Mortality-MAP Analysis

1 Mortality-Mapping and Richardson-Lucy Deconvolution

Predicting cases from observed deaths can be framed as a deconvolution problem, where the observations (COVID-19 deaths) are convoluted signals of underlying parameters (COVID-19 cases that result in death) with known transition probabilities between the two states (the distribution of time from COVID-19 symptom onset to death). There has been research on deconvolution in signal processing, and our work, along with the deconvolution research for the 1918 flu [1], relies on research on deconvolution for Positron Emission Tomography [2]. In [2], it is shown that a Richardson-Lucy (RL) algorithm, or expectation maximization (EM) algorithm, converges to the maximum likelihood estimate of the unknown parameters. It can be shown that each iteration for \( mMAP \) is the same as the EM step presented in [2]. First note that each iteration step (ignoring the right-censoring for now) in \( mMAP \) is

\[
C_{d}^{(i)}(t) = \sum_{\tau=t+1}^{t_{\text{max}}} D(\tau) \cdot \frac{p(T = (\tau - t)) \cdot \frac{C_{d}^{(i-1)}(\tau)}{\sum_{t=1}^{\tau-1} p(T = (\tau - s)) \cdot \frac{C_{d}^{(i-1)}(s)}{\sum_{t=1}^{\tau-1} p(T = (\tau - s)) \cdot C_{d}^{(i-1)}(s)}}}{\sum_{s=1}^{t_{\text{max}}} p(T = (\tau - s)) \cdot C_{d}^{(i-1)}(s)} \tag{1}
\]

\[
= C_{d}^{(i-1)}(t) \sum_{\tau=t+1}^{t_{\text{max}}} D(\tau) \cdot \frac{p(T = (\tau - t))}{\sum_{s=1}^{t_{\text{max}}} p(T = (\tau - s)) \cdot C_{d}^{(i-1)}(s)} \tag{2}
\]

[2] discusses this problem in the context of reconstructing positron emission tomography (PET) images. They denote \( n^*(d) \) as the observed count of data in tube \( d \), \( \lambda(b) \) as the unobserved count of data in box \( b \), and \( p(b, d) \) as the probability of transmission from \( B \) to tube \( d \). They demonstrate that the expectation-maximization algorithm to solve the maximum likelihood estimate for the \( \lambda(b)'s \) involves this step:

\[
\lambda^{\text{new}}(b) = \lambda^{\text{old}}(b) \frac{D}{\sum_{d=1}^{D} n^*(d) p(b, d)} \tag{3}
\]

where \( D \) and \( B \) are the total numbers of tubes and boxes, respectively. Note that this step is the same as step (2) in \( mMAP \), replacing \( n^*(d) \) with observed deaths \( D(\tau) \), \( \lambda(b) \) with unobserved cases \( C(t) \), and \( p(b, d) \) with the transition probabilities \( p(T = (\tau - t)) \). Therefore, while both methods are presented differently, they rely on the same iterative process. The mathematical justifications provided for RL deconvolution in [2], specifically that the likelihood monotonically increases and that it is concave (so EM converges to a global maximum), can therefore be used to justify \( mMAP \).
2 Simulated and Empirical Validation

To validate mMAP, it was analyzed using simulated and real death data until June 7 from six countries: United States, Italy, Spain, Germany, Japan and South Korea. Fig A compares mMAP predicted cases with reported cases. To visually scale the reported cases, the following equation is used:

\[
\text{reported-scaled} = \frac{\sum \text{predicted}}{\sum \text{reported}} \cdot \text{reported}
\]

While the scales differ, the trends of predicted cases generally follow the trends of reported cases. The later increase of reported cases as opposed to predicted cases in most countries (except Germany) could be a result of increasing case detection around the start of April. As testing increases, we would expect to see more of a relative increase in reported cases than in reported deaths (because we would likely be picking up more of the less severe cases), which would cause the reported cases to increase more steeply than mMAP predictions.

In Fig B, the deaths for each country are simulated from the reported cases. Deaths are stochastically simulated from the reported cases using an sCFR of 0.013 and the same distribution from symptom onset to death used in the rest of this paper [3]. From the simulated deaths, mMAP predicts the original cases. This figure demonstrates that mMAP recovers the original cases reasonably well.

Both plots offer validation that mMAP can successfully predict the trend of the reported cases. However, these plots do not demonstrate if the scale of mMAP predictions are on target, as this is influenced by the under-reporting of deaths and the sCFR.

References


Figure A: $mMAP$ predictions compared to reported cases.

Figure B: Simulated $mMAP$ predictions compared to reported cases.