#### L03 Reproduction numbers and their estimation<sup>1</sup>

#### Michael Höhle<sup>1</sup>

<sup>1</sup>Department of Mathematics, Stockholm University, Sweden

\*\* m hoehle

#### STA427 FS2021

Statistical Methods in Infectious Disease Epidemiology Epidemiology, Biostatistics and Prevention Institute University of Zurich, Switzerland



#### Outline

- 1 Effective Reproduction Number
- 2 Implementation and Results
  - Outbreak simulation
  - ullet R(t) for Switzerland
- Outlook

#### Overview

- Effective Reproduction Number
- 2 Implementation and Results
  - Outbreak simulation
  - R(t) for Switzerland
- Outlook

#### Outline

- Effective Reproduction Number
- Implementation and Results
- Outlook

#### Basic Reproduction Number

 In a previous lecture we considered the basic reproduction number  $R_0$ , which was given as<sup>2</sup>

$$R_0 = \beta/\gamma S(0)$$

- R<sub>0</sub> is the mean number of direct offspring one infectious case generates in a completely susceptible population without intervention measures
- However, once an epidemic unfolds susceptibles are depleted. Furthermore, interventions or behavioral changes aim at reducing  $\beta$ ,  $\gamma$  or the amount of susceptibles ( $\rightarrow$  vaccination).
- In order to assess public health interventions the basic reproduction number is not useful (Delamater, Street, Leslie, Yang, and Jacobsen 2019).

 $<sup>^2\</sup>beta$  was the infectious contact rate,  $\gamma$  the mean duration of the infectious period and S(0) is the number of susceptibles at time zero  $(S(0) \approx N)$ .

#### Effective Reproduction Number (1)

- We define the time-varying effective reproduction number R(t) as a time varying quantity denoting the average number of secondary cases generated by one case at calendar time t
- If contact and recovery rates do not vary with time we get

$$R(t) = R_0 \times \frac{S(t)}{S(0)}$$

Interventions could be reflected by extending this to

$$R(t) = (1 - c(t)) \times R_0 \times \frac{S(t)}{S(0)}$$

- The derivations in this lecture will follow Fraser (2007), who does not explicitly consider depletion of susceptibles
- Let the **transmissibility** function  $\beta(t,\tau)$  be a function of calendar time t and time since infection  $\tau$
- Dependence on  $\tau$  could, e.g., be due to time-dependence of viral shedding in the host, but also changes in contact behaviour after infection, depletion of susceptibles
- Renewal equation

$$I(t) = \int_0^\infty \beta(t, \tau) I(t - \tau) d\tau \tag{1}$$

## Effective Reproduction Number (3)

• Case reproduction number

$$R_c(t) = \int_0^\infty \beta(t+ au, au) d au$$

Interpretation: Average number of individuals an infected at time t can expect to infect

Instantaneous reproduction number

$$R(t) = \int_0^\infty \beta(t, \tau) d\tau$$

Interpretation: Average number of individuals an infected at time t is expected to infect should conditions remain unchanged

#### Effective Reproduction Number (4)

• Assuming the factorization

$$\beta(t,\tau) = \phi_1(t)\phi_2(\tau),$$

Fraser (2007) shows that by assuming  $\int_0^\infty \phi_2(\tau) d\tau = 1$  we get

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

• Here  $w(\tau)$  denotes the distribution of the new infections as a function of time since infection, i.e. the generation time distribution.

#### Effective Reproduction Number (5)

Inserting (2) into (1) yields:

• Instantaneous reproduction number

$$R(t) = \frac{I(t)}{\int_0^\infty I(t-\tau)w(\tau)d\tau}$$

Case reproduction number

$$R_c(t) = \int_0^\infty R(t+\tau)w(\tau)d\tau$$

#### Discretised Instantaneous Reproduction Number (1)

• If only discrete observations for time intervals  $t_i \equiv [t_i, t_{i+1}]$  are available, then one would use a discretized version of the instantaneous reproduction number

$$R(t_i) = \frac{I_i}{\sum_{j=0}^n w_j I_{i-j}}$$

• Note: the  $0 \le w_i \le 1$  now denote the probability mass function of a discretized version of the generation time, i.e.  $\sum_{i=0}^{n} w_i = 1$ .

Casting this into a distributional framework, e.g.,

$$I_i|I_{i-1},I_{i-1},\ldots \sim Po\left(R(t_i)\times \sum_{j=0}^n w_jI_{i-j}\right)$$

allows the consideration of  $R(t_i)$  as a parameter to be estimated in a statistical model

• This also allows the construction of confidence intervals or credibility regions for  $R(t_i)$ .

#### Smoothed Instantaneous Reproduction Number

- Estimate R(t) using discrete data is subject to fluctuations and it is therefore in general not recommended to just use one time point (Cori, Ferguson, Fraser, and Cauchemez 2013)
- $\bullet$  Suggested improvement: Compute instantaneous reproduction number over an interval of  $\tau$  days as follows

$$R_{\tau}(t_i) = \frac{\sum_{k=i-\tau+1}^{i} I_k}{\sum_{k=i-\tau+1}^{i} \sum_{j=0}^{n} w_j I_{k-j}}$$

• Note that this is slightly different from just using the mean of the  $\tau$  computed R(t) values, i.e.

$$\overline{R}_{ au}(t) = rac{1}{ au} \sum_{k=i- au+1}^{i} R(t_{i-k})$$

#### Superspreading

- So far we have considered the *mean* number of offspring one infectious case generates
- This view ignores the variability in the offspring distribution, e.g.,
   the larger the variation, the greater the probability of extinction
- Lloyd-Smith, Schreiber, Kopp, and Getz (2005) analysed that the 2002-2004 SARS-CoV-1 epidemic was driven by a small number of events where one case directly infected a large number of secondary cases, whereas many other cases did not infect any secondary cases
- If we can identify settings where super-spreading occurs, one can reduce the mean number of offspring substantially

#### Outline

- Effective Reproduction Number
- 2 Implementation and Results
  - Outbreak simulation
  - R(t) for Switzerland
- Outlook

## Outbreak simulation (1)

- Consider a growth model and denote by  $y_t$  the expected number of new symptom onsets we observe on day t.
- Let  $(w_1, \ldots, w_M)'$ , denote the probability mass function of the generation time distribution, i.e.  $P(GT = i) = w_i$  for  $i = 1, 2, \ldots, M$ .
- Assume that the expected number of cases can be described by the homogeneous linear difference equation

$$y_t = \sum_{i=1}^{M} R_c(t-i)w_i y_{t-i}, \quad t = 2, 3, \dots$$

• Note: We use the case reproduction number and ignore terms on the RHS when  $t-M \le 0$ .

## Outbreak simulation (2)

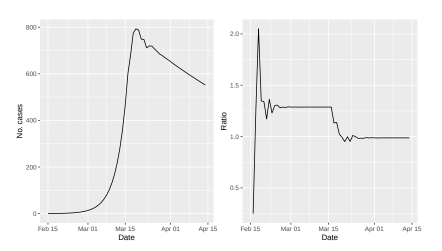
- Somewhat arbitrarily we fix  $y_1 = 1$  and conceptually denote by t = 1 the 15th of February 2020 in calendar time.
- To simulate a COVID-19 like outbreak with lockdown type intervention we use

$$R_e(t) = \left\{ egin{array}{ll} 2.5 & \mbox{if } t \leq 2020\mbox{-}03\mbox{-}15 \ 0.95 & \mbox{otherwise} \end{array} 
ight.$$

• We use the following generation time distribution

```
## 0 1 2 3 4 5 6 7
## 0.0 0.1 0.1 0.2 0.2 0.2 0.1 0.1
```

## Outbreak simulation (3)



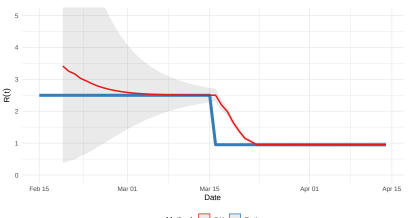
# EpiEstim (1)

- An implementation of the instantaneous reproduction estimate is available in the R package EpiEstim (Cori, Ferguson, Fraser, and Cauchemez 2013)
- Input is a 'data.frame' containing the column 'dates' and 'l' (for the incidence each day)
- It also computes credibility regions for R(t) and allows one to address uncertainty in the estimation of the generation time distribution

# EpiEstim (2)

```
library(EpiEstim)
# Rename data.frame columns to names handled by the EpiEstim pkg.
out_epiestim <- out %% rename(I = y, dates = Date) %>% select(dates, I)
# Estimate the instantaneous reproduction number
res <- EpiEstim::estimate_R(out_epiestim, method = "non_parametric_si",
                            config=make config(si distr=GT obj$GT,
                                               t_start=2:nrow(out_epiestim),
                                               t end=2:nrow(out epiestim))
# Convert result to a data.frame
rt irt df <- data.frame(Date=res$dates[res$R$t end],
                        R hat=res$R$`Mean(R)`.
                        lower=res$R$`Quantile.0.025`,
                        upper=res$R$`Quantile.0.975`,
                        Method="R(t)")
```

# EpiEstim (3)



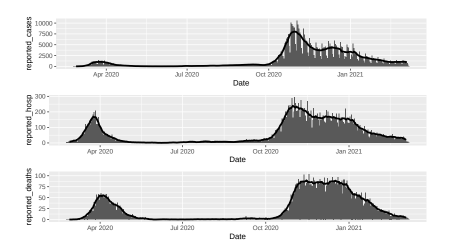
Method: R(t) Truth

## R(t) for Switzerland

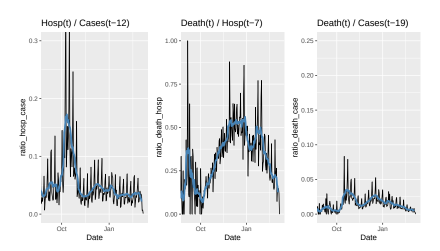
- Aim: Produce a crude R(t) for Switzerland based on the daily number of new cases as reported by the Swiss Federal Office of Public Health (FOPH)
- In order to do so, we first look a bit closer at the data as of 2021-03-10 available from https://www.covid19.admin.ch/

reported_cases	reported_hosp	reported_deaths
562364	23665	9294

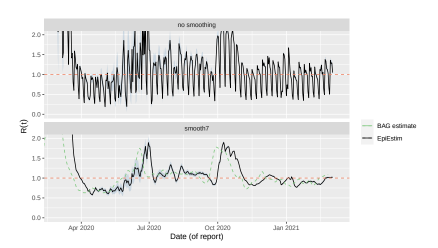
# COVID-19 Time Series for Switzerland (1)



# COVID-19 Time Series for Switzerland (2)



### Coarse R(t) Estimate (1)



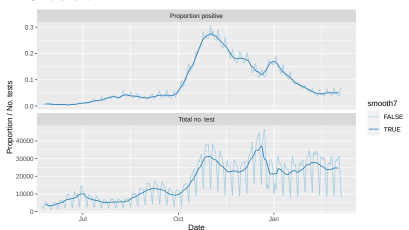
## Coarse R(t) Estimate (2)

#### Limitations:

- All cases are considered as community transmissions, but
  - there is an increase in the overall number of tests
  - some settings are tested more pro-actively than others (e.g. hospital staff)
  - imported cases are ignored (negligible from mid March)
- Serial interval
  - The analysis used an estimate from Nishiura, Linton, and Akhmetzhanov (2020) with mean 4.8 days and a standard deviation of 2.3 days
  - Difference between generation time and serial interval (infection vs. symptom onset - see Svensson (2007))
  - However, in this coarse analysis the event is actually "Day of report".
  - No specific Swiss serial interval was available to me and no estimation uncertainty is considered

## Coarse R(t) Estimate (3)

 Total number of tests and prroportion of positive tests over time in Switzerland



## Coarse R(t) Estimate (4)

- R(t) always needs context, e.g., the absolute number of (reported) cases.
- The implication of R(t) = 1.3 is very different when you have 10,000 cases or when you have 100.
- Especially when the case number is low, spikes due to outbreaks can lead to large R(t) fluctuations
- Be very careful when using R(t) graphs to assess interventions!

#### Outline

- Effective Reproduction Number
- 2 Implementation and Results
- Outlook

#### Outlook

- Good overview preprint about pitfalls when estimating R(t) by Gostic et al. (2020)
- Site estimating COVID-19 R(t) for countries around the world:
   https://epiforecasts.io/covid

#### Literature I



Cori, Anne, et al. 2013. "A New Framework and Software to Estimate Time-Varying Reproduction Numbers During Epidemics".

American Journal of Epidemiology 178 (9): 1505-1512.

doi:10.1093/aje/kwt133. eprint: http://aje.oxfordjournals.org/content/178/9/1505.full.pdf+html. http://aje.oxfordjournals.org/content/178/9/1505.abstract.



Delamater, P. L., et al. 2019. "Complexity of the Basic Reproduction Number ( $R_0$ )". 25 (1): 1. doi:10.3201/eid2501.171901.



Fraser, Christophe. 2007. "Estimating Individual and Household Reproduction Numbers in an Emerging Epidemic". PLOS ONE 2, no. 8 (): e758.



Gostic, Katelyn M, et al. 2020. "Practical considerations for measuring the effective reproductive number, Rt." PLoS Computational Biology 16, no. 12 (): e1008409. doi:10.1371/journal.pcbi.1008409.

#### Literature II



Lloyd-Smith, J. O., et al. 2005. "Superspreading and the effect of individual variation on disease emergence". <u>Nature</u> 438 (7066): 355–359. doi:10.1038/nature04153.



Nishiura, Hiroshi, Natalie M. Linton, and Andrei R. Akhmetzhanov. 2020. "Serial interval of novel coronavirus (COVID-19) infections".

International Journal of Infectious Diseases 93 (): 284–286. ISSN: 1201-9712, visited on 04/12/2020. doi:10.1016/j.ijid.2020.02.060.



Svensson, Å. 2007. "A note on generation times in epidemic models".  $\underline{\text{Math Biosci}}$  208, no. 1 (): 300–311.  $\underline{\text{doi:}}10.1016/\text{j.mbs.}2006.10.010$ .