

Simulation of the COVID-19 on the simplified social network of Slovenia: effective strategies of the virus containment

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Abstract

The paper describes the simulation of the coronavirus spread over the social network of more than 2 million nodes. The simulation is used to propose the potential strategies of the coronavirus containment to decision bodies.

1 Introduction

There are two several ways to simulate the pandemic dynamics. The most common approach is to solve a system of differential equations given some predefined parameters. These epidemic models are widely known as susceptible(S), immune (I), recovered (R), i.e. SIR models [1, 2]. Another variation of SIR models is SEIR model, which accounts also for the exposed. Complex transfer or activation functions are usually used to smoothly model external factors and to account for probability distributions

Here, we propose more computationally expensive node-based approach to simulate the virus spread. We simulate the spread over a realistic social network of more than 2 million nodes with a total of up to 50 million connections, representing the population of Slovenia and the connections of their inhabitants. A hard-to-overcome limitation of the SIR model and its variants is that the information is homogeneously spread. In reality, there are some who spread the information or virus more - so called superspreaders. Another advantage of such approach is that it allows to realistically simulate the quarantine orders of the decision bodies. Furthermore, it allows to simulate strategies for the disease containment and optimal case testing strategies.

Section 2 describes the model and its parameters. Section 3 demonstrates four different deterministic scenarios of the pandemic dynamics. Section 4 presents the results and the most likely outcome of the epidemics based on the ensemble computations which include the uncertainty of the input parameters. Section 5 discusses the potential strategies for the control of the COVID-19 pandemics. Discussion, conclusions and further outlook are given in Section 6.

2 Methodology

2.1 Social network model

The social network of the inhabitants of Slovenia is constructed based on the recent data of Statistical Office of Republic of Slovenia [3]. A total of $N = 2045795$ nodes is used in the social network. The number of k -person households is given in Table 1. There are approximately 100 elderly care centers in Slovenia with a total of approximately 20000 residents. Each elderly care center is assumed to include 8 distinct groups of 25 people.

Average household/care group has 2.5 people in Slovenia so the average number of contacts per person within household is 1.5.

Connectivity distribution in normal conditions follows power law distribution with fat tails [4], which are associated with superspreader events in pandemic dynamics. However, since all public

k persons in household	number of k -person households
1	269898
2	209573
3	152959
4	122195
5	43327
6	17398
7	6073
8	3195
25	100 care centers with 8 groups each

Table 1: Households distribution in Slovenia.

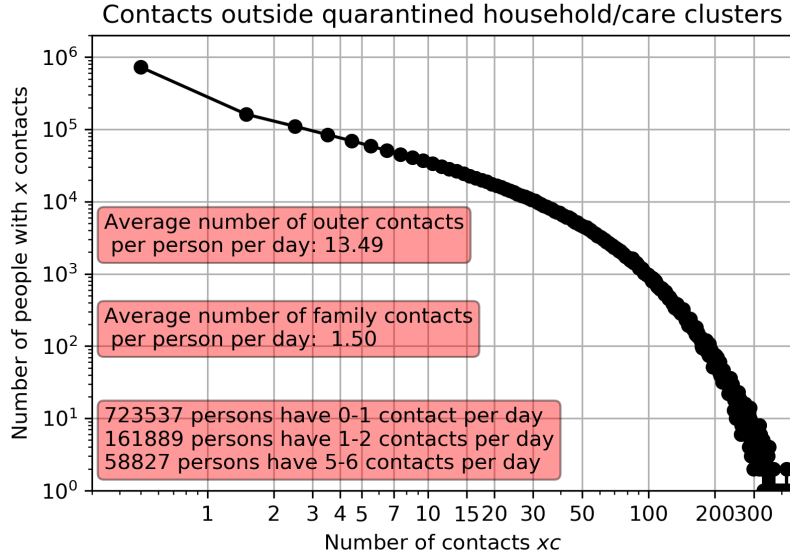


Figure 1: Number distribution by the number of contacts in social network.

events are canceled, those fat tails are cut off [5] and the topology of the social network changes drastically. In quarantine conditions, a reasonable assumption is to model the connectivity, i.e. the number of contacts per person using the gamma probability distribution, which is essentially an exponential distribution

$$p(x; k, \theta) = \frac{1}{\Gamma(k)\theta^k} x^{k-1} e^{-\frac{x}{\theta}}. \quad (1)$$

In this study, we used $k = 0.3$ and $\theta = 22.5$ for the initial setup, which gives an average number of 13.5 outer contacts per person per day (Figure 1). Together with 1.5 family contact per day, the total number of contacts per person per day is 15. Here, we assume that the average contact number is the same for each age group, despite studies showing seniors have reduced number of contacts already in normal conditions [6]. Another study shows that Italians have on average almost 20 contacts per day, Germans around 13.5, so 15 contacts per day is a reasonable guess for Slovenia [7].

Technically, we connect the graph in the following way

1. numbers of contacts for each node are randomly drawn from Gamma distribution (1). If node i has $x_i = 0.33$ contacts per day, it means that it will have 0 contacts 2/3 of the time and 1 contact 1/3 of the time of the simulation

2. for each node i we randomly assign the connections to x_i other nodes, where x_i is the number of contacts of node i . However, the assignments do not have equal probability. Node j which has x_j contacts is picked as a neighbour with probability $x_j N(x_j)/T$, where $N(x_j)$ is the number of nodes with x_j contacts and T is the total number of contacts in the network, two times the number of connections. Sampling over Gamma distribution (1) gives us a distribution of $N(x) = p(x)N$. When picking the neighbours, we actually sample the same Gamma distribution times x , i.e.

$$p_n(x) = p(x; k, \theta)x = \frac{1}{\Gamma(k)\theta^k} x^k e^{-\frac{x}{\theta}} \propto p(x; k+1, \theta) \quad (2)$$

3. The shape of the social network is changing at every time step of the simulation to account for people's mobility. (a) The number of contacts of node i is fixed (randomly jumps between $\lfloor x_i \rfloor$ and $\lceil x_i \rceil$ based on the value of x_i). For example, if a node has 0.33 contacts per day, 1 contact is picked with probability 1/3 and 0 contacts with probability 2/3. (b) The social network is rewired at every time step to account for superspreaders mobility. Note that rewiring might not be a good choice for those with low number of contacts.

2.2 Virus spread parameters

Reproduction number R_0 A basic reproduction number, R_0 , only provides the info on the average dynamics of transmission, however it is crucial to understand what settings drive the virus spread. Different methodologies produced different results, however the majority of reported numbers is within 2 and 3. Here, we use median reported R_0 from a number of studies, as well as its median confidence intervals, i.e. $R_0 = 2.68$, with 95% confidence interval [2, 3.9]. This approach is surely not the optimal one, since we are trading accuracy for precision. The published R_0 values as well as our deduced R_0 distribution is shown in Figure 2. The optimal log-normal distribution should match the following conditions:

- $\text{CDF}(R_0^L; \mu, \sigma, \Delta x) = 0.05$,
- $\text{CDF}(R_0^H; \mu, \sigma, \Delta x) = 0.95$,
- $\text{median}(\text{CDF}) = R_0$,

where CDF stands for log-normal cumulative distribution function, i.e.

$$\text{CDF}(x; \mu, \sigma, \Delta x) = \frac{1}{2} + \frac{1}{2} \text{erf} \left[\frac{\ln(x - \Delta x) - \mu}{\sqrt{2}\sigma} \right] \quad (3)$$

and median being $\exp(\mu)$. Then, we define a quadratic cost function, which includes all the above criteria, and by minimizing it, we obtain the optimal parameters for log-normal distribution: $\Delta x = 0.36$, $\sigma = 1.14$, $\exp \mu = 1.54$.

Attack rate In general, the basic reproduction number, R_0 can be decomposed into the secondary attack rate times the number of contacts. The secondary attack rate (SAR) is defined as the probability that an infection occurs among susceptible people within a specific group (i.e. household or other close contacts). The measure can provide an indication of how social interactions relate to transmission risk. We can further decompose the R_0 into the household risk of infection and outer risk of infection (following [12])

$$R_0 = SAR_h N_h + SAR_c N_c, \quad (4)$$

where SAR_h and SAR_c are secondary attack rates within household and outside household (outer contacts) and N_h and N_c are the numbers of risk contacts made. The study of Liu et al. [12] suggest SAR_h value of 35% (95% CI 27-44%).

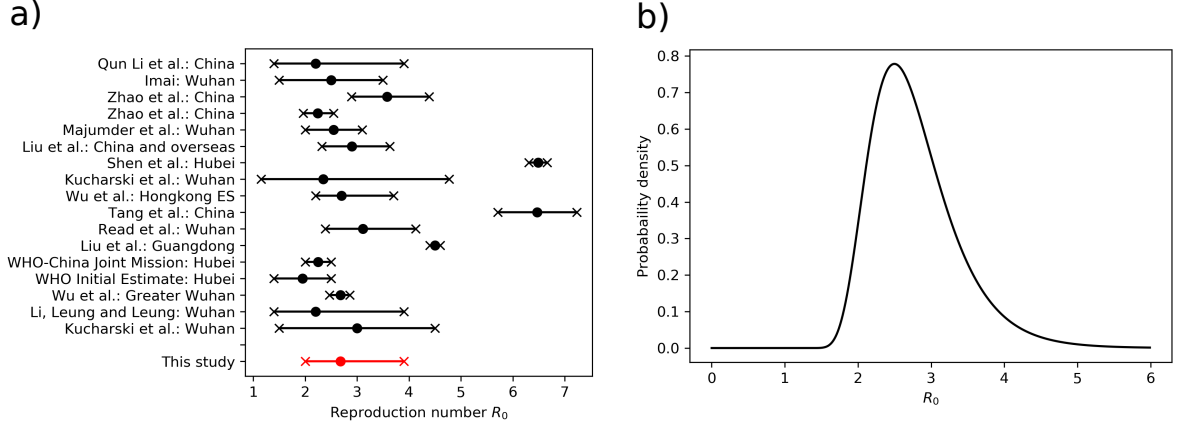


Figure 2: a) Basic reproduction number R_0 , reported in number of studies, which can be found in Wu et al. [8], Kucharski et al. [9], Li et al. [10], Liu et al. [11] and references therein. b) Log-normal probability density function of the basic reproduction number, used for ensemble simulations.

SAR_h is almost normally distributed with mean 35% and $2\sigma \approx 8.5$. The distribution of R_0 is given in the previous paragraph. It holds: $SAR_c = (R_0 - SAR_h * N_h)/N_c$. This gives a transmission efficiency of $SAR_c = 0.16$. Figure 3 shows probability distributions of secondary attack rates as used in the ensemble of simulations.

Given the infectious period of $T_{inf} \approx 10$ days (incubation period $T_{inc} \approx 5$ days minus 2 days + another week, check subsection 2.3), we can assume that the daily risk of getting infected from a certain household member is $SAR_{h,daily}$ where $1 - (1 - SAR_{h,daily})^{10} = SAR_h$ and

$$SAR_{h,daily} = 1 - \exp\left(\frac{1 - SAR_h}{T_{inf}}\right) \quad (5)$$

being equal 4.2% (3.1-5.6%). Similarly, we compute $SAR_{c,daily} = 1.7\%$.

Some studies [e.g. 13] have concentrated only on the symptomatic secondary attack rates and have shown relatively smaller numbers: 0.45% (CI=0.12%-1.6%) among all close contacts and 10.5% (CI=0.12%-1.6%) among household members. However, these numbers cannot reproduce the reported R_0 between 2 and 3.9 with realistic number of contacts. Another study shows similar attack rates we use in this study [14].

2.3 Disease and hospitalization parameters

Case fatality ratio The baseline case fatality ratio (CFR), fatality ratio among all positively tested, is assumed 1.38% (CI 1.23-1.53%) [15, 16]. Dividing deaths-to-date by cases-to-date leads to a biased estimate of CFR, called naive CFR (nCFR) as the delays from confirmation of a case to death is not accounted for, as well as due to under-reporting of cases. The reported numbers agree with recently published study for symptomatic case fatality ratio in China [17].

Infection fatality and hospitalisation ratios Infection fatality ratio (IFR) estimates are based on the study from Verity et al. [15] and are taken to be 0.9% with 95% confidence interval 0.4% to 1.4% (95% CI is 2σ for normal distribution). These estimates are consistent with IFR on Princess Diamond Cruise ship and were used also in Ferguson's Imperial College report [18]. However, the countries vastly vary in demography. The study was performed for China with median age of 37.4 years. Slovenia has a median age of 44.5 years. The study also found an increasing infection fatality profile in age (Table2). These profile is used to determine the effective infection fatality rate for Slovenia. Performing a weighted average, we use the total IFR of 1.16% (CI 0.63-2.22%). Analogously, we compute the average hospitalisation rate of 6.37% (95% CI

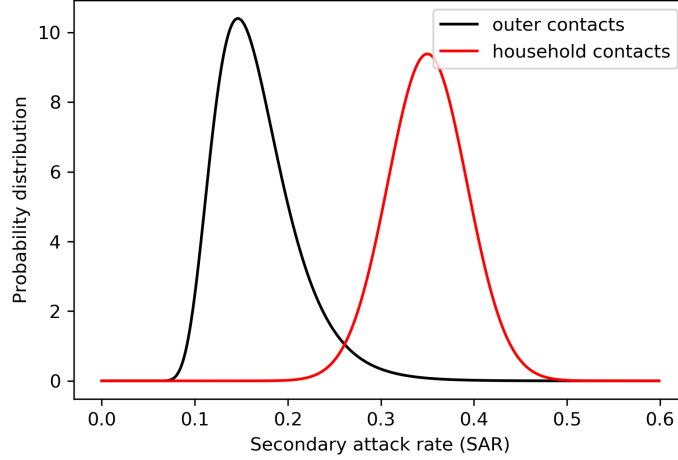


Figure 3: Probability distribution of secondary attack rates for household contacts and outer contacts.

Age group	IFR (95% CI)	IHR (95% CI)	Ratio of total population in SLO
0 to 9	0.0016% (0.000185,0.0249)	0% (0,0)	0.102
10 to 19	0.007% (0.0015,0.050)	0.04% (0.02, 0.08)	0.093
20 to 29	0.031% (0.014,0.092)	1.1% (0.62, 2.1)	0.102
30 to 39	0.084% (0.041,0.185)	3.4% (2.1, 7.0)	0.140
40 to 49	0.16% (0.076,0.32)	4.3% (2.5, 8.7)	0.146
50 to 59	0.60% (0.34,1.3)	8.2% (4.9, 16.7)	0.145
60 to 69	1.9% (1.1,3.9)	11.8% (7.0, 24.0)	0.135
70 to 79	4.3% (2.5,8.4)	16.6% (9.9, 33.8)	0.083
80+	7.8% (3.8,13.3)	18.4% (11.0, 37.6)	0.054

Table 2: Estimates of the proportion of all infections that would be hospitalised (IHR) or fatal (IFR) by age group. Disease data from Verity et al. [15]. Population data from PopulationPyramid.net [19].

3.8-13%). Using the minimization procedures, we obtain parameters of log-normal distribution which best fits both values and their 95% confidence interval (Figure 4).

Incubation period - infection to illness onset Mean incubation period is taken to be 5.0 days (95% CI 4.2-6.0), while the 95th percentile of the distribution was 10.6 days (95% CI 8.5-14.1) and 99th percentile 15.4 days (99% CI 11.7-22.5) [20]. Similar numbers were reported in an earlier study which included less patients [10, 21]. Log-normal distribution is used among for incubation period among nodes. However, the parameters of the lognormal distribution remain fixed due to numerical instability of their computation. Thus, all ensemble members have the same log-normal distribution of incubation period. Incubation period distribution and other outcome parameters are shown in Figure 5.

Infectious period The infectious period starts $T_{start} = 2$ days before incubation and likely ends around day $T_{end} = 7$ from symptoms onset [22], i.e. $T_{inc} - T_{start} + T_{end} \approx 10$ days, where average incubation period lasts 5 days. Note that none of the interval boundaries are known exactly, however several cases are known where infected transferred the virus before developing

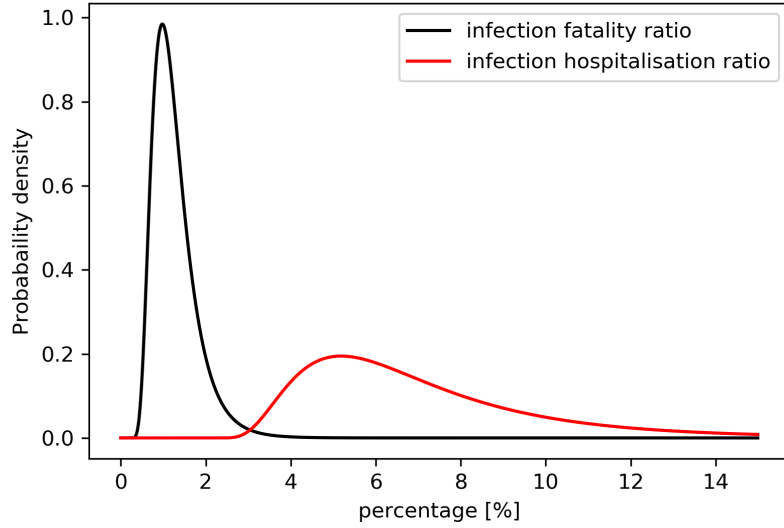


Figure 4: Infection fatality ratio distribution and infection hospitalisation ratio distribution for ensemble simulations.

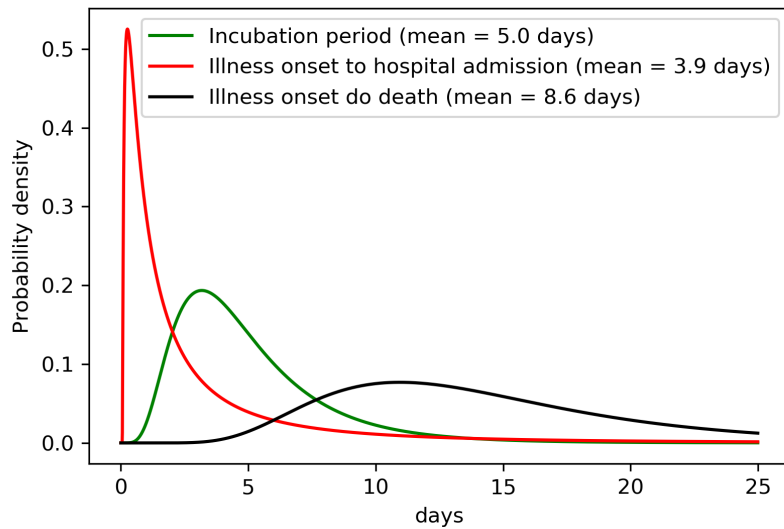


Figure 5: Incubation period, illness onset to hospitalisation and illness onset to death distribution among COVID-19 patients.

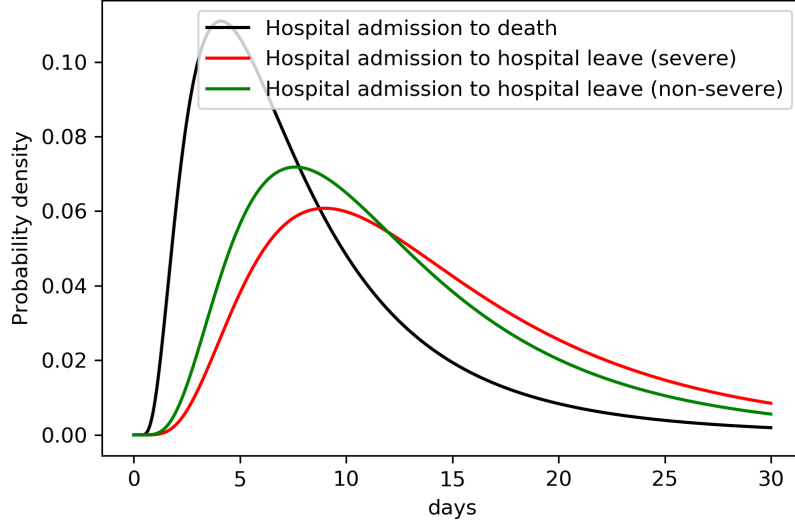


Figure 6: Mean distribution of hospital admission to death, hospital admission to hospital leave for severe and for non-severe illness.

symptoms. Thus, we randomly perturb $T_{start} = \mathcal{N}(3, 0.5)$ and $T_{end} = \mathcal{N}(2, 1)$. The infectious period fall in line with study of Bi et al. [14], Figure S2.

Illness onset to hospital admission We take mean numbers from the study of Linton et al. [20] and its distribution: mean is 3.9 days, median 1.5 days, 5% percentile at 0.2 days and 95% percentile at 14 days. Since we now understand the severity of the illness, only the distribution of data for living patients is accounted for. In China, those who died waited longer to visit the doctor.

Illness onset to death Mean 14.5 days, median 13.2 days, 5th percentile 6.5, 95th percentile 26.8 [20]. Similar numbers were reported by Russell et al. [16].

Hospital admission to death Mean 8.6 days, median 6.7 days, 5th percentile 2.2, 95th percentile 20.5, 99th percentile 32.6 days [20]. Shown in Figure 6

Hospital admission to hospital leave Hospital admission to recovery is on average longer than hospital admission to death. While the full recovery is important for economy, hospitalisation length is more important for the state of healthcare system. The median hospitalisation length is 11 days (95% CI 10-13) for non-severe cases and 13 days for severe (95% CI 11-17) [20]. Both are log-normally distributed. For ensemble computations, their medians are further log-normally distributed according to their respective confidence intervals.

Similar numbers: mean (larger than median for lognormal distribution) hospital length of stay and ICU length of stay are 11 days (95% CI 7-14) and 8 days (95% CI 4-12) were reported by Zhou et al. [23].

Other parameters - review!? Fatality ratio of severe cases in need of intensive care is 50%. Fatality ratio of severe cases without intensive care is normally distributed with mean of 90% and $\sigma = 5\%$. Fatality ratio of severely ill without oxygen is 10% with $\sigma = 5\%$.

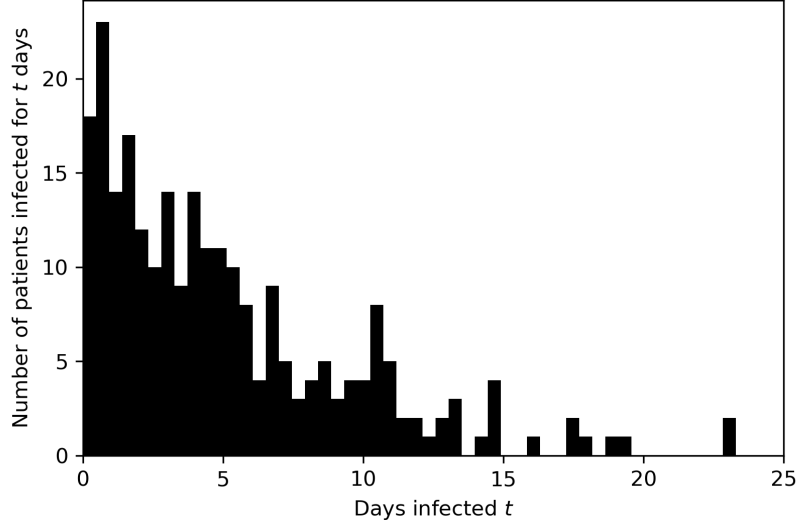


Figure 7: Distribution of 89 infected people on March 12, 2020, in Slovenia, by the time-length of their infection.

2.4 Initial condition

The initial condition for the simulation is defined for March 12, 2020. To that day, there were 89 symptomatic cases who tested positive, 8 days after first positive case, which implies an anomalously low doubling time of $\tau = 1.23$ days. This number is case specific as there was winter holiday in Slovenia at the end of February and beginning of March, when lots of people went skiing to Italy (including Lombardy). Most of the initial cases were imported [??]. Other studies typically suggest a doubling time of around 5 days (95% CI 4.3 - 6.2) in the initial uncontrolled stage of the epidemic expansion [25]. However, Abbott et al. [26] report smaller values of around 3.5 days in most of Western Europe. Thus, our choice is doubling time of $T_{double} = 4.2$ days, normally distributed with $\sigma = 0.5$.

Different numbers of actually infected people were suggested in the media reports, ranging from 5 to 10 times the number of reported positive cases. Given the average incubation period of 5 days + (1 day for visit) and somewhat smaller doubling period of 3.5 days, factor $2^{\frac{T_{inc}}{T_{double}}} = 2.7$ applies. Furthermore, the proportion of asymptomatic cases is around 18% based on the data from Diamond Princess Cruise Ship [27] and around 33% based on the more recent study [28]. There have been reports from Iceland, who undergo vast testing that nearly 50% of positively tested are asymptomatic [??]. We opt for 40%, normally distributed with std. of 10%. This means that around 400 people were infected in Slovenia on March 12. Factor 4.5 is however smaller than the assumption of Kucharski et al. that only 16% of onsets are known <https://cmmid.github.io/topics/covid19/current-patterns-transmission/wuhan-early-dynamics.html>.

Based on the exponential growth in initial stage and incubation period, we randomly generate the infection length of the patients with exponential distribution with shape factor of 4.9, so that 89 people have infection for more than 5 days and develop symptoms. Initial distribution of 89 infected people by the time-length of their infection is shown in Figure 7. Note that in reality, due to many imported cases, the actual distribution may be different.

2.5 Limitations

- Nodes do not have age as property.

- Contacts are not distributed by age. Older have less connections.
- ...

3 Scenarios

Draw four deterministic plots for different numbers of connections.

4 Results

4.1 The effect of superspreaders

5 Strategy

The main strategic points would be:

- Restriction of travels, which prevents community-to-community spread [29]
- Reduction of large events, e.g. conferences [29]
- widespread testing and inclusion of recovered back into society
- tracking of potential secondary and tertiary contacts and strict isolation in special facilities [25]. Isolate them from their families. Telecommunication data or mobile app data would be needed .
- Largely reduce the number of contacts. Complete lockdown. Strategy to avoid people mixing in supermarkets. Special online orders and takeaway.
- Recent study (under review) has shown, that contact tracing is successful only for the case of low R_0 [30]. For example, around 50% (70%) of cases would ne to be traced for $R_0 = 1.5(2.5)$ if we are to controll the outbreak.
- The survival ability of SARS coronavirus in human specimens and their environments is relatively strong, however UV and heating (such as in cars exposed to sun, playgrounds etc.) dan efficiently eliminate the viral infectivity [31] and reduce the environmental transmission.

5.1 Isolation

We can define illness onset (developed symptoms) to isolation function, similarly as in Hellewell et al. [30]. Two parameters are important: percentage of tracked and the delay. Isolation efficiency is assumed 100%. In the case of short delay, we track and isolate potentially infected before they develop symptoms, but in the case of long delay, we only isolate them when they start developing symptoms.

5.2 Isolation of tertiary infections

More progressive tracking.

6 Conclusions

Accurate models of the real-world social networks are needed to realistically simulate the topological dynamics of the epidemics. Similarly to Numerical Weather Prediction models [32, 33], the real connectivity data obtained by postprocessing of the phone traces, should be rapidly assimilated into the virus spread prognostic model [e.g. 34]. This would allow 1) to estimate the critical virus

spread parameters, needed for accurate forecasts of the pandemics, and 2) to better prepare a strategy of the virus containment, potentially saving thousands of lives, minimizing the economic damage and enhancing people’s mobility.

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Code availability

The model is freely available at <https://github.com/zaplotnik/korona>. The core program is written in Python 2.7 and requires standard `scipy`, `numpy` and `matplotlib` libraries. Computationally critical parts of the program are written in Fortran. Python bindings are created using F2PY [35]

```
f2py -c generate_connections.f90 -m generate_connections
```

Fortran random number generator (`random.f90`) for Fortran is taken from Allan Miller’s Fortran Software repository <https://jblevins.org/mirror/amiller/>.