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April 22, 2020

Abstract

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18 Introduction

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38 Models and Methods

39 **Data.** We use daily confirmed case data $\mathbf{X} = (X_1, \dots, X_T)$ from several different countries. These
 40 incidence data summarize the number of individuals X_t tested positive for SARS-CoV-2 RNA (using
 41 RT-qPCR) at each day t . Data was retrieved from **REFS** for the following regions: Wuhan, China;
 42 Austria; ...

43 **SEIR model.** We model SARS-CoV-2 infection dynamics by following the number of susceptible
 44 S , exposed E , reported infected I_r , and unreported infected I_u individuals in a population of size N .
 45 This model distinguishes between reported and unreported infected individuals: the reported infected
 46 are those that have enough symptoms to eventually be tested and thus appear in daily case reports, to
 47 which we fit the model.

48 Susceptible (S) individuals become exposed due to contact with reported or unreported infected
 49 individuals (I_r or I_u) at a rate β_t or $\mu\beta_t$. The parameter $0 < \mu < 1$ represents the decreased transmission
 50 rate from unreported infected individuals, who are often subclinical or even asymptomatic. The
 51 transmission rate β_t may change over time t due to behavioral changes of both susceptible and infected
 52 individuals. Exposed individuals, after an average incubation period of Z days, become reported
 53 infected with probability α_t or unreported infected with probability $(1 - \alpha_t)$. The reporting rate α_t may
 54 also change over time due to changes in human behavior. Infected individuals remain infectious for an
 55 average period of D days, after which they either recover, or becomes ill enough to be quarantined. They
 56 therefore no longer infect other individuals, and therefore the model does not track their frequency.
 57 The model is described by the following equations:

$$\begin{aligned}
 \frac{dS}{dt} &= -\beta_t S \frac{I_p}{N} - \mu\beta_t S \frac{I_u}{N} \\
 \frac{dE}{dt} &= \beta_t S \frac{I_p}{N} + \mu\beta_t S \frac{I_u}{N} - \frac{E}{Z} \\
 \frac{dI_r}{dt} &= \alpha_t \frac{E}{Z} - \frac{I_r}{D} \\
 \frac{dI_u}{dt} &= (1 - \alpha_t) \frac{E}{Z} - \frac{I_u}{D}.
 \end{aligned} \tag{1}$$

59 This model is inspired by Li et al.¹ and Pei and Shaman², who used a similar model with multiple
 60 regions and constant transmission β and reporting rate α to infer COVID-19 dynamics in China and
 61 the continental US, respectively.

Likelihood function. The *expected* number of new reported infected individuals on day t is

$$Y_t = \alpha_t E(t)/Z.$$

We define \tilde{Y}_t to be the cumulative expected number of reported infected individuals up to day t ,

$$\tilde{Y}_t = \sum_{i=1}^t Y_i$$

As mentioned above, X_t is the number of confirmed cases in day t . Then,

$$\tilde{X}_t = \sum_{i=1}^t X_i$$

is the cumulative number of confirmed cases until day t (with $X_0 = 0$). We assume that reported infected individuals yet to be confirmed, i.e. individuals in \tilde{Y}_t , are confirmed and therefore appear in

the daily case report of day t with probability p_t , which may change over time (note that t is a specific date, and not the elapsed time since infection). Therefore, we assume that the number of confirmed cases in day t is binomially distributed,

$$X_t \sim \text{Bin}(n_t, p_t),$$

where n_t is the *realized* number of reported infected individuals yet to appear in daily reports by day t . Given \tilde{X}_{t-1} , we assume n_t is Poisson distributed,

$$(n_t | \tilde{X}_{t-1}) \sim \text{Poi}(\tilde{Y}_t - \tilde{X}_{t-1}), \quad n_1 \sim \text{Poi}(Y_1).$$

62 Therefore, $(X_t | \tilde{X}_{t-1})$ is a binomial conditioned on a Poisson, which reduces to a Poisson with

$$63 \quad (X_t | \tilde{X}_{t-1}) \sim \text{Poi}((\tilde{Y}_t - \tilde{X}_{t-1})p_t), \quad X_1 \sim \text{Poi}(Y_1 p_1). \quad (2)$$

Therefore, for given vector θ of model parameters

$$\theta = \left(Z, D, \mu, \{\beta_t\}, \{\alpha_t\}, \{p_t\}, S(0), E(0), I_r(0), I_u(0) \right),$$

64 which also includes the initial conditions (state at $t = 0$), it is possible to compute the expected number
 65 of exposed $\{E(t)\}_{t=1}^T$ and number of new infections $\{Y_t\}_{t=1}^T$ for each day. Then, since \tilde{X}_{t-1} is a function
 66 of X_1, \dots, X_{t-1} , we can use Eq. (2) to write the probability of the confirmed case data $\mathbf{X} = (X_1, \dots, X_T)$
 67 as

$$68 \quad \mathbb{L}(\theta | \mathbf{X}) = P(\mathbf{X} | \theta) = P(X_1 | \theta)P(X_2 | X_1, \theta) \cdots P(X_T | X_1, \dots, X_{T-1}, \theta). \quad (3)$$

69 This defines our *likelihood function* for the parameter vector θ given the data \mathbf{X} .

71 Discussion

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90 **Acknowledgements**

91 This work was supported in part by the Israel Science Foundation 552/19 (YR) and XXX/XX (Alon Rosen)

92 **References**

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