

Estimating the time-varying reproduction number of SARS-CoV-2 using national and subnational case counts

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Abstract

Background: Interventions are now in place worldwide to reduce transmission of the novel coronavirus. Assessing temporal variations in transmission in different countries is essential for evaluating the effectiveness of public health interventions and the impact of changes in policy.

Methods: We use case notification data to generate daily estimates of the time-dependent reproduction number in different regions and countries. Our modelling framework, based on open source tooling, accounts for reporting delays, so that temporal variations in reproduction number estimates can be compared directly with the times at which interventions are implemented.

Results: We provide three example uses of our framework. First, we demonstrate how the toolset displays temporal changes in the reproduction number. Second, we show how the framework can be used to reconstruct case counts by date of infection from case counts by date of notification, as well as to estimate the reproduction number. Third, we show how maps can be generated to clearly show if case numbers are likely to decrease or increase in different regions. Results are shown for regions and countries worldwide on our website (<https://epiforecasts.io/covid/>) and are updated daily. Our tooling is provided as an open-source R package to allow replication by others.

Conclusions: This decision-support tool can be used to assess changes in virus transmission in different regions and countries worldwide. This allows policymakers to assess the effectiveness of current interventions, and will be useful for inferring whether or not transmission will increase when interventions are lifted. As well as providing daily updates on our website, we also provide adaptable computing code so that our approach can be used directly by researchers and policymakers on confidential datasets. We hope that our tool will be used to support decisions in countries worldwide throughout the ongoing COVID-19 pandemic.

Keywords: Covid-19, SARS-CoV-2, surveillance, forecasting, time-varying reproduction number

Introduction

The coronavirus disease 2019 (COVID-19) pandemic that emerged in December 2019 has since spread to over 100 countries in every continent except Antarctica. While some information on the progress of an outbreak in a given country can be gained from the reported numbers of confirmed cases and deaths, these numbers can obscure changes in the underlying dynamics of the outbreak due to delays between infection and the eventual reporting of a case or death. Accounting for the delays from infection to symptom onset, and delays from symptom onset to hospital admission, diagnostic testing or potential death, followed by further delays until data are recorded in official statistics, requires the use of specific statistical methods for handling

right-truncated data [1–3] and the creation of a “nowcast” [4,5] (an estimate of the current number of newly infected or symptomatic cases).

Another method for tracking the progress of an outbreak is measuring changes in the time-varying reproduction number (effective reproduction number), which represents the average number of secondary infections generated by each new infectious case [6–8]. This approach can be advantageous compared to monitoring numbers of newly reported or symptomatic cases since, in principle, reproduction number estimates reflect variations in transmission intensity. Due to the delays in disease progression, recorded numbers of newly notified or symptomatic cases will increase or decrease for a period after transmissibility has reduced or increased, respectively. Monitoring changes in the time-varying reproduction can account for this delay and reveals variations in transmissibility that are not obvious from reported case numbers. Changes in the time-varying reproduction number can also quantify the impact of public health interventions [7,9]. Discerning whether or not current interventions are reducing transmission effectively using case notification data alone is challenging, since, case numbers may still be increasing while transmission is declining. Tracking the reproduction number over time may also be useful when relaxing interventions for the same reasons.

This paper details the methods we have developed for nowcasting and forecasting global time-varying reproduction number estimates, which are presented on a regularly updated website (<https://epiforecasts.io/covid/>). We first estimate cases by date of infection based on reported cases, accounting for right truncation and uncertainty in the reporting delay and incubation period. We then estimate the time-varying reproduction number and use an ensemble of time series models to forecast future changes in the reproduction number by extrapolating underlying temporal trends. We then reverse the process of estimating the reproduction number from cases by date of infection to derive forecasts of future reported cases by date of infection. Our estimates use reports of confirmed cases at the national, or sub-national, level that are extracted from publicly available repositories. This work builds on previously published tools [10,11] by adapting them for use on the currently available data. This overcomes some of the limitations of more naive implementations that derive estimates for the reproduction number directly from numbers of reported cases without adjusting (or with only partial adjustment) for the delay from infection to symptom onset or from onset to notification. The code that creates and updates the website is open source, allowing policymakers and researchers to run analyses in private repositories using confidential data. The methods outlined in this paper and corresponding code base are under development, and new versions of this live article will be released alongside changes to the methods to create a record of the methodology used throughout the pandemic.

Methods

Data

We use daily counts of confirmed cases reported by the European Centre for Disease Control for all analyses conducted at the national level [12,13]. To estimate the delay from symptom onset to reporting (once confirmed with a positive laboratory test), we use all cases from a publicly available linelist for which onset and notification dates are available [13,14]. This linelist combines all known linelist data from over 100 countries and at the time of writing has 4,132 entries with both an onset date and a notification date. Countries are only included in the reported estimates if at least 60 cases have been reported in a single day. This restriction reduces the likelihood of spurious estimates for countries with limited transmission or case ascertainment.

For sub-national analyses, the source of the data is reported on the respective page on our website. The data are fetched from government departments or from individuals who maintain a data source if no official data are available. Subnational entities within countries are only reported if at least 40 cases have been reported in a single day. A lower limit is possible for sub-national compared to national data due to more consistent case reporting in the source datasets.

All analyses described below are run independently for each national or subnational entity under consideration.

Adjusting for reporting delays

To estimate the reporting delay with appropriate uncertainty, we fit exponential and gamma distributions to 100 subsampled bootstraps (each with 250 samples drawn with replacement) of the delay between symptom onset and case notification, accounting for left and right censoring occurring in the data as each date is rounded to the nearest day and truncated to the maximum observed delay. We fit each model in the statistical modelling program *stan* [15] and compared to goodness-of-fit of each distribution to the data by comparing the approximate leave-one-out cross-validation information criterion (LOOIC) [16].

The distribution that gave the lowest LOOIC was selected as the most appropriate and 10 samples of the fitted distribution parameters were then drawn per bootstrap (giving 1000 in total). For a given country, we used sample i from the posterior distribution of delay distribution parameters, Θ_i , to draw a sample of delays, d_i , to transform each observed notification date, c_i , into a sample onset date, o_i , as follows:

$$o_i = c_i - d_i,$$

where $d_i \sim \exp(\Theta_i)$ or $\text{gamma}(\Theta_i)$

This resulted in 1000 date of onset samples for each confirmed case. For countries/regions with high case loads (more than 10,000 reported cases) the sampling step is approximated using the probability density function of the reporting delay.

Adjusting for right-truncation of notification dates

When moving from notification dates to onset dates it is important to consider that the total number of confirmed cases lags behind the number of cases that have onset, since there is a delay between onset occurring and the case being counted upon notification. To account for this right truncation, we used binomial upscaling to increase the estimated numbers of case onsets close to the present. After transforming the observed notification dates to onset dates and tallying case onset numbers by day, we then drew a sample of the number of case onsets that occurred but have not yet been confirmed.

If t is the last date on which cases were reported then the number of onsets on day $t - j$, denoted as o_{t-j} , is regarded as the result of a Bernoulli trial from the total true number of cases with symptom onsets on day $t - j$. o_{t-j} is then distributed according to a negative binomial distribution as follows:

$$o_{t-j}^* \sim \text{NegBin}(n = o_{t-k} + 1, p = F(t - j, \Theta_i))$$

where $F(t - j, \Theta_i)$ is the cumulative distribution function for the given delay distribution. It gives the proportion of onset cases from $t - j$ days ago that are expected to have been confirmed over the j days from that time until the present. The final numbers of case onsets that were used to estimate the time varying reproduction numbers for day t are consequently given by $o_t + o_t^*$. As our approach could not fully reconstruct unreported cases without bias we truncated our results and did not use estimates from the last 3 days. To prevent spurious estimates we truncated the allowed amount of upscaling to be less than 10 times the reported cases on any given day.

Adjusting for the delay between onset and infection

We repeated the steps outlined above for adjusting for the delay from report to symptom onset to account for the delay between symptom onset and infection by replacing the bootstrapped report delay distribution with an incubation period distribution with a mean of 5 days [17]. Uncertainty from this distribution was propagated through the model by sampling the distributions parameters assuming they were both normally distributed.

Estimating the time-varying reproduction number

We used the *EpiEstim* R package [6,10,18,19] to fit a model that estimated the time-varying reproduction number from the daily number of infections and an uncertain generation time with a mean of 3.6 days (sd: 0.7 days) and a standard deviation of 3 days (sd: 0.8 days). The generation time estimate was derived using the data and method of [20] modified to use the incubation period from [17]. The instantaneous reproduction number represents the number of secondary cases arising from an individual showing symptoms at a particular time, assuming that conditions remain identical after that time, and is therefore a measure of the instantaneous transmissibility (in contrast to the case reproduction number - see Fraser (2007) [8] for a full discussion). We used a gamma prior for the reproduction number with mean 2.6 and standard deviation 2. This is based on early estimates for the basic reproduction number (R_0) from the initial stages of the outbreak in Wuhan [21,22] with long tails to allow for differences in the reproduction number between countries. Our approach can also be used to account for imported cases where data is available [7].

We incorporated uncertainty in the generation time distribution by providing *EpiEstim* with 1000 samples each derived using a different sample of the log mean and log standard deviation of the assumed log normal distribution. We evaluated the reproduction number by assuming that it is constant over a backwards looking sliding time window [6]. We evaluated window lengths from 1 to 7 days, running *EpiEstim* separately for each window choice. The optimal time-varying window was selected by first estimating the one day ahead number of cases implied by each time-varying reproduction number estimate [23] and then scoring this nowcast against the observed number of cases using the ranked probability score (RPS) score [24,25]. For each sample the window with the lowest RPS score was selected at each time point.

The estimates of the time-varying reproduction number at each time point were combined over 1000 samples, using the optimal window for each, to give a credible interval that incorporates uncertainty from the delay from case onset to notification, the incubation period and the generation time.

Estimated change in daily cases

We defined the estimated change in daily cases to correspond to the proportion of reproduction number estimates for the current day that are below 1 (the value at which an outbreak is in decline). It was assumed that if less than 5% of samples were subcritical then an increase in cases was definite, if less than 20% of samples were subcritical then an increase in cases was likely, if more than 80% of samples were subcritical then a decrease in cases was likely and if more than 95% of samples were subcritical then a decrease in cases was definite. For countries/regions with between 20% and 80% of samples being subcritical we could not make a statement about the likely change in cases (defined as unsure).

As another metric of outbreak progression, we estimated the rate of spread (r) using a quasipoisson regression model [26]. The R^2 value of the regression fit was then used to assess the goodness-of-fit. In order to account for potential changes in the rate of spread over the course of the outbreak we used a 7-day sliding window to produce time-varying estimates of the rate of spread and the corresponding R^2 . The doubling time was then estimated by calculating $\ln(2)^{\frac{1}{r}}$ for each estimate of the rate of spread.

The effect of changes in testing procedure

The results presented here are sensitive to changes in COVID-19 testing practices and the level of effort put into detecting COVID-19 cases, e.g. through contact tracing. For example, if numbers of incident infections remain constant but a country begins to find and report a higher proportion of cases, then an increasing value of the reproduction number will be inferred. This is because all changes in the number of cases are attributed to changes in the number of infections resulting from previously reported cases, and are not assumed to be a result of improved testing and surveillance. On the other hand, if a country reports a lower proportion of cases because a lower number of tests are performed (which can happen if reagents required for testing are no longer available, for example) or the surveillance system captures a lower proportion of infections, then the model will attribute this to a drop in the reproduction number that may not be a true reduction.

In order for our estimates to be unbiased not all cases have to be reported, but the level of testing effort (and therefore the proportion of detected cases) must be constant [27]. This means that, whilst a change in testing effort will initially introduce bias, this will be reduced over time as long as the testing effort remains consistent from this point onwards.

Countries may also change the focus of their surveillance over the course of the outbreak. They may initially focus on identifying travellers returning from areas of known COVID-19 transmission and performing contact tracing on the contacts of known cases. As the outbreak evolves this may change to passive surveillance at hospitals. Here, the case definition may also change from tests based on polymerase chain reaction (PCR) to diagnoses based on symptoms and computed tomography (CT) scans. In the future, different kinds of COVID-19 tests may be deployed that could influence results, such as tests that detect both active and past infections.

Forecasting the reproduction number and case counts by date of infection

We forecast the time-varying effective reproduction number over a 14 day time horizon using the best performing ensemble of time series models [11] as assessed by iteratively fitting to a subsample of the estimated effective reproduction number estimates for each region [28]. Performance was assessed using CRPS scores, interval scores, PIT calibration, bias and sharpness with an ensemble being preferred that minimised the CRPS score whilst being calibrated, unbiased and as sharp as possible over the full time horizon [29–31]. The reproduction number forecast was then transformed into a case forecast using the renewal equation and a Poisson distribution of cases [6,19]. These forecasts are indicative only and should not be considered with a weight equal to the real-time estimates. Changes in contact rates, mobility, and public health interventions are not accounted for which may lead to significant inaccuracy.

Reporting

We report the median and 90% highest density credible intervals for all measures with 50% and 90% high density regions shown in figures. The analysis was conducted independently for all regions and is updated regularly as new data becomes available. As our credible intervals do not capture the proportion of cases that have been upscaled (when correcting for right truncation), we represent this in figures using translucency. This is presented as our confidence in the estimates which we define as the proportion of symptom onsets that are expected to have been reported by the date of estimation. All results are available in the source repository in a comma-separated values file.

Results

Daily updated estimates of the time-varying reproduction number, epidemic doubling time, and rate of spread at the national level are given for more than 90 countries on our website. New countries are being added as data become available. An example plot created on the 23rd of May 2020, showing the numbers of cases by date of infection for Austria and the inferred time-varying reproduction number, is shown in Figure 1.



Figure 1: Confirmed cases by date of report (bars; top) and their estimated date of infection (ribbon; top) and time-varying reproduction number (bottom) in Austria. The light and dark blue ribbons show the 90% and 50% credible interval, respectively. The estimates were generated on the 23rd May 2020. Due to the delay between being infectious and becoming a confirmed case, the estimates lag behind the present. Confidence in the estimated values is indicated by translucency with increased translucency corresponding to increased uncertainty deriving from right-truncation of reported cases in the estimate. The forecast (red) is coloured differently from the nowcast (blue) to indicate the drop in reliability when trying to forecast future cases.

Sub-national breakdowns can highlight differences in how the outbreak is progressing within a country (Figure 2) and are currently provided for 6 countries (Italy, Germany, United Kingdom, United States of America, Brazil, India). For example, on the 23rd of May 2020 we estimated that cases were likely decreasing in every country in the United Kingdom except Northern Ireland where the estimate was classified as unsure.



Figure 2: Estimates of numbers of cases that will be newly infected on the date the estimate was made and that will end up being reported (top panel), and the time-varying reproduction number (bottom panel) across different nations/regions of the United Kingdom. Estimates were produced on 23rd of May 2020. Nations/regions with fewer than 40 confirmed cases reported on a single day are not included in the analysis.

We present the nowcasting results on a map to effectively visualise regional differences in transmission (Figure 3). This helps identify areas where intervention policies are more effective at reducing transmission than others, which can inform decision-making going forwards.



Figure 3: A map of the estimates of the expected change in daily cases at the state level for the United States of America. Estimates were produced on 23rd of May 2020. Regions with fewer than 40 confirmed cases reported on a single day are not included in the analysis (light grey).

The methodology and toolset described here has also been used separately to produce estimates of the time-varying reproduction number at the state level in Australia, an example of how researchers and policymakers can apply the methods to their own data [32]. This has an added benefit of researchers being able to use generation time and delay distributions derived from local data and not our global estimates. The authors also use this tooling with confidential hospital admissions data to generate estimates of the time-varying reproduction number for policymakers in the United Kingdom.

Discussion

We provide a centralised resource, which generates comparable daily estimates of the time-varying reproduction number and a daily nowcast of the number of cases newly infected derived using a standardised method. The estimates are free of any hypotheses about the impact of interventions, since they are derived only from reported case counts and an estimate of the generation time. We explicitly account for the delay between infection and case notification and include all sources of quantifiable uncertainty. This resource may be useful for policymakers to track the progression of the COVID-19 outbreak and evaluate the effectiveness of intervention measures. As new data become available, we will include sub-national estimates for additional countries, and provide additional support for public health agencies or researchers interested in applying our methods to their data. We routinely utilise our own tooling to provide estimates of the reproduction number in the United Kingdom for policymakers using confidential hospital admissions data.

There are several advantages associated with our approach. Firstly, reported case counts are the only data required, which allows our approach to be used in a wide variety of contexts. Secondly, we apply the same methodology to all countries. This means that estimates can be compared without having to consider differences in the underlying methodology (even if differences in testing should still be accounted for as discussed below). Finally, we have constructed our approach using open source tools and all of our code, raw data, and results are available online. This means our approach can be applied by others to non-public data and be fully evaluated by end users.

Our approach is also subject to several limitation. Firstly, the model requires that the proportion of infections that are notified is constant. In other words, it requires consistency in the focus of the surveillance method, level of effort spent on testing, and case definition. Yet it is often the case that the level of under-reporting in a country changes over the course of an outbreak [27]. However, it should be noted that any changes in

surveillance testing procedures will only bias the estimates temporarily if they begin to remain consistent again after they have changed. How long the bias remains in the reproduction number estimates will depend on the serial generation time and delay distributions, as well as the maximum window size used in the reproduction number estimation process.

In addition, the model is limited by how representative the delay that we use from infection to notification distribution is for a given location. As there is limited data to assess this, we estimate a bootstrapped global delay distribution using the combined data from every country. In particular, the delay from onset to notification can especially impact the upscaling of cases by date of onset that accounts for cases that have onset but not yet been reported. If the true delay from onset to notification for a given country is shorter than our global delay, then we will overestimate onset case numbers, and vice versa for true delays longer than the distribution we used. Additionally, estimates of the reporting delay distribution are known to be biased early in an epidemic and may vary over time [33]. However, our use of a bootstrapped subsampling approach mitigates these issues by allowing multiple delay distributions based on the observed data to be considered at the cost of increasing uncertainty in our estimates.

Our model is also limited by the data available to us. For example, the publically available linelists contain little data on the importation status of cases. This means that cases counts may be biased upwards by attributing imported cases to local transmission. This bias is particularly problematic when case counts are low. Unfortunately, in the absence of data, this issue can only be explored via scenario analysis. However, if and when data on the importation of cases is available, our approach (via *EpiEstim* [7]) supports adjusting for imported cases.

As more data becomes available, future work should look to refine the distributions used for generation time, incubation period, and the report delay. There is also the potential to extend the present model to account for overdispersion in the number of secondary infections [34] and changes in the delay from onset to notification over the course of an outbreak. Finally, there is scope to explore how outbreak dynamics that differ among particular sub-populations, such as high-risk COVID-19 patients, can bias overall reproduction number estimates.

Our approach, providing real-time estimates of the reproduction number, serves as a valuable tool for decision makers looking to track the course of COVID-19 outbreaks. The nowcasts explicitly account for delays, using the same methodology across all countries and sub-national regions. These reproduction number estimates can be used during the initial stages of an outbreak to ascertain the likely outbreak trajectory if no interventions have been implemented. They can also provide real-time feedback on whether transmission is decreasing following a particular intervention, or whether it is increasing following the relaxing or lifting of current intervention measures. We hope that our website and the related toolkit will provide a valuable resource for devising strategies to contain COVID-19 outbreaks worldwide.

Data availability

Latest data: <https://github.com/epiforecasts/covid>

Archived data at the time of publication: <https://doi.org/10.5281/zenodo.3841818>

License: MIT

Software availability

Development

- Website (Front-end): <https://github.com/epiforecasts/covid>
- *EpiNow* R package (R estimation, data processing, visualisation and reporting): <https://github.com/epiforecasts/EpiNow>

- *EpiSoon* R package (forecasting and case prediction from R trajectories): <https://github.com/epiforecasts/EpiSoon>
- *NCoVUtils* R package (data aggregation and processing): <https://github.com/epiforecasts/NCoVUtils>

Archived at the time of publication

- Website: <https://doi.org/10.5281/zenodo.3841818>
- *EpiNow* R package [35]: <https://doi.org/10.5281/zenodo.3833806>
- *EpiSoon* R package [28]: <https://doi.org/10.5281/zenodo.3833807>
- *NCoVUtils* R package [13]: <https://doi.org/10.5281/zenodo.3833808>

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References

- 1 Linton NM, Kobayashi T, Yang Y *et al*. Incubation period and other epidemiological characteristics of 2019 novel coronavirus infections with right truncation: A statistical analysis of publicly available case data. *Journal of clinical medicine* 2020;**9**.
- 2 Cori A, Donnelly CA, Dorigatti I *et al*. Key data for outbreak evaluation: Building on the ebola experience. *Philosophical transactions of the Royal Society of London Series B, Biological sciences* 2017;**372**.

- 3 Mizumoto K, Kagaya K, Zarebski A *et al.* Estimating the asymptomatic proportion of coronavirus disease 2019 (covid-19) cases on board the diamond princess cruise ship, yokohama, japan, 2020. *Eurosurveillance : bulletin Europeen sur les maladies transmissibles = European communicable disease bulletin* 2020;**25**.
- 4 Donker T, Boven M van, Ballegooijen WM van *et al.* Nowcasting pandemic influenza a/h1n1 2009 hospitalizations in the netherlands. *European journal of epidemiology* 2011;**26**:195–201.
- 5 Kasstele J van de, Eilers PHC, Wallinga J. Nowcasting the number of new symptomatic cases during infectious disease outbreaks using constrained p-spline smoothing. *Epidemiology (Cambridge, Mass)* 2019;**30**:737–45.
- 6 Cori A, Ferguson NM, Fraser C *et al.* A new framework and software to estimate time-varying reproduction numbers during epidemics. *American Journal of Epidemiology* 2013;**178**:1505–12. doi:10.1093/aje/kwt133
- 7 Thompson RN, Stockwin JE, Gaalen RD van *et al.* Improved inference of time-varying reproduction numbers during infectious disease outbreaks. *Epidemics* 2019;**29**:100356. doi:https://doi.org/10.1016/j.epidem.2019.100356
- 8 Fraser C. Estimating individual and household reproduction numbers in an emerging epidemic. *PloS one* 2007;**2**:e758.
- 9 Camacho A, Kucharski A, Aki-Sawyer Y *et al.* Temporal changes in ebola transmission in sierra leone and implications for control requirements: A real-time modelling study. *PLoS currents* 2015;**7**.
- 10 Cori A. *EpiEstim: Estimate time varying reproduction numbers from epidemic curves*. 2019. <https://CRAN.R-project.org/package=EpiEstim>
- 11 Shaub D, Ellis P. *ForecastHybrid: Convenient functions for ensemble time series forecasts*. 2020.
- 12 Disease Prevention EC for, Control. Download today’s data on the geographic distribution of covid-19 cases worldwide. 2020. www.ecdc.europa.eu/en/publications-data/download-todays-data-geographic-distribution-covid-19-cases-worldwide
- 13 Abbott S, Hellewell J, Munday JD *et al.* NCoVUtils: Utility functions for the 2019-ncov outbreak. - 2020;-. doi:10.5281/zenodo.3635417
- 14 Xu B, Gutierrez B, Hill S *et al.* Epidemiological data from the nCoV-2019 outbreak: Early descriptions from publicly available data. <http://virological.org/t/epidemiological-data-from-the-ncov-2019-outbreak-early-descriptions-from-publicly-available-data/337>
- 15 Stan Development Team. RStan: The r interface to stan. 2020.<http://mc-stan.org/>
- 16 Vehtari A, Gelman A, Gabry J. Practical bayesian model evaluation using leave-one-out cross-validation and waic. *Stat Comput* 2016;**1**–20. doi:10.1007/s11222-016-9696-4
- 17 Lauer SA, Grantz KH, Bi Q *et al.* The incubation period of coronavirus disease 2019 (covid-19) from publicly reported confirmed cases: Estimation and application. *Annals of Internal Medicine* 2020;**172**:577–82.
- 18 R Core Team. *R: A language and environment for statistical computing*. Vienna, Austria.: R Foundation for Statistical Computing 2019. <https://www.R-project.org/>
- 19 Wallinga J, Teunis P. Different epidemic curves for severe acute respiratory syndrome reveal similar impacts of control measures. *American Journal of Epidemiology* 2004;**160**:509–16. doi:10.1093/aje/kwh255
- 20 Ganyani T, Kremer C, Chen D *et al.* Estimating the generation interval for coronavirus disease (covid-19) based on symptom onset data, march 2020. *Eurosurveillance* 2020;**25**.
- 21 Imai N, Cori A, Dorigatti I *et al.* Report 3: Transmissibility of 2019-nCoV. <https://www.imperial.ac.uk/media/imperial-college/medicine/sph/ide/gida-fellowships/Imperial-2019-nCoV-transmissibility.pdf>
- 22 Abbott S, Hellewell J, Munday J *et al.* The transmissibility of novel coronavirus in the early stages of the 2019-20 outbreak in wuhan: Exploring initial point-source exposure sizes and durations using scenario analysis. *Wellcome open research* 2020;**5**:17.

- 23 Nouvellet P, Cori A, Garske T *et al.* A simple approach to measure transmissibility and forecast incidence. *Epidemics* 2018;**22**:29–35. doi:<https://doi.org/10.1016/j.epidem.2017.02.012>
- 24 Gneiting T, Raftery AE. Strictly proper scoring rules, prediction, and estimation. *Journal of the American Statistical Association* 2007;**102**:359–78. doi:10.1198/016214506000001437
- 25 Jordan A, Krüger F, Lerch S. Evaluating probabilistic forecasts with scoringRules. *Journal of Statistical Software* 2019;**90**. doi:10.18637/jss.v090.i12
- 26 Park SW, Champredon D, Weitz JS *et al.* A practical generation-interval-based approach to inferring the strength of epidemics from their speed. *Epidemics* 2019;**27**:12–8. doi:<https://doi.org/10.1016/j.epidem.2018.12.002>
- 27 Russell TW. Using a delay-adjusted case fatality ratio to estimate under-reporting. https://cmmid.github.io/topics/covid19/severity/global_cfr_estimates.html
- 28 Abbott S, Hellewell J, Bosse N *et al.* *EpiSoon: Forecast cases using reproduction numbers*. 2020. <https://epiforecasts.io/EpiSoon>
- 29 Jordan A, Kruger F, Lerch S. Evaluating probabilistic forecasts with scoringRules. *Journal of Statistical Software* 2019;**90**:1–37. doi:10.18637/jss.v090.i12
- 30 Bosse N. *Scoringutils: A collection of proper scoring rules and metrics to assess predictions*. 2020. <https://github.com/epiforecasts/scoringutils>
- 31 Funk S, Camacho A, Kucharski AJ *et al.* Assessing the performance of real-time epidemic forecasts: A case study of ebola in the western area region of sierra leone, 2014-15. *PLoS computational biology* 2019;**15**:e1006785–17.
- 32 Price DJ, Shearer FM, Meehan MT *et al.* Early analysis of the australian covid-19 epidemic. 2020;1–22.
- 33 Britton T, Scalia Tomba G. Estimation in emerging epidemics: Biases and remedies. *Journal of the Royal Society, Interface* 2019;**16**.
- 34 Endo A, Abbott S, Kucharski AJ *et al.* Estimating the overdispersion in covid-19 transmission using outbreak size outside china [version 1; peer review: 1 approved]. *Wellcome open research* 2020;**5**.
- 35 Abbott S, Hellewell J, Munday J *et al.* *EpiNow: Estimate realtime case counts and time-varying epidemiological parameters*. 2020. <https://github.com/epiforecasts/EpiNow>