Excess death in relation to Covid-19

15 de May, 2020

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1 Problem

As of May 15th 2020, the global death toll of the unfolding COVID-19 outbreak stands at 302 493 (*citation needed*). The global and national official death toll figures have drawn much attention and sparked vivid debate because they are at the center of the evaluation and comparison of the public health responses of national and local governments.

Among the controversies, there is first a debate on the severity of COVID-19 when compared to the yearly flu outbreak. Brazil's President Jair Bolsonaro refered the COVID-19 as a 'little flu' and refused to implement in his country the drastic lockdown measures that many other countries have enforced (citation needed).

Second, the limits of testing in terms of tests availability and accuracy have led many observers to point at a likely underreporting of deaths due to the novel virus. Moreover, deaths that might be indirectly due to the COVID-19 crisis because of, for example, a collapse of the health system, are not counted in official figures. On April 26th 2020, the Financial Times headlined that global coronavirus death toll could be 60% higher than reported (*citation needed*).

Finally, differences in testing and reporting policies across countries but also regions have casted more doubts on the veracity of the reporting figures. Belgium has reported the highest number of deaths per 100 000 inhabitants but Belgian officials also say they are counting in a way that no other country in the world is

Table 1: Number of deaths by week in the Netherlands

week	X2019	X2018	X2017	X2020	average
2	3262	3359	3637	3364	3637
3	3150	3364	3487	3152	3487
4	3178	3322	3626	3041	3626
5	3143	3403	3574	3158	3574
6	3185	3513	3446	3189	3446
7	3252	3660	3417	3195	3417

currently doing: counting deaths in hospitals and care homes, but also including deaths in care homes that are suspected, not confirmed, as COVID-19 cases (citation needed).

There is a need of a rigorous estimation of the excess mortality in the weeks of the outbreak. A direct week by week comparison of the observed number of deaths to historical averages as done by the Financial Times analysts is a first approach but is deeply limited as it fails to consider the variance of the number of deaths across years.

In this report, we propose a Bayesian approach to estimate the excess mortality in the outbreak weeks through relative risk. Our model intends to provide parts of the answer to the following questions:

- Is mortality significantly higher than usual in weeks of the outbreak?
- If confirmed, is there significant excess mortality on top of the reported COVID deaths?

2 Data

2.1 Description

We gathered weekly data of the total number of deaths for weeks 2 to 17 in the following countries and following years:

• Norway: years 2014 to 2020

 \bullet Netherland: years 2017 to 2020

 \bullet Belgium: years 2009 to 2020

• Germany: years 2016 to 2020

• Switzerland: years 2015 to 2020

• Italy: years 2015 to 2020

• France: years 2010 to 2020

• England and Wales: years 2010 to 2020

The data have been retrieved from national statistic institutes of the respective countries. For each country the data were transformed to a similar structure than can be visualized below.

2.2 Exploratory data analysis

2.2.1 Visualization

Fig.1 and Fig.3 show the weekly deaths over weeks 1 to 17 for each of the countries and over the years. Most countries show the same pattern with a large peak between weeks 10 and 16 in 2020 which clearly stands out

of from other years. Notably, Switzerland show a peak in the same weeks in 2020 but it is not larger than other peaks observed in previous years. Norway shows no peak, and on a contrary it shows a trough, which might be the number of deaths for an incomplete week.

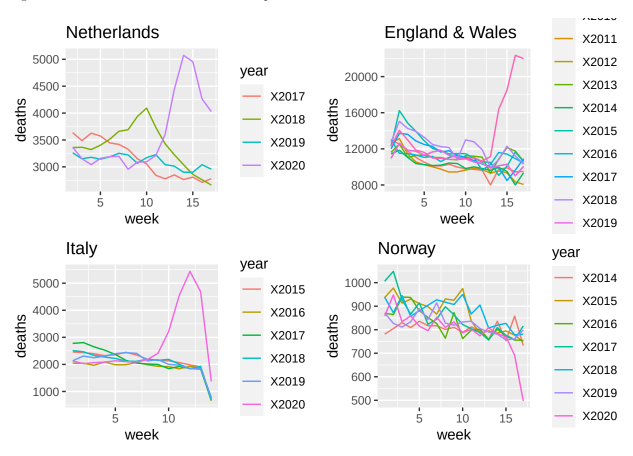


Figure 1: Weekly deaths by year and country

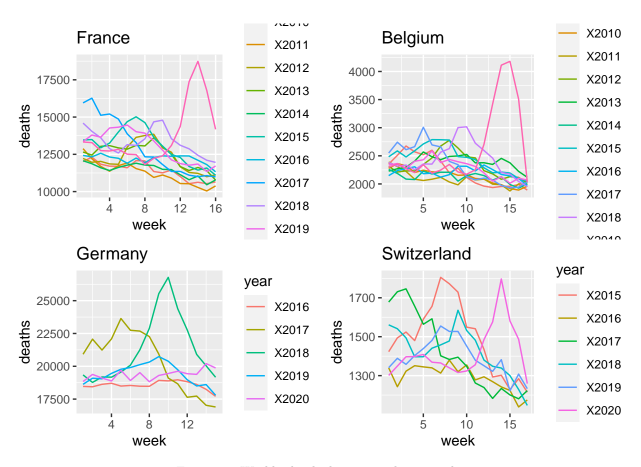


Figure 2: Weekly deaths by year and country bis

2.2.2 Overdispersion

Our data is characterized by much larger mean than variance. For example, Table 2 shows the weekly means and variance for Belgium.

Table 2: Means and variances of weekly deaths in France

3 Model

Mean

Variance

3.1 Excess mortality

3.1.1 Statistical model

We approach the modelling of excess mortality by estimating the relative risk θ understood as the ratio of weekly mortality during the COVID-19 outbreak and the weekly mortality in non-outbreak times.

$$RelativeRisk_i = \theta_i = \frac{mortality_{COV,i}}{mortality_{noCOV,i}} = \frac{\frac{O_{COV,i}}{N}}{\frac{O_{noCOV,i}}{N}} = \frac{O_{COV,i}}{O_{noCOV,i}}$$

where $O_{COV,i}$ is the observed number of deaths in week i of COVID outbreak, $O_{noCOV,i}$ the number of deaths for the same week in non-outbreak times and N is the population assumed stable across years.

The number of deaths O is a count variable. For such variable, two commonly used statistical models are the Poisson model (1) and the negative binomial model (2) that we parametrize with location and scale parameters.

(1)
$$O_{COV,i} \sim \mathcal{P}(\lambda), \ E(O_{COV,i}) = Var(O_{COV,i}) = \lambda$$

(2)
$$O_{COV,i} \sim \mathcal{NB}(\mu, \phi), \ E(O_{COV,i}) = \mu, \ V(O_{COV,i}) = \mu + \frac{\mu^2}{\phi}$$

A major difference between the two is that the Poisson model assumes the mean is equal to the variance while the negative binomial model has one more parameter that allows for overdispersion.

We have:

$$O_{COV,i} = O_{noCOV,i}\theta_i \Rightarrow E(O_{COV,i}) = E(O_{noCOV,i})\theta_i = E_i\theta_i$$

where E_i is the expected number of deaths in week i in non-outbreak times. For each week i, we estimate E_i as the historical average of number of deaths in year prior to 2020.

We therefore define the following statistical models:

(1)
$$O_{COV,i} \sim \mathcal{P}(E_i \theta_i)$$

(2)
$$O_{COV,i} \sim \mathcal{NB}(E_i\theta_i, \phi)$$

In those models, a value of θ_i larger than 1 can be interpreted as excess mortality in week i with respect to non-outbreak times and a value smaller than 1 as reduced mortality. This approach to modelling mortality is handy as the estimated value is standardized to the expected number of deaths and can therefore be compared between weeks and countries.

From the exploratory data analysis, we see that there is a clear time dependence in the weekly mortality. We then introduce a time structured random effect θ_t in our models. We use a autoregressive structure of order 1 as time structure. Our models are then as follows:

(1)
$$O_{COV,i} \sim \mathcal{P}(E_i \theta_i \theta_{t,i})$$

(2)
$$O_{COV,i} \sim \mathcal{NB}(E_i\theta_i\theta_{t,i},\phi)$$

where:

$$\theta_{t,i} = \alpha + \beta \theta_{t,i-1}$$

3.1.2 Bayesian model

We propose hierarchical models for our Bayesian estimation of θ and θ_t . Poisson model (1):

$$(O_{COV,2},...,O_{COV,17}|(\theta_2,\theta_{t,2}),...,(\theta_{17},\theta_{t,17})) \sim \prod_{i=2}^{17} \mathcal{P}(E_i\theta_i\theta_{t,i})$$

with parameters priors:

$$(\log(\theta_2), ... \log(\theta_{17})) \sim \mathcal{N}(0, \sigma)$$

$$\forall i, \log(\theta_{t,i}) = \alpha + \beta \log(\theta_{t,i-1}) + \epsilon, \ \epsilon \sim \mathcal{N}(0, \sigma_t)$$

where σ , α , β and σ_t are hyperparameters with hyperpriors:

$$\sigma \sim \mathcal{U}(a_{\sigma}, b_{\sigma})$$

$$\alpha \sim \mathcal{N}(\mu_{\alpha}, \sigma_{\alpha})$$
$$\beta \sim \mathcal{U}(a_{\beta}, b_{\beta})$$
$$\sigma_{t} \sim \mathcal{N}(\mu_{\sigma_{t}}, \sigma_{\sigma_{t}})$$

Similarly the negative binomial model (2) is:

Poisson model (1):

$$(O_{COV,2},...,O_{COV,17}|(\theta_2,\theta_{t,2}),...,(\theta_{17},\theta_{t,17})) \sim \prod_{i=2}^{17} \mathcal{P}(E_i(\theta_i\theta_{t,i}))$$

with parameters priors:

$$(\log(\theta_2), ..., \log(\theta_{17})) \sim \mathcal{N}(0, \sigma)$$

$$\forall i, \log(\theta_{t,i}) = \alpha + \beta \log(\theta_{t,i-1}) + \epsilon, \ \epsilon \sim \mathcal{NB}(E_i \theta_i \theta_{t,i}, \phi)$$

$$\phi \sim \mathcal{U}(a_{\phi}, b_{\phi})$$

where σ , α , β and σ_t are hyperparameters with hyperpriors:

$$\sigma \sim \mathcal{U}(a_{\sigma}, b_{\sigma})$$

$$\alpha \sim \mathcal{N}(\mu_{\alpha}, \sigma_{\alpha})$$

$$\beta \sim \mathcal{U}(a_{\beta}, b_{\beta})$$

$$\sigma_t \sim \mathcal{N}(\mu_{\sigma_t}, \sigma_{\sigma_t})$$

3.2 Excess mortality on top of the reported COVID deaths

To model excess mortality on top of reported COVID deaths, we use the same approach as for modelling of excess mortality during the COVID-19 outbreak with respect to non-outbreak times. We estimated the relative risk θ understood as the ratio of weekly mortality during the COVID-19 outbreak and the weekly expected mortality in non-outbreak times. We now adjust the expected mortality E_i to account for COVID deaths.

$$RelativeRisk_i = \theta_i = \frac{O_{COV,i}}{O_{noCOV,i} + D_i}$$

where $O_{COV,i}$ is the observed number of deaths in week i of COVID outbreak, $O_{noCOV,i}$ the number of deaths for the same week in non-outbreak times and D_i is the reported number of deaths due to COVID in week i.

$$\tilde{E}_i = E_i + D_i$$

4 Results

We have:

4.1 Poisson model

We started by estimating the above Poisson model in a simpler form, without the time dependence. The model resulted insuitable despite having tried numerous variations of the hierarchical model: changing the priors and levels of the multilevel model. Indeed, as can be observed in Fig. 3 for Belgium, the model adequately captured the change in relative risk in peak weeks but failed to adequately reflect variance with extremely narrow confidence intervals. We obtained similar results for the Netherlands, England & Wales, Italy and Germany. As a consequence, we abandonned the model.

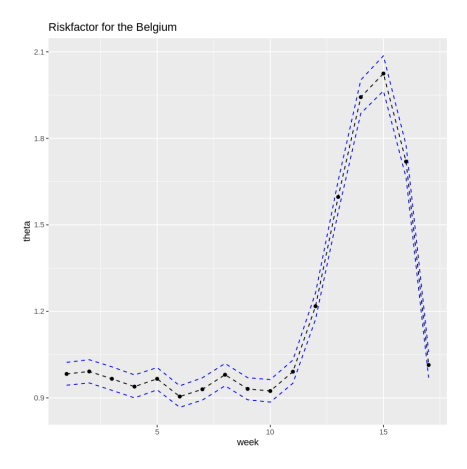


Figure 3: Poisson model estimates for Belgium

4.2 Negative binomial model

The negative binomial model proved to be much more suited to our data. This is not surprising givent the overdispersion we observed in the exploratory data analysis.

Reading the data

Extracting the data for the Netherland

Extracting data England and Wales

Extracting data Italy

Extracting the data for Norway

###Setting up the stan lists

Fit the stan models

There is a problem with divergency in our model meaning means and medians could be unreliable.

To explore this a bit more the pairs() function suggested by R are used.

5 Funtions

Funciton for outputgraphs

Function for creating the data frame with thetas and the bounds for the credible interval. #Results

5.1 Netherland

5.2 England and Wales

5.3 Italy

5.4 Norway

notice: negative risk in week 17!! Consistent with dat a:) (norway goes from 2-17 in weeks)

6 Future work

In the final report we will address the following points:

- convergence issues,
- implementation and obtention of results for the excess deaths on top of reported deaths,
- exploring the possiblity of an additional layer in our hierarchical negative binomail model by gathering the different countries into the same model,
- validation of the model,
- complete the EDA with missing countries