Monitoring the spread of COVID-19 by estimating reproduction numbers over time

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Contents

1	Introduction	1
2	Derivation of the estimator	2
3	Specifics of COVID-19	3
4	Validation on simulated data	4
5	Application to real data	7
6	Sensitivity analysis	7
7	Discussion and Outlook	7
A	Derivation of confidence intervals	7
В	Derivation of the infectivity profile for the SEIR-model	9

1 Introduction

As the Coronavirus Disease 2019 (COVID-19) threatens humanity, unprecedented measures to stop its spread have been adopted around the globe. In many countries, schools have closed and curfews have been imposed. Given the enormous burden these measures place on the economy, sooner or later they have to be relaxed. This raises important questions for policymakers and public health specialists. How large is the effect of these measures? Do they effectively stop the spread of COVID-19? What will happen if restrictions get relaxed? And in the future, how can we see whether the epidemic is getting out of hands again?

To answer these questions, one needs to know how fast the epidemic is growing. In epidemiology, this is measured by the **reproduction number**, i.e. the mean number of people someone who got infected will infect in the course of time. Its **critical value** clearly is 1: for larger values the epidemic will grow, for smaller values it will diminish.

Since conditions may change in the future, e.g. when countermeasures are introduced or lifted, this may change. We therefore follow Fraser (2007) and consider what he calls the *instantaneous* reproduction number R(t) at time t, and for which he suggests the estimator

$$\hat{R}(t) = \frac{I(t)}{\sum_{\tau=1}^{\infty} w(\tau)I(t-\tau)}$$
(1)

where I(t) is the number of incident cases at time t and w specifies the so-called **infectivty profile** which is assumed to be known. To the best of our knowledge, this estimator has first been published by Fraser and others in (Grassly et al.; 2006).

We explain the probabilistic model behind this estimator following (Cori et al.; 2013, Web Appendix 1) in Section 2. In addition, we analytically derive asymptotic confidence intervals (with details given in Appendix A) which are simple to compute. Here, we differ from Grassly et al. (2006) who use computationally more elaborate resampling techniques, namely the bootstrap, to obtain confidence intervals; Cori et al. (2013) on the other hand take a Bayesian approach, assuming a certain gamma prior distribution for R(t).

The estimator and corresponding confidence intervals are validated on simulated data in Section 4. Then, we apply this methodology to real data from Germany in Section 5, followed by a sensitivity analysis in Section 6. Finally, the results are summarised in Section 7, also discussing difficulties with this approach.

In order to continuously monitor the spread of COVID-19, a designated website has been created where the results of our analysis are shown and updated daily. It is available at https://stochastik-tu-ilmenau.github.io/COVID-19/ in English for all affected countries based on the data from (Johns Hopkins University Center for Systems Science and Engineering; 2020) as well as in German for Germany and its federal states based on the data from (Robert Koch-Institut; 2020) at https://stochastik-tu-ilmenau.github.io/COVID-19/germany. We note that a similar analysis using the Bayesian approach of (Cori et al.; 2013) was presented by Abbott et al. (2020). However, as of 06/04/2020, the analysis appears not to have been updated since 19/03/2020.

2 Derivation of the estimator

The following is an adaptation of the modelling in (Fraser; 2007) and (Cori et al.; 2013, Web Appendix 1).

Time is taken to be discrete, i.e. we consider days $t \in \mathbb{Z}$, since the spread of the epidemic shows a strong intraday variability (e.g., there are fewer infections during the night when people are at sleep), and the time scales of incubation and infectious period are on the order of days. Also, cases are reported on a daily basis.

The number of **incidences**, i.e. newly infected cases, at day t will be given as I(t). The **infection** age of an infected person in days, i.e. the number of day elapsed since the infection, is denoted by $\tau \in \mathbb{N}_0$.

The spread of the epidemic depends strongly on the time-dependent transmissibility $\beta(t,\tau) \geq 0$ specifying the expected number of susceptible individuals an infectious person at infection age τ , a so-called **primary case**, will infect at time t. The transmissibility is in particular affected by the **contact rate**, i.e. the mean number of people an infected person meets per day, and the **infectiousness** of the primary case. The former is addressed by **non-pharmaceutical interventions** such as school closures and curfews, the latter is a virological feature of the disease. Therefore we make a crucial **structural assumption**, namely that they separate:

$$\beta(t,\tau) = R(t) w(\tau) \tag{2}$$

where $R(t) \geq 0$ denotes the (instantaneous) **reproduction number** at time t of transmission, i.e. when the **secondary case** gets infected by the primary case, and $w(\tau) \in [0,1]$ specifies the **infectivity profile** at infection age τ . This models the belief that contact rates change over time but the infectiousness of the primary case depends only on τ . This is debatable: when rules for isolation or quarantine change are loosened, e.g. because hospital capacities are exhausted, β will change differently for different values of τ ; we will reiterate this point in Section 7. It also shows

in the fact that any constant factor may be alternatively incorporated into R or w. The latter is therefore standardised such that

$$\sum_{\tau=0}^{\infty} w(\tau) = 1, \tag{3}$$

i.e. w is a **probability distribution** which can be interpreted as follows: for a fixed time t randomly pick a pair of individuals where the first one is a primary case that got infected at time t, in turn infecting the second one later; $w(\tau)$ is the probability that the second case got infected at time $t+\tau$, i.e. at infection age τ of the primary case. w is assumed to be **known**; see Section 3 on how we model it for COVID-19.

In a **stochastic model** for the dynamics of the epidemic, I(t) is given as the number of successful transmissions from an infectious person to someone who is susceptible to the disease. Assuming that for each possible transmission succeeds independently (thus ignoring the possibility of multiple infections) with a probability corresponding to β , and if there are many possible transmissions, I(t) is—by the law of small numbers—approximately **Poisson distributed** conditional on the past. The intensity of this Poisson distribution is equal to

$$\mathbf{E}(I(t) | I(t-1), \dots) = \sum_{\tau=1}^{\infty} \beta(t, \tau) I(t-\tau) = R(t) \sum_{\tau=1}^{\infty} w(\tau) I(t-\tau).$$
 (4)

Here, transmissions on the same day are ruled out, i.e. w(0) = 0, which is a realistic assumption since the incubation period will be at least one day.

The last equation suggests the **estimator** $\hat{R}(t)$ for R(t) given in (Fraser; 2007, Equation (9)),

$$\hat{R}(t) = \frac{I(t)}{\sum_{\tau=1}^{\infty} w(\tau)I(t-\tau)}.$$
(5)

Note that the **case reproduction number** $R_c(t)$, i.e. the expected number of people a primary case infected at time t will infect, is given by, cf. (Fraser; 2007, Equations (2) and (8)),

$$R_c(t) = \sum_{\tau=1}^{\infty} \beta(t+\tau,\tau) = \sum_{\tau=1}^{\infty} R(t+\tau) w(\tau).$$
 (6)

This is of course difficult or even impossible to estimate as it depends on future contact rates, i.e. on countermeasures that will be imposed. However, assuming that conditions remain the same in the future, i.e. R(s) = R(t) for s > t, we obtain R(t) again, cf. (Fraser; 2007, Equation (3)),

$$\sum_{\tau=1}^{\infty} R(t+\tau)w(\tau) = R(t)\sum_{\tau=1}^{\infty} w(\tau) = R(t).$$
 (7)

This explains why R(t) is called reproduction number.

For large intensities, i.e. if the conditional expectation in Equation (4) is large, the distribution of $\hat{R}(t)$ can be well approximated by a Gaussian distribution, with small standard errors. From this, asymptotic confidence intervals can be derived, see Section A. If q denotes the $(1-\frac{\alpha}{2})$ -quantile of the standard normal distribution then

$$\left[\hat{R}(t) - q\sqrt{\frac{\hat{R}(t)}{\sum_{\tau=1}^{\infty} w(\tau)I(t-\tau)}}, \hat{R}(t) + q\sqrt{\frac{\hat{R}(t)}{\sum_{\tau=1}^{\infty} w(\tau)I(t-\tau)}}\right]$$
(8)

is an (asymptotic) $(1 - \alpha)$ -confidence interval for R(t). Note that in practice 10 or more incident cases should suffice for the asymptotics to be reliable.

3 Specifics of COVID-19

As COVID-19 is a new disease, first being described at the end of 2019, its virological features have not yet been conclusively determined. Nonetheless, we tried to choose parameter in agreement with the current state of research.

For comparisons, we note that in a population without any countermeasures, the **basic reproduction number** R_0 is believed to be given by some value between 2.4 and 4.1 (Read et al.; 2020).

The **incubation time**, i.e. the time from infection until symptom onset, ranges from 1 to 14 days with a mean of 5 to 6 days; the virus can be detected from 1 to 2 days before symptom onset for up to 7 to 12 days in moderate cases, and even up to two weeks in severe cases (Organisation; 2020). We therefore may indeed assume that w(0) = 0.

For modelling the infectivity profile w, it is important to realise that it is not proportional to the amound of viral specimens that can be detected in an infected person's sputumm, say. Indeed, since severe cases are very likely to be hospitalised and thus strictly isolated, the probability of infecting someone more than 12 days are infection is very low. Similarly, before symptom onset the probability for transmission might be very low since no sputum is distributed.

The infectivity profile is therefore set to start with 0 on the first day after infection with a linear increase up to day 4, remaining constant up to day 6 and decaying linearly again until being 0 at day 11; see Figure 1.

In Section 6 we discuss the effect this choice has on the analysis.

4 Validation on simulated data

To validate the estimator, we simulate a **stochastic SEIR** (a.k.a. Kermack-McKendrick) **model**. To be more precise, we consider a discrete-time Markov chain describing a population of n=1 million people with each individual being in one of four states: **susceptible**, i.e. not yet infected; **latent**, i.e. infected but not yet infectious; **infectious**; or **recovered** and thus immune. We start at time 0 with 100 latent individuals, all others initially being susceptible. At each time step, a susceptible person becomes infected if it meets an infectious person which happens independently with probability p_S , a latent person becomes infectious with probability p_I , and an infectious person recovers with probability p_R ; otherwise an individual remains in its state.

This results in incubation times, i.e. times spent in the latent state, which are geometrically distributed with mean $\frac{1}{p_I}$; for this to be 3, we set $p_I = \frac{1}{3}$. Similarly, the infectious period is geometrically distributed with mean $\frac{1}{p_R}$ which we would like to be 4, so we set $p_R = \frac{1}{4}$. The corresponding infectivity profile w is then given by the convolution of these two geometric convolutions. It can be calculated analytically, see Appendix B for details; the result is shown in Figure 2. Note that w(1) = w(0) = 0 since it takes at least one day to become latent and another one to become infectious in this model.

The basic reproduction rate is then given by $R_0 = \frac{np_S}{p_R}$ since an infected person on average infects np_S individuals per day (if all were susceptible) for $\frac{1}{p_R}$ days on average. In order to simulate an epidemic with $R_0 = 2.5$, we set $p_S = \frac{R_0 p_R}{n}$ accordingly.

Over time, the reproduction number changes naturally because more people recover and become immune: R(t) is R_0 times the proportion of susceptible individuals at that time. In addition, we assume that countermeasures have been imposed at time 30, resulting in R(t) being 0.7 times the proportion of susceptibles afterwards, and that measures have been relaxed at time 50, resulting in R(t) being 1.3 times the proportion of susceptibles thereafter.

Figure 3 shows one simulation run. The resulting estimates $\hat{R}(t)$ and pointwise 95%-confidence intervals ($\alpha = 5\%$ as usual) can be compared with the true reproduction rate R(t) in Figure 4.

The simulation has been repeated 100 times, and for each time point the proportion of confidence intervals containing the true reproduction number has been determined, see Figure 5. They appear not quite to have the desired nominal coverage but given that they are only asymptotic confidence intervals, and modelling errors are typically much larger they appear acceptable in practice.

These simulations demonstrate how well the estimator is able to detect changes in the reproduction number. From a practical viewpoint, this is an overly optimistic result. In fact, Equation (4) and consequently the estimator $\hat{R}(t)$ in Equation (5) are based on the number of newly infected cases. But **infection dates** are rarely known. Instead, cases are reported when they are tested with a positive test result. In our simple simulation, one should therefore consider the **newly infectious**

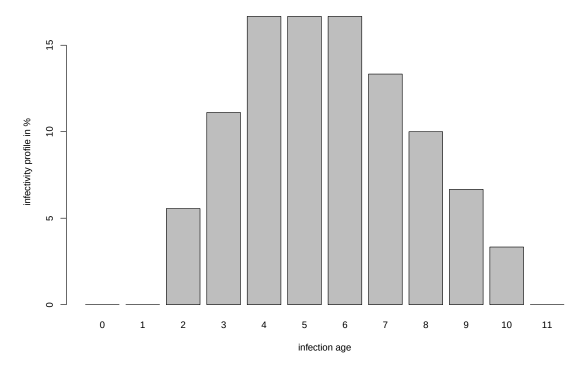


Figure 1: Modelled infectivity profile w.

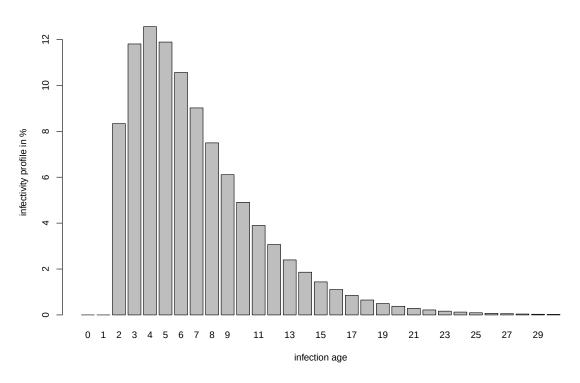


Figure 2: Computed infectivity profile w corresponding to the simulation.

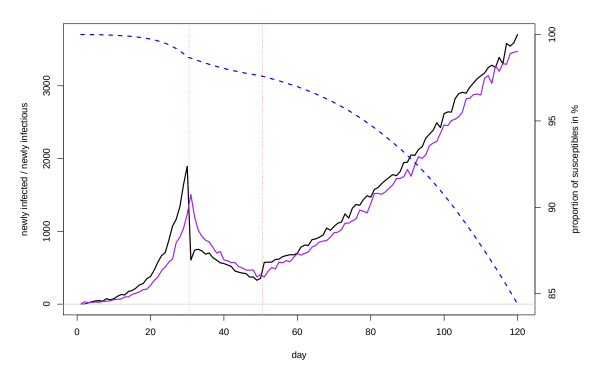


Figure 3: One simulation of the SEIR model; black solid line: newly infected, purple solid line: newly infectious; dashed blue line: proportion of susceptibles; vertical red dotted lines: intervention times.

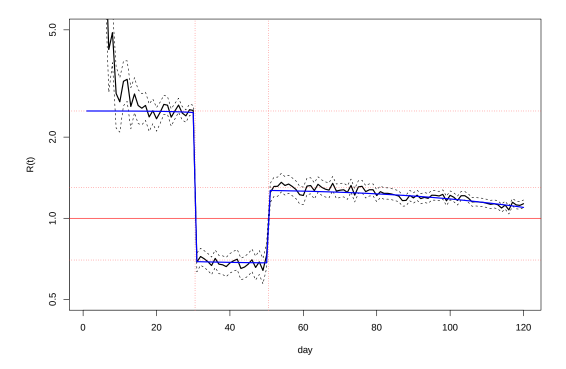


Figure 4: One simulation of the SEIR model; black solid line: $\hat{R}(t)$; black dashed lines: pointwise 95%-confidence intervals; blue solid line: R(t); vertical red dotted lines: intervention times; horizontal red lines: corresponding reproduction numbers without decrease in susceptibles taken into account.

cases at day t as input data I(t) for the estimator. Note that their increase lags behind the one of the **newly infected cases**, i.e. the **newly latent cases**, by the incubation time, see Figure 3 where they lag behind by about 1 day, the mode of the incubation time distribution.

We use a naïve approach to deal with this **reporting delay**: we shift the estimator back by 1 day. The result is shown in Figure 6 where the jump in R(t) leads only to a rapid change of $\hat{R}(t)$, approaching the true value R(t) exponentially fast, though.

5 Application to real data

- the **reporting delay** is chosen as the median incubation period plus 2, i.e. 7 days.

6 Sensitivity analysis

7 Discussion and Outlook

incubation period

structural assumption, isolation, medical treatment

infectivity profile vs amount of infectious material

weekday effects

reporting chain

infected vs reported

underreporting -; deaths, testing

mean, individual, stratification, regional effects get averaged

imported cases

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A Derivation of confidence intervals

Starting from Equation (4), the conditional expectation of $\hat{R}(t)$ given the past is

$$\mathbf{E}(\hat{R}(t) \mid I(t-1), \dots) = \frac{\mathbb{E}(I(t) \mid I(t-1), \dots)}{\sum_{\tau=-1}^{\infty} w(\tau) I(t-\tau)} = R(t). \tag{9}$$

Therefore, $\hat{R}(t)$ is **unbiased**,

$$\mathbf{E}\,\hat{R}(t) = R(t)\,,\tag{10}$$

and the **conditional variance** of $\hat{R}(t)$ is given by

$$\mathbf{Var}(\hat{R}(t) | I(t-1), \dots) = \frac{R(t)}{\sum_{\tau=1}^{\infty} w(\tau) I(t-\tau)}.$$
 (11)

An application of Slutsky's lemma gives an asymptotic $(1 - \alpha)$ -confidence interval for R(t): if q denotes the $(1 - \frac{\alpha}{2})$ -quantile of the standard normal distribution it is given by

$$\left[\hat{R}(t) - q\sqrt{\frac{\hat{R}(t)}{\sum_{\tau=1}^{\infty} w(\tau)I(t-\tau)}}, \hat{R}(t) + q\sqrt{\frac{\hat{R}(t)}{\sum_{\tau=1}^{\infty} w(\tau)I(t-\tau)}}\right]. \tag{12}$$

Note that (approximate) coverage is always guaranteed conditionally on the past, and hence also without conditioning.

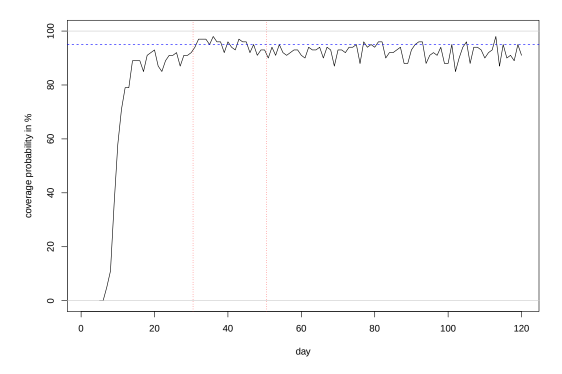


Figure 5: Estimated coverage probability based on 100 simulations (black solid line); horizontal blue dahed line: nominal coverage (95%); vertical red dotted lines: intervention times.

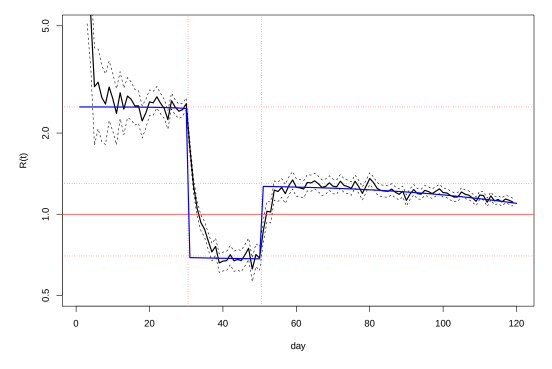


Figure 6: Estimator based on the newly infectious of one simulation of the SEIR model shifted by 1 day; black solid line: $\hat{R}(t)$; black dashed lines: pointwise 95%-confidence intervals; blue solid line: R(t); vertical red dotted lines: intervention times; horizontal red lines: corresponding reproduction numbers without decrease in susceptibles taken into account. This is to be compared with Figure 4 where the estimator is based on the newly infected cases.

B Derivation of the infectivity profile for the SEIR-model

Both latent period and infectious period are geometrically distributed with parameters $p_I = \frac{1}{t_I}$ and $p_R = \frac{1}{t_R}$, respectively. We need to compute the convolution (summing over time s of getting infectious). For $\tau > 1$ and assuming $p_I > p_R$ (the other cases are similar), we obtain

$$w(\tau) = \sum_{s=1}^{\tau-1} p_I (1 - p_I)^{s-1} (1 - p_R)^{\tau-1-s} = p_I (1 - p_R)^{\tau-2} \sum_{s=0}^{\tau-2} \left(\frac{1 - p_I}{1 - p_R}\right)^s$$

$$= p_I (1 - p_R)^{\tau-2} \frac{1 - \left(\frac{1 - p_I}{1 - p_R}\right)^{\tau-1}}{\frac{p_I - p_R}{1 - p_R}} = p_I (1 - p_R)^{\tau-1} \frac{1 - \left(\frac{1 - p_I}{1 - p_R}\right)^{\tau-1}}{p_I - p_R}$$

$$= p_I \frac{(1 - p_R)^{\tau-1} - (1 - p_I)^{\tau-1}}{p_I - p_R}.$$
(13)

References

- Abbott, S., Hellewell, J., Munday, J. D., Chun, J. Y., Thompson, R. N., Bosse, N. I., Chan, Y.-W. D., Russell, T. W., Jarvis, C. I., CMMID nCov working group, Flasche, S., Kucharski, A. J., Eggo, R. and Funk, S. (2020). Temporal variation in transmission during the COVID-19 outbreak. https://cmmid.github.io/topics/covid19/current-patterns-transmission/global-time-varying-transmission.html.
- Cori, A., Ferguson, N. M., Fraser, C. and Cauchemez, S. (2013). A New Framework and Software to Estimate Time-Varying Reproduction Numbers During Epidemics, *American Journal of Epidemiology* **178**(9): 1505–1512. https://doi.org/10.1093/aje/kwt133.
- Fraser, C. (2007). Estimating individual and household reproduction numbers in an emerging epidemic, *PLOS ONE* **2**(8): 1–12. https://doi.org/10.1371/journal.pone.0000758.
- Grassly, N. C., Fraser, C., Wenger, J., Deshpande, J. M., Sutter, R. W., Heymann, D. L. and Aylward, R. B. (2006). New strategies for the elimination of polio from India, *Science* 314(5802): 1150–1153. https://science.sciencemag.org/content/314/5802/1150.
- Johns Hopkins University Center for Systems Science and Engineering (2020). 2019 Novel Coronavirus COVID-19 (2019-nCoV) Data Repository by Johns Hopkins CSSE. https://github.com/CSSEGISandData/COVID-19.
- Organisation, W. H. (2020). Report of the WHO-China Joint Mission on Coronavirus Disease 2019 (COVID-19). https://www.who.int/publications-detail/report-of-the-who-china-joint-mission-on-coronavirus-disease-2019-(covid-19).
- Read, J. M., Bridgen, J. R., Cummings, D. A., Ho, A. and Jewell, C. P. (2020). Novel coronavirus 2019-nCoV: early estimation of epidemiological parameters and epidemic predictions. medRxiv, https://doi.org/10.1101/2020.01.23.20018549.
- Robert Koch-Institut (2020). RKI COVID19. dl-de/by-2-0, https://npgeo-corona-npgeo-de.hub.arcgis.com/datasets/dd4580c810204019a7b8eb3e0b329dd6_0.