1 Estimating the infection fatality risk of COVID-19 in New York City, March 1–May 16, 2020 Wan Yang^{1*} Sasikiran Kandula² Mary Huynh³ Sharon K. Greene⁴ Gretchen Van Wye³ Wenhui 2 Li,³ Hiu Tai Chan,³ Emily McGibbon,⁴ Alice Yeung,⁴ Donald Olson,⁵ Anne Fine,⁴ Jeffrey Shaman² 3 4 ¹Department of Epidemiology, Mailman School of Public Health, Columbia University; 5 ²Department of Environmental Health Sciences, Mailman School of Public Health, Columbia University; ³Bureau of Vital Statistics, New York City Department of Health and Mental Hygiene; 6 7 ⁴Bureau of Communicable Disease, New York City Department of Health and Mental Hygiene; 8 ⁵Bureau of Equitable Health Systems, New York City Department of Health and Mental Hygiene. 9 *correspondence to: wy2202@cumc.columbia.edu (WY)

10

11 Abstract

12 During March 1-May 16, 2020, 191,392 laboratory-confirmed COVID-19 cases were diagnosed 13 and reported and 20,141 confirmed and probable COVID-19 deaths occurred among New York 14 City (NYC) residents. We applied a network model-inference system developed to support the City's pandemic response to estimate underlying SARS-CoV-2 infection rates. Based on these 15 16 estimates, we further estimated the infection fatality risk (IFR) for 5 age groups (i.e. <25, 25-44, 17 45-64, 65-74, and 75+ years) and all ages overall, during March 1–May 16, 2020. We estimated 18 an overall IFR of 1.45% (95% Credible Interval: 1.09-1.87%) in NYC. In particular, weekly IFR was 19 estimated as high as 6.1% for 65-74 year-olds and 17.0% for 75+ year-olds. These results are 20 based on more complete ascertainment of COVID-19-related deaths in NYC and thus likely 21 more accurately reflect the true, higher burden of death due to COVID-19 than previously 22 reported elsewhere. It is thus crucial that officials account for and closely monitor the infection 23 rate and population health outcomes and enact prompt public health responses accordingly as 24 the pandemic unfolds. 25 **Key words:** COVID-19; infection fatality risk; age-specific; reporting rate; New York City 26

27 Introduction

28 The novel coronavirus SARS-CoV-2 emerged in late 2019 in China and subsequently spread to

29 200+ other countries. As of June 26, 2020, there were over 9.47 million reported COVID-19

cases and over 484.2 thousand deaths worldwide.¹ As the pandemic continues to unfold and 30 31 populations in many places worldwide largely remain susceptible, understanding the severity, 32 in particular, the infection fatality risk (IFR), is crucial for gauging the full impact of COVID-19 in 33 the coming months or years. However, estimating the IFR of COVID-19 is challenging due to the 34 large number of undocumented infections, fluctuating case detection rates, and inconsistent 35 reporting of fatalities. Further, the IFR of COVID-19 could vary by location, given differences in 36 demographics, healthcare systems, and social construct (e.g., intergenerational households are 37 the norm in some societies whereas older adults commonly reside and congregate in long-term 38 care and adult care facilities in others). Most IFR estimates thus far have come from data recorded in China, the Diamond Princess cruise ship, and France.²⁻⁵ Yet the IFR in the United 39 40 States—the country currently reporting the largest number of cases—remains unclear.

41

42 New York City (NYC) reported its first case on March 1, 2020, in a traveler, and quickly 43 became the epicenter in the United States. By May 16, 2020, there were 191,392 diagnosed 44 cases and 20,131 deaths reported in NYC (Table 1). During the pandemic, the NYC Department 45 of Health and Mental Hygiene (DOHMH) and the Mailman School of Public Health at Columbia 46 University have been collaborating in generating real-time model projections in support of the 47 City's pandemic response. Our latest model-inference system uses a network model to simulate SARS-CoV-2 transmission in the City's 42 United Hospital Fund neighborhoods.⁶ The model is 48 run in conjunction with the Ensemble Adjustment Kalman Filter (EAKF)⁷ and fit simultaneously 49 50 to case and mortality data for each of the 42 neighborhoods while accounting for under-51 detection, delay from infection to case reporting and death, and changing interventions (e.g., 52 social distancing). In this study, we apply this network model-inference system to estimate the 53 IFR for 5 age groups (i.e. <25, 25-44, 45-64, 65-74, and 75+ years) and all ages overall, from 54 March 1 to May 16, 2020. In the process, we also estimate reporting rates—i.e. the fraction of 55 infections documented as confirmed cases—and the cumulative infection rate by May 16, 2020. 56

57 Methods

58 Data

59 Laboratory-confirmed COVID-19 cases reported to the NYC DOHMH were aggregated by week 60 of diagnosis and age group (<1, 1-4, 5-14, 15-24, 25-44, 45-64, 65-74, and 75+ years) for each of the 42 United Hospital Fund neighborhoods⁶ in NYC, according to the patient's residential 61 62 address at time of report. The mortality data, from deaths registered and analyzed by the NYC 63 DOHMH, combined confirmed and probable COVID-19-associated deaths. Confirmed COVID-19-64 associated deaths were defined as those occurring in persons with laboratory-confirmed SARS-65 CoV-2 infection; and probable COVID-19 deaths were defined as those with COVID-19, SARS-66 CoV-2, or a similar term listed on the death certificate as an immediate, underlying, or contributing cause of death but did not have laboratory-confirmation of COVID-19.⁸ Due to 67 privacy concerns, mortality data were aggregated to 5 coarser age groups (<18, 18-44, 45-64, 68 69 65-74, and 75+ years) for each neighborhood by week of death. To match with the age grouping 70 for case data, we used the citywide fraction of deaths occurring in each of the five finer age 71 groups (i.e. <1, 1-4, 5-14, 15-24, 25-44) to apportion deaths in the <18 and 18-44 year age 72 categories. For this study, case and mortality data were both retrieved on May 22, 2020. 73 74 The mobility data, used to model changes in COVID-19 transmission rate due to public

health interventions implemented during the pandemic (e.g., social distancing), came from SafeGraph^{9, 10} and contained counts of visitors to locations in each zip code based on mobile device locations. The released data were anonymized and aggregated in weekly intervals. We spatially aggregated these data to the neighborhood level.

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This study was classified as public health surveillance and exempt from ethical review
 and informed consent by the Institutional Review Boards of both Columbia University and NYC
 DOHMH.

83

84 Network transmission model

85 The network model simulated intra- and inter neighborhood transmission of COVID-19 and
 86 assumed susceptible-exposed-infectious-removed (SEIR) dynamics, per the following equations:

$$\begin{cases} \frac{dS_i}{dt} = -S_i \sum_{j=1}^{j=42} b_j \beta_{city} c_{ij} I_j / N_j \\ \frac{dE_i}{dt} = S_i \sum_{j=1}^{j=42} b_j \beta_{city} c_{ij} I_j / N_j - \frac{E_i}{Z} \\ \frac{dI_i}{dt} = \frac{E_i}{Z} - \frac{I_i}{D} \\ \frac{dR_i}{dt} = \frac{I_i}{D} \end{cases}$$

87

88

89 where S_i , E_i , I_i , R_i , and N_i are the numbers of susceptible, exposed (but not yet infectious), 90 infectious, and removed (either recovered or deceased) individuals and the total population, 91 respectively, from a given age group (described below) in neighborhood *i*. β_{city} is the citywide 92 transmission rate, which incorporated seasonal variation as observed for OC43, a beta-93 coronavirus in humans from the same genus as SARS-CoV-2.¹¹ To allow differential transmission 94 in each neighborhood, we included a multiplicative factor, b_i , to scale neighborhood local 95 transmission rates. *Z* and *D* are the latency and infectious periods, respectively (Table S1). 96

97 The matrix [c_{ii}] represents changes in contact rates over time and connectivity among 98 neighborhoods and was computed based on mobility data. Briefly, changes in contact rates 99 (either intra or inter neighborhoods) for week-t were computed as a ratio of the number of 100 visitors during week-t to that during the week of March 1, 2020 (the first week of the pandemic 101 in NYC when there were no interventions in place), and further scaled by a multiplicative factor 102 m_1 ; m_1 was estimated along with other parameters. To compute the connectivity among the 103 neighborhoods, we first divided the inter-neighborhood mobility by the local mobility (this gave 104 a relative measure of connectivity; e.g., if two neighborhoods are highly connected with lots of 105 individuals traveling between them, inter-neighborhood mobility would be closer to 1 and 106 much lower than 1 otherwise); we then scaled these relative rates by a multiplicative factor m_2 , 107 which was also estimated along with other parameters.

108

109 Observation model

110 To account for delays in diagnosis and reporting, we included a time-from-infectious-to-case-111 reporting (i.e., diagnosis) lag, drawn from a gamma distribution with a mean of T_m and standard 112 deviation (SD) of T_{sd} days. To account for under-detection, we included a case reporting rate (r), 113 i.e. the fraction of infections (including subclinical or asymptomatic infections) reported as 114 cases. To compute the model-simulated number of new cases per week, we multiplied the 115 model-simulated number of infections per day (including those from the previous weeks) by the 116 reporting rate, and further distributed these simulated cases in time per the distribution of 117 time-from-infectious-to-case-reporting. We then aggregated the daily lagged, reported cases to 118 weekly totals for model inference (see below). Similarly, to compute the model-simulated 119 deaths per week, we multiplied the simulated-infections by the IFR and then distributed these 120 simulated deaths in time per the distribution of time-from-infectious-to-death, and aggregated 121 these daily numbers to weekly totals. For each week, the reporting rate (r), the mean (T_m) and 122 standard deviation (T_{sd}) of time-from-infectious-case-reporting, and the IFR were estimated 123 based on weekly case and mortality data. The distribution of time-from-diagnosis-to-death was 124 based on observations of n=15,686 COVID-19 confirmed deaths in NYC (gamma distribution 125 with mean = 9.36 days and SD = 9.76 days; Table S1).

126

127 Parameter estimation

128 To estimate model parameters (b_i , β_{citv} , Z, D, m_1 , m_2 , T_m , T_{sd} , r, and IFR, for i=1,...,42) and state 129 variables (S_i , E_i , and I_i , for i=1,...,42) for each week, we ran the network-model stochastically 130 with a daily time step in conjunction with the EAKF and fit to weekly case and mortality data 131 from the week starting March 1 to the week ending May 16, 2020. The EAKF uses an ensemble 132 of model realizations (n=500 here), each with initial parameters and variables randomly drawn 133 from a prior range (see Table S1). After model initialization, the model ensemble was integrated 134 forward in time for a week to compute the model-simulated number of cases and deaths for 135 that week; these prior estimates were then combined with the observed cases and deaths for the same week to compute the posterior per Bayes' theorem.⁷ The posterior distribution of 136 137 each model parameter/variable was updated for that week at the same time.⁷ This parameter 138 estimation process was done separately for each of the eight age groups (i.e. <1, 1-4, ..., and

75+). To account for stochasticity in model initiation, we ran the parameter estimation process
independently 10 times. Results for each age group were combined from these 10 runs (each
with 500 realizations). To combine estimates of reporting rate and IFR for <25 year-olds or all
ages overall, we weighted the age-group specific estimates by the fraction of estimated
infections from each related group.

145 **Results**

The model-inference system was able to recreate the case and mortality time series for each age group and all ages overall (Fig. 1). For most age groups, confirmed cases peaked during the week of March 29 and the mortality rate peaked about one week later than the case rate, due to the time-lag from severe infection to death (Fig. 1).

150

151 There were, however, substantial under-detection of infections, variations by age group, 152 and fluctuations of case reporting rates over time, in part due to changing testing criteria (e.g., 153 testing was restricted to severely ill patients in the early phase due to material shortages in 154 testing equipment and personal protective equipment). The estimated reporting rate for all 155 ages overall started at a low level of 2.3% [median; 95% credible interval (CrI): 0.4–4.5%; same 156 below] in the week of March 1; it increased to 21.4% (95% Crl: 14.4–31.3%) in the week of 157 March 15 and stayed at similar levels afterwards (Fig 2F). The estimated reporting rate was 158 highest for the two oldest age groups and substantially lower for younger age groups (Fig 2 D 159 and E vs. Fig 2 A-C). During the last week of this study (i.e., May 10, 2020), we estimated that 160 25.9% (95% Crl: 15.1–41.8%) of infections among 65-74 year-olds and 34.2% (95% Crl: 23.0– 161 49.9%) among 75+ year-olds were reported; in comparison, only 10.6% (95% Crl: 7.4–17.7%) of 162 infections among <25 year-olds and 16.9% (95% Crl: 13.1–26.8%) among 25-44 year-olds were 163 reported.

164

After accounting for the case reporting rate, the epidemic peak for new infections
 occurred one week sooner during the week of March 22, 2020 for <45 year-olds and all ages
 combined (Fig 2 A-B and F). This was coincident with the timing of public health interventions in

168 NYC – public schools in NYC were closed on March 16, 2020 and a citywide stay-at-home order was imposed starting the week of March 22, 2020.¹² Tallied over the entire study period, the 169 estimated overall cumulative infection rate was 15.8% (95% Crl: 11.4–24.4%) by May 16, 2020. 170 171 However, the estimated cumulative infection rate varied substantially by age group. 172 Specifically, 25-44 and 45-64 year-olds had the highest cumulative infection rates, at 20.1% 173 (95% Crl: 14.4–30.0%) and 20.7% (95% Crl: 15.7–28.0%), respectively; 65-74 and 75+ year-olds 174 had the second highest cumulative infection rates, at 14.9% (95% CrI: 11.2–22.8%) and 12.9% 175 (95% Crl: 9.9–19.9%); and <25 year-olds had the lowest cumulative infection rate (8.0%; 95% 176 Crl: 5.1–16.7%). Of note, these estimates, albeit with large uncertainties, are in line with reported measures from serology surveys (e.g., 19.9% positive in NYC, as of May 1, 2020, likely 177 from testing of 25-64 year-olds^{13, 14}). In addition, the spatial variation estimated by our model-178 inference system¹⁵ was in line with reported measures (i.e., highest in the Bronx and lowest in 179 180 Manhattan^{13, 14}). This consistency with independent serology survey data provides some 181 independent validation of our model estimates.

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During March 1 - May 16, 2020, a total of 20,141 COVID-19 deaths (15,723 confirmed and 4,418 probable) and 191,392 COVID-confirmed cases were reported in NYC. The crude confirmed case fatality risk was thus 8.22%. After accounting for changing case reporting rates and excluding the first three weeks (i.e., March 1-21, 2020) with zero or few reported deaths for which model estimates were less accurate, we estimate that the overall IFR, including both confirmed and probable deaths, was 1.45% (95% CrI: 1.09–1.87%) during March 22 – May 16, 2020.

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Examining estimates by age group, estimated IFR was lowest in young age groups. The average IFR was 0.011% (95% CrI: 0.005–0.016%) for <25 year-olds, increased by ~10 fold to 0.12% (95% CrI: 0.077–0.15%) for 25-44 year-olds, and by another 7 fold to 0.94% (95% CrI: 0.74–1.21%) for 45-64 year-olds (Fig. 3 A-C). These estimates were similar to IFRs reported for China for corresponding age groups.³ However, the estimated IFR for the two oldest age groups was much higher than the younger age groups and about twice as high as rates reported

for these age groups in China.^{3, 4} The average IFR was 4.67% (95% Crl: 3.21–6.66%) for 65-74
year-olds and 13.83% (95% Crl: 9.65–17.78%) for 75+ year-olds. In addition, the estimated IFR
fluctuated substantially over time for these two elderly groups. For 65-74 year-olds, estimated
IFR was 6.10% (95% Crl: 4.90–7.57%) during the week of April 5, 2020 but decreased to 3.79%
(95% Crl: 1.68–6.90%) during the week of May 10, 2020 (Fig 3D). For 75+ year-olds, IFR was
estimated to be 16.99% (95% Crl: 13.15–20.11%) during the week of April 5, 2020 but
decreased to 9.77% (95% Crl: 4.53–14.81%) during the week of May 10, 2020 (Fig 3F).

204

205 **Discussion**

206 In light of the large uncertainties in IFRs for COVID-19 due to under-ascertainment of cases, we 207 have used a model-inference system, developed to support the pandemic response in NYC, to 208 estimate local IFRs. During March 1–May 16, 2020, NYC recorded the largest numbers of 209 COVID-19 cases and deaths in the US and perhaps worldwide. Despite public health efforts to 210 slow the pandemic, e.g. by social distancing, and to ramp up healthcare capacity, over 20,000 211 lives were lost from COVID-19 in a short span of two months. Based on this large number of 212 deaths, the estimated overall IFR was 1.45% in NYC. This estimate included both confirmed and 213 probable COVID deaths. If only COVID confirmed deaths were included, given that 78.0% of 214 deaths among the total were laboratory-confirmed, the estimated overall IFR would be around 215 1.1%. Both estimates were higher than previously reported for elsewhere (e.g., about 0.7% in both China³ and France⁵). Importantly, NYC has nosologists who review all death certificates 216 217 and record deaths into a unified electronic reporting system rapidly. This mortality surveillance 218 infrastructure and enhanced nosology thus allow more rapid and complete death reporting in 219 NYC. As such, our estimates here likely more accurately reflect the underlying fatality risk of 220 COVID-19 infection. Further, given the likely stronger public health infrastructure and healthcare systems in NYC than many other places,¹⁶ the higher IFR estimated here suggests 221 222 that mortality risk from COVID-19 may be higher in the United States and likely other countries 223 as well than previously reported. Of note, despite the large surge in cases and hospitalizations, 224 through quick expansion of healthcare systems, most hospitals in NYC were able to meet 225 patient care demand during the two-month period. As many jurisdictions in the United States

are considering re-opening after months of social distancing, it is crucial that officials account for and closely monitor the infection rate and health outcomes including hospitalizations and mortality and take prompt public health responses accordingly.

229

230 While the IFR estimated here was similar to those previously reported elsewhere for 231 younger age groups, we found that IFRs for individuals 65 years and older in NYC were about 232 twice as high as prior reports.³ These higher IFRs may be in part due to differences in 233 population characteristics, in particular, the prevalence of underlying medical conditions such as diabetes mellitus, chronic lung disease, and cardiovascular disease.^{17, 18} Regardless, 234 235 estimated weekly IFR was as high as 6.1% for 65-74 year-olds and 17.0% for 75+ year-olds. 236 These dire estimates highlight the severity of COVID-19 in elderly populations and the 237 importance of infection prevention in congregate settings. Thus, early detection and adherence 238 to infection control guidance in long-term care and adult care facilities should be a priority for 239 COVID-19 response as the pandemic continues to unfold.

240

241 In this study, we incorporated multiple data sources, including age-grouped, spatially 242 resolved case and mortality data as well as mobility data, to calibrate our model-inference 243 system. Of note, the timing of the COVID-19 pandemic varied substantially among NYC 244 neighborhoods. For instance, peak mortality rates occurred up to 4 weeks apart among the 42 245 neighborhoods. Fitting the model-inference system simultaneously to these diverse case and 246 mortality time series thus enabled better constraint of key model parameters (e.g., case 247 reporting rate and IFR). However, we note there remain large uncertainties in model estimates. 248 A full assessment of COVID-19 severity will require comprehensive serology surveys of the 249 population by age group and neighborhood, given the large heterogeneity of infection rates 250 across population segments and space. In addition, we only included deaths that were lab-251 confirmed or explicitly coded as related to COVID-19. A recent study reported that excess deaths in NYC during about the same period could be more than 24,000.⁸ Further, recent 252 253 studies have reported severe sequelae of COVID-19 in children, i.e. Multi-system Inflammatory

- Syndrome in Children. Thus, it is important to monitor health outcomes in younger age groups
 post-infection as the pandemic unfolds, despite the low IFRs noted to date.
- 256

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- 269

270 **Conflict of Interest:**

- JS and Columbia University disclose partial ownership of SK Analytics. JS discloses consulting for
 BNI. Other authors declare no conflict of interest.
- 273

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Table 1. Summary estimates. Cases and deaths were reported during March 1 – May 16, 2020. Crude case fatality risk (CFR) was computed as the proportion of persons with confirmed COVID-19 illness who died. Cumulative infection rates, median (95% CrI), show percentages of population, for each age group or all ages overall, estimated to have been infected by May 16, 2020. IFR, median (95% CrI), was estimated here, and averaged over March 22 - May 16, 2020; we excluded estimates during March 1-21, 2020, because estimates were less accurate for these earliest weeks when zero or few deaths were reported.

	Confirmed	Confirmed and	Estimated cumulative	
Age	Cases	Probable Deaths	infection rate (%)	Estimated IFR (%)
<25	14692	40	8 (5.1, 16.7)	0.011 (0.005, 0.016)
25-44	60474	688	20.1 (14.4, 30)	0.12 (0.077, 0.15)
45-64	69839	4457	20.7 (15.7, 28)	0.94 (0.74, 1.21)
65-74	23875	4866	14.9 (11.2, 22.8)	4.67 (3.22, 6.66)
75+	22512	10090	12.9 (9.9 <i>,</i> 19.9)	13.83 (9.65, 17.78)
all	191392	20141	15.8 (11.4, 24.4)	1.45 (1.09, 1.87)

Figures

Figure 1. Model fit. Black boxes show model estimates of cases per 100,000 population and grey boxes show model estimates of mortality rates; thick horizontal lines and box edges show the median, 25th, and 75th percentiles; vertical lines extending from each box show 95% Crl. Blue dots indicate observed incidence rates and red dots show observed mortality rates.

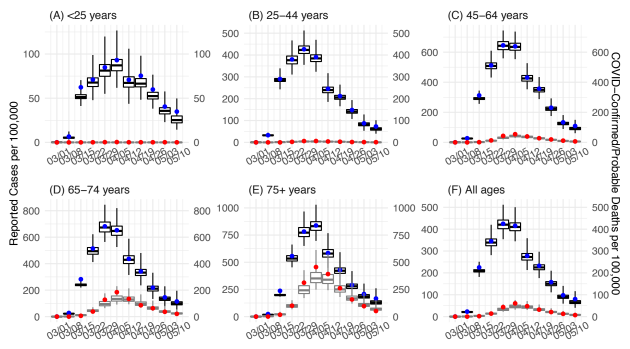
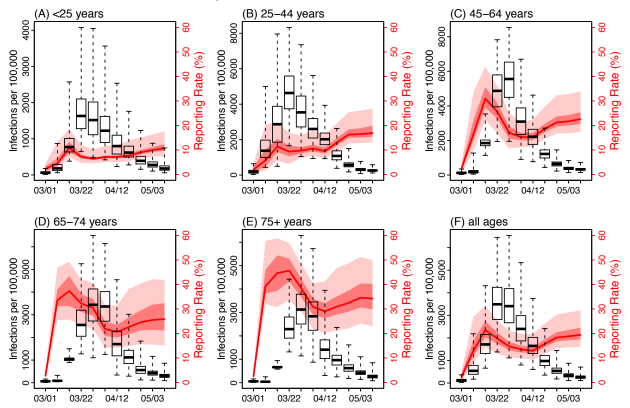


Figure 2. Estimated infection and case reporting rates over time. Boxplots show estimated infection rates. Red lines show the estimated median case reporting rate with surrounding areas show the 50% and 95% CrI. x-axis shows the first day of each week (mm/dd) from the week of March 1 to the week of March 10, 2020.



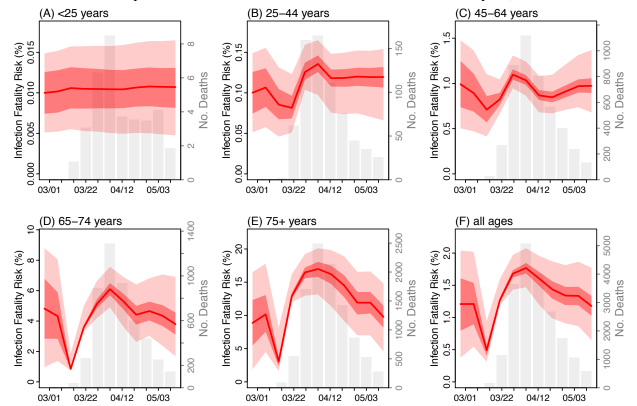


Figure 3. Estimated infection fatality risk. Red lines show the estimated median IFR with surrounding areas indicating the 50% and 95% CrI. For comparison, the grey bars show the number of deaths reported for each week from the week of March 1 to May 10, 2020.

Supplementary Material

Table S1. Prior ranges for main model parameters and variables. The spatial, temporal, and age resolution of each parameter or variable, estimated in the model-inference system, is specified in the column "Resolution". Note posterior parameter estimates can extend outside the specified prior ranges.

Parameter/variable	Symbol	Resolution	Prior range	Source/rationale
Initial exposed	<i>E</i> (t=0)	neighborhood- and age-	300 – 8000 total citywide,	Large uncertainties, used very
		group specific, estimated	scaled by population size for	wide range
		for the beginning of the	each age group and	
		Week of March 1, 2020	neighborhood	
Initial infectious	/(t=0)	neighborhood- and age-	150 – 4000 total citywide,	Assumed to be half the initial
		group specific, ,	scaled by population size for	exposed
		estimated for the	each age group and	
		beginning of the Week of	neighborhood	
		March 1, 2020		
Initial susceptible	<i>S</i> (t=0)	neighborhood- and age-	N - E - I	Assumed all were susceptible
		group specific, estimated		except for those initially
		for the beginning of the		exposed/infectious
		Week of March 1, 2020		

Population size in	Ν	neighborhood- and age-	N/A	NYC intercensal population
each age group and		group specific		estimates for 2018 ¹
neighborhood				
Citywide	B _{city}	Citywide, age-group	[0.5, 1] per day overall; scaled	Based on R_0 estimates of around
transmission rate		specific, estimated for	by contact rate for each age	1.5-4 for SARS-CoV-2 ³⁻⁵
		each week	group based on contact data	
			from the POLYMOD study ²	
Scaling of	b _i	neighborhood- and age-	[0.8, 1.2] for age groups under	Around 1; larger variation for
neighborhood		group specific, estimated	65 years; [0.5, 1.5] for age	elderly groups based on data
transmission rate		for each week	groups 65 or older	
Latency period	Ζ	Citywide, age-group	[2, 5] days	Incubation period: 5.2 days (95%
		specific, estimated for		CI: 4.1, 7) ³ ; latency period is likely
		each week		shorter than the incubation period
Infectious period	D	Citywide, age-group	[2, 5] days	Time from symptom onset to
		specific, estimated for		hospitalization: 3.8 days (95% CI:
		each week		0, 12.0) in China, ⁶ plus 1-2 days
				viral shedding before symptom

onset. We did not distinguish symptomatic/asymptomatic infections.

Multiplicative	m_1	Citywide, age-group	[1, 2] for <1 year; [0.5, 1.5] for	Initial model testing showed
factor for mobility		specific, estimated for	three age groups 1-24 years;	transmission rates for younger age
		each week	[0.1, 1.5] for age group 25-44;	groups were more sensitive to
			[1, 2.5] for age groups 45 or	changes in mobility whereas the
			older	two oldest age groups were not
				sensitive to mobility. For age
				groups with contact rates lower
				than the average (based on the
				POLYMOD study ²), we raised the
				diagonal elements in the mobility
				matrix to the power of the relative
				contact rate (<1) to account for
				insensitivity of transmission rate in
				these age groups to mobility.

Multiplicative	<i>m</i> ₂	Citywide, age-group	[0.5, 2]	Likely around 1 but with large
factor for		specific, estimated for		uncertainties
neighborhood		each week		
connectivity				
Mean of time from	T _m	Citywide, age-group	[3, 8] days	From a few days to a week from
viral shedding to		specific, estimated for		symptom onset to diagnosis/
diagnosis		each week		reporting, ⁶ plus 1-2 days of viral
				shedding (being infectious) before
				symptom onset
Standard deviation	T _{sd}	Citywide, age-group	[1, 3] days	To allow variation in time to
(SD) of time from		specific, estimated for		diagnosis/reporting
viral shedding to		each week		
diagnosis				
Reporting rate	r	Citywide, age-group	Starting from [0.001, 0.05] at	Large uncertainties
		specific, estimated for	time 0 and allowed to increase	
		each week	over time using space re-	
			probing ⁷	
Infection fatality		Citywide, age-group	[5, 15]×10 ⁻⁴ for ages under 25;	Based on previous estimates ⁸ but
risk (IFR)		specific, estimated for	[5, 15]×10 ⁻³ for ages 25-44; [5,	extend to have wider ranges
		each week	15]×10 ⁻² for ages 45-64; [0.01,	

		0.1] for ages 65-74; [0.02, 0.2]	
		for ages 75+;	
Time from	Citywide	Gamma distribution with mean	Based on <i>n</i> =15,686 COVID-19
diagnosis to death		of 9.36 days and SD of 9.76	confirmed deaths in NYC as of May
		days	17, 2020.

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