
Transmission Dynamics and Effectiveness of Control Measures during COVID-19 Surge, Taiwan, April–August 2021

Andrei R. Akhmetzhanov, Hao-Yuan Cheng, Natalie M. Linton, Luis Ponce, Shu-Wan Jian, Hsien-Ho Lin

An unprecedented surge of COVID-19 cases in Taiwan in May 2021 led the government to implement strict nationwide control measures beginning May 15. During the surge, the government was able to bring the epidemic under control without a complete lockdown despite the cumulative case count reaching >14,400 and \geq 780 deaths. We investigated the effectiveness of the public health and social measures instituted by the Taiwan government by quantifying the change in the effective reproduction number, which is a summary measure of the ability of the pathogen to spread through the population. The control measures that were instituted reduced the effective reproduction number from 2.0–3.3 to 0.6–0.7. This decrease was correlated with changes in mobility patterns in Taiwan, demonstrating that public compliance, active case finding, and contact tracing were effective measures in preventing further spread of the disease.

SARS-CoV-2, the pathogen causing COVID-19, began infecting humans in Wuhan, China, in December 2019. Within 1 year, SARS-CoV-2 spread to nearly all countries, and >178 million infections and 3.7 million deaths were reported by April 2021. Taiwan, an island with 23.8 million inhabitants, reported only slightly more than 1,000 cases by April 2021, despite being located close to the original epicenter of the COVID-19 outbreak. At that time, most infections confirmed in Taiwan were acquired abroad, and <10% were acquired locally.

The subsequent emergence of more transmissible SARS-CoV-2 variants led to multiple introductions

from those traveling to and from Taiwan, initiating cryptic transmissions in the capital city of Taipei and its surroundings in April 2021. Newly detected clusters of the virus led to an explosive growth in cases, and daily reported case numbers reached 200 by mid-May. The sudden increase in cases prompted the government to implement stricter control measures to prevent disease spread, and those measures proved effective in bringing the epidemic under control by the end of July. Those preventive measures included restricting public movement, enforcing compulsory shortening of business hours, implementing work-from-home for nonessential businesses, banning in-restaurant dining, and canceling social and religious gatherings. By October 2021, Taiwan was again reporting 0 cases daily.

The initial clusters of infections in 2021 were linked to international pilots and flight crew members, but the major epidemic hotspots were identified as owners and visitors of tea houses, which are landmarks in some districts of Taipei. Although tea houses in Taipei typically offer tea and other refreshments during the day, some also conduct business in the evening, when the potential for activities that increase risk for the transmission of SARS-CoV-2 (e.g., close physical contact) is greater and timely detection of infections can be hindered (1–3). In nightlife districts across the city, patrons and staff of tea houses and other establishments often are unwilling to share contact and travel histories with public health officials. Outside of Taipei and New Taipei City, clusters of infections were frequently linked to factories or other production sites, affecting vulnerable social groups such as migrant workers. Some initial clusters were linked to local markets and initiated by vendors traveling to the Taipei area for commercial purposes.

Author affiliations: National Taiwan University College of Public Health, Taipei, Taiwan (A.R. Akhmetzhanov, L. Ponce, H.-H. Lin); Taiwan Centers for Disease Control, Epidemic Intelligence Center, Taipei (H.-Y. Cheng, S.-W. Jian); Hokkaido University Graduate School of Medicine, Sapporo, Japan (N.M. Linton)

DOI: <http://doi.org/10.3201/eid2810.220456>

The effective reproduction number, R_t , has played a pivotal role in evaluating the effectiveness of various public health and social measures (PHSMs) during the COVID-19 pandemic (4–6). R_t is defined as the average number of secondary transmissions caused by a primary case at a given time while interventions, existing immunity, or other mediating factors are present. During the pandemic, R_t was used frequently as a data point to inform decision- and policy-making processes, because the value of R_t relative to the threshold value of 1 can be interpreted as an indicator for when PHSMs should be implemented, strengthened, or relaxed (7,8). Among the various PHSMs that might be used, stay-at-home orders, cancelling leisure activities, and restaurant-based interventions were found to be largely ineffective in curbing COVID-19 transmission in the United States (9). In contrast, strong social distancing, school closures, and widespread mask-wearing were found to be quite effective in mitigating the spread of COVID-19 in both the United States and elsewhere (10–12). One study found that only strict (complete) lockdowns could curb the spread of infections and reduce R_t to <1 (5). However, responses to the virus in Taiwan and Japan demonstrate that less extreme measures (i.e., without the implementation of a complete lockdown) were sufficient in preventing a wide, rampant spread of COVID-19 during the epidemic and returning daily counts to an acceptable level (<10 cases). The government's response to the surge of COVID-19 cases in Taiwan that began in May 2021 presents a striking example of how public compliance with such less extreme preventive measures successfully quelled a burgeoning epidemic wave.

Among various possible ways to estimate R_t , the instantaneous reproduction number based on the method of Cori et al. (13) has often been used during the COVID-19 pandemic to describe current epidemiologic situations (14,15) or to forecast future incidence (16). Predicting the real-time R_t value and accounting for covariates has been recognized as an important step toward the future real-time monitoring of disease spread in different countries (17–21).

Taiwan reported extremely low numbers of confirmed COVID-19 cases in 2020, offering an example of a relatively efficient prevention strategy against the spread of SARS-CoV-2 (22). The government instituted a 4-level system to efficiently contain and mitigate COVID-19 epidemics (Appendix Table 1, <https://wwwnc.cdc.gov/EID/article/28/10/22-0456-App1.pdf>). Before April 2021, the largest cluster of locally acquired infections had only 22 confirmed cases (23). Of the various factors contributing to Taiwan's early

pandemic success, the key components were strict border control, public compliance with untargeted PHSMs (e.g., mask-wearing, proactive case finding, and contact tracing), and use of digital technologies, such as QR codes (24). However, the increased transmissibility of subsequent SARS-CoV-2 variants and low levels of vaccine coverage posed significant challenges for COVID-19 containment in Taiwan in 2021. We investigated the effectiveness of the public health and social measures instituted by the Taiwan government during the 2021 COVID-19 surge by quantifying the change in R_t .

Methods

Data Collection

We retrieved line list data from publicly available sources and Taiwan Centers for Disease Control reports (25). The combined dataset from these sources contained de-identified case records, including information on symptom onset date (when available), case confirmation date, confirmed date of death, level of severity of the infection (asymptomatic/mild, moderate, severe), and information on residency. The 3 categories of disease severity (mild, moderate, severe) were assigned in accordance with the World Health Organization definition (26). We extracted mobility metrics from community reports provided by Google (27). The 6 metrics used fell into the following categories: "grocery and pharmacy," "parks," "residential," "retail and recreation," "transit stations," and "workplaces." We quantified each metric by a daily change in the median mobility when compared with the baseline median for the 5-week period January 3–February 6, 2020.

Estimating Epidemiologic Parameters

We fitted time intervals from symptom onset to case confirmation, onset to severe disease, onset to death, and onset to report of death (as well as from death to report of death) to a mixture of 3 distributions (gamma, Weibull, and log-normal) (23). We then fitted the serial interval distribution to left-shifted gamma, Weibull, and log-normal distributions (to account for negative values). We estimated all parameters within a Bayesian framework, using a doubly censored likelihood with right truncation and Markov chain Monte Carlo simulations (28,29). To improve convergence of the mixture model, we set the mean and SDs to be common to the 3 distributions, as has been proposed for Bayesian model averaging (M. Keller et al., unpub. data, <https://doi.org/10.48550/arXiv.1711.10016>). We estimated the reporting delay, which is the time from symptom onset to

case confirmation, under 2 scenarios: when the distribution was unchanged over time, and when the parameters of the distribution were varied in time (30).

We estimated R_t using date of symptom onset and date of infection (13,31). When R_t was classified by date of symptom onset, the expected case count on day t was proportional to R_t and a convolution of case counts on previous days with the serial interval distribution. When R_t was classified by date of infection, the formula had a more complicated form and contained a double convolution, involving the incubation period and profile of infectiousness (31,32) (Appendix). Because some case records did not contain information on symptom onset date, we back-projected those cases from the date the case was confirmed to a presumptive date of symptom onset, using a time-varied distribution of the reporting delay.

Results

Epidemiologic Situation

Little to no local transmission of SARS-CoV-2 was reported in Taiwan before April 2021. Vaccine coverage was also arbitrarily low (<1%) at that time. There were, however, multiple clusters of infections during the latter half of April 2021, followed by a wave of COVID-19 cases at the beginning of May 2021 (Figure 1, panel A). A total of 14,442 cases associated with the epidemic wave were confirmed by August 25, 2021, including 5,029 (34.8%) persons who were asymptomatic at the time of testing and 3,093 (21.4%) persons recognized as having severe disease. Among patients requiring hospitalization, 238 (1.6%) had nonsevere pneumonia, 2414 (16.7%) had severe pneumonia, and 441 (3.1%) had acute respiratory distress syndrome (Table). A total of 779 persons (5.4%) died during the epidemic wave. Most (701, 90%) of the deceased patients had known underlying chronic conditions. Eight additional deaths among patients in the study population were unrelated to SARS-CoV-2 infection.

The median age of persons with confirmed cases was 51 years; 23.9% were ≥ 65 years of age, 51.9% 45–64 years of age, 18.3% 18–44 years of age, and 5.9% <18 years of age. Only 0.4% of patients <18 years were categorized as having moderate disease, and half (50.3%) of these younger patients were reported as asymptomatic at the time of testing. In contrast, 46.6% of those ≥ 65 years of age experienced moderate symptoms, and 22.3% were asymptomatic at the time of testing. The median age of patients who died was 72 years, and 79.8% of deaths were reported among those ≥ 65 years of age. Men accounted for most deaths (63.5%). Geographically, a substantial portion

of the infections (1,874 cases, 13.0% of the total) were confirmed among residents of Wanhua District in Taipei (Figure 1, panel B).

The median time from date of symptom onset to date of case confirmation was estimated at 3.0 days (95% CI 0.7–11.9 days). The time required for disease progression from symptom onset to severe disease was an average of 7.7 days (95% CI 2.1–28.5 days). Death was observed, on average, 13.3 days after symptom onset (95% CI 1.1–92.4 days). Deaths were reported an average of 3.5 days thereafter (95% CI 1.0–12.3 days).

R_t and Efficiency of PHSMs

When quantifying R_t by date of symptom onset, we noted that the value remained relatively stable, with values of ≈ 2 –3 before the surge of COVID-19 cases reported around May 10, 2021 (Figure 2, panel A). We estimated the median posterior value of R_t to exceed 3 during the first week of May, likely because of cryptic community transmission; confirmed cases with symptom onset in the first week of May had prolonged reporting delays of nearly 10 days (Figure 2, panel B, orange line), and later cases generally had shorter reporting delays of ≈ 3 –4 days. The reporting delay quantified by date of case confirmation peaked around May 16 (Figure 2, panel B, gray line). The test-positivity rate for SARS-CoV-2 also reached its highest around the same dates (Figure 2, panel B, blue line). These results indicate that cases with earlier symptom onset dates had longer reporting delays compared with subsequent cases and serve as an indicator of persistent cryptic transmission of SARS-CoV-2 in the community between the end of April and the beginning of May 2021.

Next, we quantified the effective reproduction number by infection date and tied it to PHSMs (Figure 2, panels C, D). Taiwan adopted a 4-tier system of restrictions ranging from level 1 at the lowest to level 4 at the highest (Appendix Table 1). Level 2 restrictions began on May 11, 2021; level 3 restrictions began in Taipei and New Taipei City on May 15 and then expanded to the rest of Taiwan on May 19. Level 3 measures were further strengthened on May 29. We estimated the posterior mean R_t in the early stage of the epidemic—before level 2 restrictions began—at 2.85 (95% CI 2.51–3.26). Implementation of level 2 measures on May 11 was followed by a slightly decreased mean of 2.40 (95% CI 1.99–2.86), and level 3 measures in Taipei City and New Taipei City on May 15 further decreased the mean value to 1.59 (95% CI 1.30–1.90). Nonetheless, these measures were insufficient to bring the R_t consistently below 1. Only

after level 3 measures were expanded to all of Taiwan on May 19 did the mean R_t decrease to below 1 (0.86 [95% CI 0.76–0.95]). R_t then dropped even further

when those measures were strengthened on May 29 by prohibiting dine-in services and setting up a work-from-home order (0.65 [95% CI 0.57–0.74]).

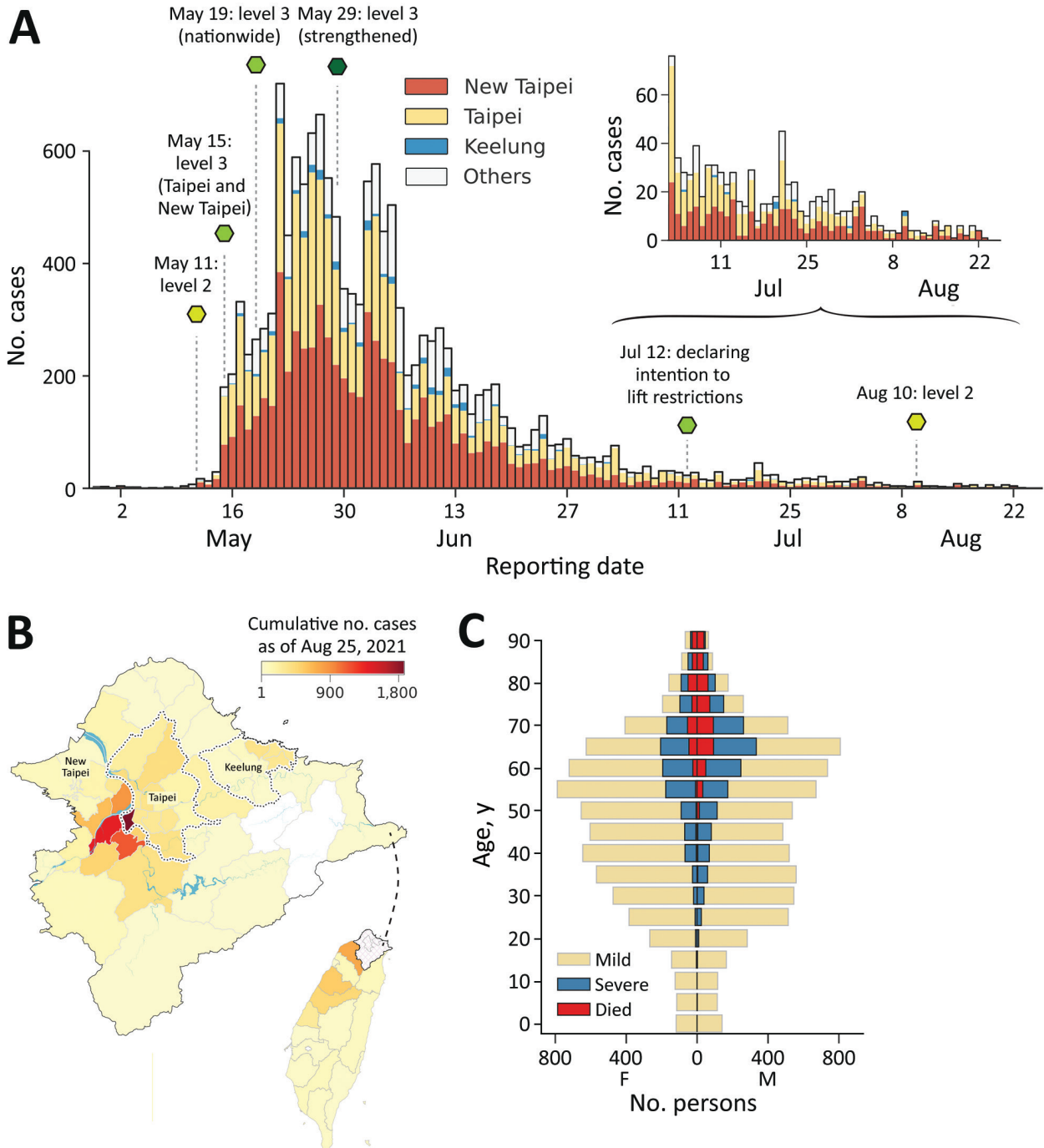


Figure 1. Epidemic wave of COVID-19 in Taiwan, April–August 2021. A) Epidemiologic curve of confirmed COVID-19 cases by reporting date, stratified by geographic area. Dashed lines and hexagons indicate timing and description of major public health and social measures; variation in hexagon colors shows relative strictness of measures, ranging from light to dark green. B) Geographic distribution of cases. The colormap indicates the cumulative number of cases confirmed by August 25, 2021, at district level for Taipei, New Taipei City, and Keelung and at county level for all other areas (indicated in gray in panel A). Inset shows location of enlarged area in Taiwan. C) Age pyramid of confirmed cases specified by known severity status or death. Age and spatial distribution of fatalities is shown in Appendix Figure 4 (<https://wwwnc.cdc.gov/EID/article/28/10/22-0456-App1.pdf>)

Table. Demographic and clinical characteristics of persons with confirmed COVID-19 cases, by geographic region, Taiwan, April 23, 2021–August 25, 2021*

Characteristic	No. (%)			
	Taiwan	Taipei	New Taipei City	Other counties
Age group				
<17	845 (5.9)	225 (4.6)	410 (6.0)	210 (7.7)
17–34	2,660 (18.4)	642 (13.2)	1,168 (17.0)	850 (31.0)
15–64	7,489 (51.9)	2,629 (54.2)	3,656 (53.3)	1,204 (44.0)
>64	3,448 (23.9)	1,354 (27.9)	1,620 (23.6)	474 (17.3)
Sex				
F	7,149 (49.5)	2,502 (51.6)	3,387 (49.4)	1,260 (46.0)
M	7,293 (50.5)	2,348 (51.6)	3,467 (49.4)	1,478 (54.0)
Severity				
Mild/asymptomatic	11,349 (78.6)	3,807 (78.5)	5,309 (77.5)	2,233 (81.6)
Severe	3,093 (21.4)	1,043 (21.5)	1,545 (22.5)	505 (18.4)
Known to be symptomatic				
No	5,037 (34.9)	1,710 (35.3)	2,193 (32.0)	1,134 (41.4)
Yes	9,405 (65.1)	3,140 (64.7)	4,661 (68.0)	1,604 (58.6)
Total	14,442	4,850 [33.6]	6,854 [47.5]	2,738 [19.0]

*Parentheses indicate a columnwise fraction of cases within each group. Brackets indicate a rowwise proportion of cases.

These estimates prompted our further investigation into why the initial set of level 3 measures implemented on May 15 for Taipei and New Taipei City and on May 19 nationwide were insufficient to bring R_t substantially below 1. We estimated R_t by infection date using 2 different functions of time. First, R_t was modeled by a piecewise constant function of time with equidistant time windows (e.g., 5 or 7 days). Second, the change in R_t was correlated with the observed change in community mobility across 6 different community metrics (see Methods).

When we modeled R_t using a piecewise constant function of time, we observed a pattern similar to that of R_t by date of symptom onset, except that the pattern was time-lagged (compare Figure 3, panel A, and Figure 2, panel A). The temporal pattern also resembled the change in various mobility metrics over time (compare Figure 3, panel A, and Figure 3, panel B). However, the posterior mean of R_t did not increase after July 12, even though some mobility metrics previously recognized as important for explaining the transmission potential of COVID-19 (17) (e.g., retail and recreation, transit stations, and workplaces) continued to increase over time. To address this contradiction, we theorized that the basic reproduction number (R_0) changed over time. The time-variability of R_0 represented the proxy measure of changing contact rate of infected and susceptible individuals over time and served as an indicator of PHSMs, including the voluntary changes in public behavior (33). When we defined it by a monotonically decreasing sigmoidal function over time, the corresponding model fit the data better. We compared a model with a time-varied R_0 with a model with a constant R_0 using a “leave-one-out” information criteria (LOOIC), which is used in Bayesian frameworks for model selection

(34). The model with a time-varied R_0 had a lower median LOOIC value (884.2) compared with that of the model that used a constant R_0 (899.6) (Appendix Figure 5). The fit resulted in the change point of R_0 on approximately July 19, and R_0 decreased from a median of 3.17 at the beginning of the epidemic to 1.72 at the end of the epidemic (defined as August 14), a 46% reduction.

We additionally investigated the association of different mobility metrics with R_t . The model with only 3 mobility metrics showed a fairly indistinguishable data fit compared to models with 4 to 6 mobility metrics, and the difference in LOOIC values was <2 ($\Delta\text{LOOIC} \leq 1.56$). By sequentially fitting the models with 1, 2, and 3 metrics, we identified that the most significant metrics describing the individual mobility were transit stations, workplaces, and grocery stores and pharmacies (Figure 3, panel D).

We investigated counterfactual scenarios wherein level 3 measures had been implemented either earlier or later than the actual May 15 date (Figure 4). If the level 3 measures had been delayed by just 3 days, the size of the epidemic on August 14 likely would have been double that of the baseline scenario (23,900 cases [95% CI 7,900–61,500]) vs. 12,500 cases [95% CI 4,000–29,800]) or the actual case count (14,400). Beginning level 3 measures 3 days earlier likely would have resulted in only 6,400 cases (95% CI 2,200–15,600) (Appendix Figure 6). Varying the date of level 3 implementation revealed a nonlinear, exponential-like relationship whereby a longer delay would accelerate the increase in the final epidemic size.

Discussion

In this study, we analyzed the spread of SARS-CoV-2 in Taiwan during April–August 2021 and quantified

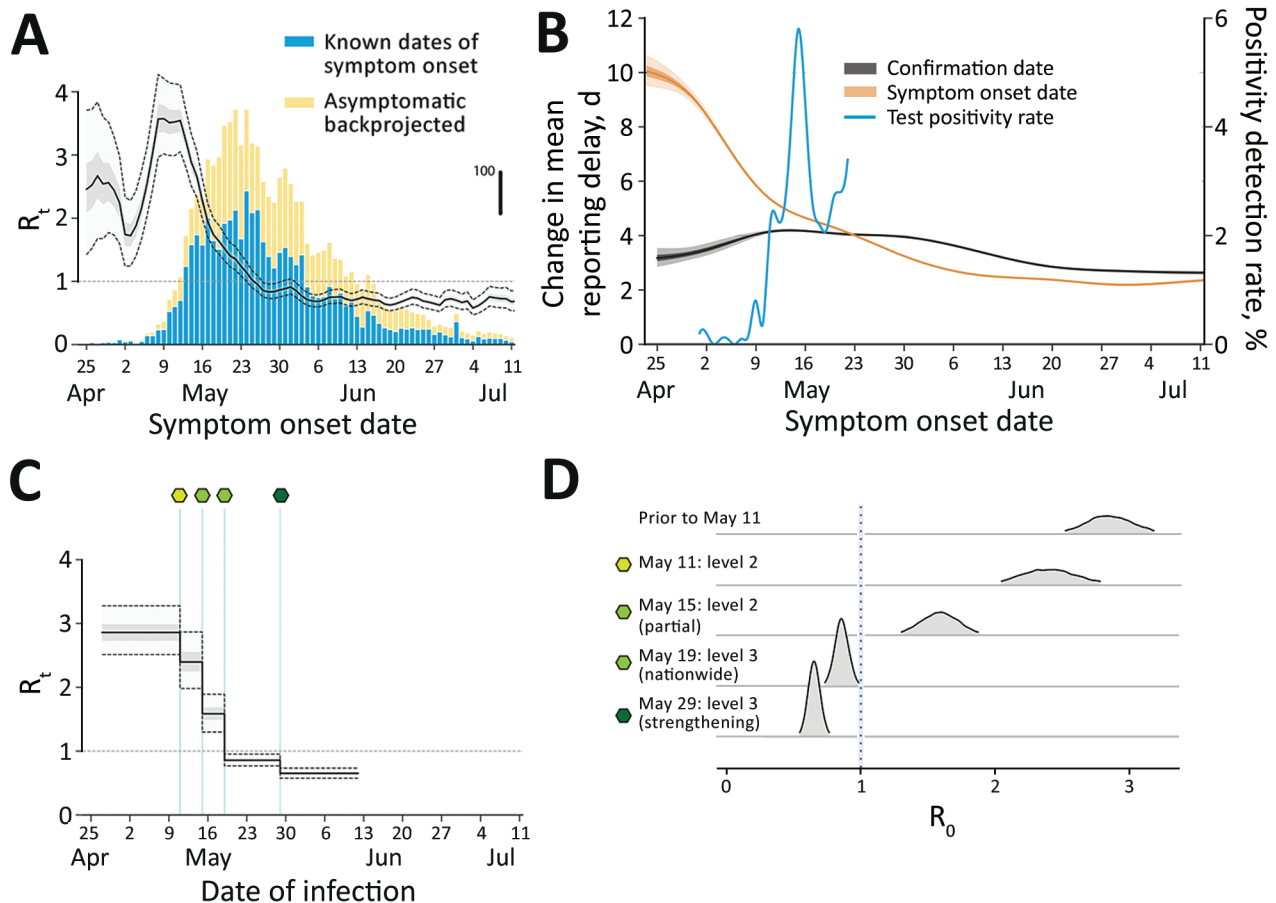


Figure 2. Comparison of R_t inferred by infection date with R_t by symptom onset date during epidemic wave of COVID-19 in Taiwan, April–August 2021. **A**) R_t by infection date (overlay) is notably shifted to the left compared with symptom onset date. Black line indicates mean; light gray shading indicates interquartile range; dotted lines indicate 95% CI. Bars indicate the nowcasted daily incidence of COVID-19 cases; vertical scale is indicated by thick black line on the right. **B**) Change in the mean reporting delay, which is the time between symptom onset date and confirmation date, over time, characterized by either the date of symptom onset (orange) or by confirmation date (black). Dark gray shading indicates IQR; light gray shading indicates 95% CI. The blue line indicates the test positivity rate that peaked around May 16 (axis on the right). **C**, **D**) The estimated R_t by date of infection, linked to public health and social measures (green-shaded hexagons, as defined in panel **D**). R_t , effective reproduction number.

the effectiveness of PHSMs implemented by the government. Initial COVID-19 cases had longer reporting delays, and there was a higher test-positivity rate at the beginning of the outbreak (Figure 1). Shortening of the reporting delay over time (Figure 2, panel **B**) indicated better management of the outbreak in later periods. Our results also showed that implementing stricter PHSMs on May 29, 2021 (Appendix Table 2), was followed by R_t falling below 1. We conclude that the timing of introduction of PHSMs by the government was judicious, and postponement by ≥ 3 days would have likely more than doubled the final size of the outbreak.

Because the number of cases grows exponentially at the beginning of an outbreak, delaying PHSMs by just 3 days can lead to a significant increase in the

disease burden and can double the final epidemic size. Given the indications that the healthcare system of Taiwan was close to being overwhelmed with COVID-19 patients in mid-May, the actual timing of level 3 measures on May 15 likely prevented an even larger healthcare crisis. Although an earlier introduction of PHSMs could have substantially improved the situation, the low case numbers might have caused some public misunderstanding regarding the necessity of strict prevention measures when there was no evidence of escalating case counts. It is fortunate that the government of Taiwan acted in accordance with the country's 4-level COVID-19 alert system criteria by implementing stricter PHSMs as soon as possible (Appendix Table 2).

The April–August 2021 epidemic wave was the first such large-scale wave seen in Taiwan. Using

Bayesian statistical inference of the effective reproduction number by date of infection, we were able to attribute reductions in R_t to the implementation of PHSMs and estimate their effectiveness. The value of R_t only fell below 1 (95% CI 0.57–0.74) consistently after the PHSMs were further strengthened. We base this result, however, largely on model assumptions, so the association might be confounded by behaviors not accounted for in the models.

Even assuming only 1 in 5 COVID-19 cases was confirmed, the cumulative number of cases would have reached fewer than 100,000 cases, according to our models. In 2022, however, Taiwan experienced a much larger outbreak associated with the Omicron variant, during which the total number of confirmed cases exceeded 4 million. Given Omicron’s higher transmissibility and greater capacity for evading immunity, coupled with pandemic fatigue and high vaccination coverage of the Taiwanese population (80.2% for the

second dose and 60.1% for the booster dose as of May 2, 2022), the government chose to relax PHSMs such as proactive case finding and contact tracing in mid-May 2022. As a result, a direct comparison between the pandemic situation in 2021 we have described and the 2022 Omicron wave is not possible.

Using mobility metrics, which are the proxies of contact rates in different settings, was unable to completely capture the temporal change in R_t . However, the additional assumption of a simultaneous decrease in R_0 at later stages of the epidemic adequately explained the observed dynamics. This decrease could likely be a result of higher efficiency in terms of case finding and contact tracing when the number of cases was significantly lower compared with the efficiency in gathering that information at the peak of the epidemic wave.

In regard to study limitations, we did not distinguish fully asymptomatic infections from those that were asymptomatic at the time of testing but became

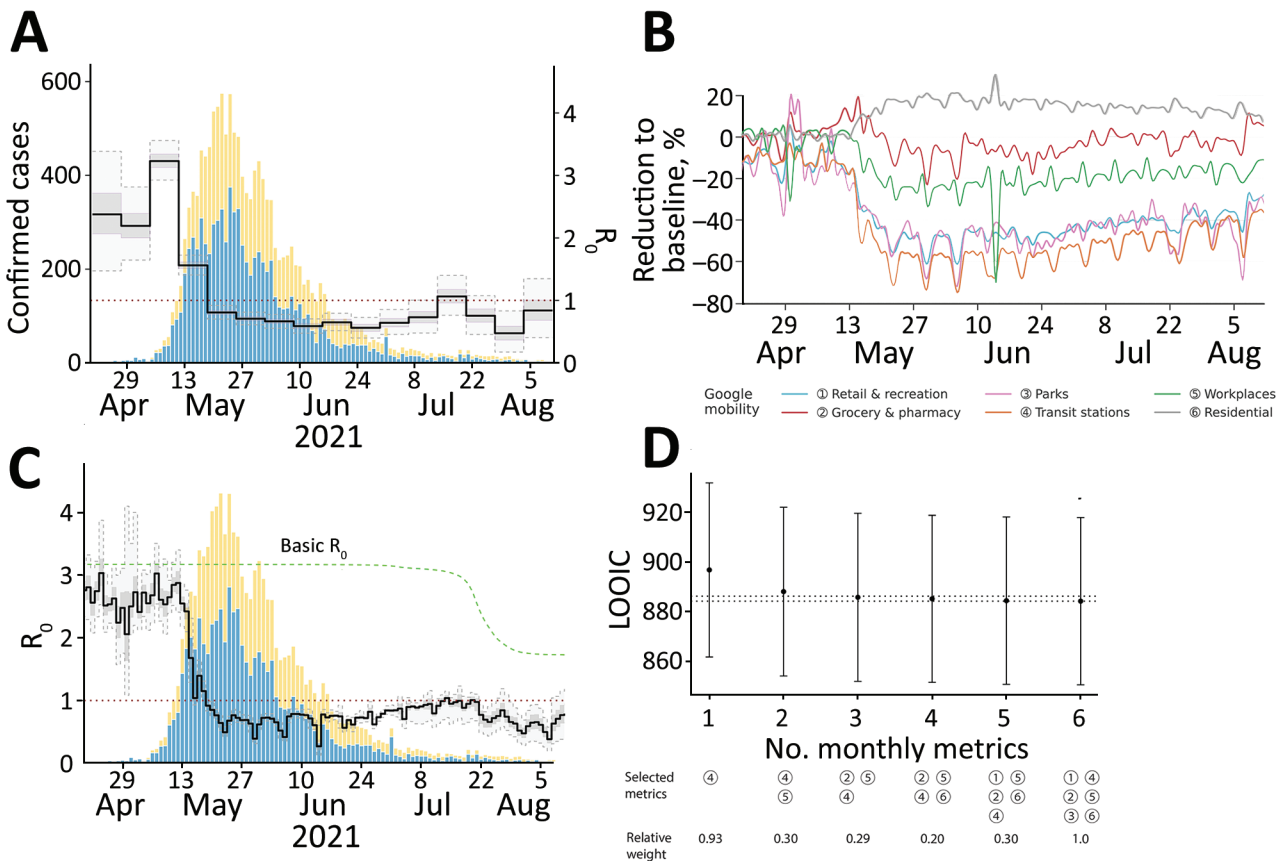


Figure 3. R_t by infection date and its link to mobility patterns for epidemic wave of COVID-19 in Taiwan, April–August 2021. A) The change in R_t modeled by a piecewise constant function with a 7-day time window. B) The R_t inferred based on monotonically decreasing basic reproduction number (green) and 6 mobility metrics. C) The temporal dynamics of mobility metrics. D) Comparison of different models based on LOOIC values under a restricted number of mobility metrics (numbers defined in panel B). The legend indicates the set of metrics with highest probability of selection shown by relative weight. Dashed lines contain the region where the change in LOOIC values does not exceed 2 from the minimum, implying a relatively equivalent fit to the data; error bars indicate SD. The blue and yellow bars in A and C are the same as in Figure 2, panel A. LOOIC, leave-one-out information criteria; R_t , effective reproduction number.

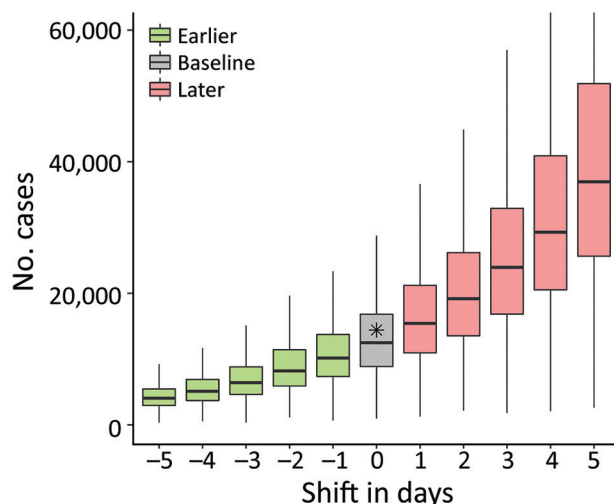


Figure 4. Epidemic wave of COVID-19 in Taiwan, April–August 2021. Model shows impact on epidemic size (by August 14, 2021) of a delay in implementing level 3 prevention measures (Appendix Figure 2, <https://wwwnc.cdc.gov/EID/article/28/10/22-0456-App1.pdf>) or of implementing them earlier. Horizontal line within boxes indicate medians; box tops and bottoms indicate interquartile ranges; whiskers indicate 95% CIs. Gray box indicates baseline scenario; asterisk indicates observed data.

symptomatic later. We also did not account for age, sex, and spatial structures in our framework for estimating R_t ; including those factors could have provided more insight into the transmission dynamics. We also did not distinguish high-risk and low-risk transmission venues in our statistical model, nor did we account for the contribution of superspreading events. We noted, with interest, that the Alpha variant was not the only variant detected among the locally acquired infections during the investigation period. An outbreak associated with the Delta variant also was reported in June 2021, which surfaced in Pingtung County in the south of Taiwan and was contained within 2 weeks. The cluster originated from 2 travelers who returned to Taiwan from Peru and involved a total of 17 cases.

In 2021, Taiwan's pandemic response demonstrated that, despite low levels of vaccine coverage, containment and elimination of COVID-19 remained feasible. The timely introduction of PHSMs helped Taiwan to avoid healthcare system collapse, and the PHSM strategies employed serve as an example for future outbreaks of emerging and re-emerging infectious diseases. In the case of SARS-CoV-2, the continued evolution of the virus toward higher transmissibility and immune evasion poses a continued threat. It is clear from the 2022 Omicron waves in Taiwan and elsewhere that high levels of vaccine coverage, although offering

protection against severe disease, are insufficient in preventing transmission. PHSMs beyond vaccination might become necessary again for future SARS-CoV-2 epidemic waves.

Acknowledgments

The authors are grateful to Jenny Wu, Chih-Chan Lan, and Tzu-You Lin for helpful discussions and assistance in data collection and Taiwan public health authorities and institutions for surveillance, laboratory testing, epidemiological investigations, and data collection. A.R.A. thanks Katherine Thielges for editing the first draft of this manuscript.

This study was approved by the Research Ethics Committee of National Taiwan University (202203HM021). A.R.A. was supported by the National Science and Technology Council, Taiwan (NSTC #111-2314-B-002-289).

About the Author

Dr. Akhmetzhanov is an assistant professor in the Global Health Program and Institute of Epidemiology and Preventive Medicine of the College of Public Health, National Taiwan University, Taiwan. His research interests include the epidemiology and prevention of infectious disease outbreaks.

References

- Kang CR, Lee JY, Park Y, Huh IS, Ham HJ, Han JK, et al.; Seoul Metropolitan Government COVID-19 Rapid Response Team (SCoRR Team). Coronavirus disease exposure and spread from nightclubs, South Korea. *Emerg Infect Dis*. 2020;26:2499–501. <https://doi.org/10.3201/eid2610.202573>
- Takaya S, Tsuzuki S, Hayakawa K, Kawashima A, Okuhama A, Kanda K, et al. Nightlife clusters of coronavirus disease in Tokyo between March and April 2020. *Epidemiol Infect*. 2020;148:e250. <https://doi.org/10.1017/S0950268820002496>
- Oshitani H; Expert Members of The National COVID-19 Cluster Taskforce at The Ministry of Health, Labour and Welfare, Japan. Cluster-based approach to coronavirus disease 2019 (COVID-19) response in Japan—February–April 2020. *Jpn J Infect Dis*. 2020;73:491–3. <https://doi.org/10.7883/yoken.JJID.2020.363>
- Pan A, Liu L, Wang C, Guo H, Hao X, Wang Q, et al. Association of public health interventions with the epidemiology of the COVID-19 outbreak in Wuhan, China. *JAMA*. 2020;323:1915–23. <https://doi.org/10.1001/jama.2020.6130>
- Flaxman S, Mishra S, Gandy A, Unwin HJT, Mellan TA, Coupland H, et al.; Imperial College COVID-19 Response Team. Estimating the effects of non-pharmaceutical interventions on COVID-19 in Europe. *Nature*. 2020;584:257–61. <https://doi.org/10.1038/s41586-020-2405-7>
- Di Domenico L, Sabbatini CE, Boëlle PY, Poletto C, Crépey P, Paireau J, et al. Adherence and sustainability of interventions informing optimal control against the COVID-19 pandemic. *Commun Med (Lond)*. 2021;1:57. <https://doi.org/10.1038/s43856-021-00057-5>
- Li Y, Campbell H, Kulkarni D, Harpur A, Nundy M,

- Wang X, et al.; Usher Network for COVID-19 Evidence Reviews (UNCOVER) group. The temporal association of introducing and lifting non-pharmaceutical interventions with the time-varying reproduction number (R) of SARS-CoV-2: a modelling study across 131 countries. *Lancet Infect Dis*. 2021;21:193–202. [https://doi.org/10.1016/S1473-3099\(20\)30785-4](https://doi.org/10.1016/S1473-3099(20)30785-4)
8. Brauner JM, Mindermann S, Sharma M, Johnston D, Salvatier J, Gavenčiak T, et al. Inferring the effectiveness of government interventions against COVID-19. *Science*. 2021;371:eabd9338. <https://doi.org/10.1126/science.abd9338>
 9. Yang B, Huang AT, Garcia-Carreras B, Hart WE, Staid A, Hitchings MDT, et al.; UFCOVID Interventions Team. Effect of specific non-pharmaceutical intervention policies on SARS-CoV-2 transmission in the counties of the United States. *Nat Commun*. 2021;12:3560. <https://doi.org/10.1038/s41467-021-23865-8>
 10. Chernozhukov V, Kasahara H, Schrimpf P. The association of opening K-12 schools with the spread of COVID-19 in the United States: County-level panel data analysis. *Proc Natl Acad Sci U S A*. 2021;118:e2103420118. <https://doi.org/10.1073/pnas.2103420118>
 11. Dighe A, Cattarino L, Cuomo-Dannenburg G, Skarp J, Imai N, Bhatia S, et al. Response to COVID-19 in South Korea and implications for lifting stringent interventions. *BMC Med*. 2020;18:321. <https://doi.org/10.1186/s12916-020-01791-8>
 12. Ku D, Yeon C, Lee S, Lee K, Hwang K, Li YC, et al. Safe traveling in public transport amid COVID-19. *Sci Adv*. 2021;7:eabg3691. <https://doi.org/10.1126/sciadv.abg3691>
 13. Cori A, Ferguson NM, Fraser C, Cauchemez S. A new framework and software to estimate time-varying reproduction numbers during epidemics. *Am J Epidemiol*. 2013;178:1505–12. <https://doi.org/10.1093/aje/kwt133>
 14. Abbott S, Hellewell J, Thompson RN, Sherratt K, Gibbs HP, Bosse NI, et al. Estimating the time-varying reproduction number of SARS-CoV-2 using national and subnational case counts [version 2, peer review: 1 approved with reservations]. *Wellcome Open Res*. 2020;5:112. <https://doi.org/10.12688/wellcomeopenres.16006.2>
 15. Röst G, Bartha FA, Bogya N, Boldog P, Dénes A, Ferenci T, et al. Early phase of the COVID-19 outbreak in Hungary and post-lockdown scenarios. *Viruses*. 2020;12:708. <https://doi.org/10.3390/v12070708>
 16. Abbott S, Hellewell J, Sherratt K, Gostic K, Hickson J, Badr HS, et al. EpiNow2: estimate real-time case counts and time-varying epidemiological parameters. 2020 [cited 2022 Jul 15]. <https://zenodo.org/record/4088545>
 17. Jung SM, Endo A, Akhmetzhanov AR, Nishiura H. Predicting the effective reproduction number of COVID-19: inference using human mobility, temperature, and risk awareness. *Int J Infect Dis*. 2021;113:47–54. <https://doi.org/10.1016/j.ijid.2021.10.007>
 18. Knock ES, Whittles LK, Lees JA, Perez-Guzman PN, Verity R, FitzJohn RG, et al. Key epidemiological drivers and impact of interventions in the 2020 SARS-CoV-2 epidemic in England. *Sci Transl Med*. 2021;13:eabg4262. <https://doi.org/10.1126/scitranslmed.abg4262>
 19. Leung K, Wu JT, Leung GM. Real-time tracking and prediction of COVID-19 infection using digital proxies of population mobility and mixing. *Nat Commun*. 2021;12:1501. <https://doi.org/10.1038/s41467-021-21776-2>
 20. Lin Y, Yang B, Cobey S, Lau EHY, Adam DC, Wong JY, et al. Incorporating temporal distribution of population-level viral load enables real-time estimation of COVID-19 transmission. *Nat Commun*. 2022;13:1155. <https://doi.org/10.1038/s41467-022-28812-9>
 21. Rüdiger S, Konigorski S, Rakowski A, Edelman JA, Zernick D, Thieme A, et al. Predicting the SARS-CoV-2 effective reproduction number using bulk contact data from mobile phones. *Proc Natl Acad Sci U S A*. 2021;118:e2026731118. <https://doi.org/10.1073/pnas.2026731118>
 22. Summers J, Cheng HY, Lin HH, Barnard LT, Kvalsvig A, Wilson N, et al. Potential lessons from the Taiwan and New Zealand health responses to the COVID-19 pandemic. *Lancet Reg Health West Pac*. 2020;4:100044. <https://doi.org/10.1016/j.lanwpc.2020.100044>
 23. Akhmetzhanov AR, Jung SM, Cheng HY, Thompson RN. A hospital-related outbreak of SARS-CoV-2 associated with variant Epsilon (B.1.429) in Taiwan: transmission potential and outbreak containment under intensified contact tracing, January–February 2021. *Int J Infect Dis*. 2021;110:15–20. <https://doi.org/10.1016/j.ijid.2021.06.028>
 24. Wang CJ, Ng CY, Brook RH. Response to COVID-19 in Taiwan: big data analytics, new technology, and proactive testing. *JAMA*. 2020;323:1341–2. <https://doi.org/10.1001/jama.2020.3151>
 25. Taiwan Centers for Disease Control. COVID-19 (SARS-CoV-2 Infection) [cited 2022 Jul 11]. <https://www.cdc.gov.tw>
 26. World Health Organization. COVID-19 clinical management: living guidance. 2021 Jan 25 [cited 2022 Jul 15]. <https://www.who.int/publications/i/item/WHO-2019-nCoV-clinical-2021-1>
 27. Google Inc. COVID-19 community mobility reports. [cited 2022 Jul 15]. <https://www.google.com/covid19/mobility/Taiwan>
 28. Reich NG, Lessler J, Cummings DAT, Brookmeyer R. Estimating incubation period distributions with coarse data. *Stat Med*. 2009;28:2769–84. <https://doi.org/10.1002/sim.3659>
 29. Linton NM, Kobayashi T, Yang Y, Hayashi K, Akhmetzhanov AR, Jung SM, et al. Incubation period and other epidemiological characteristics of 2019 novel coronavirus infections with right truncation: a statistical analysis of publicly available case data. *J Clin Med*. 2020;9:538. <https://doi.org/10.3390/jcm9020538>
 30. van de Kasstele J, Eilers PHC, Wallinga J. Nowcasting the number of new symptomatic cases during infectious disease outbreaks using constrained P-spline smoothing. *Epidemiology*. 2019;30:737–45. <https://doi.org/10.1097/EDE.0000000000001050>
 31. Nakajo K, Nishiura H. Estimation of $R(t)$ based on illness onset data: An analysis of 1907–1908 smallpox epidemic in Tokyo. *Epidemics*. 2022;38:100545. <https://doi.org/10.1016/j.epidem.2022.100545>
 32. He X, Lau EHY, Wu P, Deng X, Wang J, Hao X, et al. Temporal dynamics in viral shedding and transmissibility of COVID-19. *Nat Med*. 2020;26:672–5. <https://doi.org/10.1038/s41591-020-0869-5>
 33. Delamater PL, Street EJ, Leslie TF, Yang YT, Jacobsen KH. Complexity of the basic reproduction number (R_0). *Emerg Infect Dis*. 2019;25:1–4. <https://doi.org/10.3201/eid2501.171901>
 34. Vehtari A, Gelman A, Gabry J. Practical Bayesian model evaluation using leave-one-out cross-validation and WAIC. *Stat Comput*. 2017;27:1413–32. <https://doi.org/10.1007/s11222-016-9696-4>

Address for correspondence: A.R. Akhmetzhanov, National Taiwan University, College of Public Health, No. 17 Xuzhou Rd. Zhongzheng District, Taipei 10055, Taiwan; email: akhmetzhanov@ntu.edu.tw

Transmission Dynamics and Effectiveness of Control Measures during a COVID-19 Surge, Taiwan, April–August 2021

Appendix

Estimation of the serial interval

Estimation of the serial interval was performed by selecting symptomatic infector-infectee pairs from a series of epidemiological investigations conducted by Taiwan Centers for Disease Control (CDC) from the beginning of 2020 through March 2021 (i.e. preceding the epidemic wave under analysis). They were assigned as “certain” or “probable” pairs depending on the strength of the available evidence supporting the epidemiological linkage and directionality of transmission (accurately identifying who was the infector and who was the infectee). Three shifted distributions (gamma, Weibull, and lognormal) were fitted to data on the time between onset of symptoms of infectors and onset of symptoms of infectees for 33 certain infector-infectee pairs and the best-fit distribution was identified using the Bayesian mixture model. In that setting the best-fit distribution was the one with the maximum posterior probability. Inclusion of 25 probable infector-infectee pairs did not significantly alter the inferred estimates. The complete list of infector-infectee pairs used for estimation of the serial interval is provided (Appendix Table 1).

Estimation of within-individual time distributions

The distribution of time intervals from one event of interest to another, including:

- symptom onset to case confirmation,
- symptom onset to severe disease,
- symptom onset to death,
- symptom onset to report of death, and
- death to report of death

were fitted to a mixture of three unimodal (gamma, Weibull, and lognormal) distributions within the framework of a Bayesian mixture model. Each of the three component distributions was defined by the probability density function $f_l(o; \theta)$ ($l = 1, 2, 3$) that was given a two-dimensional parameter vector $\theta = (\mu, \nu)$, where μ was the mean and ν was the coefficient of variation (CV). Both parameters were common to all three component distributions.

Suppose that the time of the first event for a case k ($k \in K$) was denoted by o_k and the time of the second event was denoted by c_k . When both times were recorded in the dataset, they were given by their calendar dates, \mathcal{O}_k and \mathcal{C}_k , respectively. Both times, o_k and c_k , were assigned to uniform priors of 1-day:

$$o_k \sim \text{Uniform}(\mathcal{O}_k, \mathcal{O}_k + 1 \text{ day}) \quad (1)$$

$$c_k \sim \text{Uniform}(\max(\{o_k, \mathcal{C}_k\}), \mathcal{C}_k + 1 \text{ day}) \quad (2)$$

where $\mathcal{O}_k \leq \mathcal{C}_k$ was held for all aggregated records, and $o_k < c_k$ was imposed for simplicity for $\mathcal{O}_k = \mathcal{C}_k$. Because of a non-zero probability of viral RNA detection in a sample from a presymptomatic individual, the condition $o_k < c_k$ can be invalid in general. However, when the alternative model with shifted distributions was verified, we found that it did not improve the data fit, and therefore chose to use the simpler model version.

When the date of the first event, \mathcal{O}_k , was not recorded, the case was asymptomatic at the time of testing, \mathcal{T}_k . According to the data collection procedures of the Taiwan CDC, the date of testing \mathcal{T}_k is backlogged by 1 day or occurs on the same day as the reporting day \mathcal{R}_k if confirmation of the case represents social significance defined based on expert opinion within the agency. Because the severity status of all confirmed cases was regularly updated by the Taiwan CDC, we noticed that some cases with an initially unknown symptom onset date \mathcal{O}_k became severe at later dates, denoted as \mathcal{S}_k . Suppose that two conditions are additionally met: $\mathcal{T}_k \leq \mathcal{C}_k$ and $\mathcal{R}_k < \mathcal{S}_k < \mathcal{C}_k$. The prior (1) can then be replaced with the following prior valid for that subset of cases:

$$o_k \sim \text{Uniform}(\mathcal{T}_k = \mathcal{R}_k - 1 \text{ day}, \mathcal{S}_k + 1 \text{ day}) \quad (3)$$

Otherwise, the cases were omitted from the analysis.

The likelihood was defined by the weighted sum of three components:

$$L_{\Sigma}(\theta; D = \{o_k, c_k\}) = \sum_{l=1}^3 w_l L_l(\theta; D) \quad (4)$$

$$L_l(\theta; D) = \prod_{k \in K} \frac{f_l(c_k - o_k; \theta)}{F_l(T - o_k; \theta)} \quad (5)$$

where $F_l(o; \theta)$ is the cumulative distribution function of $f_l(o; \theta)$, and T is the date of the latest release of data by Taiwan CDC, set as 2 p.m. on of August 25, 2021. The probability of selecting the best-fit distribution could be then given by categorical sampling from the three components with relative probabilities $w_l L_l / L_{\Sigma}$ ($l = 1, 2, 3$).

Time-varied reporting delay

The effective reproduction number, R_t , and case-fatality ratio are often estimated retrospectively when an epidemic has already been declared over, but they can also be estimated in real-time. However, real-time estimation presents unavoidable challenges to researchers (1,2). First, snapshots of count data are right-truncated respective to the time of the latest update. In other words, cases that have already been infected but not yet confirmed are not recorded in the data. Unless these truncated cases are accounted for—e.g., by nowcasting— R_t will always show a decreasing trend toward zero just before the latest data cutoff date. Second, a substantial fraction of case records may not contain the necessary information, such as date of symptom onset, as that information might not be required for initial reporting of a case (particularly if the confirmed case definition is based primarily on a positive laboratory test) and thus might only be collected retrospectively. COVID-19 cases can be completely asymptomatic, pre-symptomatic at the time of testing and not followed up later, or have unknown symptom status (e.g., owing to patient noncooperation or a deteriorating health condition). Because missing symptom onset dates can be backprojected from reporting dates, identification of the reporting delay distribution becomes an important quantity to estimate. Hence, we estimated a time-varied reporting delay distribution that allowed us to backproject the symptom onset date for cases with unknown presence of symptoms and to nowcast cases that were not yet reported in Taiwan’s COVID-19 epidemic wave in mid-2021.

In our framework, the coefficient of variation (CV) of the reporting delay ν was set constant, but the mean reporting delay μ varied over multiple lengthscales similarly to previous work (3). Two lengthscales were used, each characterized by two parameters: W , which defined a long-range lengthscale and ω , which defined a short-range lengthscale. Their implementation

in our statistical framework is described in more detail below. The baseline values for W and ω were set to 10 days and 7 days, respectively (Appendix Figure 1). The sensitivity analysis included other values, $W = \{5 \text{ days}, 15 \text{ days}\}$ and $\omega = \{3 \text{ days}, 11 \text{ days}\}$, demonstrating relative robustness (Appendix Figure 2). A comparison with a piecewise constant reporting delay was also performed (Appendix Figure 3).

For the long-range variation in the reporting delay controlled by the parameter W , the temporal dynamics of its mean were modeled using a cubic B-spline, $\mu(t) = \mu_{\kappa,\theta}(t)$, such that:

$$\mu_{\kappa,\theta}(t) = \sum_{n=1}^N \vartheta_n B_{n,\kappa}(t, \theta) \quad (6)$$

where $\kappa = 3$ is the degree of the B-spline, ϑ_n ($n = 1, \dots, N$) are B-spline coefficients, $\theta = \{\theta_m, m = 0, \dots, M\}$ is a knot sequence composed of equidistant time points $\theta_m = Tm/M$, and $N = M + \kappa - 1$ is the total number of the basis spline functions (4). The sensitivity of the B-spline function was controlled by the parameter W . The number of splits M was pre-defined as $M = \lfloor T/W \rfloor + 1$, where the operator $\lfloor T/W \rfloor$ was the floor division of T on W . The choice was made to ensure that the distance between the nearest knots, $\delta = T/M$, was not larger than W and had a finite lower bound proportional to W , $W(1 - 1/M) \leq \delta \leq W$.

The likelihood for a short-range variation in the reporting delay controlled by the parameter ω was defined as follows:

$$L_l(\theta_t = \{\mu_{\kappa,\theta}(t), \nu\}, \{o_k, c_k \mid k \in K(t)\}) = \prod_{k \in K(t)} \frac{f_l(c_k - o_k, \theta_t)}{F_l(T - o_k, \theta_t)} \quad (7)$$

where the set $K(t)$ consisted of all cases whose symptom onset date O_k (or confirmation date C_k) was within the time window ω : $t - \omega \leq O_k \leq t + \omega$ (or $t - \omega \leq C_k \leq t + \omega$), o_k and c_k are the inferred symptom onset and confirmation time:

$$o_k \sim \text{Uniform}(O_k, O_k + 1 \text{ day}),$$

$$c_k \sim \text{Uniform}(\max(\{o_k, C_k\}), C_k + 1 \text{ day}),$$

as for all records $O_k \leq C_k$. The distribution l was chosen among the gamma, Weibull, and lognormal distributions ($l = 1, 2, 3$). The overall likelihood was present by a sum of three components:

$$L_\Sigma(\theta, D = \{O_k, C_k\}) = \sum_{l=1}^3 w_l \prod_t L_l(\theta_t, \{o_k, c_k \mid k \in K(t)\}) \quad (8)$$

Because the reporting delay could be specified either by the date of symptom onset \mathcal{O}_k (a “forward-looking” reporting delay) or by the date of confirmation \mathcal{C}_k (a “backward-looking” reporting delay according to notations (5)), the two possible reporting delay distributions could lead to different results. The forward reporting delay, indicated by “(o)” in our work, e.g., $f^{(o)}$, describes the situation in which the mean reporting delay was estimated using the left-end symptom onset date. By comparison, the backward reporting delay, indicated by “(c)”, e.g., $f^{(c)}$, describes the situation in which the mean reported delay was estimated using the right-end confirmation date. Depending on the concrete application, one form can be preferred over another and would allow different interpretations. For instance, when cases with unknown symptom onset dates were backprojected to their symptom onset dates, the form $f^{(c)}$ could be appropriately used. When the cases were nowcasted to a given confirmation date to predict the number of cases that had not yet been reported, the form $f^{(o)}$ was appropriate. Park and colleagues (5) argued that the backward-looking delay distribution is prone to a bias toward lower values when an exponential growth in cases is observed during an epidemic, whereas the forward-looking delay distribution is likely to be representative of the unbiased (intrinsic) delay distribution.

Thus, the reporting delay distribution was modeled based on a mixture of three unimodal distributions (gamma, Weibull, and lognormal). The plausibility of each distribution was determined by its relative weight w_l in the likelihood L_Σ , when the probability of selecting the distribution l was equal to $w_l L_l / L_\Sigma$ (see equations (7) and (8)). The temporary change in the reporting delay was modeled by a time-varied mean reporting delay, whereas the CV of the reporting delay was kept unchanged. The alternative formulation, where the standard deviation (SD) was used instead of the CV and kept constant, did not fit the data, so it was omitted from our analysis. The time variation of the mean reporting delay μ was modeled at two timescales: for the long timescale, the change in $\mu(t)$ was defined by a cubic B-spline function, while for the short timescale the reporting delay distribution was adjusted locally using the likelihood (7). The resulting fit was obtained by combining the two timescales and quantified either by the symptom onset date, denoted by the upper index (o), or by the confirmation date, denoted by the upper index (c).

Effective reproduction number by date of symptom onset

To estimate the effective reproduction number by the date of symptom onset, one needs to know whether the case is asymptomatic or symptomatic. If the case is symptomatic, the symptom onset date should be recorded. However, the extracted dataset was composed of two types of case records—some records contained the symptom onset date and the confirmation date, while others had a blank symptom onset date. For the latter, the asymptomatic status was not definitive, and the record could not differentiate truly asymptomatic and pre-symptomatic cases. The unknown symptom onset dates were backprojected from the case confirmation dates using the reporting delay distribution $f^{(c)}$.

Let i_t be the daily counts of cases with known symptom onset dates at day of symptom onset t , and y_t be the daily counts of cases with unknown symptom onset dates at day of confirmation t . The expected daily count i_t^* that combines both was defined as follows:

$$i_t^* = i_t + \sum_{\tau>0} y_{t+\tau} f_{l,\tau}^{(c)}(\theta_{t+\tau}) \quad (9)$$

where $f_{l,\tau}^{(c)}(\theta_{t+\tau})$ is a discretized reporting delay distribution, $f_{l,\tau}^{(c)}(\theta_t) = F_l^{(c)}(\tau, \theta_t) - F_l^{(c)}(\tau - 1, \theta_t)$ for $\tau = 1, 2, \dots$, and the function $F_l^{(c)}(\circ, \theta_t)$ is the cumulative distribution function of the reporting delay $f_l^{(c)}(\circ, \theta_t)$.

To account for right truncation, the expected counts i_t^* were nowcasted according to (6):

$$i_t^{nc} = \frac{i_t^*}{F_l^{(o)}(T-t, \theta_t)} \quad (10)$$

where the function $F_l^{(o)}(\circ, \theta_t)$ is a cumulative distribution function of the forward reporting delay distribution $f_l^{(o)}(\circ, \theta_t)$.

The effective reproduction number by date of symptom onset, $R_t^{(o)}$, was estimated using the renewal process written within the negative binomial likelihood:

$$i_t^{nc} \sim \text{NegBinom}(\text{mean} = E(i_t^{nc}), \text{overdisp.} = \phi_{\{1\dots 4\}})$$

$$E(i_t^{nc}) = R_t^{(o)} \sum_{\tau=1}^{t-1} i_{t-\tau}^{nc} \cdot g_\tau \quad (11)$$

where g_τ is the serial interval distribution estimated for Taiwan consisting of the best-fit lognormal distribution with a mean (\pm SD) of 4.6 ± 3.4 days, which is in line with previous

reports (7,8). The overdispersion parameter ϕ followed one of the four functional forms: (i) $\phi_1 \equiv \text{const}$, which is a commonly used form of the negative binomial distribution, also known as NB2, (ii) $\phi_2 \equiv \text{const} \cdot E(i_t^{nc})$, which resembles a quasi-Poisson distribution, also known as NB1, (iii) $\phi_3 \equiv \text{const} \cdot (E(i_t^{nc}))^{1/2}$, which presents an intermediate situation between (i) and (ii), and (iv) $\phi_4 \gg 1$, which is a limiting case leading to the Poisson likelihood. Green and colleagues (9) previously indicated a better fit of the data when negative binomial likelihood (i) is used instead of Poisson likelihood (iv). However, other studies (10,11) have shown that the overdispersion parameter ϕ given by (iii) is a better choice than (i) or (ii). In our framework, all four likelihoods (i)–(iv) were implemented within the Bayesian mixture model with a similar likelihood as in (8). The best-fit configuration was directly selected from the results of statistical inference.

Because the nowcasted counts i_t^{nc} (10) are expected values and are not integer counts of incidence, the negative binomial likelihood (11) was replaced with the gamma likelihood that matched the first and the second moments (12):

$$i_t^{nc} \sim \text{Gamma}(\text{shape} = \alpha_t, \text{rate} = r_t) \quad (12)$$

where $r_t = \phi_{\{1\dots4\}} / (\phi_{\{1\dots4\}} + E(i_t^{nc}))$ and $\alpha_t = E(i_t^{nc}) \cdot r_t$.

Finally, the effective reproduction number was smoothed by calculating the centered rolling average over a 5-day time window (13), which implied that the value $R_t^{(o)}$ on day t was estimated using five sequential values of the nowcasted counts on days $\{t - 2, t - 1, t, t + 1, t + 2\}$.

Effective reproduction number by date of infection

The number of infections j_t on day t was generated by nowcasted cases with symptom onset dates on either side of that day and was proportional to the effective reproduction number by the date of infection $R_t^{(i)}$. Framed within the renewal process framework (14), the equation holds the form:

$$j_t = R_t^{(i)} \sum_u i_{t-u}^{nc} \cdot \lambda_u \quad (13)$$

where λ_u denotes the profile of infectiousness, which is the probability of a secondary transmission from a primary case at time u measured with respect to the symptom onset date.

Notably, u covers both positive and negative integers. Negative integers imply pre-symptomatic transmission, which is characteristic of SARS-CoV-2 infection. The profile of infectiousness λ_u was previously estimated using a gamma distribution shifted 12.3 days to the left that peaked at time zero and indicated 44% of pre-symptomatic infections (15).

Because newly infected cases j_t experience symptom onsets that are time-lagged by the incubation period, we can write:

$$E(i_t^{nc}) = \sum_{\tau=1}^{t-1} j_{t-\tau} h_{\tau} \quad (14)$$

where h_{τ} defines the incubation period fitted by a lognormal distribution with a mean of 5.2 days (16). Combining equations (13)–(14) results in the renewal equation previously derived by Nakajo and Nishiura (14):

$$E(i_t^{nc}) = \sum_{\tau=0}^t R_{t-\tau}^{(i)} h_{\tau} \sum_{u=-x}^{t-\tau} i_{t-\tau-u}^{nc} \lambda_u \quad (15)$$

where $x = 13$ days is the least integer for the shift of 12.3 days. The observation model was implemented using a negative binomial likelihood function analogous to equation (13), however, $E(i_t^{nc})$ was replaced by a double sum (15).

The effective reproduction number was first modeled by a piecewise constant function of time t . The change points were set to every 7 days, which was a rough approximation of both the generation time (17–19) and a calendar week. Two alternative approaches were also explored. In the first approach, the change in the effective reproduction number was attributed to PHSMs (Appendix Table 2). In this case, the effective reproduction number remained constant during the intervening time periods but changed abruptly when the PHSMs were implemented. In the second approach, the effective reproduction number was correlated with a change in mobility metrics and involved a sigmoidal, monotonically decreasing change in the baseline (basic) reproduction number, for example, owing to behavioral changes or more efficient case finding and contact tracing toward the end of the epidemic in August–September 2021. The effect of overall mobility reduction was modeled by a combination of mobility metrics in different settings. In this case, the effective reproduction number changed on the log scale according to the following equation:

$$\log R_t^{(i)} = \log R_{0,t} + \sum_{p=1}^P \beta_p X_t^{(p)} \quad (16)$$

where the intercept $\log R_{0,t}$ is the logarithm of the baseline reproduction number given by a sigmoidal function over time (20):

$$R_{0,t} = R_0 \left(\eta + \frac{1-\eta}{1+\exp(\xi(t-t_{sw}-v))} \right) \quad (17)$$

where each of five parameters, R_0 , η , v , ξ , and t_{sw} , were assigned to a weakly informative prior. The parameter R_0 defined the baseline reproduction number at the beginning of the epidemic. The parameter η measured the reduction in transmission during the later stage of the epidemic, $\eta \sim \text{Beta}(2.5, 4)$, v measured the delay in the reduction of the baseline reproduction number, $v \sim \text{Exponential}(1/5)$, ξ measured the slope of the sigmoidal function, which was restricted to be negative, $\xi \sim \text{Uniform}(0.5, 1.5)$, and t_{sw} defined the change point of the baseline reproduction number during late stage of the epidemic, $t_{sw} \sim \text{Uniform}(\text{July 1, 2021, August 25, 2021})$. The variable $X_t^{(p)}$ defined the p th mobility metric on day t across the total $P = 6$ metrics.

The restricted number of mobility metrics $P_{\min} \leq P$ was identified by comparing the “leave-one-out information criterion” LOOIC values for the respective models with the total P_{\min} metrics and $C_p^{P_{\min}}$ combinations among all P metrics. The LOOIC is commonly used in Bayesian frameworks for model selection (21). Specifically, the fit of the model with only P_{\min} mobility metrics was compared to the fit of other models by selecting the model with the smallest LOOIC value. The negative binomial likelihood used for the data fit was analogous to (11) and was sequentially replaced by the gamma likelihood (12).

Counterfactual scenarios

The statistical model for the effective reproduction number based on mobility metrics was applied to predict the incidence of COVID-19 under different counterfactual scenarios. Compared with the baseline scenario shown in Appendix Figure 1, panel A, the counterfactual scenarios described situations in which a set of Level 3 measures, initially introduced on May 15, 2021, was shifted either to the right or to the left by y days. This was modeled by accordingly shifting the mobility patterns starting from May 15, 2021. When the mobility patterns were shifted to the left (i.e., earlier implementation of PHSMs), the mobility metrics during the period of (May 15, 2021 – y days) to (May 15, 2021 – 1 day) were replaced with the shifted mobility metrics starting from May 15. Owing to the shift, the final ($y - 1$) days before the ending time T

remained unfilled. They were replaced with zeros because they had a negligible effect on the results of the counterfactual scenario. When the mobility metrics were shifted to the right (i.e., a later implementation of PHSMs), the unfilled gap in metrics covering the time period from May 15, 2021 through (May 15, 2021 – $(y - 1)$ days) was filled with mobility readings randomly imputed from a 2-week period between May 1 and May 14. For instance, if the gap was 3 days ($y = 3$), the mobility metrics for those 3 days were imputed by randomly selecting 3 days from May 1–14. The result of each counterfactual scenario was expressed as a cumulative number of infections observed as of August 14, 2021. This cutoff point was chosen because it marked a 3-month period after the initial Level 3 implementation on May 15.

The following protocol was used to perform computer simulations. The seeded number of infections j_0 and the seeding time t_0 were chosen based on a grid search and calculation of either the root-mean-square error (RMSE) or Dawid–Sebastiani score (DSS) (22,23). Prior to the seeding time, all infections were set to zero. Because the transmission chains observed in April 2021 were thoroughly investigated by Taiwanese authorities, we assumed that the seeding time was between April 5 and April 20, 2021 and the number of seeded infections was unlikely to exceed five. The RMSE value was derived by comparing the simulated incidence \tilde{i}_t characterized by the symptom onset date and the nowcasted incidence i_t^{nc} according to the following formula:

$$RMSE(i_t^{nc}, \tilde{i}_t) = \left(\sum_{t=t_{min}}^{t_{max}} \frac{(i_t^{nc} - \tilde{i}_t)^2}{K} \right)^{1/2} \quad (18)$$

$$DSS(i_t^{nc}, \tilde{i}_t) = \left(\frac{i_t^{nc} - \mu_{\mathcal{P}_t}}{\sigma_{\mathcal{P}_t}} \right)^2 + 2 \log \sigma_{\mathcal{P}_t} \quad (19)$$

where $K = t_{max} - t_{min} + 1$ is the total number of data points from t_{min} to t_{max} covering the observed time period. t_{min} was set to April 21, 2021 (1 day after the end of the seeding time period), t_{max} was set to August 14, 2021, as discussed earlier. The DSS value representing a proper scoring rule was calculated by comparing the nowcasted counts with the posterior predictive distribution \mathcal{P}_t . Denoting the mean of the posterior distribution as $\mu_{\mathcal{P}_t}$ and the standard deviation as $\sigma_{\mathcal{P}_t}$, we can write:

The model with smaller values of DSS or RMSE was preferred over the others.

To predict the case counts \tilde{i}_t and their posterior distribution \mathcal{P}_t for a given counterfactual or baseline scenario, the following two-step procedure was implemented. In the first step, the number of infections \tilde{j}_t was obtained using the renewal process and the Poisson count model:

$$\tilde{j}_t \sim \text{Poisson}(E(\tilde{j}_t))$$

$$E(\tilde{j}_t) = R_t^{(i)} \sum_{\tau=1}^{t-1} \tilde{j}_{t-\tau} \cdot \tilde{g}_\tau \quad (20)$$

for any $t > t_0$. Otherwise, $\tilde{j}_{t=t_0} = j_0$ and $\tilde{j}_{t < t_0} = 0$. The effective reproduction number $R_t^{(i)}$ was defined by equation (16) and included modified mobility metrics according to the chosen counterfactual scenario. The distribution \tilde{g}_τ defined the generation time distribution that was previously estimated by the gamma distribution with a mean (\pm SD) of 5.7 ± 1.7 days (18). Currently, there is no evidence that the generation time intervals were substantially different between the Alpha variant and the wild-type strain of SARS-CoV-2 (19). In the second step, the simulated case counts \tilde{i}_t were obtained by combining multinomial samples from the days preceding t , where the size parameter was equal to the number of infections and the probability distribution was given by the incubation period h :

$$\tilde{i}_t = \sum_{u>0} z_t^{(t-u)}$$

$$z_{\{s+1, s+2, \dots\}}^{(s)} \sim \text{Multinomial}(\text{size} = \tilde{j}_s, \text{prob.} = \{h_\tau, \tau = 1, 2, \dots\}) \quad (21)$$

Summary

Our framework has several important implications to advance the methods used for the real-time estimation of epidemiological parameters. First, we estimated a time-varied reporting delay distribution that measured the time between symptom onset and case confirmation. Similar to previous works (3,24,25), we performed statistical inference within a Bayesian framework and conducted Bayesian nowcasting of cases not yet reported. Our approach was based on using cubic B-splines, which is less computationally demanding than Gaussian processes (3). Our method shows a similar performance to approximating the reporting delay by a piecewise constant function, but it forces the coefficient of variation to remain constant (Appendix Figure 3). It appeared that the method was robust to varying the parameters (Appendix Figure 2). Second, we statistically inferred the effective reproduction number according to the date of infection, as proposed by Nakajo and Nishiura (14). By using a Bayesian model, we

demonstrated reliance on both pointwise and Bayesian posterior estimates while expanding the original methodology. Our methods improve the real-time estimation techniques and can be used to build accurate risk assessments and to precisely monitor the disease spread at different locations in the future.

Technical details

All model parameters were estimated by using Markov chain Monte Carlo methods within a Bayesian framework. Statistical inference was performed using the *Stan* programming language (*cmdStan* version 2.28.2). Data processing, analysis and presentation of the results were performed using *R* (version 4.1.2) and *Python* (version 3.9) with other base packages involved. The DSS and RMSE values were calculated using the *R* package *scoringutils* (26). The code scripts necessary for the replication of our results is available at:

<https://github.com/aakhmetz/COVID19-Reff-Taiwan-2021>.

References

1. Nishiura H, Klinkenberg D, Roberts M, Heesterbeek JA. Early epidemiological assessment of the virulence of emerging infectious diseases: a case study of an influenza pandemic. *PLoS One*. 2009;4:e6852. [PubMed https://doi.org/10.1371/journal.pone.0006852](https://doi.org/10.1371/journal.pone.0006852)
2. Munayco C, Chowell G, Tariq A, Undurraga EA, Mizumoto K. Risk of death by age and gender from CoVID-19 in Peru, March-May, 2020. *Aging (Albany NY)*. 2020;12:13869–81. [PubMed https://doi.org/10.18632/aging.103687](https://doi.org/10.18632/aging.103687)
3. Hawryluk I, Hoeltgebaum H, Mishra S, Miscouridou X, Schnekenberg RP, Whittaker C, et al. Gaussian process nowcasting: application to COVID-19 mortality reporting. Preprint at *arXiv*. 2021 <https://arxiv.org/abs/2102.11249>
4. Kharratzadeh M. Splines in Stan. *Stan Case Studies* 2017, 4. https://mc-stan.org/users/documentation/case-studies/splines_in_stan.html
5. Park SW, Sun K, Champredon D, Li M, Bolker BM, Earn DJD, et al. Forward-looking serial intervals correctly link epidemic growth to reproduction numbers. *Proc Natl Acad Sci U S A*. 2021;118:e2011548118. [PubMed https://doi.org/10.1073/pnas.2011548118](https://doi.org/10.1073/pnas.2011548118)
6. Tsuzuki S, Lee H, Miura F, Chan YH, Jung SM, Akhmetzhanov AR, et al. Dynamics of the pneumonic plague epidemic in Madagascar, August to October 2017. *Euro Surveill*. 2017;22:2–7. [PubMed https://doi.org/10.2807/1560-7917.ES.2017.22.46.17-00710](https://doi.org/10.2807/1560-7917.ES.2017.22.46.17-00710)

7. Nishiura H, Linton NM, Akhmetzhanov AR. Serial interval of novel coronavirus (COVID-19) infections. *Int J Infect Dis.* 2020;93:284–6. [PubMed https://doi.org/10.1016/j.ijid.2020.02.060](https://doi.org/10.1016/j.ijid.2020.02.060)
8. Ryu S, Kim D, Lim JS, Ali ST, Cowling BJ. Serial interval and transmission dynamics during SARS-CoV-2 Delta variant predominance, South Korea. *Emerg Infect Dis.* 2022;28:407–10. [PubMed https://doi.org/10.3201/eid2802.211774](https://doi.org/10.3201/eid2802.211774)
9. Greene SK, McGough SF, Culp GM, Graf LE, Lipsitch M, Menzies NA, et al. Nowcasting for real-time COVID-19 tracking in New York City: an evaluation using reportable disease data from early in the Pandemic. *JMIR Public Health Surveill.* 2021;7:e25538. [PubMed https://doi.org/10.2196/25538](https://doi.org/10.2196/25538)
10. Counotte MJ, Althaus CL, Low N, Riou J. Impact of age-specific immunity on the timing and burden of the next Zika virus outbreak. *PLoS Negl Trop Dis.* 2019;13:e0007978. [PubMed https://doi.org/10.1371/journal.pntd.0007978](https://doi.org/10.1371/journal.pntd.0007978)
11. Nouvellet P, Bhatia S, Cori A, Ainslie KEC, Baguelin M, Bhatt S, et al. Reduction in mobility and COVID-19 transmission. *Nat Commun.* 2021;12:1090. [PubMed https://doi.org/10.1038/s41467-021-21358-2](https://doi.org/10.1038/s41467-021-21358-2)
12. Li M, Dushoff J, Bolker BM. Fitting mechanistic epidemic models to data: A comparison of simple Markov chain Monte Carlo approaches. *Stat Methods Med Res.* 2018;27:1956–67. [PubMed https://doi.org/10.1177/0962280217747054](https://doi.org/10.1177/0962280217747054)
13. Gostic KM, McGough L, Baskerville EB, Abbott S, Joshi K, Tedijanto C, et al. Practical considerations for measuring the effective reproductive number, Rt. *PLOS Comput Biol.* 2020;16:e1008409. [PubMed https://doi.org/10.1371/journal.pcbi.1008409](https://doi.org/10.1371/journal.pcbi.1008409)
14. Nakajo K, Nishiura H. Estimation of R(t) based on illness onset data: An analysis of 1907-1908 smallpox epidemic in Tokyo. *Epidemics.* 2022;38:100545. [PubMed https://doi.org/10.1016/j.epidem.2022.100545](https://doi.org/10.1016/j.epidem.2022.100545)
15. He X, Lau EHY, Wu P, Deng X, Wang J, Hao X, et al. Temporal dynamics in viral shedding and transmissibility of COVID-19. *Nat Med.* 2020;26:672–5. [PubMed https://doi.org/10.1038/s41591-020-0869-5](https://doi.org/10.1038/s41591-020-0869-5)
16. Li Q, Guan X, Wu P, Wang X, Zhou L, Tong Y, et al. Early transmission dynamics in Wuhan, China, of novel coronavirus–infected pneumonia. *N Engl J Med.* 2020;382:1199–207. [PubMed https://doi.org/10.1056/NEJMoa2001316](https://doi.org/10.1056/NEJMoa2001316)

17. Ganyani T, Kremer C, Chen D, Torneri A, Faes C, Wallinga J, et al. Estimating the generation interval for coronavirus disease (COVID-19) based on symptom onset data, March 2020. *Euro Surveill.* 2020;25:2000257. [PubMed https://doi.org/10.2807/1560-7917.ES.2020.25.17.2000257](https://doi.org/10.2807/1560-7917.ES.2020.25.17.2000257)
18. Lau, Y.C., Tsang, T.K., Kennedy-Shaffer, L., Kahn, R., Lau, E., et al. Joint estimation of generation time and incubation period for coronavirus disease (Covid-19). *J. Infect. Dis.* 2021;jiab424.
19. Linton NM, Akhmetzhanov AR, Nishiura H. Correlation between times to SARS-CoV-2 symptom onset and secondary transmission undermines epidemic control efforts. *medRxiv.* 2021 Aug 31. <https://doi.org/10.1101/2021.08.29.21262512>
20. Grinsztajn L, Semenova E, Margossian CC, Riou J. Bayesian workflow for disease transmission modeling in Stan. *Stat Med.* 2021;40:6209–34. [PubMed https://doi.org/10.1002/sim.9164](https://doi.org/10.1002/sim.9164)
21. Vehtari A, Gelman A, Gabry J. Practical Bayesian model evaluation using leave-one-out cross-validation and WAIC. *Stat Comput.* 2017;27:1413–32. <https://doi.org/10.1007/s11222-016-9696-4>
22. Dawid AP, Sebastiani P. Coherent dispersion criteria for optimal experimental design. *Ann Stat.* 1999;27:65–81. <https://doi.org/10.1214/aos/1018031101>
23. Funk S, Camacho A, Kucharski AJ, Lowe R, Eggo RM, Edmunds WJ. Assessing the performance of real-time epidemic forecasts: A case study of Ebola in the Western Area region of Sierra Leone, 2014-15. *PLOS Comput Biol.* 2019;15:e1006785. [PubMed https://doi.org/10.1371/journal.pcbi.1006785](https://doi.org/10.1371/journal.pcbi.1006785)
24. Mallela A, Neumann J, Miller EF, Chen Y, Posner RG, Lin YT, et al. Bayesian Inference of state-level COVID-19 basic reproduction numbers across the United States. *Viruses.* 2022;14:157. [PubMed https://doi.org/10.3390/v14010157](https://doi.org/10.3390/v14010157)
25. McGough SF, Johansson MA, Lipsitch M, Menzies NA. Nowcasting by Bayesian Smoothing: A flexible, generalizable model for real-time epidemic tracking. *PLOS Comput Biol.* 2020;16:e1007735. [PubMed https://doi.org/10.1371/journal.pcbi.1007735](https://doi.org/10.1371/journal.pcbi.1007735)
26. Bosse NI, Abbott S. EpiForecasts, Funk, S. scoringutils: utilities for scoring and assessing predictions. *Zenodo* 2020. <https://doi.org/10.5281/zenodo.4618017>.

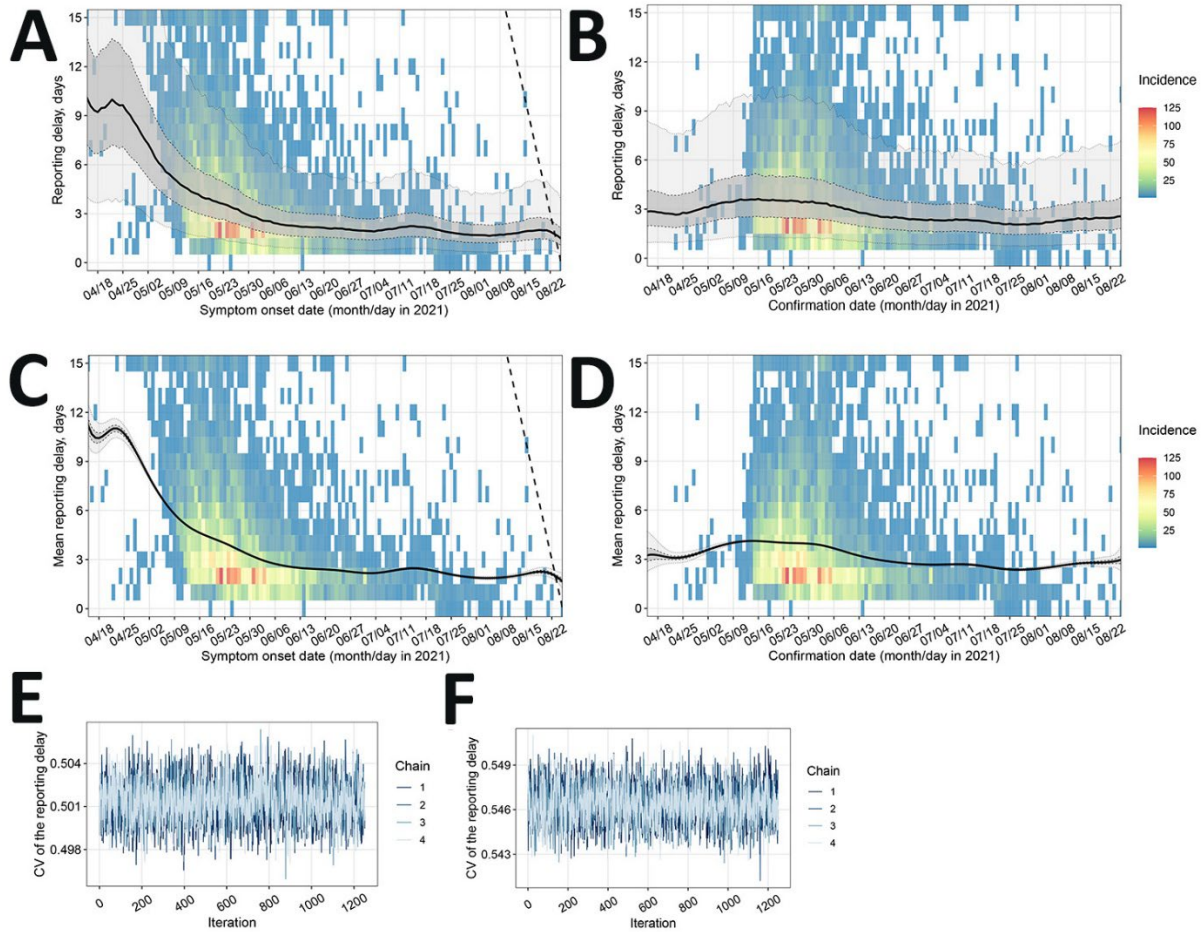
Appendix Table 1. List of infector-infectee pairs from a series of epidemiological investigations conducted by Taiwan Centers for Disease Control from the beginning of 2020 through March 2021 used for estimation of the serial interval

#	Infector National ID Case #	Infector Onset Date	Infectee National ID Case #	Infectee Onset Date	Classification*	Note
1	5	2020-01-25	8	2020-01-26	certain	
2	10	2020-01-21	9	2020-01-27	certain	
3	19	2020-01-27	21	2020-02-06	probable	family cluster
4	19	2020-01-27	22	2020-01-28	probable	family cluster
5	19	2020-01-27	23	2020-02-03	probable	family cluster
6	24	2020-01-22	25	2020-01-24	probable	family cluster
7	27	2020-02-05	28	2020-01-31	probable	family cluster, transmission chain is not clear
8	27	2020-02-05	29	2020-01-29	probable	family cluster, transmission chain is not clear
9	27	2020-02-05	30	2020-02-04	probable	family cluster, transmission chain is not clear
10	27	2020-02-05	32	2020-02-24	probable	a caretaker, transmission chain is not clear
11	34	2020-02-18	36	2020-02-18	certain	a nurse
12	34	2020-02-18	37	2020-02-25	certain	a nurse
13	34	2020-02-18	38	2020-02-25	certain	a nurse
14	34	2020-02-18	46	2020-03-03	probable	son, family cluster
15	39	2020-02-20	43	2020-03-03	certain	
16	122	2020-03-18	307	2020-03-25	certain	
17	71	2020-03-13	92	2020-03-17	certain	
18	59	2020-03-12	103	2020-03-15	certain	
19	59	2020-03-12	130	2020-03-17	certain	
20	160	2020-03-14	124	2020-03-17	probable	
21	160	2020-03-14	168	2020-03-16	probable	
22	160	2020-03-14	169	2020-03-18	probable	
23	84	2020-03-16	216	2020-03-20	certain	
24	228	2020-03-21	247	2020-03-23	certain	
25	209	2020-03-21	246	2020-03-23	certain	
26	289	2020-03-22	293	2020-03-23	certain	
27	290	2020-03-22	335	2020-03-23	certain	
28	277	2020-03-22	269	2020-03-23	certain	
29	269	2020-03-23	299	2020-03-26	certain	
30	336	2020-03-17	347	2020-03-24	certain	
31	301	2020-03-06	352	2020-03-30	certain	
32	356	2020-03-17	343	2020-03-20	certain	
33	356	2020-03-17	365	2020-03-25	certain	
34	812	2020-12-29	838	2021-01-08	certain	
35	838	2021-01-08	839	2021-01-09	certain	
36	839	2021-01-09	870	2021-01-18	certain	
37	838	2021-01-08	869	2021-01-16	certain	
38	838	2021-01-08	852	2021-01-14	certain	
39	838	2021-01-08	856	2021-01-16	certain	
40	856	2021-01-16	863	2021-01-14	probable	
41	856	2021-01-16	868	2021-01-17	probable	
42	863	2021-01-14	864	2021-01-14	probable	family cluster
43	863	2021-01-14	865	2021-01-18	probable	family cluster
44	863	2021-01-14	907	2021-01-28	probable	family cluster
45	863	2021-01-14	910	2021-01-29	probable	family cluster
46	765	2020-12-12	771	2020-12-14	certain	
47	765	2020-12-12	760	2020-12-16	certain	
48	765	2020-12-12	766	2020-12-17	certain	
49	852	2021-01-14	881	2021-01-21	certain	
50	870	2021-01-18	924	2021-02-01	probable	
51	889	2021-01-19	890	2021-01-20	probable	
52	889	2021-01-19	908	2021-01-29	probable	

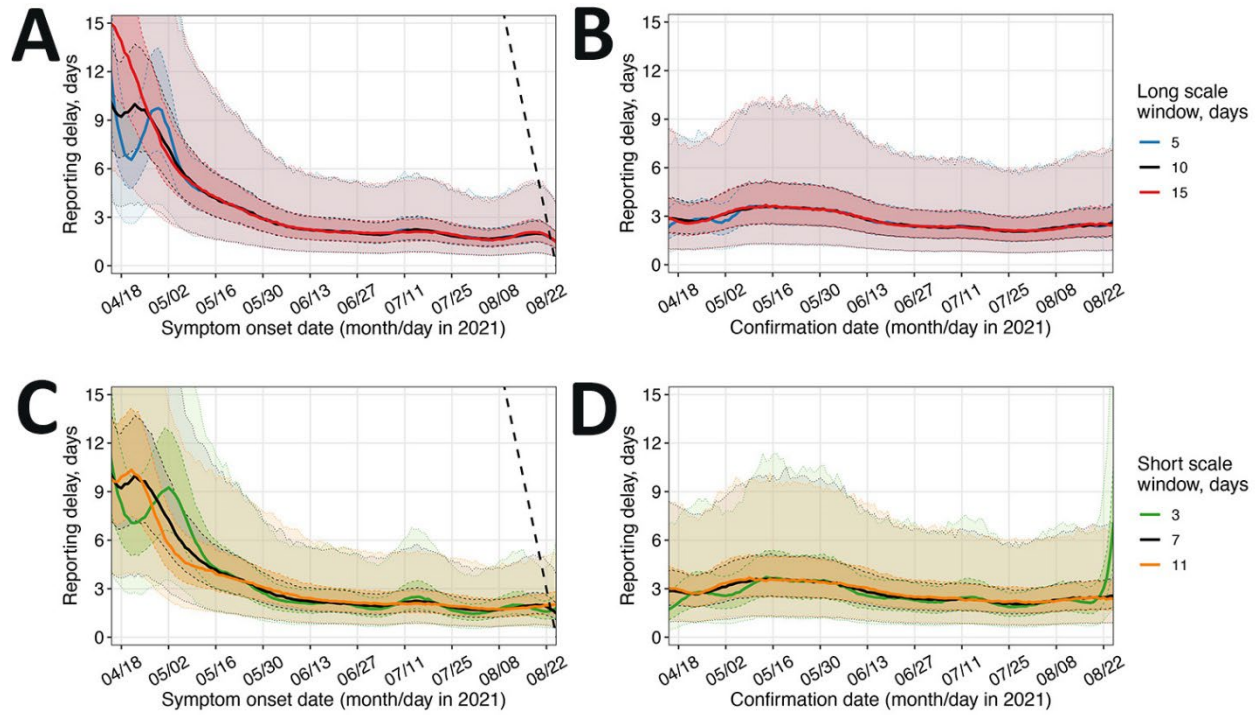
*The pair was classified as certain if the epidemiological link between an infector and an infectee could be established on a one-to-one relationship based on cumulative evidence. The pair was classified as probable if more than one possible infector could be assigned to a given infectee.

Appendix Table 2. Criteria and prevention measures of 4-level COVID-19 alert system in Taiwan

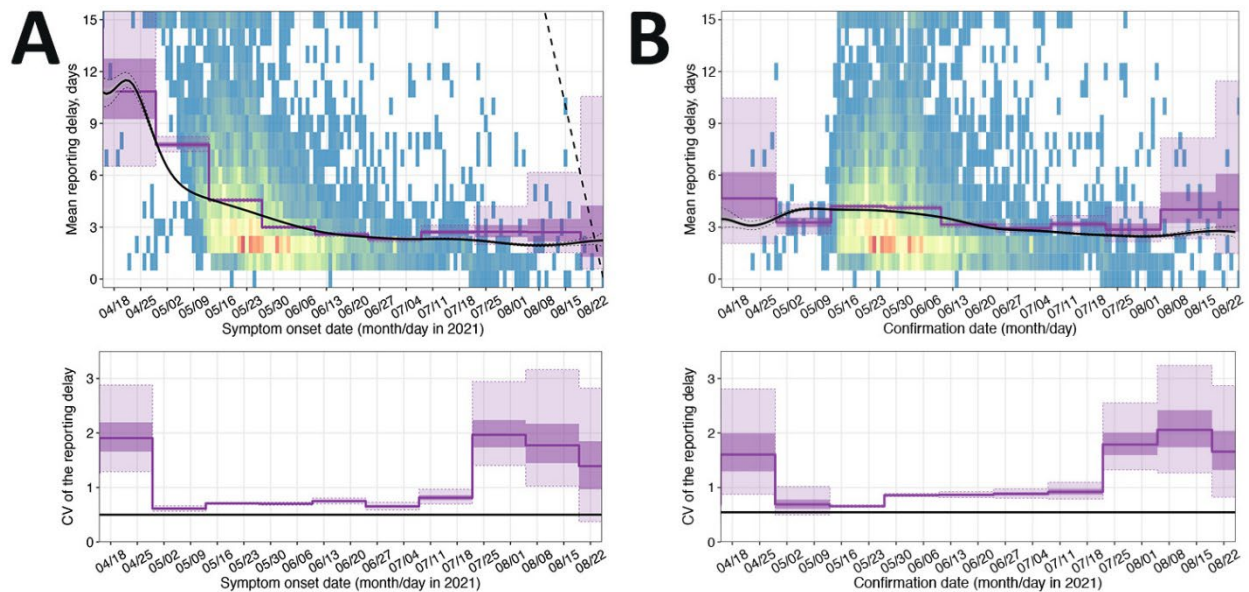
Level	Epidemic level as criteria	Prevention measures
Level 1	Sporadic community infections caused by imported cases	<ul style="list-style-type: none"> Masks must be worn all the time while using public transportation and being at crowded public venues. Recommendation to cancel or postpone non-essential gatherings that will bring people into close contact with each other. Business and public venues must implement prevention measures which include an identification-based registration system, social distancing, temperature checks, and routine disinfection.
Level 2	New local cases with unknown source of infection	<ul style="list-style-type: none"> Impose fines on those who do not obey the regulation of mask wearing. Suspend all outdoor gatherings with 500+ participants and indoor gatherings with 100+ participants. Any other gatherings must implement prevention measures including social distancing, mask wearing/partitions, an identification-based registration system, temperature checks, routine disinfection, crowd control, and crowd mobility management. Business venues must impose crowd control; those that are unable to implement necessary epidemic prevention measures must temporarily suspend operations. If necessary, entertainment related business and public venues are asked to be closed.
Level 3	3+ community clusters in one week or 10+ local cases with unknown source of infection	<ul style="list-style-type: none"> Masks must be worn at all times outdoors. Suspend all outdoor gatherings with 10+ participants and indoor gatherings with 5+ participants. Apart from essential services, which include maintenance venues, medical and public services, all other business and public venues must be closed. In the community with an ongoing active transmission where the rapid containment is required, residents must comply with COVID-19 testing protocols, do not leave the pre-defined control places and also suspend all gatherings and close schools.
Strengthened level 3		<p>Five additional measures on top of that in Level 3:</p> <ul style="list-style-type: none"> Violation of mask wearing regulation at all times will be fined. Strict inspection of entertainment related venues which have already been announced to be closed. Illegal operation, which involves operators, on-site practitioners, consumers and participants will be penalized accordingly with the law. Only take-out for all food service is possible. The enhanced crowd control is required for supermarkets and all other markets. Cancellation of all banquets for wedding ceremonies and public memorial ceremonies for funerals. Suspension of all religious gatherings. Religious venues must be closed for the public.
Level 4	Rapid increase of local cases (average number of cases more than 100 per day within 14 days) and half of them with unknown source of infection	<ul style="list-style-type: none"> Residents can leave their home only for essential activities, for example, to buy food, receive medical treatment, or to do essential work. The social distance must be maintained and wearing a mask must be done at all times outdoors. Residents must wear a mask and maintain social distance when at home. Suspension of all gatherings. Apart from essential services, which include maintenance venues, necessary medical and public services, all other venues and schools must be closed. Implementation of a lockdown policy in cities/counties or districts that reported severe outbreaks. The lockdown areas must be defined precisely and clearly, which will include the definition of restriction zones for people's entrance and exit. Residents must stay home.



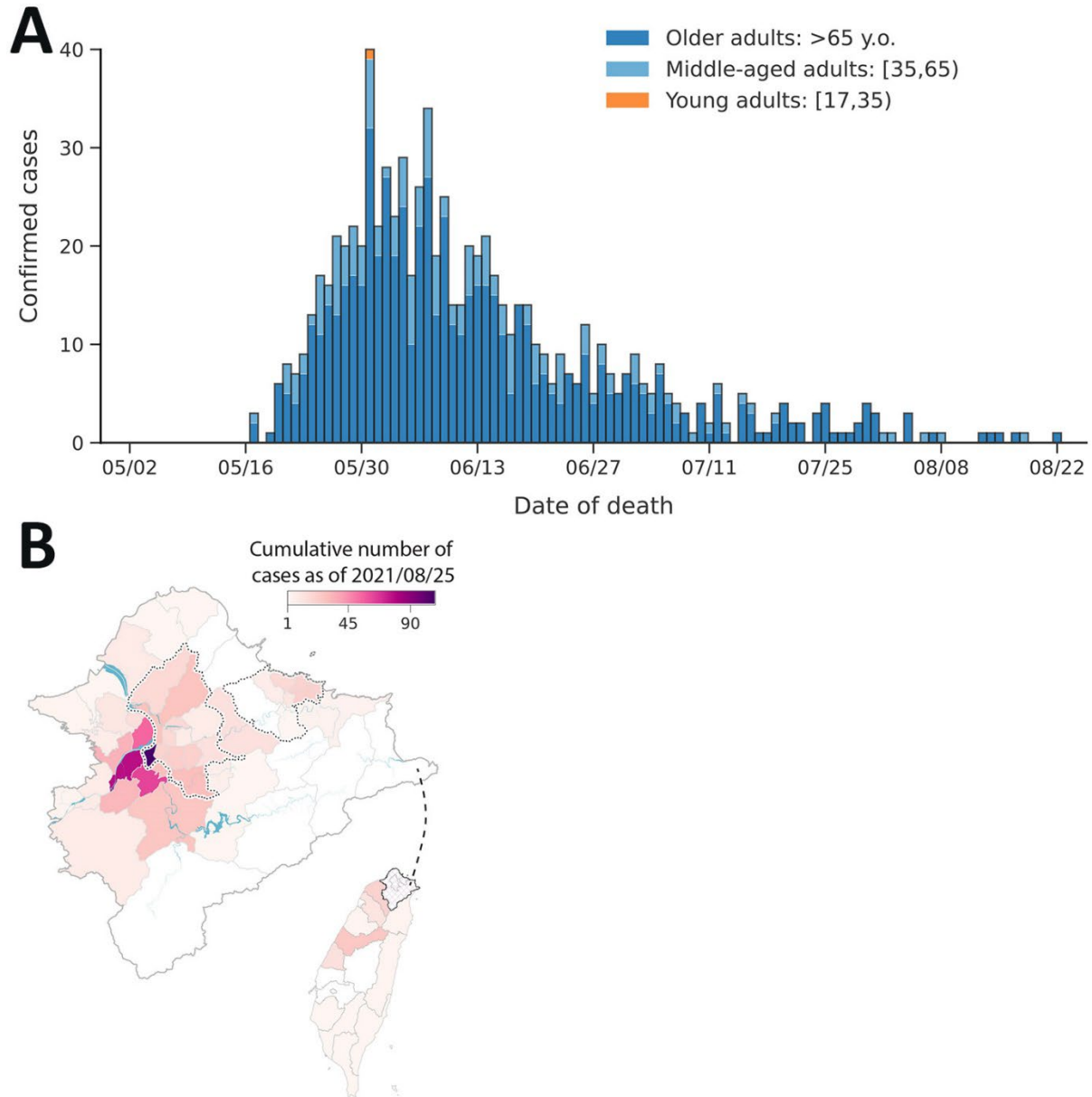
Appendix Figure 1. Data from a study of transmission dynamics and effectiveness of control measures during a COVID-19 surge, Taiwan, April-August 2021. Posterior distribution of the forward (by symptom onset date [A]) and backward (by confirmation date [B]) reporting delay, which also involved estimation of the mean (C,D) and coefficient of variation (CV) (E,F) of the reporting delay. The heatmap in the background shows the incidence of COVID-19 cases by a 2-dimensional contingency table, with horizontal axis being a date of symptom onset (A,C,E) or case confirmation (B,D,F) and with vertical axis being the observed reporting delay. The color code is indicated in the legend. CV remained constant during the epidemic, and it is shown by a trace plot of Markov chain Monte Carlo (MCMC) simulations. The dashed black line indicates the right truncation corresponding to the latest update of August 25, 2021.



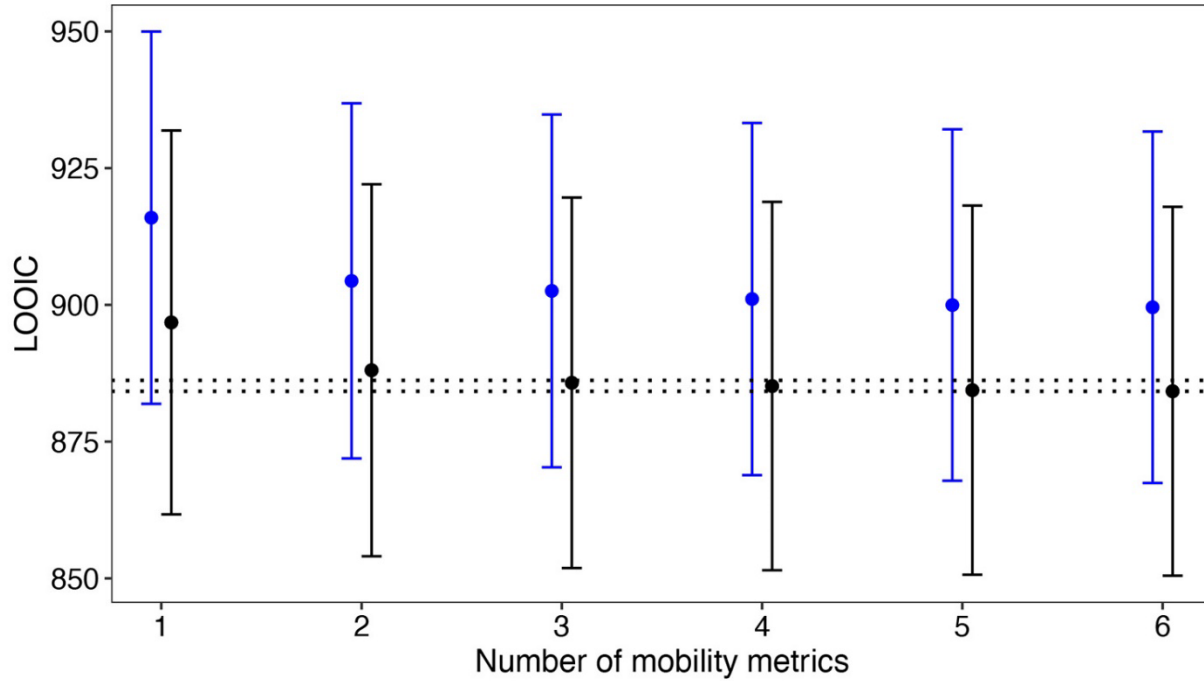
Appendix Figure 2. Data from a study of transmission dynamics and effectiveness of control measures during a COVID-19 surge, Taiwan, April-August 2021. Effect of varying the long-scale time window W (A,B) and short-scale time window ω (C,D) on the fit of the forward (A,C) and backward (B,D) reporting delay distributions. Only 1 of 2 parameters was varied, while the other was fixed at its baseline value—10 days for the long scale window and 7 days for the short-scale window—and shown in black (legend). The dashed black line in AC indicates the right truncation line respectively to the latest update date of August 25, 2021.



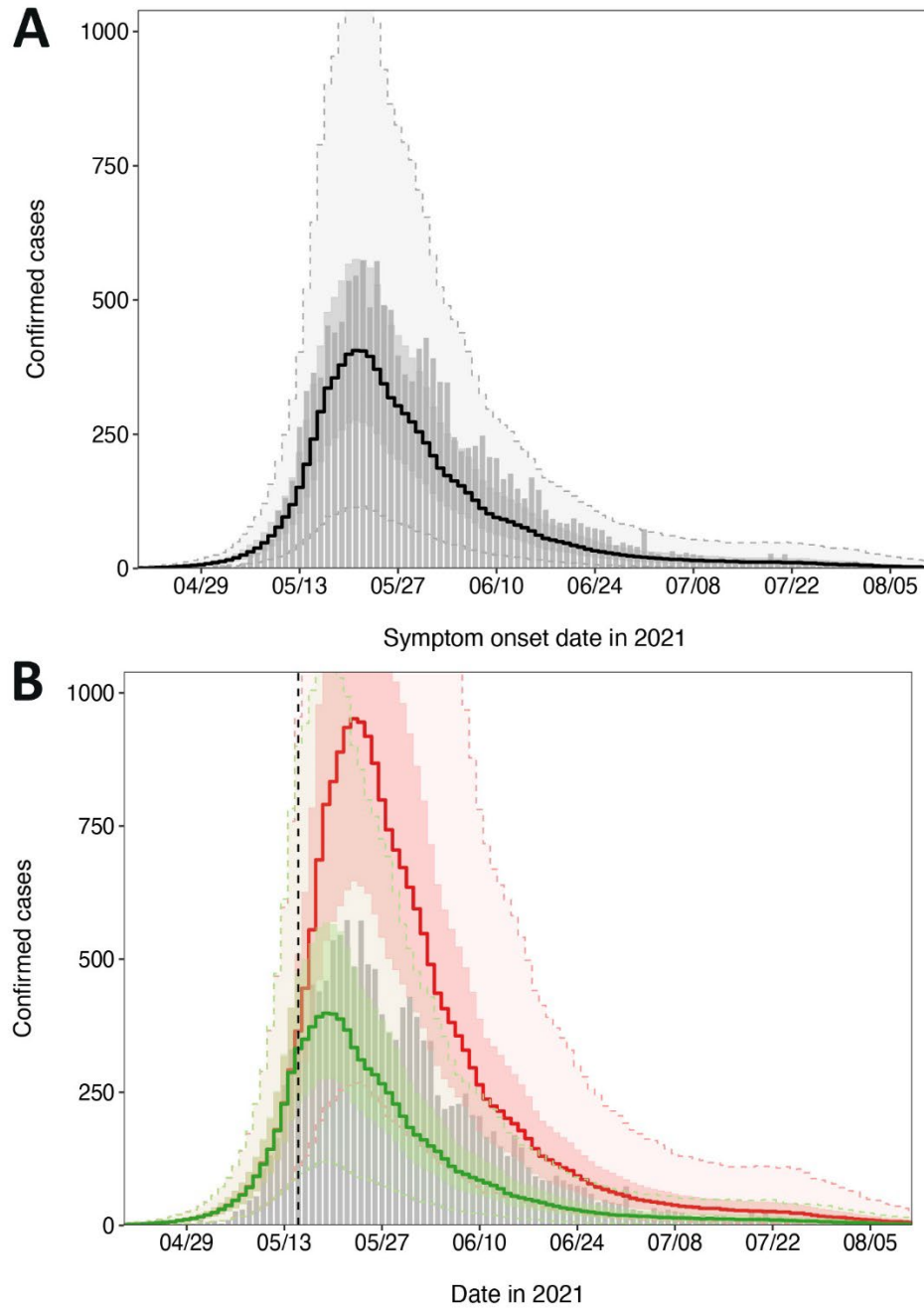
Appendix Figure 3. Data from a study of transmission dynamics and effectiveness of control measures during a COVID-19 surge, Taiwan, April-August 2021. Comparing the mean and coefficient of variation (CV) for the forward (A) and backward (B) reporting delay distributions when the mean is given by a cubic B-spline and CV is constant (black), or both the mean and CV are given by a piecewise-constant function over time with time-window of 14 days (purple). The heatmap shows the incidence of COVID-19 cases represented as a 2-dimensional contingency table, where the horizontal axis is the date of symptom onset or case confirmation and the vertical axis is the reporting delay. The color code is shown in legend. CV was constant during the epidemic and shown as a trace plot of MCMC simulations. The dashed black line in A (top panel) indicates the right truncation line respectively to the latest update date of August 25, 2021.



Appendix Figure 4. Data from a study of transmission dynamics and effectiveness of control measures during a COVID-19 surge, Taiwan, April-August 2021. Age and spatial distribution of deaths confirmed in Taiwan from May through August 25, 2021. A) Epidemiologic curve of confirmed COVID-19 death cases stratified by age group and shown by date of death. B) Geographic distribution of deaths. The colormap indicates the cumulative number of deaths reported as of August 25, 2021, at the district level for Taipei City, New Taipei City, and Keelung City, and at county level for the rest of Taiwan.



Appendix Figure 5. Data from a study of transmission dynamics and effectiveness of control measures during a COVID-19 surge, Taiwan, April-August 2021. Comparing the "leave-one-out" information criterion (LOOIC) values for 2 models when the baseline reproduction number is constant during the epidemic (blue) or modeled by a sigmoidal monotonically decreasing function of time implying one changepoint (black). The double-dashed line indicates LOOIC values exceeding the minimal LOOIC value by no more than 2 units.



Appendix Figure 6. Data from a study of transmission dynamics and effectiveness of control measures during a COVID-19 surge, Taiwan, April-August 2021. Reconstructed epidemiologic curve for baseline scenario (A) and 2 counterfactual scenarios (B) when the Level 3 measures were implemented either 3 days earlier (green) or 3 days later (red). The thick line indicated the median of the posterior distribution. The light shaded area indicates the 95% credible interval, whereas the dark shaded area shows the interquartile range of the posterior distribution. Gray bars show the observed and backprojected counts of confirmed cases by date of symptom onset. The vertical dashed line in B indicates the date of May 15, 2021, when the baseline Level 3 measures were implemented at national level.