

Accelerated Article Preview**Burden and characteristics of COVID-19 in the United States during 2020**

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Burden and characteristics of COVID-19 in the United States during 2020

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The COVID-19 pandemic disrupted health systems and economies throughout the world during 2020 and was particularly devastating for the United States, which experienced the highest numbers of reported cases and deaths during 2020^{1–3}. Many of the epidemiological features responsible for observed rates of morbidity and mortality have been reported^{4–8}; however, the overall burden and characteristics of COVID-19 in the United States have not been comprehensively quantified. Here we use a data-driven model-inference approach to simulate the pandemic at county-scale in the United States during 2020 and estimate critical, time-varying epidemiological properties underpinning the dynamics of the virus. The pandemic in the US during 2020 was characterized by national ascertainment rates that increased from 11.3% (95% credible interval (CI): 8.3 – 15.9%) during March to 24.5% (18.6 – 32.3%) during December. Population susceptibility at year's end was 69.0% (63.6 – 75.4%), indicating that roughly one third of the US population had been infected. Community infectious rates, the percentage of people harbouring a contagious infection, rose above 0.8% (0.6 – 1.0%) before the end of the year, and were as high as 2.4% in some major metropolitan areas. In contrast, the infection fatality rate fell to 0.3% by year's end.

During 2020, the United States (US) documented more COVID-19 cases and deaths than any other country in the world¹. The first COVID-19 case was identified in Washington state on January 20, 2020². Over the course of the year, three pandemic waves took place: 1) a spring season outbreak in select, mostly urban areas following the introduction of the virus to the US; 2) a summer wave that predominantly affected the southern half of the country; and 3) a fall-winter wave that remained pervasive through the spring of 2021. To understand the transmission of the virus and better control its progression in the future, it is vital that the epidemiological features that have supported these outbreaks be quantified and analyzed in both space and time.

Here we use a county-resolved metapopulation model to simulate the transmission of SARS-CoV-2 within and between the 3142 counties of the US. The model depicts both documented and undocumented infections and is coupled with an iterative Bayesian inference algorithm—the ensemble adjustment Kalman filter—which assimilates observations of daily cases in each county, as well as population movement between counties^{9,10} (see Supplementary Information, Extended Data Table 1). The Bayesian inference supports a fitting of the model to case observations and estimation of unobserved state variables (e.g. population susceptibility within a county) and system parameters (e.g. the ascertainment rate in each county). Synthetic tests indicate that the inference approach can recover key time-varying parameters across a diversity of simulation scenarios (Extended Data Fig. 1). The model fitting to observed case data captures the 3 waves of the outbreak as manifest at national scales (Fig. 1a), as well as in major metropolitan areas and at county scales (Extended Data Fig. 2). These inference results are robust to parameter settings and model

configurations (see Supplementary Information, Extended Data Figs. 3–4).

To further validate the fitting, we compared model estimates of cumulative infections to findings from US Centers for Disease Control and Prevention (CDC) seroprevalence surveys conducted at site and state levels³. The seroprevalence data, which provide an out-of-sample corroboration of the model fitting, were adjusted for the waning of antibody levels following adaptive immune response^{11,12} (see Supplementary Information, Extended Data Fig. 5). Model estimates of cumulative infected percentages are well aligned with adjusted seroprevalence estimates from the CDC 10-site survey across sites and through time (Pearson $r=0.97$, mean absolute error (MAE)=1.31%) (Fig. 1b) and are similarly well matched to adjusted estimates at the state level (Extended Data Fig. 6). In addition, the seroprevalence generated using the estimated daily infections adjusted for seroreversion also matches the observed seroprevalence, and the results are robust to assumed use of a lower sensitivity seroassay (Extended Data Fig. 6).

A critical feature of SARS-CoV-2 is its ability to infect and transmit largely from individuals never diagnosed with the virus⁴. The model structure and fitting enable estimation of the ascertainment rate, the percentage of infections confirmed diagnostically, at county scales. The national population-weighted ascertainment rate averaged for all of 2020 was 21.8% (95% credible interval (CI): 15.9 – 30.3%), similar to an estimate derived from surveys on healthcare-seeking behaviours¹³. This national ascertainment rate increased from 11.3% (8.3 – 15.9%) during March 2020 to 24.5% (18.6 – 32.3%) during December 2020 (Fig. 1c). The increase through time is a likely by-product of increasing testing capacity, a relaxation of initial restrictions on test usage, and

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increasing recognition, concern and care-seeking among the public. We additionally focus on 5 metropolitan areas within the US. Small differences in the ascertainment rate manifest across these areas; in particular, ascertainment rates for Phoenix and Miami are higher than the national average for much of the year, whereas New York City, Chicago, and Los Angeles are consistently below the national average.

At the national level, three pandemic waves are clearly evident during spring, summer and fall/winter (Fig. 1a); however, among the 5 focus metropolitan areas the structure differs with New York and Chicago experiencing strong spring and fall/winter waves but little activity during summer, Los Angeles and Phoenix undergoing summer and fall/winter waves, and Miami experiencing all 3 waves (Extended Data Fig. 2). Los Angeles County, the largest county in the US with a population of more than 10 million people, was particularly hard-hit during fall/winter. The differences in virus activity produced different cumulative infection numbers through time (Fig. 2a). Population susceptibility at the end of the year was 69.0% (63.6–75.4%) for the US, and among the focal metropolitan areas ranged from 47.6% (37.2–54.8%) in Los Angeles to 73.2% (68.3–77.8%) in Phoenix. Though there is variability among counties, a substantial portion of the US population (69.0%) had not been infected by the end of 2020; however, pockets of lower population susceptibility, which are evident in the southwest and southeast on August 1, 2020 (Fig. 2b), expanded considerably by December 31, 2020 (Fig. 2c). In particular, areas of the upper Midwest and Mississippi valley, including the Dakotas, Minnesota, Wisconsin and Iowa, are estimated to have population susceptibility below 40% as of December 31, 2020.

The structure of the outbreak is evident in both incidence and prevalence estimates (Fig. 3, Extended Data Fig. 7). Incidence indicates the daily number of newly infectious individuals—both those who will be confirmed COVID-19 cases and those whose infections will remain undocumented. The majority of infections each month are undocumented (Fig. 3a), as indicated by the low ascertainment rates (Fig. 1c). For all of 2020, an estimated 78.2% of infections in the US were undocumented. Estimates of daily prevalence provide a measure of the community infectious rate (CIR), the fraction of the population currently harboring a contagious infection. The national SARS-CoV-2 CIR was 0.77% (0.60–0.98%) on December 31 2020, indicating that roughly 1 in 130 persons was contagious (a similar percentage, 0.83% (0.52–1.26%), was estimated to be latently infected, i.e. infected but not yet contagious) (Fig. 3b). Among the 5 focal metropolitan areas, the CIR varied considerably: in mid-November, Chicago reached a CIR of 1.51% (1.27–1.82%); whereas in Miami CIR rose to 1.25% (1.03–1.53%) during July. Los Angeles was even more burdened at the end of 2020 with a CIR of 2.42% (2.05–2.86%) as of December 31, 2020 (Extended Data Fig. 7).

The model fitting enables estimation of the case fatality rate (CFR) and the infection fatality rate (IFR). Using public line-list data from the US CDC¹⁴, we estimated the distribution of time lag from case confirmation to death for each county and, using these estimates, deconvolved observed deaths to their date of case reporting¹⁵ (see Supplementary Information, Extended Data Figs. 8–9). CFR and IFR were then generated using these deconvolved death data. Both rates were highest nationally at the beginning of the spring wave: the CFR was 7.1% (4.8–9.8%) and the IFR was 0.77% (0.51–1.25%) during April (Fig. 3c). The national cumulative IFR through June 1 was 0.69% (0.47–1.04%), in line with previous studies^{5–7} (see Supplementary Information, Extended Data Fig. 2). Over the course of the year, with earlier diagnosis and treatment, improved patient care^{16–18}, and, in the case of CFR, increased reporting of mild infections, the CFR and IFR dropped to 1.29% (0.98–1.68%) and 0.31% (0.22–0.44%) by December 2020, respectively. Both rates varied by location and over time; for instance, intermediate drops of CFR and IFR began for Chicago, Phoenix and Miami during the summer wave in association with a decrease of the average age of hospitalized patients (Extended Data Fig. 8). During the winter of 2020, the CFR and IFR in most metropolitan areas increased slightly, possibly driven by greater hospitalization rates among the elderly (Extended Data Fig. 8) and

strained healthcare resources¹⁹. Overall, these findings delineate the mortality risk associated with infection broadly. The national IFR during the latter half of 2020 hovers around 0.30%, well above estimates for both seasonal influenza (<0.08%)²⁰ and the 2009 influenza pandemic (0.0076%)²¹. We note that as COVID-19 deaths are likely under-reported, our estimate of IFR could be biased low.

We further examined the change of the reproduction number, R_t , in response to changing local, reported COVID-19 case numbers in five US regions (Northeast, Southeast, Midwest, Southwest, and West) during the spring, summer, and fall/winter waves (see Supplementary Information). Results indicate that communities with increasing cases had greater reductions of virus transmission potential, R_t (Extended Data Fig. 10). However, the rate of reduction in R_t decreased over successive waves. These findings are potentially driven by a number of factors modulating the reproduction number, including changing compliance with non-pharmaceutical interventions²² and seasonal modulation of virus transmissibility²³. A more thorough analysis of this preliminary finding is needed.

The US experienced the highest numbers of confirmed COVID-19 cases and deaths in the world during 2020¹. Our findings provide quantification of the time-evolving epidemiological characteristics associated with successive pandemic waves in the US, as well as conditions at the end of the year and prospects for 2021. Critically, despite more than 19.6 million reported cases at year's end, an estimated 69% of the population remained susceptible to viral infection. Several factors will considerably alter population susceptibility in the coming months. Firstly, ongoing transmission will infect naïve hosts and continue to deplete the susceptible pool. Secondly, as more vaccine is distributed and administered, additional individuals will be protected against symptomatic infection and the IFR will decrease. Lastly, our model does not represent re-infection, either through waning immunity or immune escape; however, re-infection has been documented^{24,25}, evidence of waning antibody levels exists^{26,27}, and new variants of concern have emerged^{28,29} and will likely continue to do so. All these processes will affect population susceptibility over time and help determine when society enters a post-pandemic phase, the pattern of endemicity the virus ultimately assumes, and its long-term public health burden³⁰.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41586-021-03914-4>.

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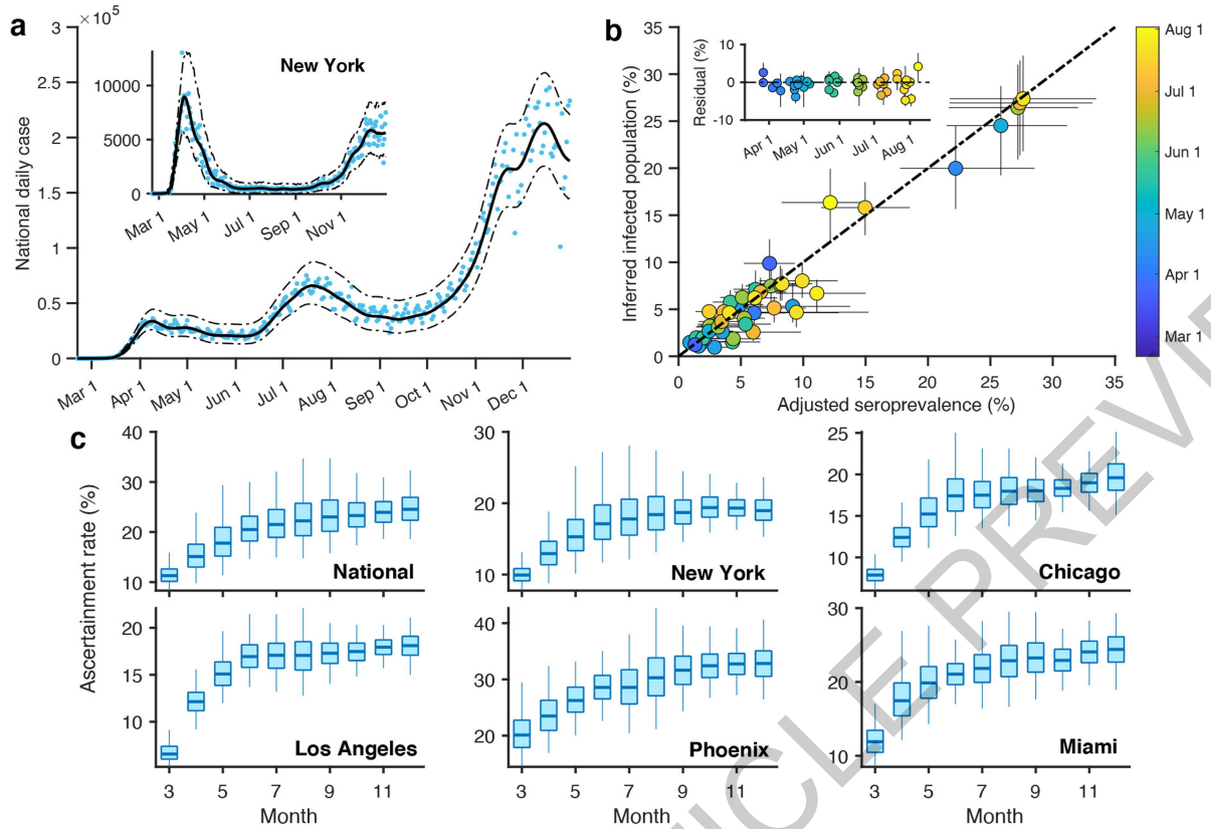


Fig. 1 | Model calibration and ascertainment rate. (a), Model fitting to daily case numbers (blue dots) in the US and the New York metropolitan area (inset). Solid and dashed lines show the median estimate and 95% CIs, respectively. (b), Comparison between inferred percentage cumulative infections and seroprevalence in 10 locations adjusted for antibody waning. The inset shows residuals of inference (inferred percentage of infected population minus adjusted seroprevalence). Centres and whiskers show medians and 95% CIs,

and color indicates the sample collection date in each location. Distributions are obtained from $n=100$ ensemble members. Details on the serological survey are provided in Supplementary Information. (c), Distributions of estimated ascertainment rate in the US and five metropolitan areas. Centre and box bounds represent median, 25%, and 75% percentiles, and whiskers show 2.5% and 97.5% percentiles. Monthly posterior estimates are presented for March to December 2020. Distributions are obtained from $n=100$ ensemble members.

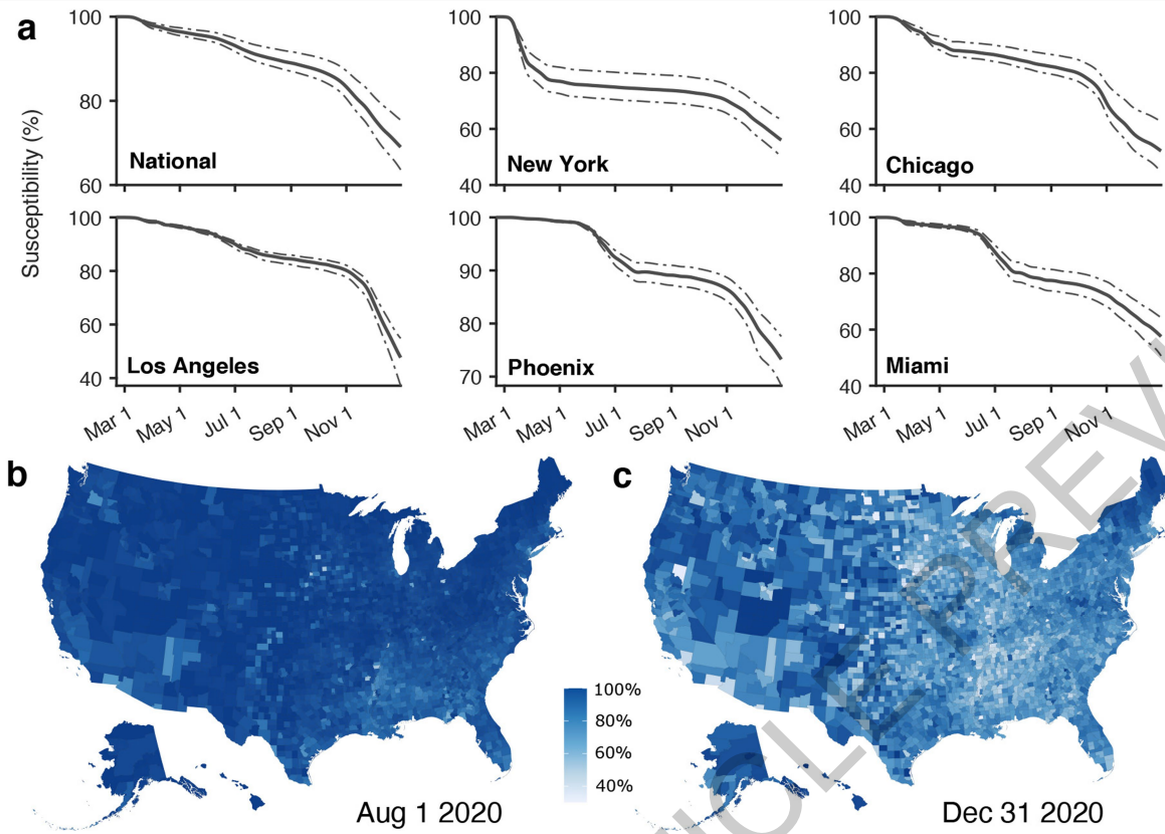


Fig. 2 | Estimates of population susceptibility. (a), Estimated evolution of susceptibility to COVID-19 in the US and five metropolitan areas. Solid and dashed lines show median estimate and 95% CIs. (b-c), Estimated susceptibility in 3,142 US counties on August 1 and December 31, 2020. Color shows median estimate.

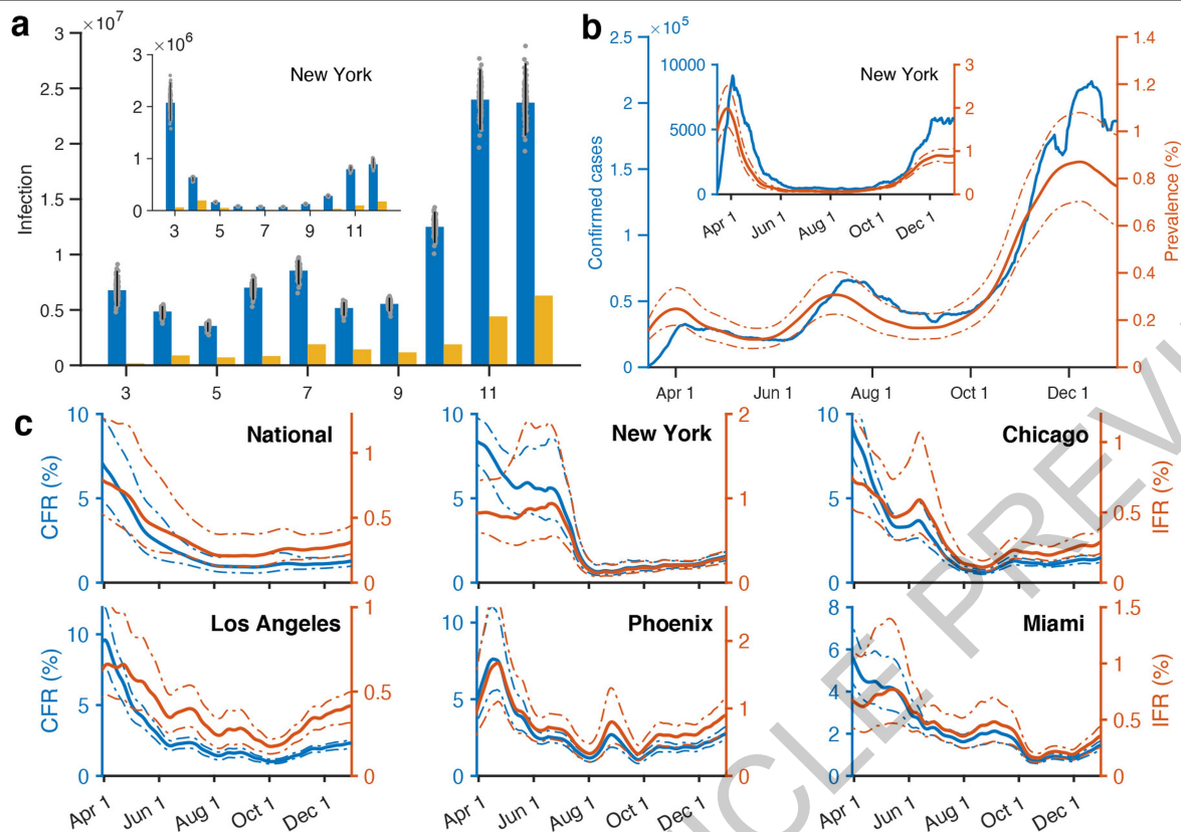


Fig. 3 | Estimated transmission and characteristics of COVID-19 in the US.

(a), Estimated monthly total infections (blue bars) and confirmed cases (orange bars) in the US and the New York metropolitan area (inset). Distributions are obtained from $n=100$ ensemble members. The blue bars represent the medians and whiskers show 95% CIs. (b), Daily confirmed cases (blue line, 7-day moving

average) and estimated prevalence of contagious infections (red line, median and 95% CIs) in the US. Result for the New York metropolitan area is shown in the inset. (c), Estimated CFR (blue lines) and IFR (red lines) in the US and five metropolitan areas. Solid and dashed lines show median estimate and 95% CIs.

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

Data Availability

The human mobility and COVID-19 surveillance data that support the findings of this study are available at GitHub (https://github.com/SenPei-CU/COVID_US_2020). The county-level COVID-19 surveillance data in the US are available at Johns Hopkins University coronavirus resource center (https://github.com/CSSEGISandData/COVID-19/tree/master/csse_covid_19_data/csse_covid_19_time_series). County-to-county commuting data were downloaded from the US Census Bureau (<https://www.census.gov/data/tables/2015/demo/metro-micro/commuting-flows-2015.html>). Human mobility data in 2020 were provided by SafeGraph (<https://safegraph.com/>), a data company that aggregates anonymized location data from numerous applications in order to provide insights about physical places, via the SafeGraph Community. To enhance privacy, SafeGraph excludes census block group information if fewer than five devices visited an establishment in a month from a given census block group. We aggregated the mobility data to county level to estimate change of inter-county mobility in 2020. Aggregated and derived data are allowed to be shared publicly by SafeGraph. Seroprevalence data were published by the US CDC (<https://www.cdc.gov/coronavirus/2019-ncov/cases-updates/>

[commercial-lab-surveys.html](https://www.cdc.gov/coronavirus/2019-ncov/cases-updates/commercial-lab-surveys.html)). The line-list datasets are available at the US CDC website (<https://data.cdc.gov/Case-Surveillance/COVID-19-Case-Surveillance-Public-Use-Data/vbim-akqf> and <https://data.cdc.gov/Case-Surveillance/COVID-19-Case-Surveillance-Public-Use-Data-with-Ge/n8mc-b4w4>). Source data are provided with this paper.

Code Availability

Custom code supporting this study is available at GitHub (https://github.com/SenPei-CU/COVID_US_2020). Source data are provided with this paper.

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Author contributions SP and JS conceived the study, SP, TKY, SK, MG performed the analysis, SP and JS drafted the manuscript, all authors revised and reviewed the manuscript.

Competing interests J.S. and Columbia University disclose partial ownership of SK Analytics. J.S. discloses consulting for BNI. All other authors declare no competing interests.

Additional information

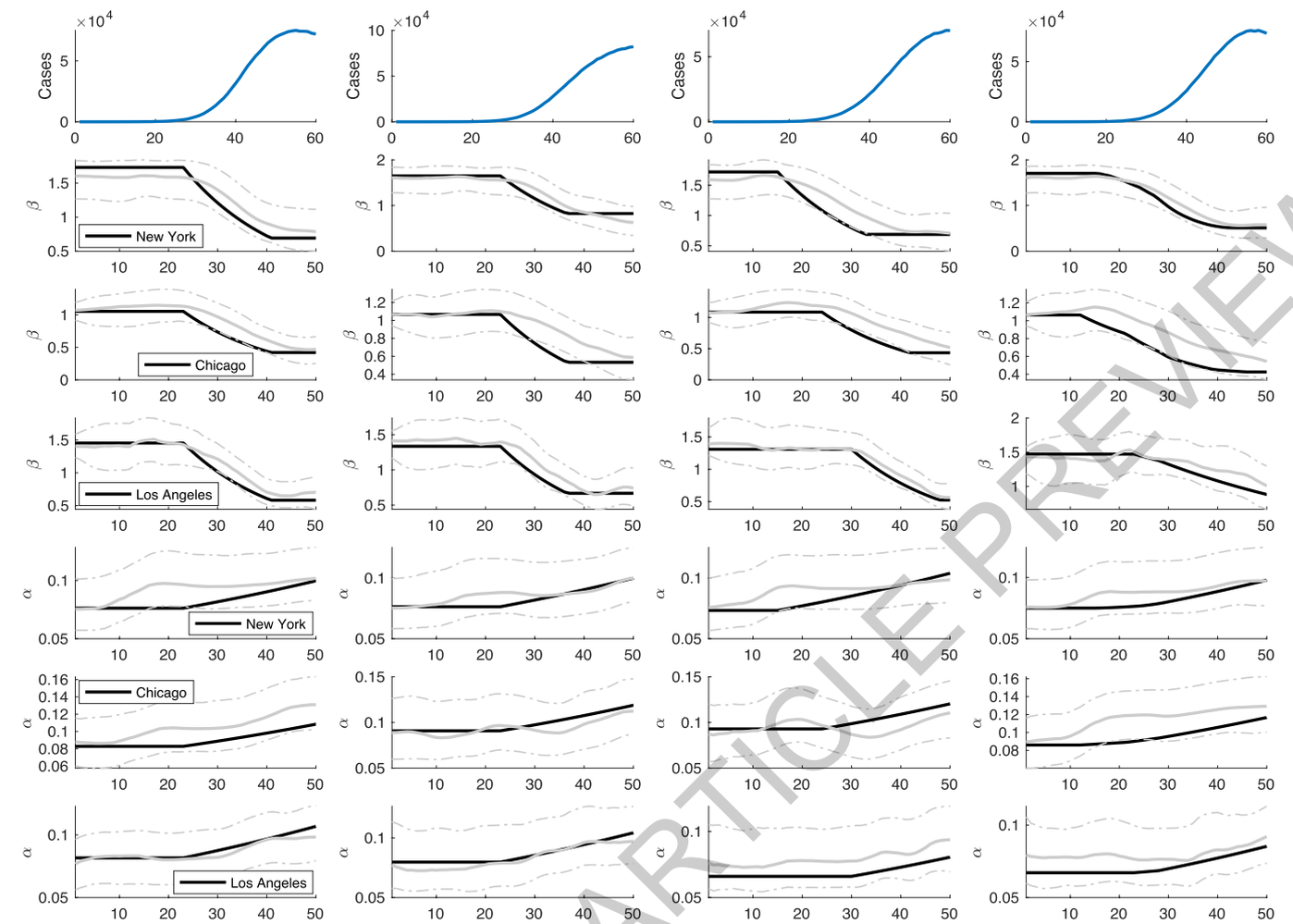
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Correspondence and requests for materials should be addressed to P.S. or J.S.

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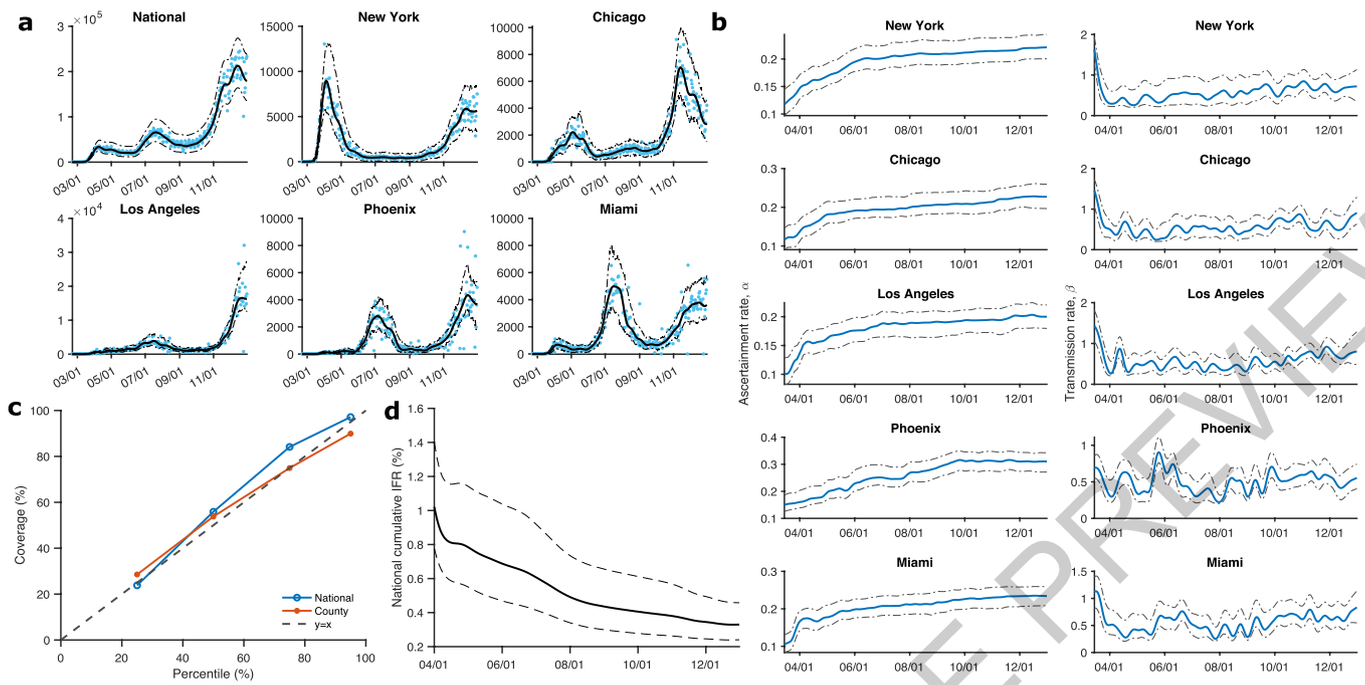
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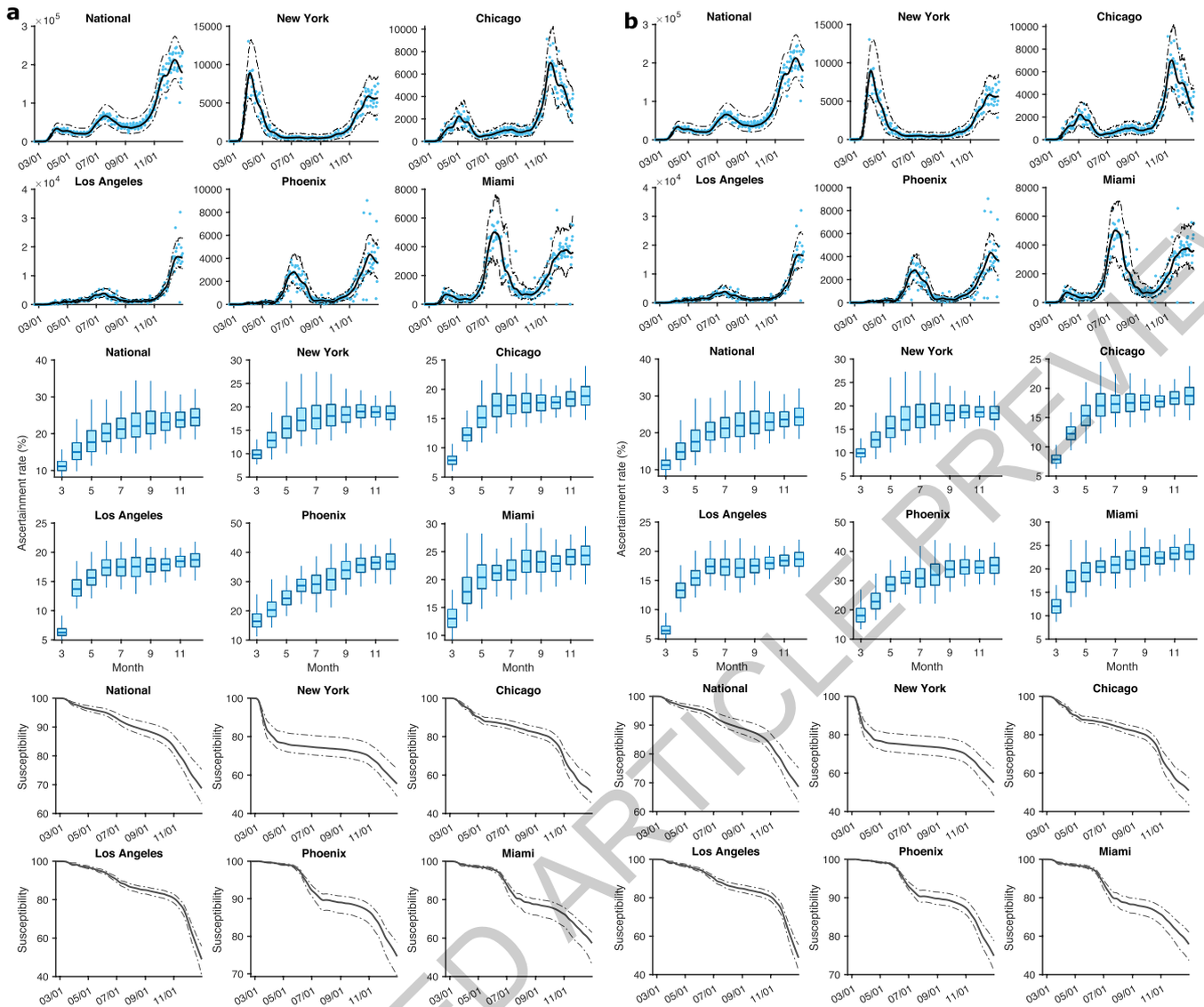
Extended Data Fig. 1 | Parameter inference for simulated outbreaks. Results are shown for three major metropolitan areas – New York, Chicago, and Los Angeles. Outbreaks were generated for 60 days using four prescribed scenarios. National daily cases are shown in the top row. Parameter estimates

for the last 10 days are not displayed as there is not enough data at the end of the time series to constrain the model. Solid and dashed lines show the median estimate and 95% CIs respectively.



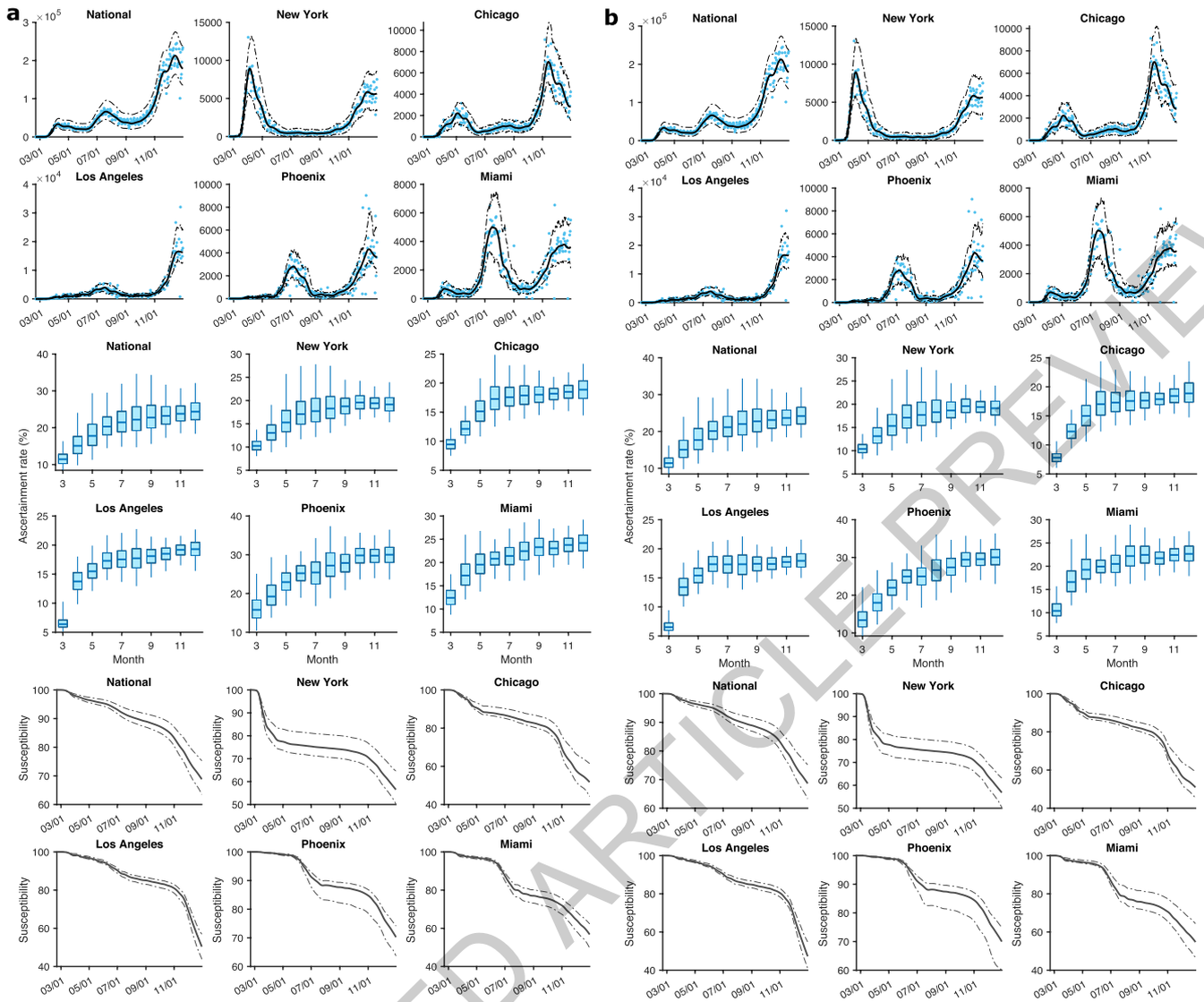
Extended Data Fig. 2 | Model fitting and inference results. (a) Model fitting to daily case numbers (blue dots) in the US and five metropolitan areas. Solid and dashed lines show the median estimate and 95% CIs respectively. (b) Estimated daily ascertainment rates (left column) and transmission rates (right column) for five metropolitan areas. Solid and dashed lines show the median estimate and 95% CIs respectively. (c) Reliability plot for model

calibration. Data points show the coverage of the 25%, 50%, 75% and 95% CIs of the posterior fitting at county and national levels. (d) The estimated national cumulative IFR in 2020. The cumulative IFR is computed using the estimated cumulative numbers of death (deconvolved) and infections prior to a given date.



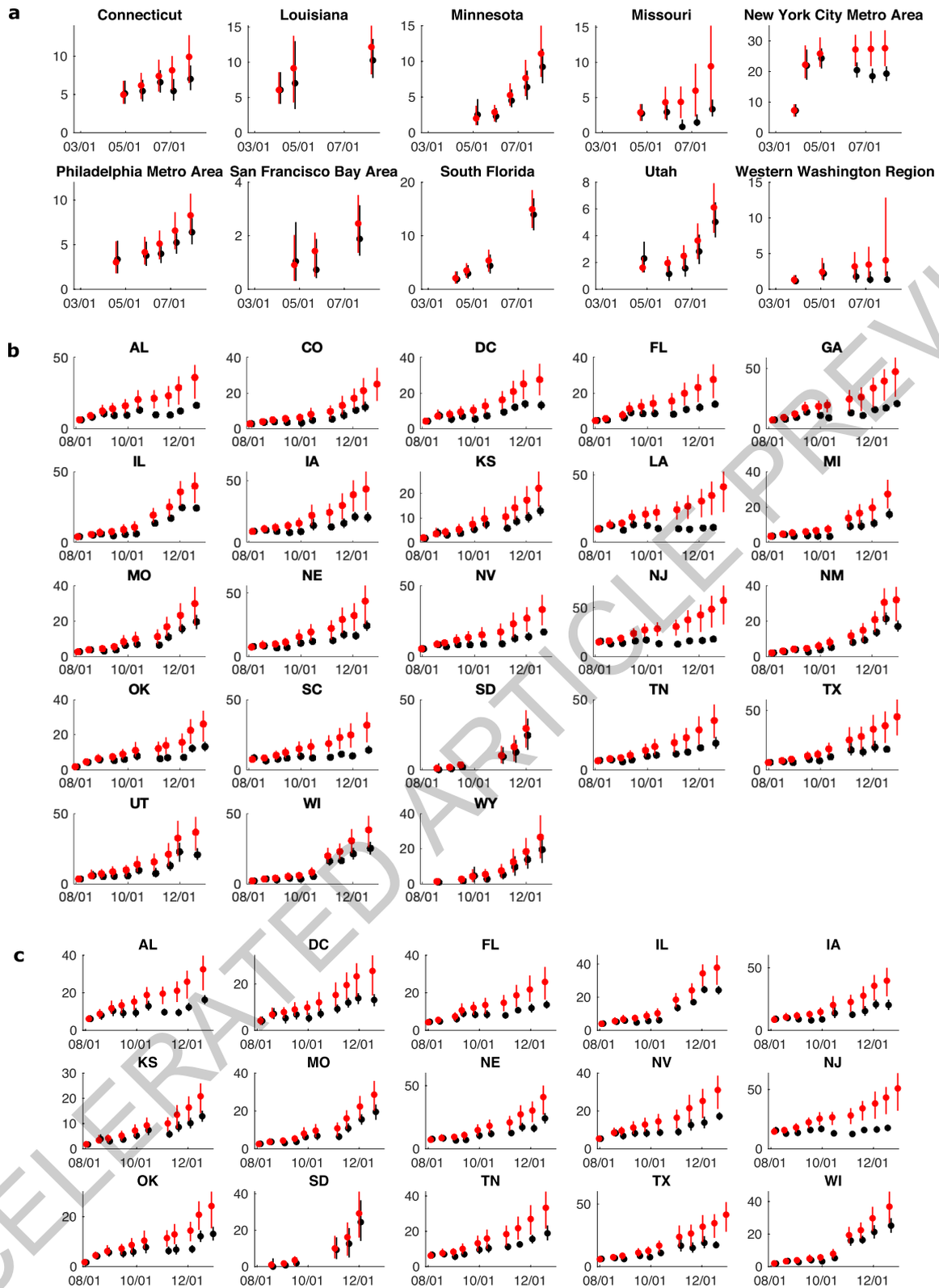
Extended Data Fig. 3 | Sensitivity analyses on inference results. (a) Inference results using fixed parameters (Z, D, μ, θ) estimated from case data prior to April 2 2020. (b) Inference results from a modified version of the transmission model in which the relative infectiousness of undocumented infections, μ , is allowed to vary over time. Fitting to case data (top two rows), estimated monthly ascertainment rate (middle two rows) and population susceptibility

(bottom two rows) are shown. Distributions are obtained from $n=100$ ensemble members. In the top two and bottom two rows, the solid line represents median, and the dash lines show 95% CIs. In the middle two rows, centre and box bounds represent median, 25%, and 75% percentiles, and whiskers show 2.5% and 97.5% percentiles.



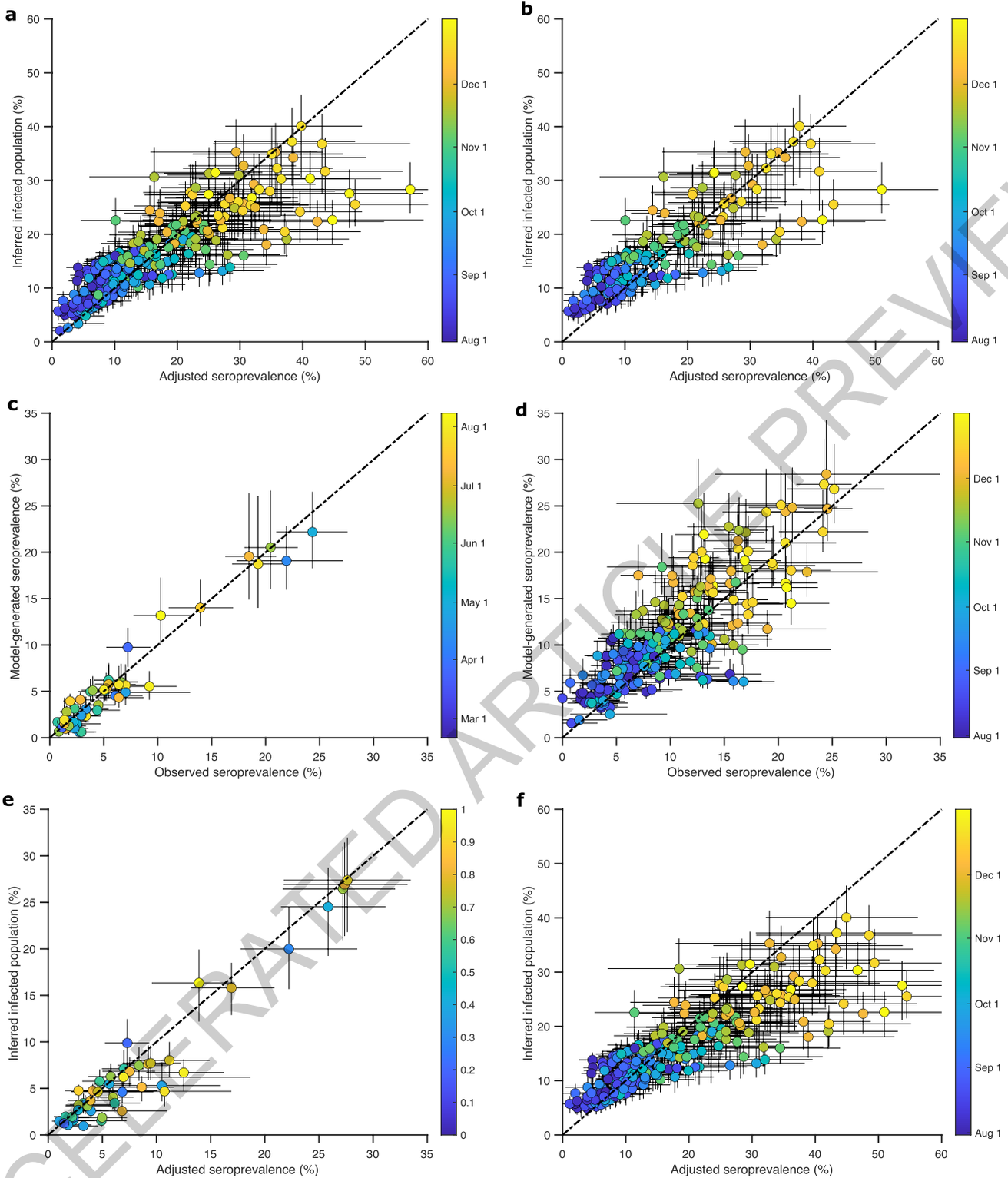
Extended Data Fig. 4 | Inference results from a modified version of the transmission model permitting movement of documented infections among counties. (a) 25% of documented infections are allowed to move among counties. (b) 50% of documented infections are allowed to move among counties. Fitting to case data (top two rows), estimated monthly ascertainment rate (middle two rows) and population susceptibility (bottom two rows) are

shown. Distributions are obtained from $n=100$ ensemble members. In the top two and bottom two rows, the solid line represents median, and the dash lines show 95% CIs. In the middle two rows, the centre and box bounds represent median, 25%, and 75% percentiles, and the whiskers show 2.5% and 97.5% percentiles.



Extended Data Fig. 5 | The reported (black) and adjusted (red) seroprevalence. (a) Results for the 10-site study. (b) and (c) show the results for state-level serological surveys obtained using a maximum monthly attenuation

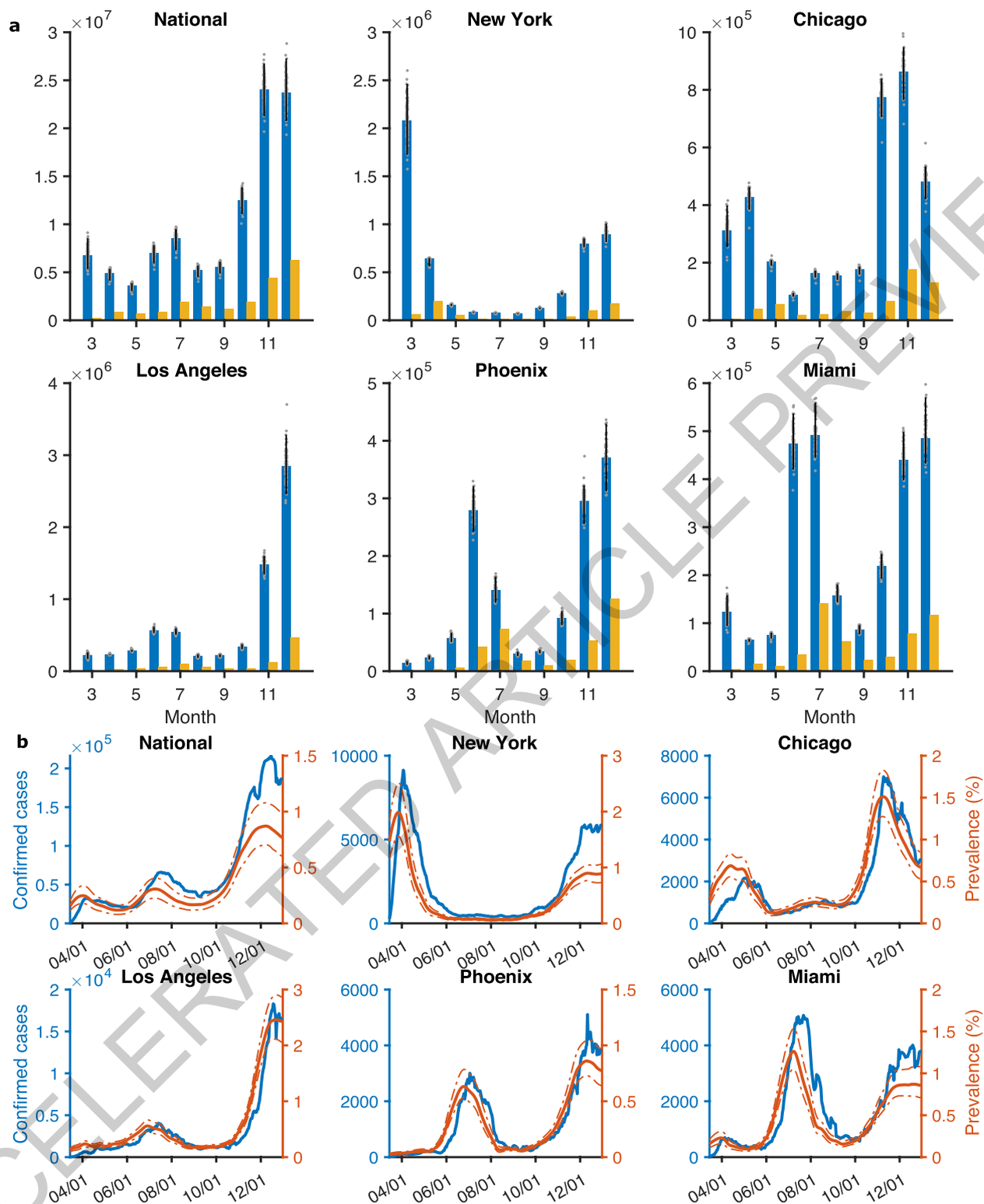
rate of 17.5% and 15%, respectively. Dots and whiskers show the median and 95% CIs respectively. Distributions are obtained from $n=1,000$ simulated seroprevalence samples.



Extended Data Fig. 6 | Validation of inference using seroprevalence data.

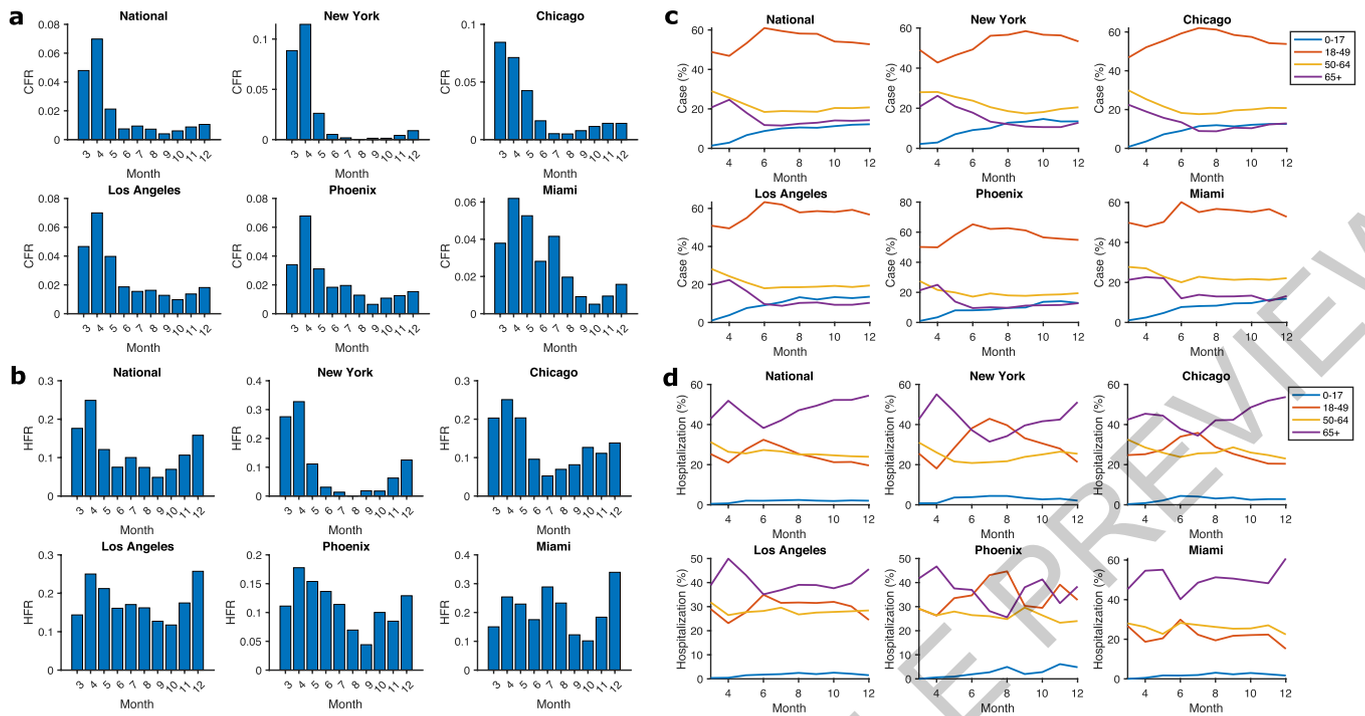
(a) – (b) Comparison between the inferred percentage of cumulative infections and seroprevalence at the state level adjusted for antibody waning. Seroprevalence data adjusted using a maximum monthly attenuation rate of 17.5% (a) and 15% (b) are included in the analysis. (c) – (d) Comparison between the model-generated seroprevalence and observed seroprevalence in 10

locations (c) and at the state level (d). (e) – (f) Comparison between the inferred percentage of cumulative infections and seroprevalence in 10 locations (e) and at the state level (f) adjusted for antibody waning using lower sensitivity and specificity. Distributions are obtained from $n=100$ ensemble members. Centre and whiskers show median and 95% CIs. Color indicates the sample collection date for each location.



Extended Data Fig. 7 | Inference results in the US and five metropolitan areas. (a) Estimated monthly total infections (blue bars) and confirmed cases (orange bars) in the US and five metropolitan areas. Distributions are obtained from $n=100$ ensemble members. The blue bars show medians and whiskers

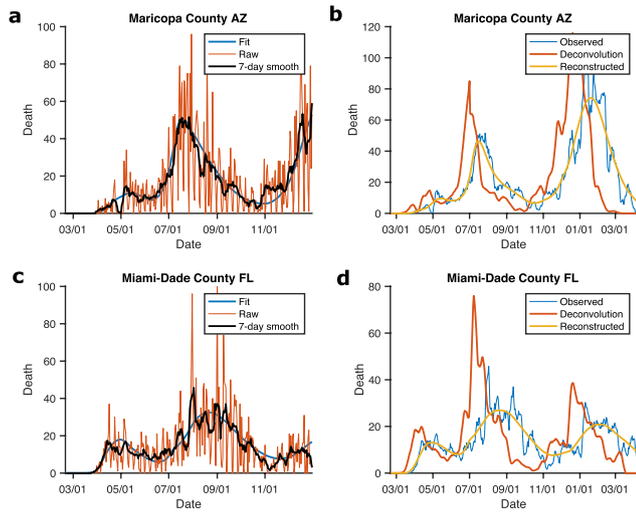
show 95% CIs. (b) Daily confirmed cases (blue line, 7-day moving average) and estimated prevalence of contagious infections (red line, median and 95% CIs) in the US and five metropolitan areas.



Extended Data Fig. 8 | Key statistics obtained from line-list data for the US and five metropolitan areas. (a) – (b) The crude monthly CFR (a) and HFR (b) obtained from line-list data for the US and five metropolitan areas. Note that due to incomplete reporting of deaths in the line-list data, these estimates are

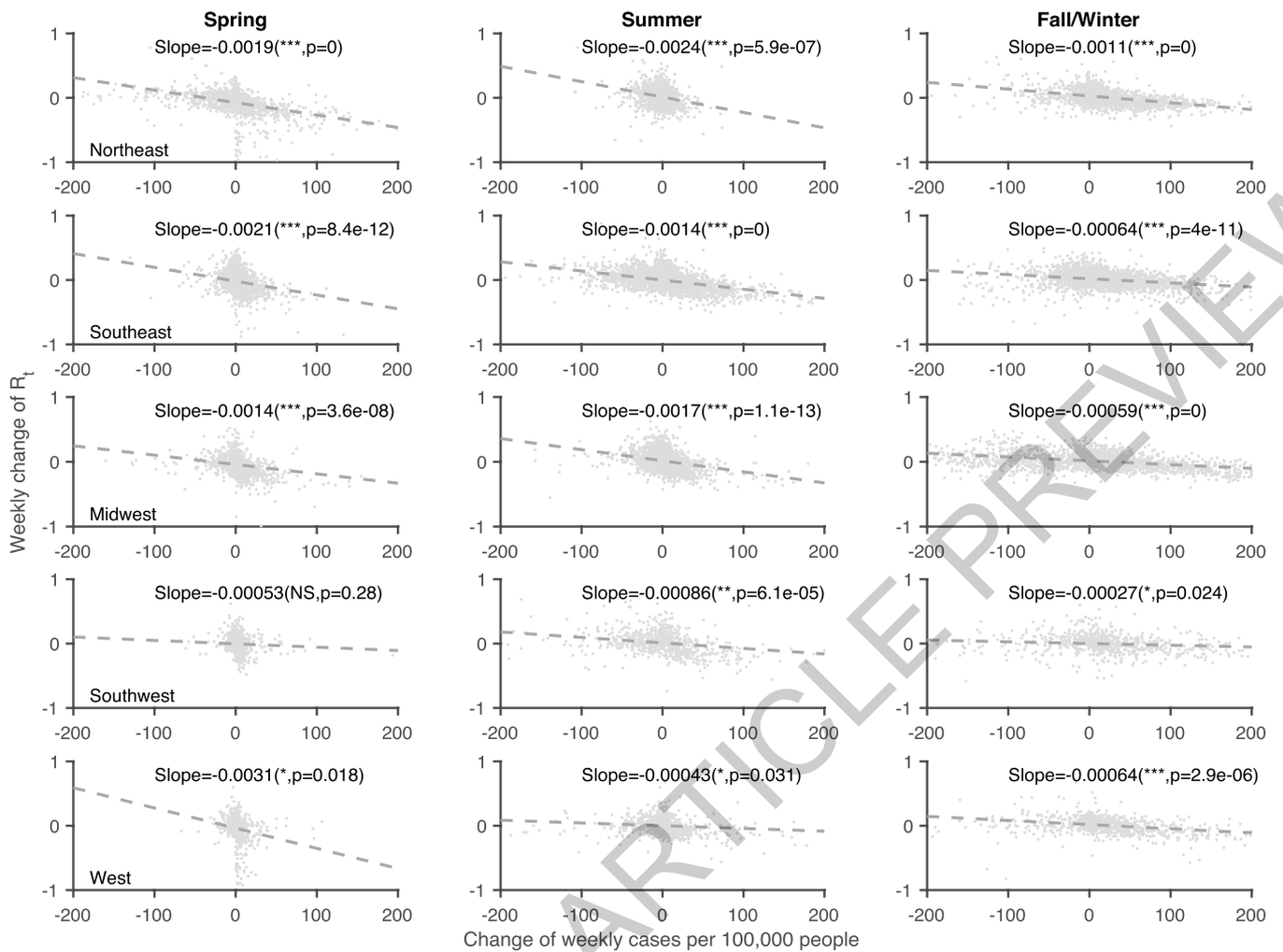
likely low. (c) – (d) The proportion of confirmed cases (c) and hospitalizations (d) in four age groups (0-17, 18-49, 50-64, 65+) in the line-list data. Data are shown monthly for the US and five metropolitan areas.

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Extended Data Fig. 9 | Estimation of the time-to-event distribution from case confirmation to death for Maricopa County AZ (a) and Miami-Dade County FL (c). Deconvolution of daily deaths using the estimated delay distributions for Maricopa County AZ (b) and Miami-Dade County FL (d).

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Extended Data Fig. 10 | Weekly change of R_t in response to the change of weekly cases per 100,000 people at county level. The analysis was performed for five US regions (Northeast, Southeast, Midwest, Southwest, West) during the spring (Feb 21 – May 31), summer (Jun 1 – Sep 15), and fall/winter (Sep 16 – Dec 31) waves. In the five US regions, 116, 162, 126, 45 and 54 counties that reported cumulative cases over 100 per 100,000 people during

all three waves and had a population over 100,000 were included in the analysis. A positive/negative change of weekly cases in the x-axis indicates increasing/decreasing community prevalence of COVID-19. The dash lines are the linear fits. The statistical significance of the slope is indicated by asterisks (two-sided t-test. ***: $p < 10^{-5}$, **: $p < 0.001$, *: $p < 0.05$; NS: not significant. P-values are reported in the legends).

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Our web collection on [statistics for biologists](#) contains articles on many of the points above.

Software and code

Policy information about [availability of computer code](#)

Data collection No software was used for data collection.

Data analysis We implemented the EAKF algorithm in MATLAB R2021a. Data analysis was performed using MATLAB R2021a. The custom codes are publicly posted at GitHub: https://github.com/SenPei-CU/COVID_US_2020.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio [guidelines for submitting code & software](#) for further information.

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- Accession codes, unique identifiers, or web links for publicly available datasets
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The human mobility and COVID-19 surveillance data that support the findings of this study are available at GitHub (https://github.com/SenPei-CU/COVID_US_2020). The county-level COVID-19 surveillance data in the US are available at Johns Hopkins University coronavirus resource center (https://github.com/CSSEGISandData/COVID-19/tree/master/csse_covid_19_data/csse_covid_19_time_series). County-to-county commuting data were downloaded from the US Census Bureau (<https://www.census.gov/data/tables/2015/demo/metro-micro/commuting-flows-2015.html>). Human mobility data in 2020 were provided by SafeGraph (<https://safegraph.com/>), a data company that aggregates anonymized location data from numerous applications in order to provide insights about physical places, via the Placekey Community (<https://placekey.io/>). To enhance privacy, SafeGraph excludes census block group information if fewer than five devices visited an

establishment in a month from a given census block group. We aggregated the mobility data to county level to estimate change of inter-county mobility in 2020. Aggregated and derived data are allowed to be shared publicly by SafeGraph. Seroprevalence data were published by the US CDC (<https://www.cdc.gov/coronavirus/2019-ncov/cases-updates/commercial-lab-surveys.html>). The line-list datasets are available at the US CDC website (<https://data.cdc.gov/Case-Surveillance/COVID-19-Case-Surveillance-Public-Use-Data/vbim-akqf> and <https://data.cdc.gov/Case-Surveillance/COVID-19-Case-Surveillance-Public-Use-Data-with-Ge/n8mc-b4w4>).

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Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

Life sciences Behavioural & social sciences Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see [nature.com/documents/nr-reporting-summary-flat.pdf](https://www.nature.com/documents/nr-reporting-summary-flat.pdf)

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size	We did not collect data from samples. In the EAKF algorithm, we used n=100 ensemble members to represent the distributions of model states, parameters, and inference outcomes.
Data exclusions	No data were excluded.
Replication	This is a modeling study without experiments. We repeated the inference multiple times and the inference results are reproducible and robust.
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