifies the number of individuals in compartment  $E$  in some set of specified nodes. As this parameter cannot be accurately observed, it is inferred.

# 4 ABC-SMC Inference

## 4.1 Algorithm description

Simple network sim being a network-based model, the likelihood is computationally intractable to derive. Therefore model-agnostic and simulation-based methods must be used in order to infer the parameters. We use a Bayesian approach and we aim at obtaining the posterior distribution of model parameters conditional on the data, using an ABC-SMC algorithm introduced in [1], which we will briefly describe.

Based on ABC (Approximate Bayesian Computation, see [2]), and inspired from sequential importance sampling and particle filters, ABC-SMC aims at reducing the running

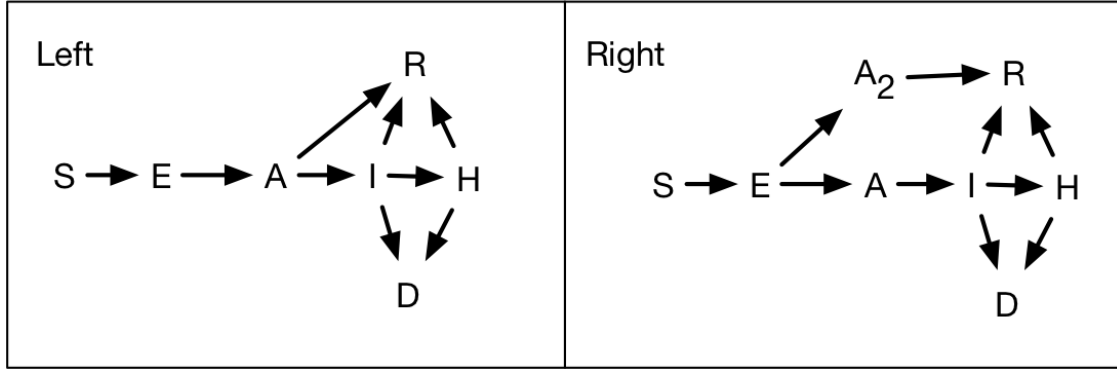


Figure 1: Two compartmental models currently available as input files to simple network sim. In both,  $S$  are susceptible individuals,  $E$  are exposed individuals who are not yet infectious,  $I$  are infectious individuals with symptoms,  $H$  are hospitalised individuals,  $R$  are recovered individuals,  $D$  are dead individuals. On the left,  $A$  are asymptomatic individuals who may or may not ever develop symptoms, whereas on the right,  $A$  are asymptomatic individuals who will eventually develop symptoms and  $A_2$  are *truly asymptomatic* individuals who will never develop symptoms

time of ABC by resampling in the population of particles accepted by the rejection mechanism, in an iterative way. Let  $\theta$  be the parameter vector to be estimated,  $\mathcal{D}$  the data, and  $\pi(\theta)$  the prior used. Our aim is to compute  $\pi(\theta|\mathcal{D}) \propto f(\mathcal{D}|\theta)\pi(\theta)$ .  $N$  particles  $\theta^1, \dots, \theta^N$  are propagated through a sequence of intermediate distributions  $\pi(\theta|d(x_0, x) \leq \epsilon_i), i = 1, \dots, T$  until they represent a sample distributed according to  $\pi(\theta|d(x_0, x) \leq \epsilon_T)$ , which will be a good approximation of the posterior  $\pi(\theta|\mathcal{D})$ . The tolerances  $\epsilon_1 > \dots > \epsilon_T \geq 0$  are chosen so that the iterated distribution converges to the real posterior. The algorithm works as follows (see [1] for more details and derivation):

```

Initialize  $\epsilon_1 > \dots > \epsilon_T \geq 0$ ;
for  $t=1, \dots, T$  do
  while Particles accepted  $< N$  do
    if  $t = 0$  then
      Sample  $\theta^{**}$  from  $\pi(\theta)$ ;
    else
      while  $\pi(\theta^{**}) = 0$  do
        Sample a particle  $\theta^*$  from the previous population  $\{\theta_{t-1}^i\}_i$  with
        normalized weights  $\{w_{t-1}^i\}_i$  and perturb the particle to obtain
         $\theta^{**} \sim K_t(\theta|\theta^*)$ , where  $K_t$  is the perturbation kernel;
      end
    end
    Simulate a candidate dataset  $x^* \sim f(x|\theta^{**})$ ;
    if  $d(x^*, x_0) \leq \epsilon_t$  then
      Accept the particle, set  $\theta_t^i = \theta^{**}$  and calculate its weight:
      
$$w_t^i = \begin{cases} 1, & \text{if } t = 0 \\ \frac{\pi(\theta_t^i)}{\sum_{j=1}^N w_{t-1}^j K_t(\theta_{t-1}^j, \theta_t^i)}, & \text{if } t > 0 \end{cases} \quad (1)$$

    end
  end
end

```

**Algorithm 1:** ABC-SMC

## 4.2 Parameter inference for simple network sim

We aim at inferring:

- The infection probability  $p$
- The initial state of the model which we assume as  $E_0(A_{u,[17,70)})$ , the number of exposed people in age group  $[17, 70)$  in every node at time 0. There are 14 such parameters
- The contact multipliers  $y_t$ , which is a constant-by-parts function with 2 changes

This is a total of 18 parameters to infer, which is quite a large problem. For starting time of the model, we use the date of 9 March 2020, 2 weeks before the first recorded death in Scotland. The perturbation kernel used is the uniform noise  $K_t(\theta, \theta^*) \sim \mathcal{U}(-\sigma, \sigma)$  where  $\sigma$  is different for every inferred parameter. The distance used is the root mean sum

of squares in the time series of deaths per healthboard,

$$d(x^*, x_0) = \sqrt{\frac{1}{\#hb * \#t} \sum_{hb,t} (model\ deaths_t^{hb} - historical\ deaths_t^{hb})^2}$$

where the time series are aggregated by weeks for increasing smoothing,  $\#hb$  is the total number of healthboards,  $\#t$  is the number of timesteps. The time series of deaths is one of the few reliable numbers (compared for example to the number of infected people, which is mostly unknown due to insufficient testing). We use  $T = 5$ ,  $N = 100$ , and the  $\epsilon$  strategy is to use the median of the distances at the previous SMC iteration, with an infinite threshold for the first iteration. Finally, the following priors are used:

- $p \sim Beta(4, 76)$  which allows to have a distribution peaked around a mean value of 5% which is our prior
- for every region  $u$ ,  $E_0(A_{u,[17,70)}) \sim LogNormal(mean = estimated\ infected, std = max(0.2*mean, 10))$ , i.e. the parameters are chosen such that we match our manually estimated numbers, with a standard deviation proportional to the mean to account for the fact that different regions have very different estimations due to different remoteness
- $y_t \sim LogNormal(mean = estimated\ infected, std = 0.2*mean)$ , with a similar logic as above

The inference results for the infection probability are shown in Figures [2] and [3]. Finally we show a run from a fitted model, in Figures [4] and [5].

## References

- [1] Tina Toni, David Welch, Natalja Strelkowa, Andreas Ipsen and Michael P.H Stumpf, Approximate Bayesian computation scheme for parameter inference and model selection in dynamical systems, J. R. Soc. Interface.6187–202 <http://doi.org/10.1098/rsif.2008.0172>
- [2] Mark A. Beaumont, Wenyang Zhang and David J. Balding, Approximate Bayesian Computation in Population Genetics, GENETICS December 1, 2002 vol. 162 no. 4 2025-2035 <https://www.genetics.org/content/162/4/2025>



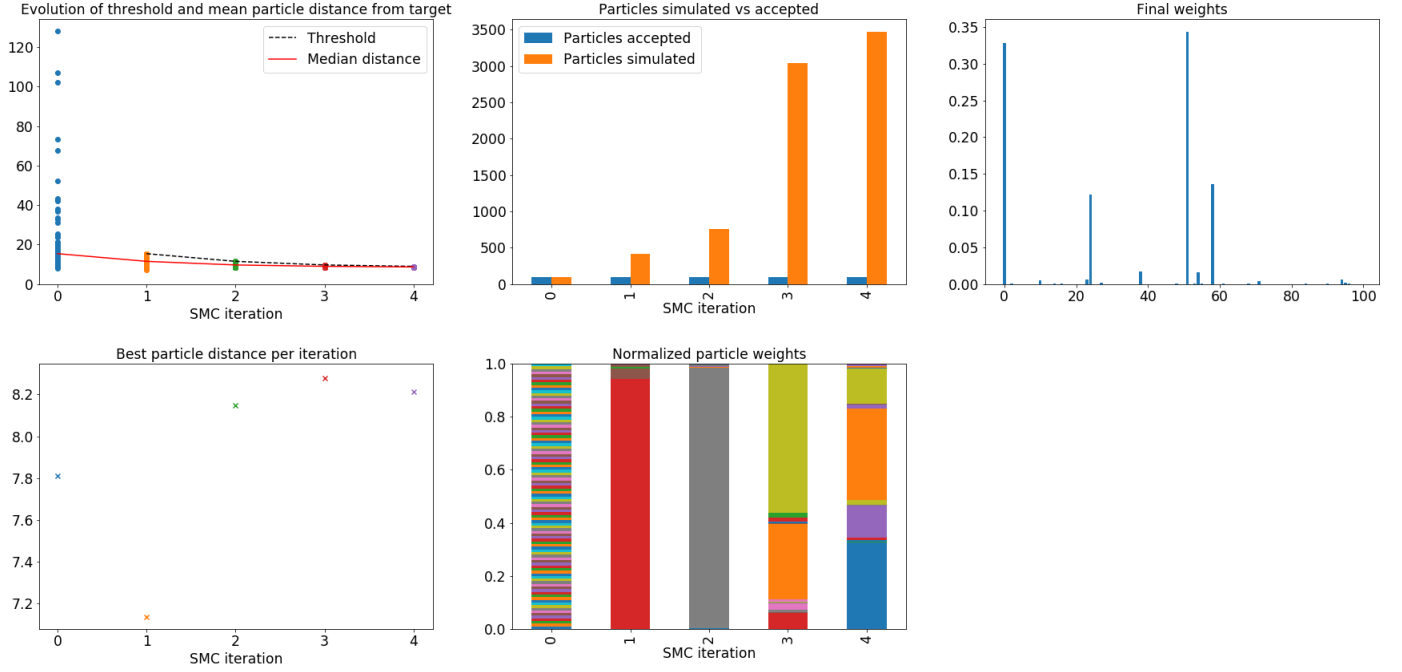


Figure 2: ABC-SMC run statistics. We notice the distances of particles quickly decrease and converge to a value of approximately 6.6, which seems to be the minimum attainable using this model parametrization. The weights start off uniformly distributed but subsequently target more particles. We also notice an increasing number of particles simulated in order to fill the required population, but remains in the thousands. The total run-time is about 1h30 on a somewhat powerful personal computer.

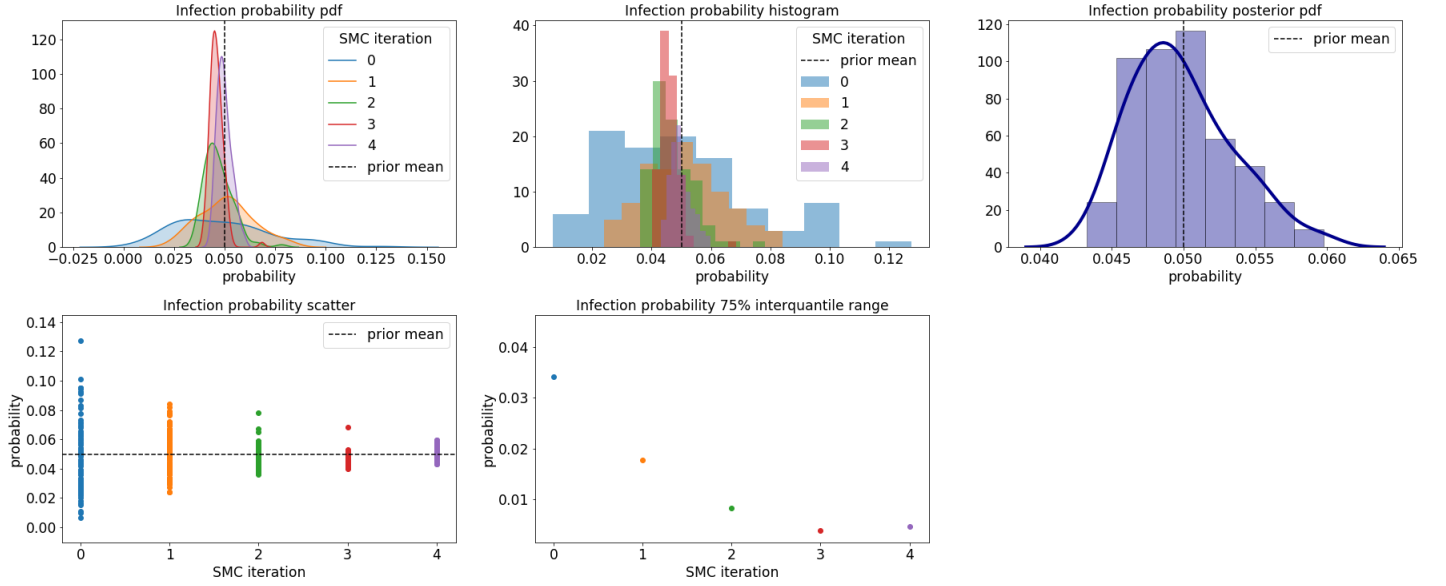


Figure 3: Inference results for infection probability. We observe that each intermediate distribution becomes more and more peaked and concentrated around what seems to be the posterior value, as evidenced by a converging interquantile range. The resulting approximation of the posterior distribution is peaking at 5.3%, very close to our prior. We observe similarly good convergence for the other inferred parameters.

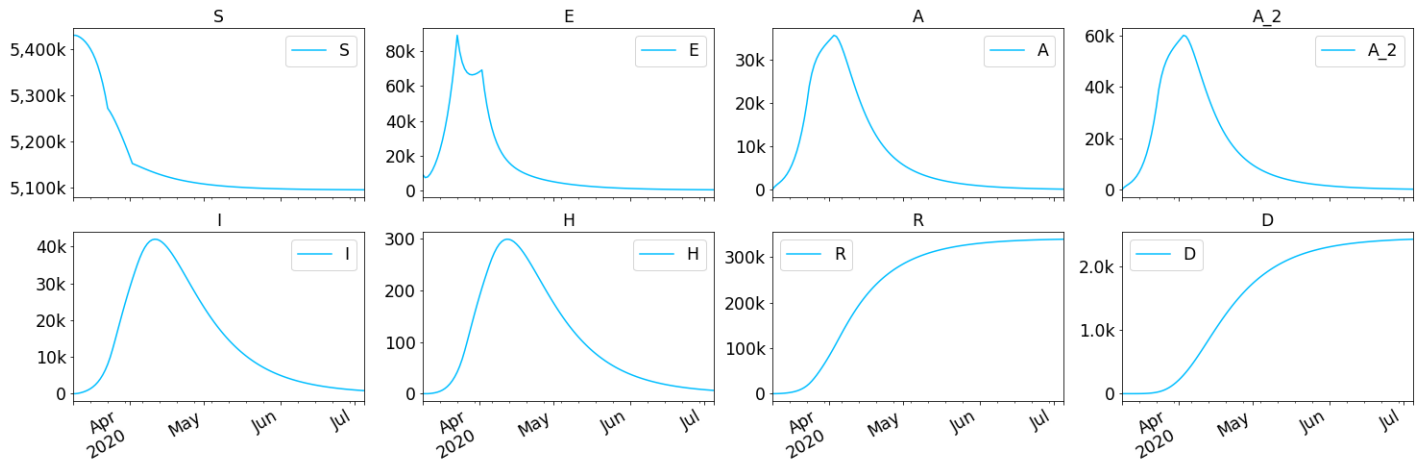


Figure 4: Evolution of different disease compartments for fitted model.

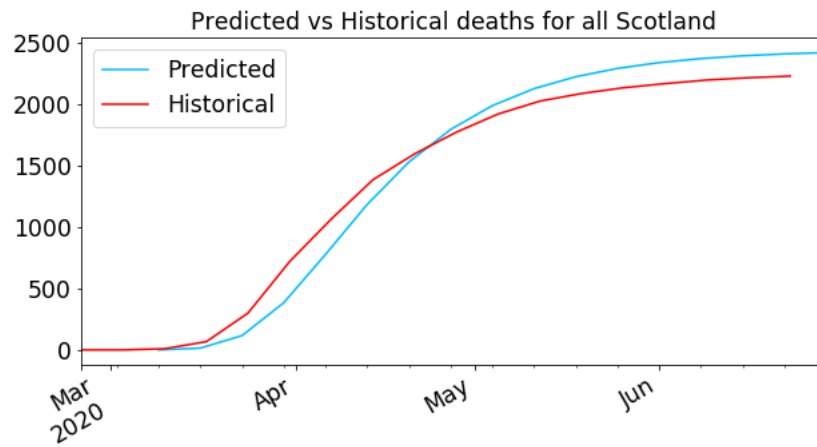


Figure 5: Fitted vs historical death time series for all Scotland.