

Improving the robustness of local R_t estimates during periods of geographical dropout

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Introduction

- General intro to R_t estimation
- Discussion of real-time
- Emphasis on “local” methodologies
- Extension to estimation across many locations simultaneously
- Not all locations have the same reporting frequency
- High-level description of the methods and a “killer figure” that shows what we do.

Results

This study could combine the algorithm maybe as a small R function that users can apply to data. There are several questions to investigate that could complete the paper.

- What is the value of combining local and global information relative to projecting forward locally? When should this work?
- How should we define the global scale? Perhaps pick the areas with a history of having similarly synched epidemics? Nearest neighbours?
- Given p locations in a region, how many can we tolerate simultaneous dropout in? What about other patterns? How long (relative to the generation time) does dropout need to be before there is no chance of correction?

Practically, we could combine some more simulated examples using a combination of the `rtestim` and `EpiFilter` code that we already have developed that can provide insights into when and how global information can supplement local data scarcity as well as provide guidelines for when this global information will not work. This could be about 2 figures. Then about 3 figures with the more impactful content that uses real examples, perhaps from the CFA pipeline or the other data sources we discussed. As baselines we can always compare to local forecasting and hold out real data to use for validation.

Discussion

Below are some ideas to describe as potential future work.

- Can we inform the additional parameter ϵ_h in some way to improve the correction? Does this require other data e.g., mobility?
- Potential to generalize the methods to more complicated incidence forecasting. View our simple procedure as a process with 2 modules, a `forecaster` and an `Rtestimator`. Then the meta procedure is
 1. Use `forecaster` to produce $\{\Lambda_{T+h,\ell}\}$ for $2 \leq h \leq H$. We do this conditional on all available information (incidence at the location, the region, or some related quantity).
 2. We can supplement these forecasts with auxiliary signals (wastewater, deaths etc.). Currently, we have a minimal `forecaster` (uses global R_t and local convolved incidence).
 3. Calculate $\hat{I}_{T+h,\ell}$ by taking the first (backward) differences of $\{\Lambda_{T+h,\ell}\}$.
 4. Use `Rtestimator` once across the pseudo time series to produce $\{R_{t,\ell}\}_{t=1}^{T+H}$.

Methods

Connecting local and global transmission

We consider $p \geq 2$ heterogeneous locations (the local scale) that compose a region of interest (the global scale). We assume that homogeneous mixing is valid in all locations. We neglect interactions among locations and do not model geographical movements of infections. As an example of this setup, if our global scale is at the state level, then our local scale may be at the county level. Our choice of scales is also determined by the availability of infection data.

We examine two periods of time (usually in units of days). In the first, which spans $1 \leq t \leq T$, we have access to the incidence of new infections (or a meaningful proxy such as reported

cases) in all locations of interest. We denote this $I_{t,\ell}$ for location ℓ at time t . The global incidence follows as the sum over locations and is written as $I_t = \sum_{\ell} I_{t,\ell}$. We also assume knowledge of the distribution of disease generation times, which is location independent and defined by probabilities ω_i such that $\sum_{i=1}^{\infty} \omega_i = 1$.

We can use the renewal transmission model to compute time-varying effective reproduction numbers at both local and global scales. At a location ℓ , this reproduction number is denoted $R_{t,\ell}$ and the corresponding global reproduction number is R_t . The renewal models are constructed as in (1) as a Poisson distribution describing intrinsic stochasticity in generating infections and the Λ terms defining the total infectiousness at a given scale.

$$\begin{aligned} I_{t,\ell} &\sim \text{Pois}(R_{t,\ell}\Lambda_{t,\ell}) \text{ with } \Lambda_{t,\ell} = \sum_{i=1}^{t-1} \omega_i I_{t-i,\ell} \quad (\text{local scale}) \\ I_t &\sim \text{Pois}(R_t\Lambda_t) \text{ with } \Lambda_t = \sum_{i=1}^{t-1} \omega_i I_{t-i} \quad (\text{global scale}) \end{aligned} \tag{1}$$

We can replace the Poisson distribution with any noise distribution of interest, but in this case it is possible to directly connect the transmission scales as $R_t = \sum_{l=1}^p \Lambda_{t,\ell} R_{t,\ell} (\sum_{m=1}^p \Lambda_{t,m})^{-1}$. Throughout this period up to T we are able to compute estimates from these models for all locations and globally. These result in local posterior distributions $\mathbb{P}(R_{t,\ell} \mid I_{1,\ell}^t)$ and the global posterior $\mathbb{P}(R_t \mid \sum_{l=1}^m I_{1,\ell}^t)$, which are directly obtained from a number of standard reproduction number estimators. Here, the notation $I_{1,\ell}^t$ means the past time series in location ℓ from times 1 to t .

The second period of interest is from times $T \leq t \leq T + H$. In this period there is dropout i.e., a loss of data in location ℓ . Consequently, we have no knowledge of $I_{T,\ell}^{T+h}$ and cannot assess the transmissibility in this location directly. We consider two options for providing estimates for location ℓ during this period. Across a horizon $1 \leq h \leq H$ we can either:

1. Project from our last local posterior $\mathbb{P}(R_{T,\ell} \mid I_{1,\ell}^T)$ to time $T + h$. This generates the baseline prediction $\mathbb{P}(R_{T+h,\ell} \mid I_{1,\ell}^T)$. For a short enough h this is expected to be reasonable. For large h it simply reproduces our prior information.
2. Borrow information from the global posteriors (or remaining locations with data) $\mathbb{P}(R_{T+h} \mid \sum_{m \neq \ell} I_{1,m}^{T+h})$. This essentially relies on some degree of correlation naturally occurring among locations due perhaps to similar drivers of spread. We do not assume knowledge of these correlations.
3. Use a mixture distribution of the above options.

Using global transmission to supplement dropout

Our main contribution lies in developing a sensible but real-time and computational simple algorithm to correct estimates in dropout locations. We propose to use the last local posterior in location ℓ as a prior and then draw information from the global posteriors (which exclude ℓ) to update this into the posterior distribution of (2).

$$\mathbb{P}\left(R_{T+1,\ell} \left| I_{1,\ell}^T, \sum_{m \neq \ell} I_{T+1,m} \right.\right) \propto \mathbb{P}\left(R_{T+1,\ell} \left| \sum_{m \neq \ell} I_{T+1,m} \right.\right) \mathbb{P}(R_{T+1,\ell} \mid I_{1,\ell}^T) \quad (2)$$

This follows due to the conditional independence of the infections in locations. Later time steps are obtained by iterating this process sequentially (with the prior reset to the dropout posterior from the last time step). We have not yet implemented this full version and so test a ‘deterministic dropout correction’ via the following sequence:

- Because location ℓ is unavailable for $t = T + 1, \dots, T + H$, we also cannot compute Λ_t for $t = T + 1, \dots, T + H$. Therefore, we calculate the **global incidence correction factor**: $\eta = 1 - \Lambda_{\ell,t}/\Lambda_t$.
- Compute global R_t by using the adjusted incidence $(I_1, \dots, I_t, I_{t+1}/\eta, \dots, I_{t+h}/\eta)$.
- Input $\{R_t\}_{t=1}^{T+H}$, $\{R_{t,\ell}\}_{t=1}^T$, $\Lambda_{\ell,t}$. For $h = 1, \dots, H$, do
 1. Predict $\hat{I}_{T+h,\ell} = R_{T+h-1}\Lambda_{T+h-1,\ell}$ (expectations from renewal models).
 2. Convolve $\Lambda_{T+h,\ell} = \sum_{i=1}^{T+h-1} \omega_i \tilde{I}_{T+h-i}$, where $\tilde{I}_j = I_j \mathbb{1}[j \leq T] + \hat{I}_j \mathbb{1}[j > T]$.
 3. Estimate local $\hat{R}_{T+h,\ell} = \epsilon_h R_{T+h} \frac{\Lambda_{T+h,\ell}}{\Lambda_{T+h+1,\ell}}$, with ϵ_h as a factor we currently set to 1.
- Given $\{\hat{I}_{\ell,t}\}$ and $\{\Lambda_{\ell,t}\}$ for $t = 1, \dots, T + H$, we recompute the local $R_{t,\ell}$ using both real ($t = 1, \dots, T$) and the estimated pseudodata ($t = T + 1, \dots, H$).