Survival-Convolution Models for Predicting COVID-19 Cases and Assessing Effects of Mitigation Strategies

Qinxia Wang¹, Shanghong Xie¹, Yuanjia Wang^{1,*}, Donglin Zeng^{2,*}

- Department of Biostatistics, Mailman School of Public Health, Columbia University, New York,
 NY, USA
- ² Department of Biostatistics, Gillings School of Public Health, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA

Correspondence*: Yuanjia Wang and Donglin Zeng yw2016@cumc.columbia.edu, dzeng@email.unc.edu

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11 Summary

Countries around the globe have implemented unprecedented measures to mitigate the coronavirus disease 2019 (COVID-19) pandemic. We aim to predict COVID-19 disease course and compare effectiveness of mitigation measures across countries to inform policy decision making using a robust and parsimonious survival-convolution model. We account for transmission during a pre-symptomatic incubation period and use a time-varying effective reproduction number (R_t) to reflect the temporal trend of transmission and change in response to a public health intervention. We estimate the intervention effect on reducing the infection rate using a natural experiment design and quantify uncertainty by permutation. In China and South Korea, we predicted the entire disease epidemic using only early phase data (two to three weeks after the outbreak). A fast rate of decline in R_t was observed and adopting mitigation strategies early in the epidemic was effective in reducing the infection rate in these two countries. The nationwide lockdown in Italy did not accelerate the speed at which the infection rate decreases. In the United States, R_t significantly decreased during a 2-week period after the declaration of national emergency, but declines at a much slower rate afterwards. If the trend continues after May 1, COVID-19 may be controlled by July 26 (CI: July 9 to August 27). However, a loss of temporal effect (e.g., due to relaxing mitigation measures after May 1) could lead to a long delay in controlling the epidemic (November 19 with less than 100 daily cases) and a total of more than 2 million cases.

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Keywords: COVID-19, survival-convolution model, time-varying effective reproduction number, mitigation measures, prediction

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36 1 Introduction

COVID-19 pandemic is currently a daunting global health challenge. The novel coronavirus was observed to have a long incubation period and highly infectious during this period ¹⁻⁴. The cumulative case number surpasses 4.1 million by May 10, with more than 1.3 million in the United States (US). It is imperative to study the course of the disease outbreak in countries that have controlled the outbreak (e.g., China and South Korea) and compare mitigation strategies to inform decision making in regions that are in the midst of (e.g., the US) or at the beginning of outbreak (e.g., South America).

Various infectious disease models are proposed to estimate the transmission of COVID19⁵⁻⁷ and investigate the impact of public health interventions on mitigating the spread 8-12.
Several studies modeled the transmission by stochastic dynamical systems 5-7,10, such as susceptible-exposed-infectious-recovered (SEIR) models 5, extended Kalman filter 13-15, and individual-based simulation models 8,9. Some models did not explicitly take into account of behavioral change (e.g., social distancing) and government mitigation strategies that can have major influences on the disease course, while other work modified the infection rate as public-health-intervention-dependent 10,12 or time-varying 7. A recent study 11 considered the disease incubation period and used a convolution model based on SEIR. A state-space susceptible-infectious-recovered (SIR) model with time-varying transmission rate 16 was developed to account for interventions and quarantines.

SEIR models can incorporate mechanistic characteristics and scientific knowledge of virus transmission to provide useful estimates of its temporal dynamics, especially when individual-level epidemiological data are available through surveillance and contact tracing. However, these sophisticated models may involve a large number of parameters and assumptions about individual transmission dynamics. Thus, they may be susceptible to perturbation of parameters and prior assumptions, yielding wide prediction intervals especially when granular individual-level data are not available. In contrast to infectious disease models, alternative statistical models are proposed to predict summary statistics such as deaths
and hospital demand under a nonlinear mixed effects model framework ¹⁷, survival analysis
has been introduced to model the occurrence of clinical events in infectious disease studies ¹⁸,
and a nonparametric space-time transmission model was developed to incorporate spatial
and temporal information for predictions at the county level ¹⁹. Nonparametric modelling or
survival models are data-driven, so parameters may not be scientifically related to disease
epidemic.

In this work, we propose a parsimonious and robust population-level survival-convolution 69 model that is based on main characteristics of COVID-19 epidemic and observed number of confirmed cases to predict disease course and assess public health intervention effect. Our 71 method models only key statistics (e.g., daily new cases) that reflect the disease epidemic 72 over time with at most six parameters, so it may be more robust than models that rely on individual transmission processes or a large number of parameters and assumptions. We construct our model based on prior scientific knowledge about COVID-19, instead of posthoc observations of the trend of disease spread. Specifically, two important facts we consider include (1) SARS-CoV-2 virus has an incubation period up to 14-21 days¹ and a patient can 77 be highly infectious in the pre-symptomatic phase; (2) infection rate varies over time and can 78 change significantly when government guidelines and mitigation strategies are implemented; 79 (3) intervention effect may be time-varying. 80

We aim to achieve the following goals. The first goal is to fit observed data to predict daily new confirmed cases and latent pre-symptomatic cases, the peak date, and the final to-tal number of cases. The second goal is to assess the effect of nationwide major interventions across countries (e.g., mitigation measures) under the framework of natural experiments (e.g., longitudinal pre-post quasi-experimental design ²⁰). Quasi-experiment approaches are often used to estimate intervention effect of a public health intervention (e.g., HPV vaccine ²¹) or a health policy where randomized controlled trials (RCTs) are not feasible. Our

third goal is to project the future trend of COVID-19 for the countries (e.g., US) amid the epidemic under different assumptions of future infection rates, including the continuation of the current trend and relaxing mitigation measures.

91 2 Methods

2.1 Data source

We used data from a publicly available database that consolidates multiple sources of official reports (World Meters[https://www.worldometers.info/coronavirus/]). We analyzed 94 two countries with a large number of confirmed cases in Asia (China, South Korea) and two 95 outside (Italy, US). Since both China and South Korea are already at the end of epidemic, 96 we used their data to test empirical prediction performance of our method. We included data 97 in the early phase of epidemic as training set to estimate model parameters and leave the 98 rest of the data as testing set for evaluation. For China, we used data up to two weeks post the lockdown of Wuhan city (January 23) as training (data from January 20 to February 100 4), and used the remaining observed data for evaluation (February 5 to May 10). Similarly, 101 for South Korea we used data from February 15 to March 4 as training and leave the rest 102 for evaluation (March 5 to May 10). Italy is the first European country confronted by a 103 large outbreak and currently has passed its peak. We estimate the effect of the nation-wide 104 lockdown in Italy (dated March 11) using 10 weeks data (February 20 to April 29). For the US, since after May 1 some mitigation measures were lifted in various states, we also 106 included about 10 weeks data (February 21 to May 1) to assess the effect of its mitigation strategies.

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2.2 Survival-Convolution Model

Let t denote the calendar time (in days) and let $N_0(t)$ be the number of individuals who 110 are newly infected by COVID-19 at time t. Let t_j denote the time when individual j is 111 infected ($t_j = \infty$ if never infected), and let T_j be the duration of this individual remain-112 ing infectious to any other individual and in the transmission chain. Let t_0 be the un-113 known calendar time when the first patient (patient zero) is infected. Therefore, at time 114 t, the total number of individuals who can infect others is $\sum_{j} I(t_j \leq t, T_j \geq t - t_j) =$ 115 $\sum_{m=0}^{C} \sum_{\{j: j \text{ is infected at } (t-m)\}} I(T_j \geq m)$, where $C = \min(t-t_0, C_1)$ with C_1 as the max-116 imum incubation period (i.e., 21 days for SARS-CoV-2) and I(E) denotes an indicator 117 function with I(E) = 1 if event E occurs and I(E) = 0 otherwise. Since the total number of 118 individuals who are newly infected at time (t-m) is $N_0(t-m)$, the number of individuals 119 who remain infectious at time t is $M(t) = \sum_{m=0}^{C} N_0(t-m)S(m)$, where S(m) denotes the proportion of individuals remaining infectious after m days of being infected, or equivalently, 121 the survival probability at day m for T_i . On the other hand, right after time t, some individuals will no longer be in the transmission chain (e.g., due to testing positive and quarantine 123 or out of infectious period) with duration $T_j = (t - t_j)$. The total number of these individuals 124 is $\sum_{j} I(t_j \le t, T_j = t - t_j) = \sum_{m=0}^{C} \sum_{j: j \text{ is infected at } (t-m)} I(T_j = m)$, or equivalently

$$Y(t) = \sum_{m=0}^{C} N_0(t-m)[S(m) - S(m+1)].$$
(1)

Therefore, (M(t) - Y(t)) is the number of individuals who can still infect others after time t. Assuming the infection rate at t to be a(t), then at time (t+1) the number of newly infected patients is a(t)[M(t) - Y(t)], which yields

$$N_0(t+1) = a(t) \sum_{m=0}^{C} N_0(t-m)S(m+1).$$
(2)

Note that a(t) is time-varying because the infection rate depends on how many close contacts an infected individual may have at time t, which is affected by public heath in-

terventions (e.g., stay-at-home order, lockdown), and saturation level of the infection in the whole population. Define $R_t = \sum_{m=0}^{C} a(t+m)S(m)$, the expected number of secondary cases infected by a primary infected individual in a population at time t while accounting for the entire incubation period of the primary case. Thus, R_t is the instantaneous time-varying effective reproduction number²² that measures temporal changes in the disease spread.

Models (1) and (2) provide a robust dynamic model to characterize COVID-19 epidemic. Equation (2) gives a convolution update for the new cases using the past numbers, while equation (1) gives the number of cases out of transmission chain at time t, and M(t) computes the number of latent pre-symptomatic cases by the end of time t. This model considers three important quantities to characterize COVID-19 transmission: the initial date, t_0 , of the first (likely undetected) case in the epidemic, the survival function of time to out of transmission, S(m), and the infection rate over calendar time, a(t).

We model infection rate a(t) as a non-negative, piece-wise linear function with knots 145 placed at meaningful event times. The simplest model consists of a constant and a single 146 linear function with three parameters (infection date of patient zero, intercept and slope 147 of a(t)). When a massive public health intervention (e.g., nation-wide lockdown) is imple-148 mented at some particular date, we introduce an additional linear function afterwards with 149 a new slope parameter. Thus, the difference in slope parameters of a(t) before and after an 150 intervention reflects its effect on reducing the rate of change in disease transmission (i.e., 151 "flattening the curve"). Since the intervention effect may diminish over time, we introduce 152 another slope parameter two weeks after intervention to capture the longer-term effect. We 153 use existing knowledge of SARS-CoV-2 virus incubation period¹ to approximate S(m) and 154 perform sensitivity analysis assuming different parameters. For estimation, we minimize a 155 loss function measuring differences between model predicted and observed daily number of 156 cases. For statistical inference, we use permutation based on standardized residuals. All 157 mathematical details are in Supplementary Material.

9 2.3 Utility of Our Model

First, with parameters estimated from data and assuming that the future infection rate remains the same trend, we can use models (1) and (2) to predict future daily new cases, the peak time, expected number of cases at the peak, when R_t will be reduced to below 1.0, and when the epidemic will be controlled (the number of daily new cases below a threshold or decreases to zero). Furthermore, our model provides the number of latent cases cumulative over the incubation period at each future date, which can be useful to anticipate challenges and allocate resources effectively.

Second, we can estimate the effects of mitigation strategies, leveraging the nature of 167 quasi-experiments where subjects receive different interventions before and after the initia-168 tion of the intervention. The longitudinal pre-post intervention design allows valid inferences 169 assuming that pre-intervention disease trend would have continued had the intervention not 170 taken place and local randomization holds (whether a subject falls immediately before or 171 after the initiation date of an intervention may be considered as random, and thus the 172 "intervention assignment" may be considered to be random). Applying this design, the in-173 tervention effects will be estimated as the difference in the rate of change of the infection 174 rate function before and after an intervention takes place. 175

Third, we study the impact of an intervention (e.g., lifting mitigation measures) that
changes the epidemic at a future date. Using permutations, we obtain the joint distribution
of the parameter estimators and construct confidence intervals (CI) for the projected case
numbers and interventions effects.

3 Results

For China, the infection rate a(t) is a single linear function (estimates in Table 1). The first community infection was estimated to occur on January 3, 17 days before the first reported

case (Table 1). Figure 1A shows that the model captures the peak date of new cases, the 183 epidemic end date, and the prediction interval contains the majority of observed number of 184 cases except one outlier (due to a change of diagnostic criteria). The reproduction number 185 R_t decreases quickly from 3.34 to below 1.0 in 14 days (Figure 2A). We only used data up to 186 February 4 to estimate our model. The observed total number of cases by May 10 is 82,901, 187 which is inside the 95% CI of the estimated total number of cases (58,415; 95% CI: (42,516, 188 133,083)). There are two outlier days (February 12, 13) with a total of 19,198 cases reported 189 in the testing set. Excluding two outliers, the observed number of cases 62,356. 190

For South Korea, Figure 1B shows that the model captures the general trend of the epidemic except at the tail area (after March 15) where some small and enduring outbreak is observed. The effective reproduction number decreases dramatically from 5.37 at the beginning of the outbreak to below 1.0 in 14 days (Figure 2B). The predicted number of new cases at the peak is 665 and the total number of predicted cases at the peak time is close to the observed total (4,300 vs 4,335). The predicted total number by March 15 is 7,816 and the observed total is 8,162.

For Italy, we model a(t) as a four-piece linear function to account for the change in 198 mitigation strategies with a knot placed at the lockdown (March 11), and two additional 199 knots at 2-week intervals (March 25, April 8) to account for time-varying intervention effect. 200 Difference on the rate of change before and after the first knot measures the immediate 201 effect of lockdown on reducing the infection rate. Change before and after the second and 202 third knot measures whether the lockdown effect can be maintained in longer term. The 203 rate of change in R_t is not significantly different before and two weeks after the lockdown 204 (Figure 2C). The reproduction number decreased from 3.73 at the beginning to 1.02 two 205 weeks post-lockdown. However, starting from the third week post-lockdown (March 26), 206 R_t stops decreasing and remains close to 1.0 until April 16. The slope of a(t) (infection 207 rate) increases by 116% to a slightly positive value after March 26 (Table 1, comparing a₂ and a_3 for Italy). This is consistent with a relatively flat trend of observed daily new cases during this period (Figure 1C). The estimated total by May 10 is 216,300 (95%CI: (214,863, 228,406)) and close to the observed total (219,070). Recent daily cases in the testing set also closely follow our predicted trend (Figure 1C).

In the US, we fit a three-piece model for a(t) with a knot on March 13 (the declaration of 213 national emergency) and an additional knot two weeks after (March 27). The predicted peak 214 date is May 3 (Figure 3A) with a total number of 1,176,915 cases by May 3, which is close 215 to the observed total (1,188,122). R_t increases during the early phase but decreases sharply 216 after the declaration of national emergency (Figure 3B) up to two weeks after. During 217 the next period (March 28 to April 10), R_t decreases at a much slower rate. If this trend 218 continues, the end of epidemic date is predicted to be July 26 (scenario 1, Figure 3A), and the 219 predicted total over the entire epidemic will be 1,626,950 (CI: (1,501,036, 1,918,602), Table 220 1). However, since states in the US are gradually lifting mitigation measures after May 1. 221 the trend of infection rate may change. We predicted epidemic control date assuming a(t)decreases slower after May 1 by 50% (scenario 2), 75% (scenario 3), and 100% (scenario 4) 223 in Table 1. Under scenario 4 where the temporal effect of mitigation measures is completely 224 lost (i.e., a(t) is a constant over time), the projected total number of cases will be more 225 than 2 million, and the epidemic cannot be controlled until November 19 (with less than 100 226 daily cases, Table 1). Assuming a case fatality rate of 6% as observed by May 10, the total 227 number of deaths would be around 120,000. 228

We show the estimated number of latent cases present on each day (i.e., including pre-symptomatic patients infected k days before but have not shown symptoms) in Supplementary Material (Figure S1). For all countries, there were a large number of latent cases around the peak time. We performed a sensitivity analysis using different distributions of S(m) assuming a delay in reporting confirmed cases. The results show that predicted daily new cases were similar under different parameters of S(m) for both US and Italy (Sup-

plementary Material Figures S2 and S3), demonstrating robustness of our method to the assumptions of S(m).

²³⁷ 4 Discussion

In this study, we propose a parsimonious and robust survival convolution model to predict 238 daily new cases of the COVID-19 outbreak and use a natural quasi-experimental design to 239 estimate the effects of mitigation measures. Our model accounts for major characteristics of 240 COVID-19 (long incubation period and highly contagious during incubation) with a small 241 number of parameters (up to six) and assumptions, directly targets prediction accuracy, and 242 provides measures of uncertainty and inference based on permuting the residuals. We allow 243 the infection rate to depend on time and modify the basic reproduction number R_0 as a 244 time-dependent measure R_t to estimate change in disease transmission over time. Thus, R_t 245 corrects for the naturally impact of time on the disease spread. Our estimated reproduction 246 number at the beginning of the epidemic ranges from 2.81 to 5.37, which is consistent with R_0 247 reported in other studies²³ (range from 1.40 to 6.49, with a median of 2.79). For predicting 248 daily new cases, our analyses suggest that the model estimated from early periods of outbreak 249 can be used to predict the entire epidemic if the disease infection rate dynamic does not change dramatically over the disease course (e.g., about two weeks data is sufficient for China and fits the general trend of South Korea). 252

Comparing the effective reproduction numbers across countries, R_t decreased much more rapidly in South Korea and China than Italy (Figure 2). In South Korea, the effective reproduction number had been reduced from 5.37 to under 1.0 in a mere 13 days and the total number of cases is low. The starting reproduction number in South Korea was high possibly due to many cases linked to patient 31 and outbreaks at church gatherings. Similarly for China, the reproduction number reduced to below 1.0 in 14 days. Italy's R_t decreased until almost reaching 1.0 on March 25, but remained around 1.0 for 3 weeks. The US followed a fast decreasing trend during a two-week period after declaring national emergency $(a_2 = -1.031)$, which is faster than the first two weeks in China $(a_1 = -0.693)$, but its R_t decreased at a much slower rate $(a_3 = -0.042)$ afterwards and was below 1.0 on May 5.

Comparing mitigation strategies across countries, the fast decline in R_t in China sug-263 gests that the initial mitigation measures put forth on January 23 (lockdown of Wuhan city, 264 traffic suspension, home quarantine) were successful in controlling the transmission speed of 265 COVID-19. Additional mitigation measures were in place after February 2 (centralized quar-266 antine and treatment), but did not seem to have significantly changed the disease course. In 267 fact, our model assuming the same infection rate trajectory after February 2 fits all observed 268 data up to May 10. A recent analysis of Wuhan's data^{24,25} arrived at a similar conclu-269 sion, and their estimated R_t closely matches with our estimates. However, their analyses 270 were based on self-reported symptom onset and other additional surveillance data, where we 271 used only widely available official reports of confirmed cases. Another mechanistic ²⁶ study confirmed the effectiveness of early containment strategies in Wuhan. 273

South Korea did not impose a nation-wide lockdown or closure of businesses, but at the very early stage (when many cases linked to patient 31 were reported on February 20) conducted extensive broad-based testing and detection (drive through tests started on February 26), rigorous contact tracing, isolation of cases, and mobile phone tracking. Our results suggest that South Korea's early mitigation measures were also effective.

Italy's initial mitigation strategies in the most affected areas reduced R_t from 3.73 to 1.92 in 20 days. To estimate the intervention effect of the nation-wide lockdown as in a natural experiment, we require local randomization and the continuity assumption. The former requires that characteristics of subjects who are infected right before or after the lockdown are similar. Since in a very short time period, whether a person is infected at time t or t+1 is likely to be random, the local randomization assumption is likely to be valid. Continuity assumption refers to that the infection rate before the lockdown would

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continue to capture the trend afterwards had the intervention not been implemented. Under this assumption, the lockdown in Italy is not effective to further reduce the transmission 287 speed (slopes of a(t) are similar before and after lockdown on March 11). There were 10,149 288 cases reported in Italy as of March 10, suggesting that the lockdown was placed after the 289 wide community spread had already occurred. Nevertheless, it is possible that without 290 the lockdown the infection rate would have had increased, i.e., the lockdown enhanced and 291 maintained the effect of quarantine for two weeks. In fact, after two weeks of lockdown, we 292 observe a loss of temporal effect so that R_t has remained around 1.0 for about 2-3 weeks 293 before it starts to decrease again. 294

For the US, R_t ranges between 2.81 and 4.50 before the declaration of national emergency 295 on March 13, but R_t declines rapidly over a two-week period after March 13. Although the 296 disease trend and mitigation strategies vary across states in the US, since the declaration 297 of national emergency, many states have implemented social distancing and ban of large gathering. The large difference before and after March 13 is likely due to states with large 299 numbers of cases that implemented state-wide mitigation measures (e.g., New York, New 300 Jersey). Our model predicted a continued decrease in R_t from March 27 to May 1 but at 301 a much slower rate (95.9% slower; Table 1, comparing a_2 and a_3 for the US). If the trend 302 continues after May 1, the first wave of epidemic will be controlled by July 26 (CI: July 9, 303 August 27). However, after May 1 many states enter a re-opening phase. If the guidelines 304 on quarantine measures are relaxed so that the effect of quarantine cannot be maintained, 305 the control date can be delayed by 32 days (50% slower decrease in the infection rate) or 306 70 days (75\% slower). If the temporal effect of quarantine measures is completely lost, the 307 predicted total number of cases is more than 2 million, with a long delay in controlling the 308 epidemic (less than 100 cases by November 19, and no new case by May, 2021). 309

Other studies reported transmission between asymptomatic individuals⁶, which is not accounted for here. However, asymptomatic individuals can only be identified and confirmed

by serological tests which are not widely available. When there is a delay in reporting some symptomatic patients, the daily reported cases are a mixture of new symptomatic cases and 313 patients presenting after having had symptoms for a few days. In this case, the average 314 number of days to testing positive may be higher than the virus incubation period of 5.2 315 days. However, as shown in our sensitivity analysis, the prediction of daily reported cases 316 was not affected by using a larger mean value for S(m), demonstrating robustness of the 317 model. Our model does not consider subject-specific covariates and focuses on predicting 318 population-level quantities. Neither have we considered borrowing information from multiple 319 countries or state-level analysis for the US, which are worthy of study in a mixed effects model 320 framework. We do not consider prediction of daily new deaths or hospitalizations. These 321 data can be included to enhance the prediction of new cases by linking the distribution of 322 time to COVID symptom onsets, hospitalization, or death. Lastly, we can consider a broader 323 class of models for infection rate a(t) to allow discontinuity in both intercepts and slopes 324 before and after an intervention under a regression discontinuity design ^{21,27}. 325

Despite these limitations, our study offers several implications. Implementing mitigation 326 measures earlier in the disease epidemic reduces the disease transmission rate at a faster speed 327 (South Korea, China). Thus for regions at the early stage of disease epidemic, mitigation 328 measures should be introduced early. Nation-wide lockdown may not further reduce the 329 speed of R_t reduction compared to regional quarantine measures as seen in Italy. In countries 330 where disease transmissions have slowed down, lifting of quarantine measures may lead to a 331 persistent infection rate delaying control of epidemic and thus should be implemented with 332 caution and close monitoring. 333

$_{\scriptscriptstyle{334}}$ Data sharing

All data and optimization codes are publicly available at [https://github.com/COVID19BIOSTAT].

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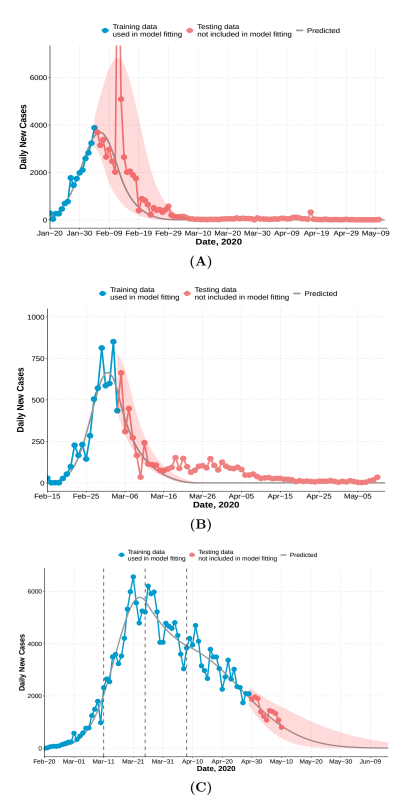


Figure 1: Observed and predicted daily new cases and 95% prediction interval (shaded). **(A)** China. Training data: January 20 to February 4; testing data: February 5 to May 10. 14,108 cases were reported on February 12 and not shown on figure. The recent cases since April are imported cases. **(B)** South Korea. Training data: February 15 to March 4; testing data: March 5 to May 10. **(C)** Italy. First dashed line indicates the nation-wide lockdown (March 11). Second and third dashed line indicates two or four weeks after. Training data: February 20 to April 29 (7 weeks after the lockdown); testing data: April 30 to May 10.

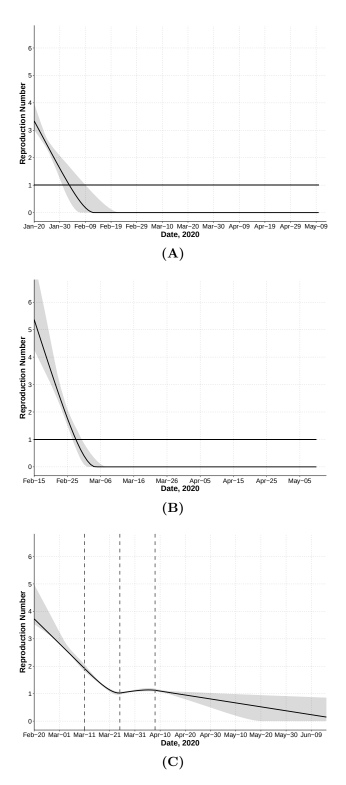


Figure 2: Effective reproduction number R_t for each country computed as the average number of secondary infections generated by a primary case at time t accounting for the incubation period of the primary case. Dashed lines indicate knots for infection rate a(t). (A) China. (B) South Korea. (C) Italy.

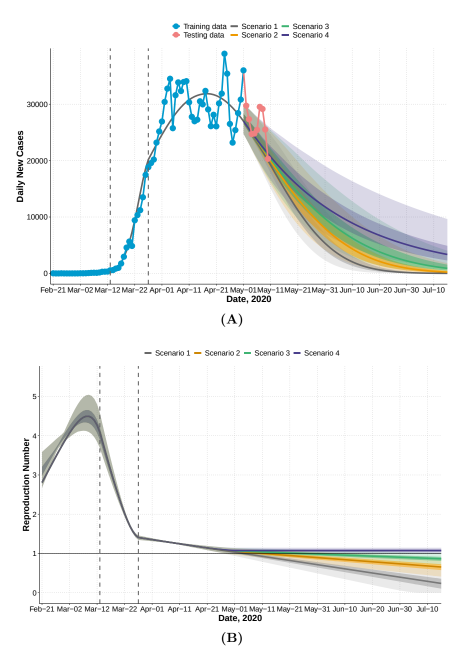


Figure 3: United States: observed and predicted daily new cases, 95% prediction intervals (lighter shaded) and 50% prediction intervals (darker shaded) under four scenarios that assume relaxation of mitigation measures occurs after May 1. Scenario 1: infection rate a(t) follows the same trend after May 1 as observed between March 27 and May 1. Scenario 2: rate of decrease of a(t) slows by 50% after May 1. Scenario 3: rate of decrease of a(t) slows by 75% after May 1. Scenario 4: rate of decrease of a(t) slows by 100% after May 1 (complete loss of temporal decreasing effect). First dashed line indicates the declaration of national emergency (March 13). Second dashed line indicates two weeks after (March 27). Training data: February 21 to May 1 (7 weeks after declaring national emergency); testing data: May 2 to May 10. (A) Observed and predicted daily new cases. (B) Effective reproduction number R_t .

Table 1: Model Estimated Parameters in Each Country

Country	Parameter or Prediction*	Estimate	95% CI
China	$t_0(d)$	Jan 3 (17)	(12, 21)**
Training data: Jan 20 to Feb 4	a_0	0.793	(0.68, 1.02)
Testing data: Feb 5 to May 10	a_1	-0.693	(-1.13, -0.42)
	Duration	44	(39, 55)
	End date	Mar 4	(Feb 28, Mar 15)
	Total	$58,\!415$	(42,516, 133,083)
South Korea	$t_0(d)$	Feb 11 (4)	(1, 7)
Training data: Feb 15 to Mar 4	a_0	1.363	(1.03, 1.98)
Testing data: Mar 5 to May 10	a_1	-1.496	(-2.39, -0.96)
	Duration	39	(37, 43)
	End date	Mar 25	(Mar 23, Mar 29)
	Total	7,977	(7,307, 10,562)
Italy	$t_0(d)$	Feb 10 (10)	(4, 11)
Training data: Feb 20 to Apr 29	a_0	0.789	(0.73, 1.10)
Testing data: Apr 30 to May 10	a_1	-0.358	(-0.68, -0.26)
	a_2	-0.372	(-0.46, -0.31)
	a_3	0.061	(0.02, 0.12)
	a_4	-0.057	(-0.12, -0.01)
	Duration	123	(103, 179)
	End date	Jun 22	(Jun 2, Aug 17)
	Total	223,410	(216,848, 257,710)
United States	$t_0(d)$	Feb 15 (6)	(1, 4)
Training data: Feb 21 to May 1	a_0	0.410	(0.34, 0.62)
Testing data: May 2 to May 10	a_1	0.526	(0.23, 0.72)
	a_2	-1.031	(-1.24, -0.86)
	a_3	-0.042	(-0.06, -0.03)
Scenario 1: Continue current [†]	Duration	156	(139, 188)
	End date	Jul 26	(Jul 9, Aug 27)
-	$_{-}$ Total	1,626,950	(1,501,036, 1,918,602)
Scenario 2: 50% slower	Duration	188	(163, 233)
after May 1	End date	Aug 27	(Aug 2, Oct 11)
G	Total	1,731,992	(1,563,122, 2,113,294)
Scenario 3: 75% slower	Duration	226	(190, 289)
after May 1	End date	Oct 4	(Aug 29, Dec 5)
C	Total	1,832,291	(1,616,574, 2,324,552)
Scenario 4: 100% slower	Duration [‡]	272 Narr 10	(201, 448)
after May 1	Control date [‡] Total [‡]	Nov 19	(Sep 9, May 13 (2021))
	Tota1,	2,084,235	(1,728,028, 3,094,518)

^{*:} t_0 is the estimated date of the first undetected community infection; d is the estimated gap days between the first undetected case and the first reported case; a_0 is the infection rate before the reported first case; a_1 , a_2 and a_3 are rates of change of a(t) in each period measured as change per 21 days; "Duration" is the number of days from the date of the first reported case to "End date"; "End date" is the date when predicted new case decreases to zero; "Total" is the total number of predicted cases by the "End date". **: CI for d. †: Scenario 1 assumes the infection rate decreases at the same rate (i.e., a_3) after May 1; Scenarios 2 to 4 assume the relaxation of quarantine measures after May 1 will lead to a slower decrease of infection rate by 50%, 75% and 100% (complete loss of temporal effect over time). †: Under scenario 4, "Duration" and "Control date" is defined by the date when the predicted daily new case is less than 100 since the distribution of new cases has an extremely long tail (the end date defined by zero new case is May 3, 2021; CI: Dec 27, 2021 to Mar 16, 2022); and "Total" is the total predicted cases by the "Control date".