

# The probability that a previously susceptible individual is infected as a function of time after exposure to SARS-CoV-2

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## Abstract

We consider an individual who, following a specific exposure to SARS-CoV-2, is symptom free and has a negative test and calculate the probability that they are in fact in the latent state or infectious at the time of the test. We parameterize the probability of actual infection as a consequence of the exposure and, in particular, compare the certainly infected and certainly uninfected cases. The results shed light on the questions of how long to isolate and when to test an individual following an exposure event.

## 1 Introduction

Throughout this work we use *infected* to mean that an individual is either in the latent state or one of the active states of the disease.

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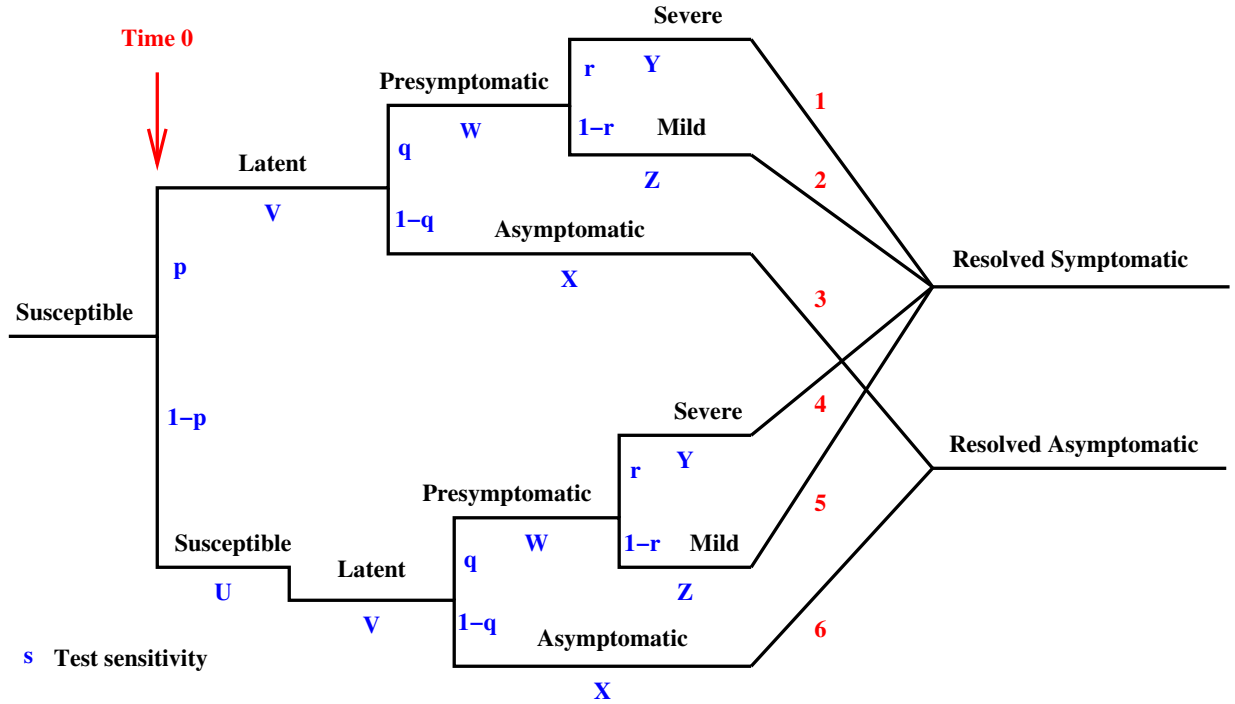
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## 2 Methods

Figure 1 gives a schematic of the possible paths through a course of infection following an exposure to SARS-CoV-2 . With probability  $p$  the exposure results in an infection and the individual enters the latent state and progresses through subsequent phases. With probability  $1 - p$  the individual remains susceptible until they become infected through general background exposure. Note that we distinguish two resolved states depending on whether or not the individual previously experienced symptoms. This is the sole distinction between these states and is introduced only as an artifice to simplify the computations described below. Probability parameters control the path taken, and the time in state variables are all assumed to be Gamma distributed.

Figure 1: Model for the course of infection. Letters representing the random variables specifying the time in each state and branching probabilities are shown in blue. The red numbers enumerate the possible paths.



For the most part, to model the effects of the original variants of SARS-CoV-2 , we chose values for the path splitting parameters and the Gamma shape and rate parameters to approximately match the values given in table 1 of Eikenberry et al. (2020). Tables 1 and 2 give details of these parameters. The columns marked *Lower* and *Upper* in table 1 give the lower and upper 2.5% quantiles of these Gamma distribution and roughly correspond to the *Likely range* column given by Eikenberry et al. (2020). The values for the time in the susceptible state were arbitrarily chosen so that the background risk of infection is 1 per 1000 per day. The current daily risk in the US is approximately 1 per 10000 per day, but we use the higher rate to better reflect the risks to active individuals, and allow for under reporting of cases. Note that the model used by Eikenberry et al. (2020) does

not have a *presymptomatic* phase. We assumed that this had a mean of 3 days with likely range of (1.0, 5.0) and reduced the times in the symptomatic phases accordingly.

Table 1: Gamma time-in-state distribution parameters and summary values for the original SARS-CoV-2 variants.

Parameter	Name	Shape	Rate	Mean	Variance	Lower	Upper
U	Sus	1.00	0.001	1000.00	1000000.00	25.32	3688.88
V	Lat	3.47	0.680	5.10	7.50	1.22	11.70
W	Asy	9.00	3.000	3.00	1.00	1.37	5.25
X	Pre	6.53	0.933	7.00	7.50	2.71	13.30
Y	Mld	2.66	0.242	11.00	45.50	1.97	27.62
Z	Svr	0.55	0.110	5.00	45.50	0.01	24.06

Table 2: Probability parameter values specifying the path through the course of infection for the original SARS-CoV-2 variants.

Parameter	Name	Value
q	Symptomatic	0.50
r	Severe	0.10

With regard to the later, omicron, SARS-CoV-2 variants, Jansen et al. (2021) estimate that the latent period is roughly 60% that for the original variants. Symptomatic periods are estimated to be reduced by a factor of 90% (Menni et al. 2022), and we have assumed that both the presymptomatic and asymptomatic periods are similarly reduced. These changes have been made by reducing both the means and standard deviations of the Gamma distributions by the appropriate factor. Table 3 gives details of these Gamma distributions. The path probability parameters have been left unchanged.

Table 3: Gamma time-in-state distribution parameters and summary values for more recent SARS-CoV-2 variants.

Parameter	Name	Shape	Rate	Mean	Variance	Lower	Upper
U	Sus	1.00	0.001	1000.00	1000000.00	25.32	3688.88
V	Lat	3.47	1.133	3.06	2.70	0.73	7.02
W	Asy	9.00	3.333	2.70	0.81	1.23	4.73
X	Pre	6.53	1.037	6.30	6.08	2.43	11.97
Y	Mld	2.66	0.269	9.90	36.86	1.78	24.86
Z	Svr	0.55	0.122	4.50	36.86	0.01	21.66

Appendix A describes methods to compute the probabilities of being in each state at times following the exposure event given the model parameter values described here. The methods have been implemented in the R statistical programming environment (R Core Team 2015), and can be downloaded from [github](#) at

<https://github.com/alun-thomas/covidisolation>.

A **shiny** application is also provided that allows the user to adjust the model and see the effects on state probabilities. This is available at

<https://alunthomas.shinyapps.io/covidisolation/>.

Observations about the individual following the exposure event will change the state probabilities. If they are observed not to have shown any symptoms, the mild and severe disease state are excluded as is the resolved symptomatic state.

In considering the effects of negative test results we distinguish between antigen tests and PCR tests. Since we assume that both test types are fully specific, they are both obligatorily negative for individuals in the susceptible and latent states, and subject to the test sensitivity, can be positive in the asymptomatic, presymptomatic, mild and severe disease states. Antigen tests are typically negative for resolved individuals, while PCR tests are typically positive for some time after resolution. PCR tests are also more sensitive. We set the sensitivity to 95% for PCR tests and 90% for antigen tests.

Appendix A also gives methods for computing state probabilities conditional on observations regarding symptoms and test results.

Since we are primarily concerned with modeling the short term following exposure, we do not model a transition from the resolved state back to susceptible, and we do not model a decrease in PCR sensitivity with time in the resolved state.

### 3 Results

Figure 2 shows the different state probabilities for an individual following an exposure with a 70% chance of resulting in an infection for the Delta SARS-CoV-2 variant. This also shows the effects on these probabilities of observing that the individual shows no symptoms, and, in addition, that they have either a negative PCR or antigen test. Figure 3 shows the same plots for the Omicron variant.

Figure 4 addresses the inference problem of whether an infection occurred following exposure given that the individual remains asymptomatic and has a negative test. The curves show the posterior probabilities in a Bayesian analysis assuming a range of prior probabilities. This is done for both antigen and PCR tests and for both Delta and Omicron variants.

To address the question of when to release an infected individual from isolation or quarantine, we compared the probabilities of infection for individuals who were certainly infected at exposure with those who were certainly not. Because we assume that the infected individual will eventually recover and, in the short term, be resistant and that the uninfected individual will be exposed to background risk, the probabilities for the former will eventually be lower than those for the latter. The point at which this occurs might be considered a guide to an appropriate time to end isolation. Figure 5 shows plots for these probabilities, and also how these are affected by observing that an individual shows no symptoms and has a negative SARS-CoV-2 test.

Figure 2: State probabilities following a 70% chance of transmission at exposure to the SARS-CoV-2 Delta variant. This shows the unconditional probabilities and those when no symptoms and negative tests are observed.

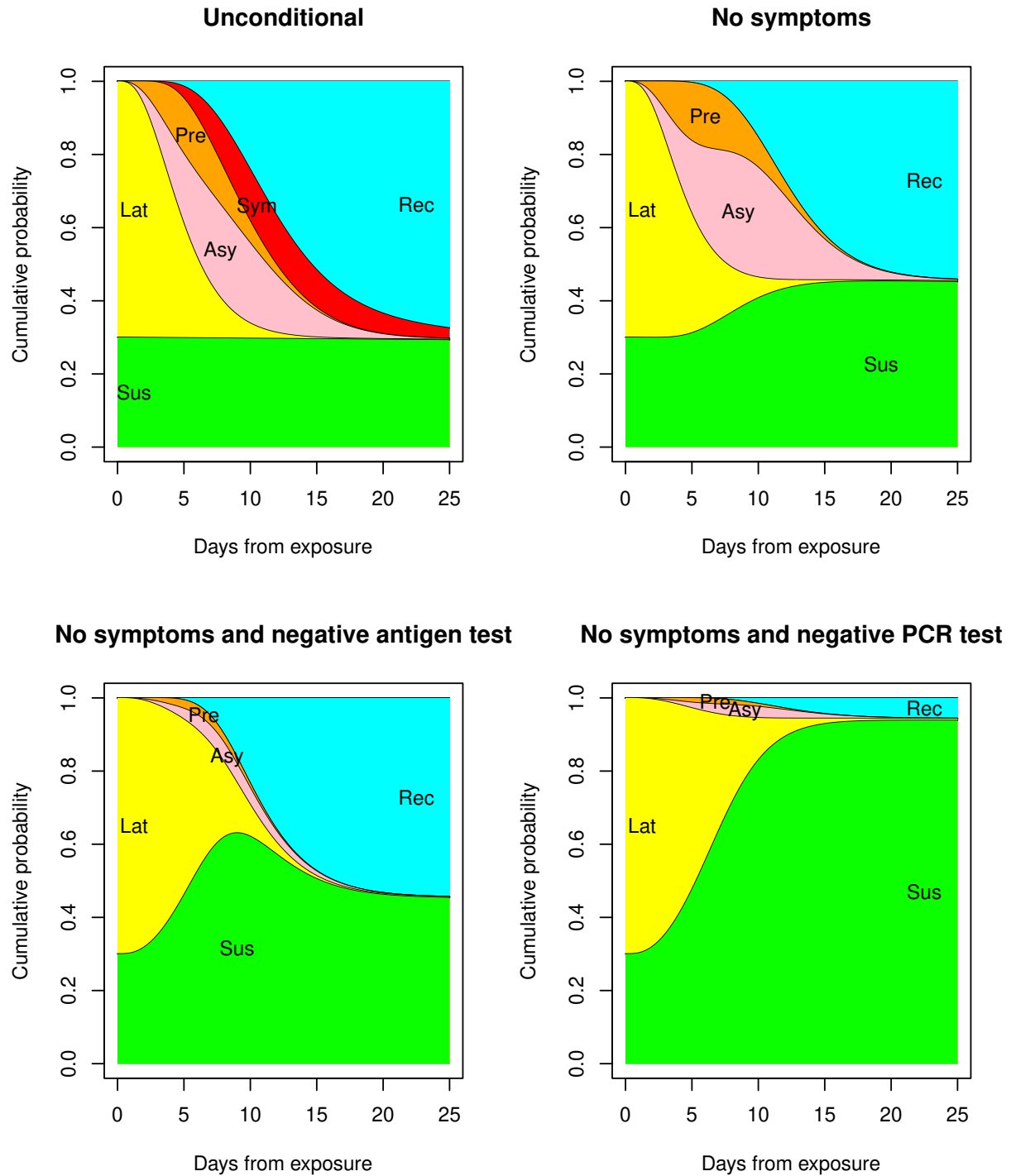


Figure 3: State probabilities following a 70% chance of transmission at exposure to the SARS-CoV-2 Omicron variant. This shows the unconditional probabilities and those when no symptoms and negative tests are observed.

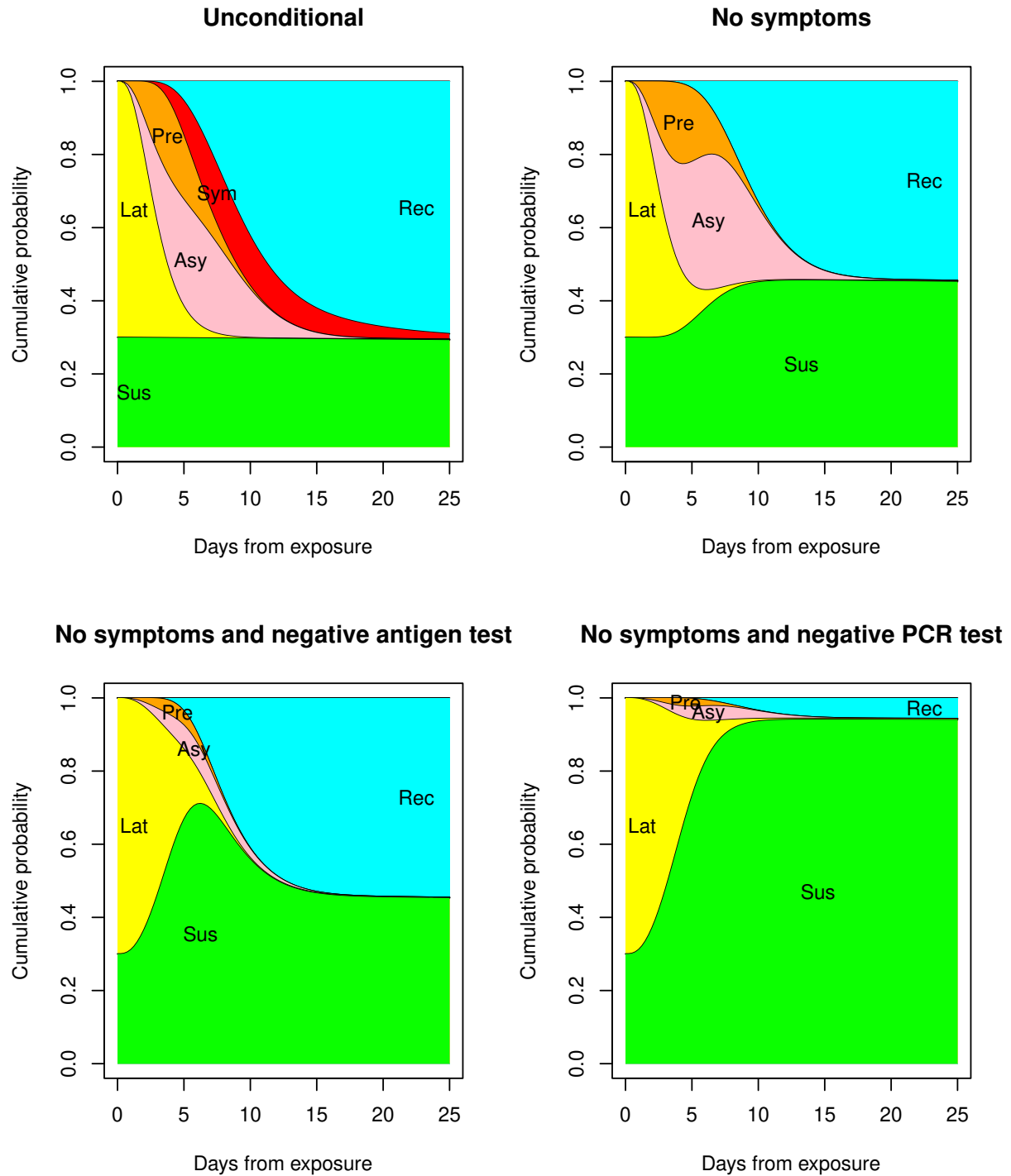


Figure 4: The posterior probability that an infection occurred at time 0 given a negative test following. The curves are given for varying prior probabilities. The vertical lines show where the posteriors are lowest.

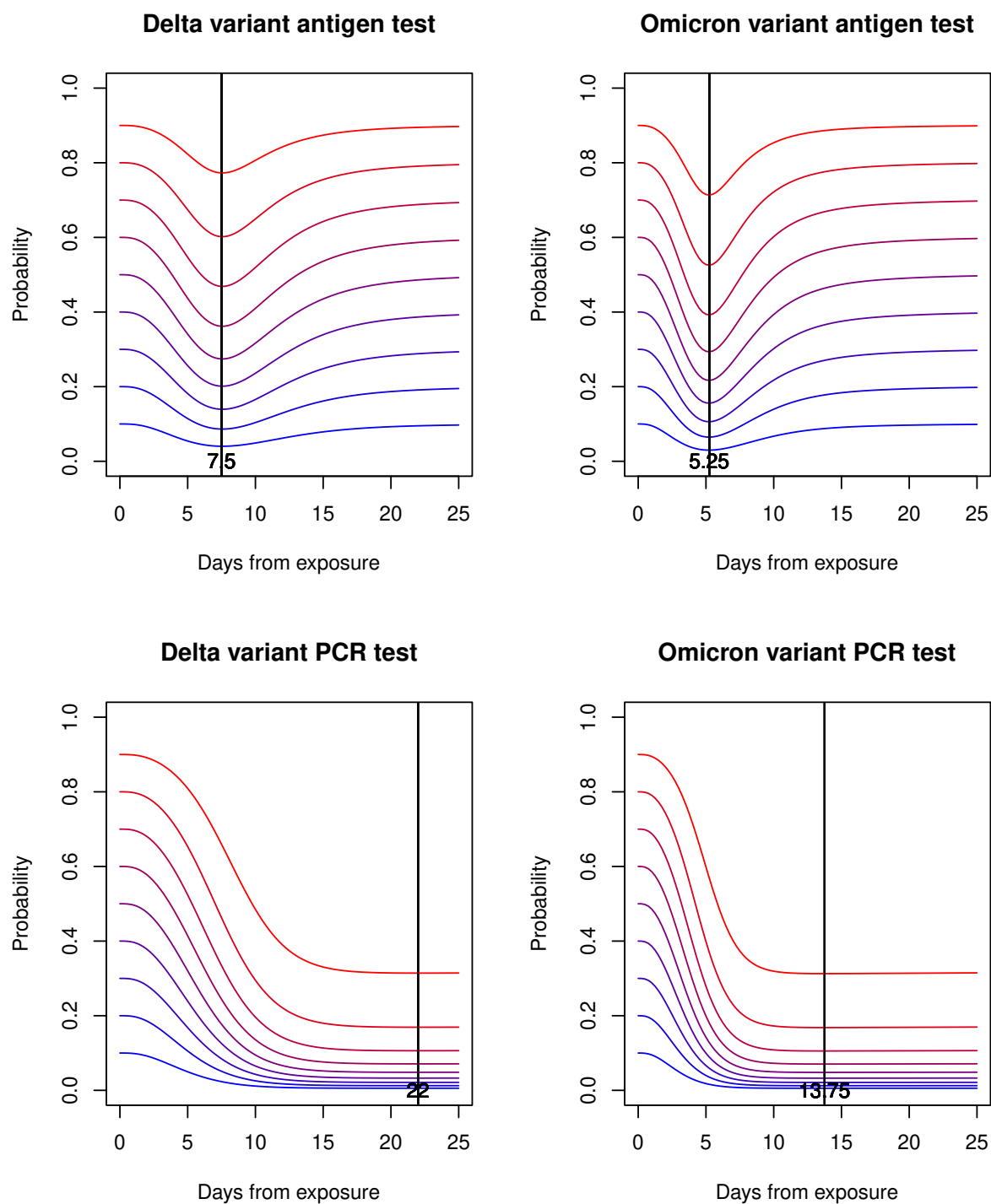
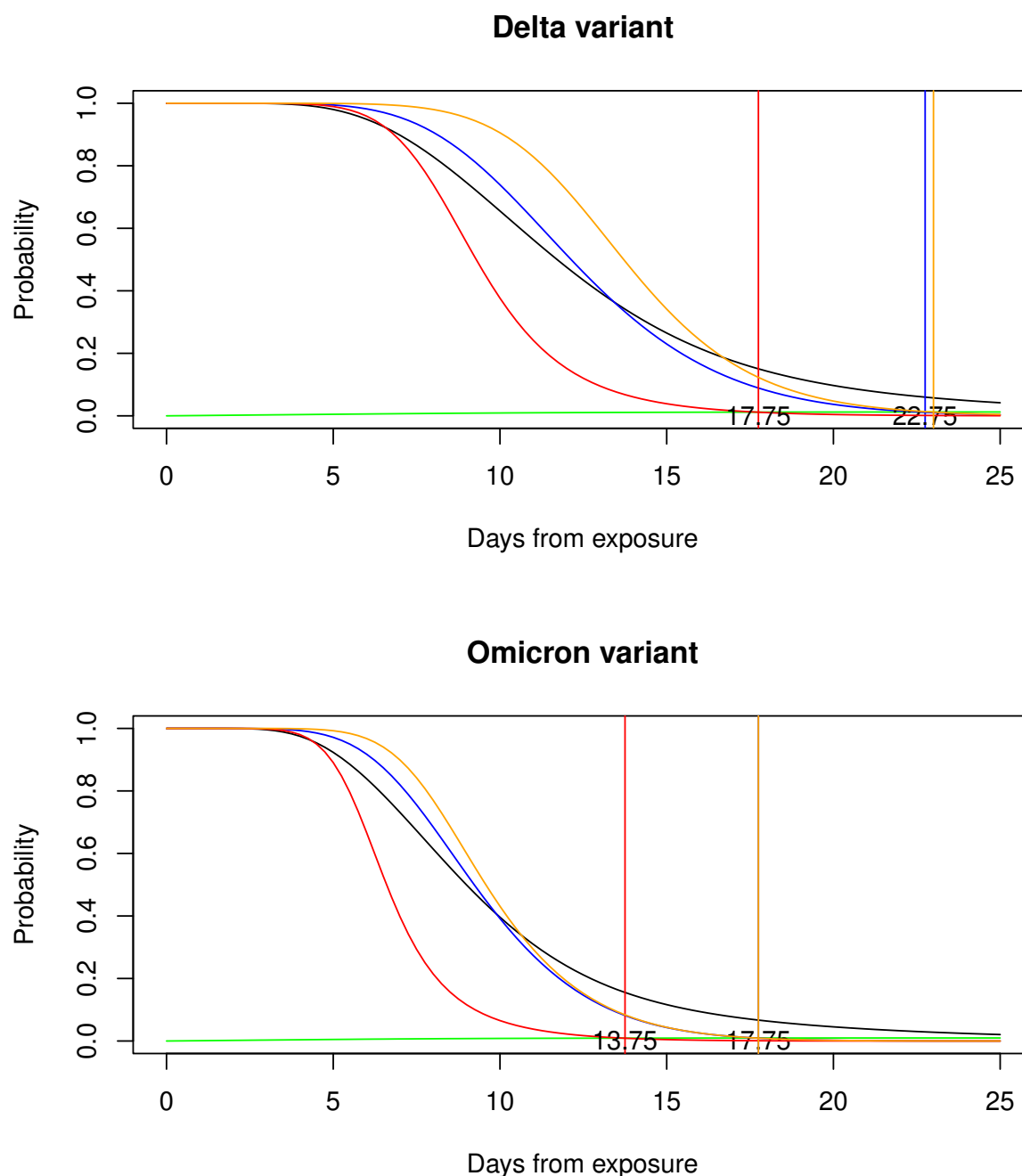




Figure 5: The probability that an individual who was certainly infected at time zero is still infectious at a given time following is shown in black. The blue line shows the probability for an individual who is asymptomatic up to the given time, the red line for an asymptomatic individual who has a negative Antigen test at the given time, the orange line for an asymptomatic individual who has a negative PCR test at the given time. The green line shows the probability for an individual who was certainly not infected at time zero, but was subsequently exposed to background risk of infection. Vertical lines in the appropriate colour show where the probabilities for certainly infected individuals first fall below those for certainly uninfected. Note that the blue and orange lines are very close in both plots.



## 4 Discussion

- PCR test is good for detecting presence of recent infection, antigen test not (see figure 4). Optimal time to run PCR test for this purpose is about 22 days for Delta, 14 days for Omicron. Antigen test is uninformative for this purpose after about 2 weeks.
- Antigen test is good for detecting end of infection, PCR test not (see figure 5). Optimal time to run Antigen test for this purpose is about 18 days for Delta, 14 days for Omicron. Shortens isolation time compared with just observing no symptoms by about 5 days for Delta and 4 for Omicron. PCR test offers very little in addition to observation of no symptoms.
- Reduction to background risk is a pretty cautious criterion, might consider higher threshold.
- Test differences are due to how they behave for individuals in recovered state, not so much because of difference in sensitivity.
- Differences between Delta and Omicron largely due to reduced latent period for latter. (can be seen in figures 2 and 3).
- Limitation: This analysis is limited to the short or medium term, before the possibility of reinfection becomes a factor.

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## A Methods and programs for probability calculations

We calculate the state probabilities following an exposure event by conditioning on the paths taken through the course of the infection. The paths, states, variable and parameter names used here are those illustrated in figure 1.

We let  $H \in \{1, \dots, 6\}$  index the path and let  $S_x \in \{\text{Sus, Lat, Asy, Pre, Mld, Svr, ResS, ResA}\}$  be the state the individual is in at time  $x$  days following the exposure event. Table 4 gives  $P(H)$ , the path probabilities derived from the splitting probabilities, and  $P(S_x|H)$ , the conditional probabilities of being in each state  $S_x$  given that path  $H$  is followed. We use  $F_\cdot$  as shorthand for the distribution function of sums of arbitrary Gamma distributed random variables. Thus, for instance,

$$\begin{aligned} F_{UVW} &= F_{U+V+W}(x) \\ &= P(U + V + W \leq x) \\ &\quad \text{where } U, V, W \sim \text{Gamma}(\bullet, \bullet) \end{aligned} \tag{1}$$

The state probabilities are then given by

$$P(S_x) = \sum_H P(S_x|H)P(H). \tag{2}$$

The specific path splitting probabilities and Gamma shape and rate parameters used here are given in tables 1, 2 and 3, but our implementation allows the user to change these values.

Table 4: The probability,  $P(H)$ , of following each possible path of progression through the infection, and the probability of being in each state at time  $x$  given the path,  $P(S_x|H)$ .

	$H$					
	1	2	3	4	5	6
$P(H)$	$pqr$	$pq(1-r)$	$p(1-q)$	$(1-p)qr$	$(1-p)q(1-r)$	$(1-p)(1-q)$

$S_x$	$P(S_x H)$					
Sus	0	0	0	$1 - F_U$	$1 - F_U$	$1 - F_U$
Lat	$1 - F_V$	$1 - F_V$	$1 - F_V$	$F_U - F_{UV}$	$F_U - F_{UV}$	$F_U - F_{UV}$
Asy	0	0	$F_V - F_{VX}$	0	0	$F_{UV} - F_{UVX}$
Pre	$F_V - F_{VW}$	$F_V - F_{VW}$	0	$F_{UV} - F_{UVW}$	$F_{UV} - F_{UVW}$	0
Mld	0	$F_{VW} - F_{VWZ}$	0	0	$F_{UVW} - F_{UVWZ}$	0
Svr	$F_{VW} - F_{VWY}$	0	0	$F_{UVW} - F_{UVWY}$	0	0
ResS	$F_{VWY}$	$F_{VWZ}$	0	$F_{UVWY}$	$F_{UVWZ}$	0
ResA	0	0	$F_{VX}$	0	0	$F_{UVX}$

To calculate the conditional probabilities given in the results section, we consider the probability of some observation  $O_x$  at time  $x$  and specify  $P(O_x|S_x)$ , the probability of the observation given the underlying state at that time. Then, using Bayes rule, we have

$$P(S_x|O_x) = \frac{P(O_x|S_x)P(S_x)}{\sum_{S_x} P(O_x|S_x)P(S_x)}. \tag{3}$$

Table 5 specifies the conditional probabilities required for the following events:

- $I_x$  indicates that the individual is uninfected (0), or infected (1), at time  $x$ ,
- $M_x$  indicates that the individuals has not (0) or has (1) exhibited symptoms up to time  $x$ ,
- $T_x$  indicates that a test at time  $x$  is negative (0) or positive (1).

We note that the states ResS and ResA differ only in the path taken to get to them and we use this convention only because it allows us to distinguish between recovered individuals who have or have not shown symptoms in the past without explicit reference to the path, thus simplifying the first column of table 5.

To address the inferential problem of whether an exposure at time 0 resulted in an infection we let  $E_0$  be an indicator for the event that an infection occurred (1) or didn't (0). We can then calculate the required likelihoods given observations  $O_x$  as

$$\begin{aligned} P(O_x|E_0 = 0) &= \sum_{S_x} P(O_x|S_x)P(S_x|p = 0) \\ P(O_x|E_0 = 1) &= \sum_{S_x} P(O_x|S_x)P(S_x|p = 1) \end{aligned} \quad (4)$$

and if  $P(E_0 = 1)$  is the prior probability that an infection occurred, the posterior given  $O_x$  is

$$P(E_0 = 1|O_x) = \frac{P(O_x|E_0 = 1)P(E_0 = 1)}{P(O_x|E_0 = 1)P(E_0 = 1) + P(O_x|E_0 = 0)P(E_0 = 0)}. \quad (5)$$

Table 5: For each state we give  $P(M_x = 0|S_x)$ , the probability of no symptoms up to time  $x$  given the state,  $P(I_x = 1|S_x)$ , the probability of being infected given the state,  $P(T_x = 0|S_x, \text{Antigen})$ , the probability of having a negative antigen test given the state, and  $P(T_x = 0|S_x, \text{PCR})$ , the probability of having a negative PCR test given the state.

$S_x$	$P(M_x = 0 S_x)$	$P(I_x = 1 S_x)$	$P(T_x = 0 S_x, \text{Antigen})$	$P(T_x = 0 S_x, \text{PCR})$
Sus	1	0	1	1
Lat	1	1	1	1
Asy	1	1	$1 - s$	$1 - s$
Pre	1	1	$1 - s$	$1 - s$
Mld	0	1	$1 - s$	$1 - s$
Svr	0	1	$1 - s$	$1 - s$
ResS	0	0	1	$1 - s$
ResA	1	0	1	$1 - s$

We wrote programs to implement the above methods using the R statistical programming environment (R Core Team 2015). These allow specifying arbitrary path splitting probabilities and arbitrary shape and rate parameters for the Gamma time in state variables. These calculations require the `pcoga()` function in the `coga` package (Hu et al. 2021, Hu et al. 2020) to calculate the density functions of sums of independent Gamma random variables. `codecoga` can be installed using the standard R command

`install.packages("coga")`, and is also available from the CRAN repository of R packages at <https://cran.r-project.org/web/packages/coga/index.html>.

The R source code for these general programs is available from github at <https://github.com/alun-thomas/covidisolation> in the file `covid.R`. The specific code necessary to produce the tables and figures in this paper are also provided in the file `isol.R` and may be a useful example for those wishing to use or extend the methods described here.

A **shiny** application for calculating the state probabilities following an exposure using these programs is available at <https://alunthomas.shinyapps.io/covidisolation/>. The application allows the user to input parameters and change the conditioning events.