Real-time estimation of the effective reproduction number CNVAC 2022 Seminar series

Sam Abbott @seabbs samabbott.co.uk





Case studies

- Who am I
- Overview
- Challenges
- Tools for estimation
- Uses
- What is left to do?
- Summary

Real-time estimation of the effective reproduction number?
Real-time analysis case studies

Who am I?

Sam Abbott @seabbs samabbott.co.uk





I'm an infectious disease researcher interested in real-time analysis, forecasting, semi-mechanistic modelling, and open source tool development.

- Research Fellow at the London School of Hygiene and Tropical Medicine
- Member of epiforecasts.io
- PhD in modelling Tuberculosis and the optimal use of the BCG vaccine
- Background in Mathematical modelling and mathematics



Sam Abbott

Sam Abbott







I'm an infectious disease researcher interested in real-time analysis, forecasting, semi-mechanistic modelling, and open source tool development. More on my research interests here.

Hi there 🁋

- . Working at the London School of Hygiene and Tropical Medicine in the Epiforecasts group;
- Crafting extensions to forecast vocs
- 14 Crafted last epinowcast . Currently working on:
- · Estimation of the test to test distribution as a proxy for generation interval distribution for the Omicron variant in England
- · Real-time estimation of the time-varying transmission advantage of Omicron in England using S-Gene Target Status as a Proxy
- · Evaluating the use of real-time sequences for short-term forecasting · Evaluating a new method for nowcasting right truncated count data

- . I'm currently working at London School of Hygiene and Tropical Medicine
- . I did my PhD at the University of Bristol
- . I use daily: R, stan
- III like to perform analysis using novel models on interesting data and generalise those approaches into software . Of m mostly active within the R Community
- Learning all about Julia and Turing.jl
- Reading all of China Miéville's work.
- Ping me about statistical modelling of infectious diseases, real-time analysis of infectious diseases, estimating transmission dynamics in realtime, and team science opportunities
- ■Reach me: sam.abbott@lshtm.ac.uk

samabbott.co.uk

Real-time estimation of the effective reproduction number?
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Overview

Sam Abbott @seabbs samabbott.co.uk

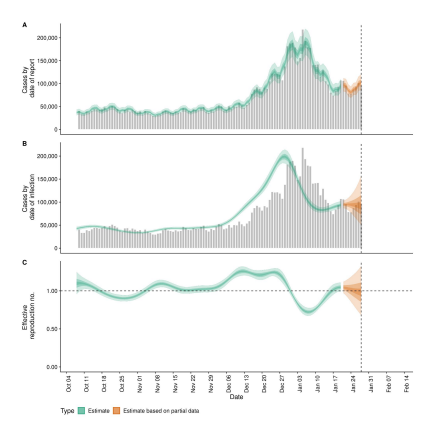




An 80% right paper before a policy decision is made is worth ten 95% right papers afterwards, provided the methodological limitations imposed by doing it fast are made clear.

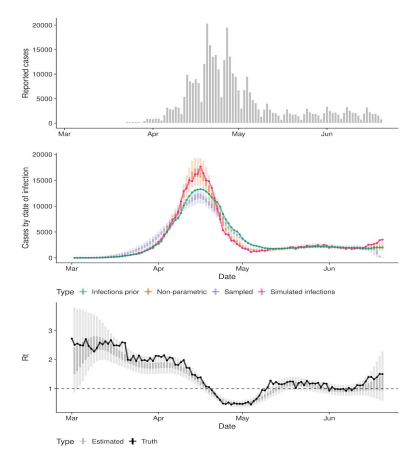
The average number of secondary infections produced by a single infected person.

- A helpful metric to track transmission.
- Difficult to estimate as depends on unobserved infections and on the interval between primary and secondary infections.
- Estimating using various methodologies since February 2020.
- Estimates submitted as part of the SPI-M consensus estimate each week.
- Estimates were also published each day for over 1000 locations since April 2020 on epiforecasts.io/covid.



Key Challenges

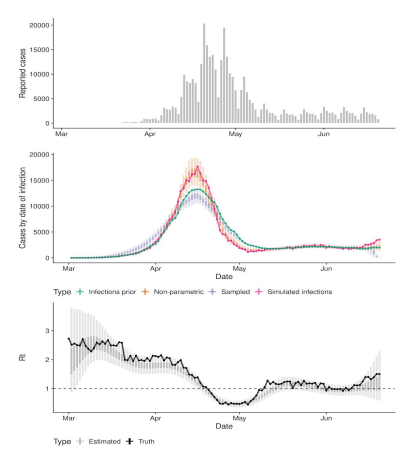
- We care about linking policy changes with changes in transmission but only observe delayed proxies like reported cases and deaths.
- Surveillance data subject to a range of difficult to account for biases.
- Estimation at scale in real-time is computationally challenging.



Sherratt et al. Phil Trans B, 2021, DOI: 10.1098/rstb.2020.0283

Objectives

- Develop a model that can be used for real-time surveillance, nowcasting, and short-term forecasting.
- Include known epidemiological structure of the infection and reporting process.
- The model should include a parameter that is referenced to the infection process and that can be used to compare disparate surveillance data sources (here the effective reproduction number).
- The model should ideally capture changes in trend as quickly as possible to make it useful for situational awareness.



Sherratt et al. Phil Trans B, 2021, DOI: 10.1098/rstb.2020.0283

EpiNow2 1.3.3.10 Home News 🖟 Functions

EpiNow2: Estimate real-time case counts and timevarying epidemiological parameters

License MIT contributors 12 PRs welcome commits since v1.3.2 85 DOI 10.5281/zenodo.5036949

This package estimates the time-varying reproduction number, growth rate, and doubling time using a range of open-source tools (Abbott et al.), and current best practices (Gostic et al.). It aims to help users avoid some of the limitations of naive implementations in a framework that is informed by community feedback and is under active development.

How do we define the reproduction number?

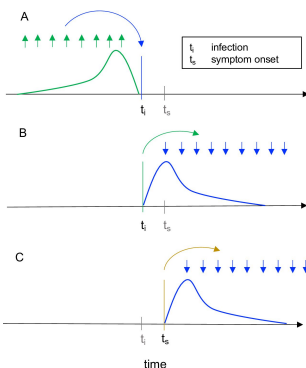
$$R_t^{\text{inst}} = \beta(t)S(t)D$$

$$R_t = \frac{I_t}{\sum_{s=1}^t I_{t-s} w_s},$$

$$R_t^{\text{case}} = \int_{u=t}^{\infty} R_u^{\text{inst}} w(u-t) du$$

$$p_{ij} = \frac{w(t_i - t_j)}{\sum_{i \neq k} w(t_i - t_k)}$$

$$R_j = \sum_i p_{ij}$$



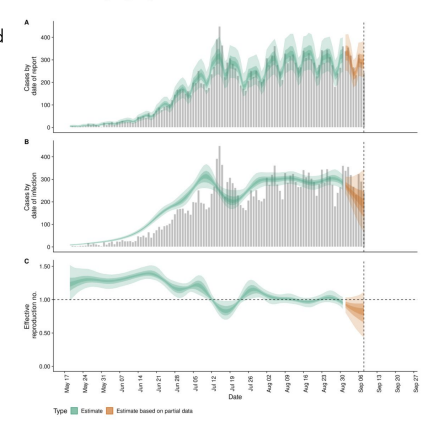
Practical considerations for measuring the effective reproductive number, R_t

Katelyn M. Gostic , Lauren McGough, Edward B. Baskerville, Sam Abbott, Keya Joshi, Christine Tedijanto, Rebecca Kahn, Rene Niehus, James A. Hay, Pablo M. De Salazar, Joel Hellewell, Sophie Meakin, James D. Munday, [...], Sarah Cobey [view all]

General approach

- Bayesian approach combining nowcasting and *R* estimation
- Uncertain generation interval estimates allowing for negative serial interval
- Latent process for estimating I_{t}
- Negative binomial reporting with multiplicative day-of-the-week effect
- R_t estimates with correlation between R_{t+1} and R_t based on Gaussian Process prior

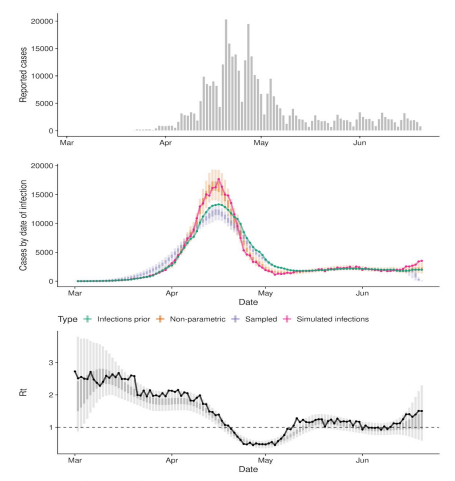
Confirmed cases, their estimated date of report, date of infection, and time-varying reproduction number estimates



The model

$$egin{aligned} \log R_t &= \log R_{t-1} + \mathrm{GP}_t \ I_t &= R_t \sum_{ au=1}^{15} w(au | \mu_w, \sigma_w) I_{t- au} \ O_t &= \sum_{ au=0}^{15} \xi_O(au | \mu_{\xi_O}, \sigma_{\xi_O}) I_{t- au} \ D_t &= lpha \sum_{ au=0}^{15} \xi_D(au | \mu_{\xi_D}, \sigma_{\xi_D}) O_{t- au} \ C_t &\sim \mathrm{NB}\left(\omega_{(t \mod 7)} D_t, \phi
ight) \end{aligned}$$

Method: doi.org/10.12688/wellcomeopenres.16006.2 Stan code: git.io/JUxRt



Type + Estimated + Truth

The model



$$egin{aligned} I_t &= I_0 \exp(rt) \ I_0 &\sim \mathcal{LN}(\log I_{obs}, 0.2) \ r &\sim \mathcal{LN}(r_{obs}, 0.2) \end{aligned}$$

20000

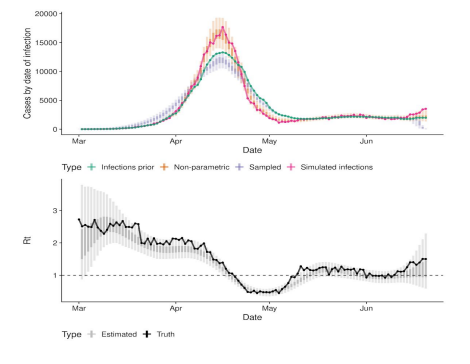
Seported cases 150000

5000

Mar

Apr

$$egin{aligned} w &\sim \mathcal{G}(\mu_w, \sigma_w) \ \xi_O &\sim \mathcal{LN}(\mu_{\xi_O}, \sigma_{\xi_O}) \ \xi_D &\sim \mathcal{LN}(\mu_{\xi_D}, \sigma_{\xi_D}) \end{aligned}$$

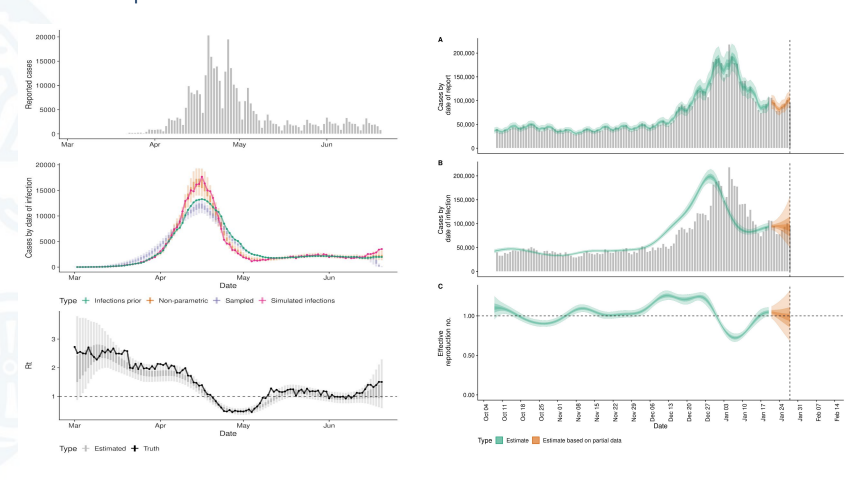


May

Jun

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Challenges

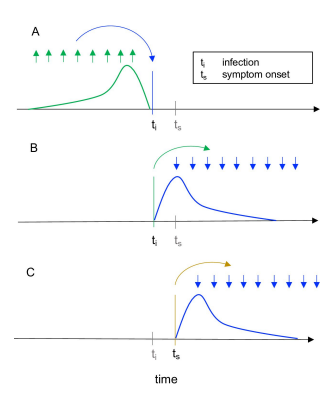
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What are we actually trying to do?

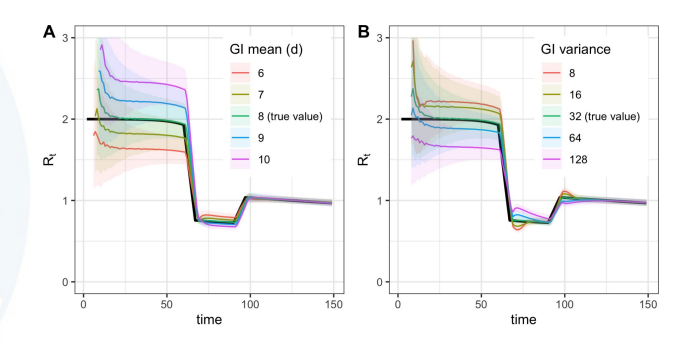
- What kind of reproduction number are we estimating (case, instantaneous etc).
- Are we trying to estimate the reproduction number for the true latent infection process or using observed data as a proxy?
- Do we want a non-parametric estimate, a forecast of future transmission, or an understanding of the impact of NPIs etc.
- What do we a priori believe about how the reproduction number and generation time evolve over time.



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Generation interval misspecification

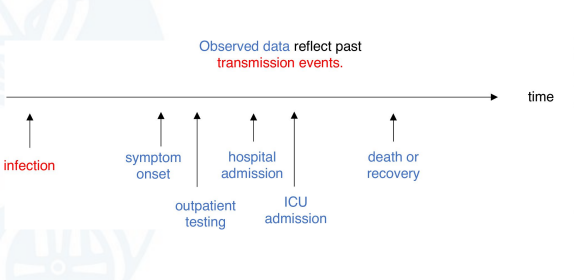


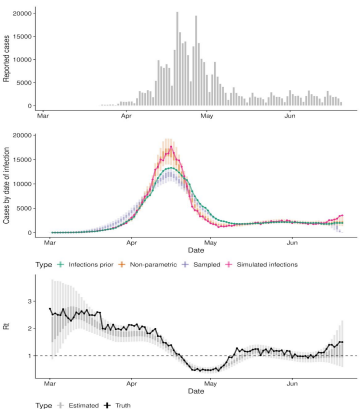
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Adjusting for delays

We observe some later proxy for infections blurred by a delay distribution.





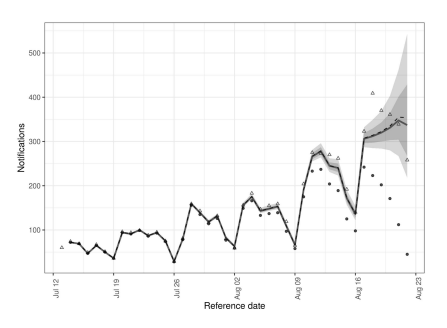
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Adjusting for right truncation

Incoming data can be delayed meaning what we currently observed isn't what we will observe.

Equally as infections are observed via a delay after reconstructing them they are also not fully observed.

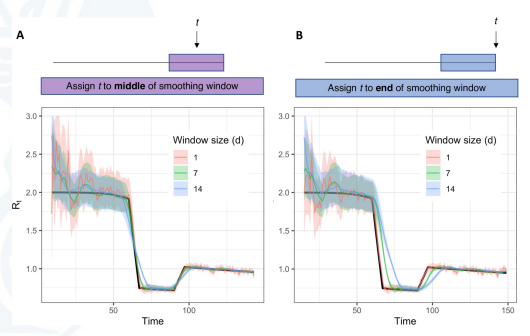


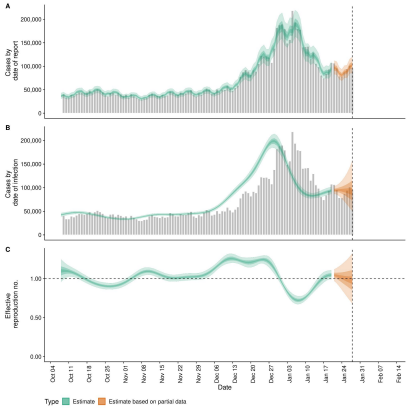
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How to smooth

How does the reproduction number evolve over time and how much of that do we want to include in our estimates?





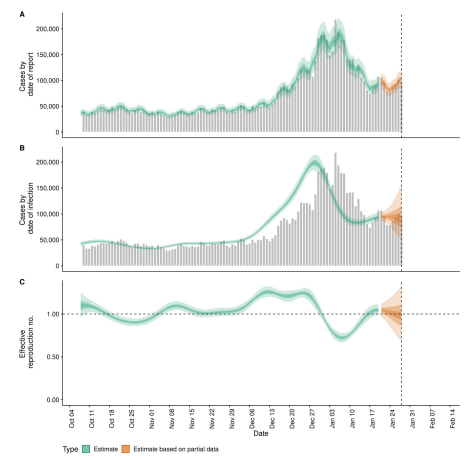
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Retrospective or real-time?

Real-time = Orange. This is expensive to care about.

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Tools for estimation

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EpiNow2 - Real-time non-parametric estimation

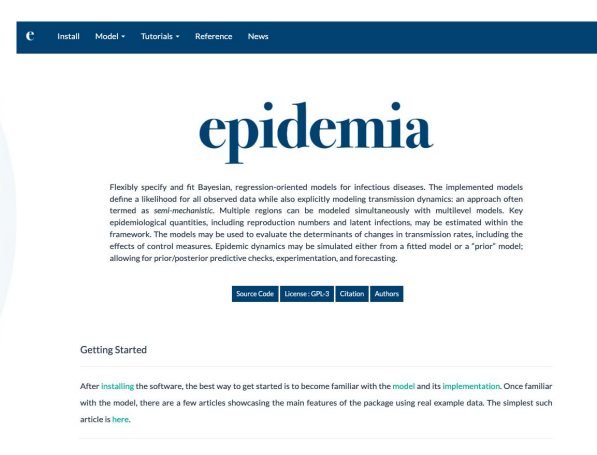
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Epidemia - Hierarchical regression + flexible model options



0

EpiEstim - The core model, well implemented

EpiEstim: Estimate Time Varying Reproduction Numbers from Epidemic Curves

Tools to quantify transmissibility throughout an epidemic from the analysis of time series of incidence as described in Cori et al. (2013) < doi:10.1093/aje/kwt133> and Wallinga and Teunis (2004) < doi:10.1093/aje/kwh255>.

Version: 2.2-4 Depends: R (≥ 2.10)

Imports: coarseDataTools (≥ 0.6-4), stats, graphics, reshape2, ggplot2, gridExtra, fitdistrplus, coda, incidence (≥ 1.7.0), scales, grDevices

Suggests: testthat, utils, vdiffr, covr, knitr, rmarkdown

Published: 2021-01-07

Published: 2021-01-

Author: Anne Cori 👩 [aut, cre], Simon Cauchemez [ctb], Neil M. Ferguson 👩 [ctb], Christophe Fraser 👩 [ctb], Elisabeth Dahlqwist 👩 [ctb], P. Alex Demarsh [ctb], Thibaut Jombart 👩 [ctb], Zhian N. Kamvar 👩 [ctb], Justin Lessler 👩 [ctb], Shikun Li [ctb], Jonathan A.

Polonsky @ [ctb], Jake Stockwin [ctb], Robin Thompson @ [ctb], Rolina van Gaalen [ctb]

Maintainer: Anne Cori ca.cori at imperial.ac.uk>
BugReports: https://github.com/mrc-ide/EpiEstim/issues
License: GPL-2 | GPL-3 | expanded from: GPL (2 2) |
URL: https://github.com/mrc-ide/EpiEstim

NeedsCompilation: no

Materials: README NEWS
In views: Epidemiology
CRAN checks: EpiEstim results

Documentation:

Reference manual: EpiEstim.pdf

Vignettes: <u>EpiEstim demonstration</u>

Downloads:

Package source: <u>EpiEstim 2.2-4.tar.gz</u>

Windows binaries: r-devel: EpiEstim 2.2-4.zip, r-release: EpiEstim 2.2-4.zip, r-oldrel: EpiEstim 2.2-4.zip

 $macOS\ binaries: \ r-release\ (arm64): \underline{EpiEstim\ 2.2-4.tgz}, r-oldrel\ (arm64): \underline{EpiEstim\ 2.2-4.tgz}, r-oldrel\ (x86_64): \underline{EpiEstim\ 2.2-4.tgz}, r-$

Old sources: <u>EpiEstim archive</u>

Reverse dependencies:

Reverse imports: <u>covid19india</u>, <u>earlyR</u>, <u>EpiLPS</u> Reverse suggests: <u>epidemia</u>, <u>projections</u>

Linking:

Please use the canonical form https://cran.r-project.org/package=EpiEstim to link to this page.

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Case studies

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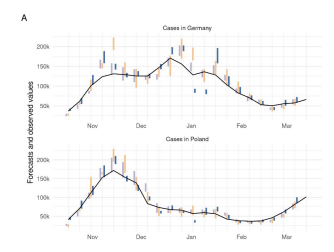


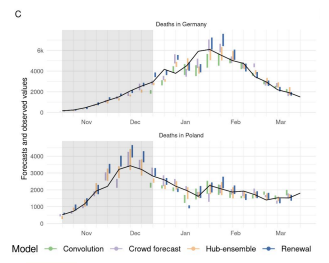


Short-term forecasts

What do we think will happen to reported metrics over the next 1-4 weeks

- Using the reproduction number model and similar discrete convolutions models.
- Submitting to the ECDC and CDC forecasting hubs weekly as well as to SPI-M 3 times a week until mid 2020.
- Performs well compared to other models but out performed by an all-model ensemble.
- Also outperformed by a human judgement model.
- All forecasts struggle to account for policy changes and behavioural changes.



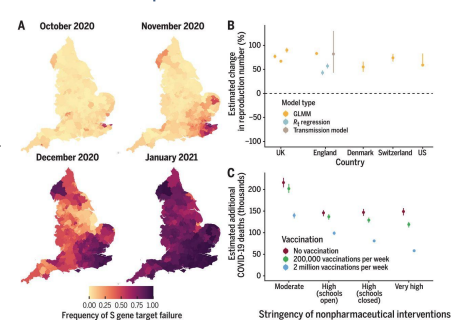


Bosse et al. medRxiv, 2020, DOI: 10.1101/2021.12.01.21266598

Estimating the transmission advantage of Alpha

How much more transmissible is Alpha than wild-type?

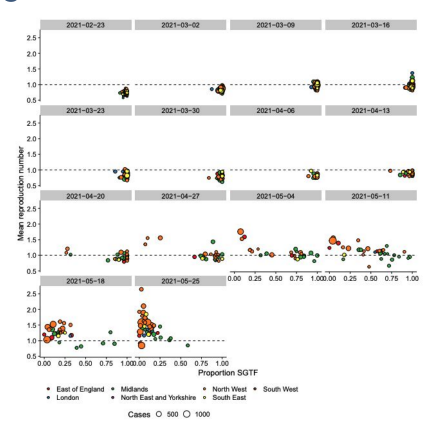
- Part of a multi-method approach by Davies et al.
- Used reproduction number estimates by Lower-Tier local authority as "data" + S-gene target failure status (SGTF) as a proxy for variant status.
- Estimated the transmission advantage using an extended regression model adjusted for confounders
- Work done between December 20th 2020 and January 1st 2021.



Estimating the transmission advantage of Delta

How much more transmissible is Delta than Alpha?

- Repurposed the approach used for Alpha using reproduction number estimates as data.
- Extended the methodology to include uncertainty for the reproduction number estimates and SGTF status.
- Estimates combined with others as apart of the SPI-M consensus statement.

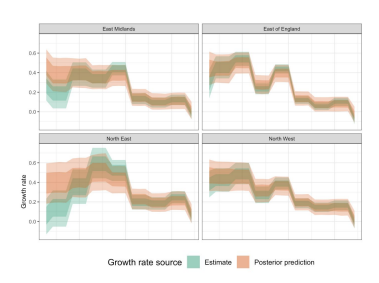


Estimating the generation time of Omicron

Is Omicron's generation time shorter than Delta's?

- Observed reduction in transmission advantage could indicate a shorter generation time.
- This is due to the relationship between the daily growth rate and the reproduction number.
- Used growth rates for Omicron and Delta to explore this and found that a shorter generation time was plausible.
- Findings supported by a study from UKHSA using household contact data.
- Results available in early January and formed part of the SPI-M consensus statement.

$$R = \left(1 + \bar{G}kr\right)^{\frac{1}{k}}$$



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What is left to do?

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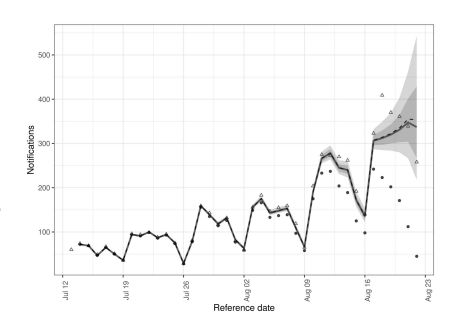




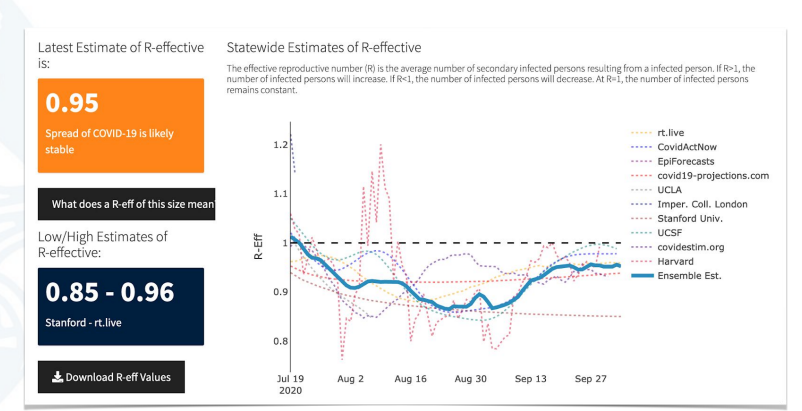
Nowcasting

What is happening now to metrics we partially observe

- International collaboration estimating 7-day hospital admissions by date of positive test in Germany (the key metric used by decision makers).
- Statistical approach is to decompose the model into forecast and reporting delay components.
- Multi-method ensemble outperforms any single model.
- Nowcasting is conceptually difficult to understand and so potentially under used.
- Open access tools, such as epinowcast, make access easier.



Validation - which method works best when



Novel data sources

Estimating epidemiological quantities from repeated cross-sectional prevalence measurements

© Sam Abbott, © Sebastian Funk

doi: https://doi.org/10.1101/2022.03.29.22273101

Estimating epidemiologic dynamics from cross-sectional viral load distributions



Authors Info & Affiliations

SCIENCE · 3 Jun 2021 · Vol 373, Issue 6552 · DOI: 10.1126/science.abh0635

Other

- Documentation, testing, case studies, modularity.
- Joint estimation of the time-varying generation time.
- Estimation and support for time-varying delays.
- Better understanding of how the reproduction number should be modelled to evolve over time.
- Hybrid models producing optimal retrospective and real-time estimates
- Linking stochastic and deterministic models
- Linking to phylodynamic methods

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Summary

Thanks to the epiforecast.io group and my collaborators. Please see individual slides for links containing more details of each case study.