<u>Dynamics of B.1.617.2 in the UK from importations, traveller-linked and non-traveller-linked transmission</u>

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Summary

- We used a deterministic approximation of a simple continuous time branching process model to combine estimates for the number of imported B.1.617.2 cases in the UK from India with local onwards transmission, then fitted this model to reported COVID-19 cases up to 17th May 2021 and B.1.617.2 sequences in COG-UK data up to 11th May 2021 to estimate importation rate, UK-based transmission, and rate of decline of non-B.1.617.2 cases in the UK. We stratified transmission so that travellers to India and non-travellers could have different values of R, with R_{traveller} ≥ R_{non-traveller} to reflected potential for early amplification that does not persist in wider community transmission.
- Based on importations, local sequences of B.1.617.2 and overall case patterns, we estimated that R_{traveller} was 3.3 (95% Crl: 1.8-10) and R_{non-traveller} was 1.7 (95% Crl: 1.5-1.8) in the UK assuming no change in generation interval. Consistent with our previous report, we predicted the majority of cases in the UK would consist of B.1.617.2 by mid-May 2021 (which should show up clearly in COG-UK data in the coming week). This corresponds to a median doubling time of 7.1 days for B.1.617.2 in the period since 1st May 2021.
- Note that these preliminary estimates of R for B.1.617.2 reflect the average level of transmission across the specific settings where this variant is currently circulating. In particular, the relatively large estimate of R_{traveller} possibly reflects higher levels of within household transmission and lower levels of vaccine coverage in specific communities. As a result, these estimates may not generalise to other areas in the UK if there are specific risk factors for elevated transmission in areas where B.1617.2 is being reported, or additional control measures being introduced or relaxed. Analysis and model structure will continue to be refined as more data become available.

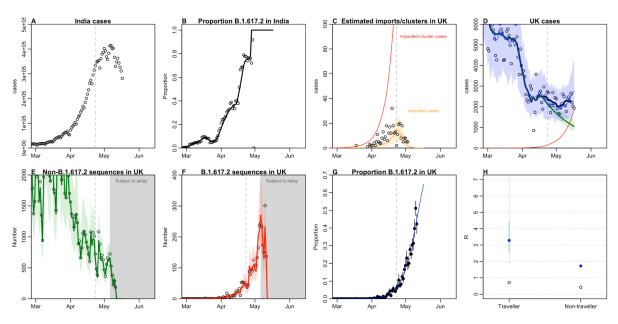
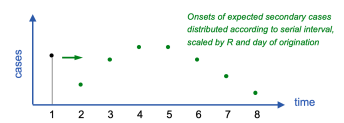


Figure 1: A) Reported cases in India. B) Proportion of reported sequences in India that are B.1.617.2, with black line showing moving average (constrained to end at 100%). C) Estimated imported cases of B.1.617.2 into the UK that contribute to onwards transmission (orange line, with 95% shaded CrI interval), reported traveller cases of B.1.617.2 in PHE Technical Report 11 (black dots); simulated imported cases and onwards transmission using maximum a posteriori (MAP) model estimate (red line). D) Reported cases in the UK. Black dots show data, black line shows 7 day centred moving average; green line shows estimated non-B.1.617.2 cases with 95% CrI; red line as in (C); blue line and shaded region shows predicted total cases in UK with negative binomial 95% CrI. E) Black dots show number of non-B.1.617.2 sequences in COG-UK data up to 11th May 2021; green line shows fitted model with 95% negative binomial CrI. Grey region shows data in the past week, which is likely subject to reporting delays. F) Black dots show number of B.1.617.2 sequences in COG-UK data up to 11th May 2021; red line shows fitted model with 95% negative binomial CrI. G) Black dots show proportion of B.1.617.2 sequences in COG-UK data up to 11th May 2021; blue line shows MAP model estimate. H) Estimate of R_{traveller} and R_{non-traveller} in model, with thick line showing 50% CrI and thin line showing 95% CrI. Dots show implied R based on contact tracing data in PHE Technical Report 11.

Methods

- We estimated imported cases of B.1.617.2 into the UK by combining two data sources: reported cases in India, proportion of sequenced cases that were B.1.617.2 in India. We then scaled these estimates by a parameter a_{import} to produce an expected number of importations over time. Details of the two data sources:
 - Reported cases in India from 1st February 2021 onwards were downloaded using from the covidregionaldata R package (Figure 1A).
 - Proportion of sequences in India were based on sequences reported in GISAID, aggregated by <u>outbreak.info</u> (Figure 1B). Note that these are based on relatively low numbers of sequences collected, which may not be representative, and assumed to converge to 100% eventually.
- We assumed that all imported infections from India ceased after the red listing on 23rd April 2021 (i.e. no leaks from hotel quarantine), and assumed a lognormal incubation period with mean 5.1 days and s.d. log(1.65) (McAloon al, BMJ Open, 2020) to estimate onsets occuring after the red list date among travellers (Figure 1).
- As a validation, we compared estimated imported to traveller cases reported during the same period (<u>PHE Technical Report 11</u>). These corresponded closely to our estimates (Figure 1C).
- To estimate overall B.1.617.2 cases resulting from initial imports, we used a deterministic approximation of a continuous time branching process model (<u>Kucharski et al. EID. 2016</u>), with the serial interval assumed to be lognormal with mean = 5.4 days and s.d. log(1.5) (<u>Rai et al. Clin Epi Glob Health. 2021</u>). The expected secondary number of cases from onsets on each day is iteratively propagated forward, with transmission depending on R and temporal pattern based on the serial interval. A schematic of this process is illustrated below:



- To estimate non-B.1.617.2 cases in the UK, we calculated the 7 day centred moving average of cases overall in the UK up to 23rd April 2021 given fluctuations in day-to-day reporting, then extrapolated forward based on the value of an exponential daily decline, a_{decline}, which was fitted.
- We estimated a vector θ of five parameters ($a_{decline}$, a_{import} , $R_{traveller}$, $R_{non-traveller}$, σ_{report}) by simulating case trajectories from the model, then calculating two likelihoods:
 - the negative binomially distributed log likelihood of sequencing the number of B.1.617.2 cases reported in reality in COG-UK data on a given day i (y_{ib}), given the mean number of B.1.617.2 onsets on each day in the simulated outbreak ($E(x_{ib})$) and overall number cases by date of specimen collection in the UK (y_{io}) and the number of cases sequenced (y_{in}). The negative binomial distribution had dispersion parameter σ_{report} . Specifically:

$$L_1(\theta) = \sum_i \log NB(x = y_{ib} \mid mu = E(x_{ib}) y_{in} / y_{io}$$
, size = $1/\sigma_{report}$)

the negative binomially distributed log likelihood of the number of overall cases reported in reality on a given day (y_{io}) , given the mean number of B.1.617.2 onsets on each day in the simulated outbreak $(E(x_{ib}))$ and estimated non-B.1.617.2 cases in the UK (y_{ic}) from 1st April 2021 onwards. We assumed the negative binomial distribution also had dispersion parameter σ_{report} . Specifically:

$$L_2(\theta) = \sum_i \log NB(x = y_{io} \mid mu = E(x_{ib}) + y_{ic}, size = 1/\sigma_{report}))$$

- We then calculated the overall log likelihood as L(θ) = L₁(θ) + L₂(θ), and estimated the parameters using MCMC (adaptive Metropolis-Hastings, implemented with the doMC R package). We estimated a daily decline of 3.0% (95% CrI: 2.0-3.8%) for non-B.1.617.2 cases (Figure 1D). This is consistent with the SPI-M consensus estimate for England on 23rd April (i.e. –5 to –1%)
- The posterior model estimates are shown in Figure 1C–F, with comparison to COG-UK data and overall cases.