Spatial and temporal dynamics of superspreading events in the 2014–2015 West Africa Ebola epidemic

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The unprecedented scale of the Ebola outbreak in Western Africa (2014–2015) has prompted an explosion of efforts to understand the transmission dynamics of the virus and to analyze the performance of possible containment strategies. Models have focused primarily on the reproductive numbers of the disease that represent the average number of secondary infections produced by a random infectious individual. However, these population-level estimates may not capture important systematic variation in the number of cases generated by infected individuals, particularly found in spatially localized transmission and superspreading events. Although superspreading features prominently in first-hand narratives of Ebola transmission, its dynamics have not been systematically characterized, hindering refinements of future epidemic predictions and explorations of targeted interventions. We used Bayesian model inference to integrate individual-level spatial information with other epidemiological data of community-based (undetected within clinical-care systems) cases and to explicitly infer distribution of the cases generated by each infected individual. Our results show that superspreaders play a key role in sustaining onward transmission of the epidemic, and they are responsible for a significant proportion (~61%) of the infections. Our results also suggest age as a key demographic predictor for superspreading. We also show that community-based cases may have progressed more rapidly than those notified within clinical-care systems, and most transmission events occurred in a relatively short distance (with median value of 2.51 km). Our results stress the importance of characterizing superspreading of Ebola, enhance our current understanding of its spatiotemporal dynamics, and highlight the potential importance of targeted control measures.

Significance

For many infections, some infected individuals transmit disproportionately more susceptibles than others, a phenomenon referred to as “superspreading.” Understanding superspreading can facilitate devising individually targeted control measures, which may outperform population-level measures. Superspreading has been described for a recent Ebola virus (EBOV) outbreak, but systematic characterizations of its spatiotemporal dynamics are still lacking. We introduce a statistical framework that allows us to identify core characteristics of EBOV superspreading. We find that the epidemic was largely driven and sustained by superspreading that were ubiquitous throughout the outbreak and that age is an important demographic predictor for superspreading. Our results highlight the importance of control measures targeted at potential superspreaders and enhance understanding of causes and consequences of superspreading for EBOV.


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study localized transmissions of EBOV and complement formal surveillance by detecting cases that did not interface with clinical care. In this work, we built an age-specific spatiotemporal framework, which allowed us to explicitly infer the probability distribution of the number of new cases generated by each infected individual (hereafter, offspring distribution). This framework was applied to the community-based EBOV case dataset and deployed to infer transmission dynamics and identify superspreaders. Specifically, we used Bayesian inferential techniques to synthesize individual-level spatial data (i.e., GPS coordinates), age data, symptoms onset time, and burial time (Study Data), and to impute unobserved infection time and transmission network (Materials and Methods and SI Text).

**Study Data**

We analyzed a community-based dataset collected from the Safe and Dignified Burials program conducted by the International Federation of Red Cross, between October 20, 2014, and March 30, 2015, in Western Area (which comprises the capital Freetown and its surrounding area) in Sierra Leone. These data contain GPS locations (collected by mobile phones) of where the bodies of 200 dead who tested positive for Ebola were collected (typically at their homes). Age, sex, time of burial (which was usually performed within 24 h of death), and symptom onset time were also recorded. Symptom onset time was reported retrospectively by next of kin.

**Results**

**Natural History Parameters.** We estimated that $R_0$ has median value 2.39, with 95% credible interval (C.I.) of [2.05, 2.84] (Fig. 1A). We also estimated the time-varying reproductive number $R_t$ (Fig. 1B). The incubation period was estimated to be 6.74 d [1.29, 16.21]. These estimates are broadly consistent with what have been reported (3, 12).

The mean of infectious period (i.e., duration from symptoms onset to death/burial) was estimated to be 3.9 d [3.75, 4.0]. Because the transmission tree and times of infection were imputed (Materials and Methods), we were also able to infer the mean generation time of EBOV, which was estimated to be 10.9 d [9.25, 13.01]. Both estimates were lower than that estimated from cases detected within the clinical care system [e.g., mean infectious period 8 d estimated for patients who received clinical care (13) and mean generation time 15.3 d estimated by the WHO (1)]. These discrepancies potentially highlight systematic differences between community-based cases and cases notified in clinical care systems, with terminal community-based cases progressing significantly more rapidly.

![Fig. 1](image1.png)\[\text{Fig. 1.} \] Estimates of reproductive number. (A) Posterior distribution of the basic reproductive number, $R_0$. (B) Posterior distribution of the weekly effective reproductive number, $R_t$. Bars represent 95% C.I., and red line connects the medians.

![Fig. 2](image2.png)\[\text{Fig. 2.} \] (A) Spatial distribution of mean number of offspring resulting from initial cases at the individual level. An infection is classified as an index case if it has a posterior probability of importation (i.e., not infected by any cases in the data) $>0.5$; otherwise, it is classified as a secondary case. Lat, latitude; Lon, longitude. (B) Distribution of mean number of offspring by different sources of infection. (C) Proportion of infected individuals who are direct and indirect descendants of the first five superspreaders (i.e., first five individuals with highest number of mean offspring; note that the choice of five is arbitrary here). “Any” includes superspreaders who were also the index cases (i.e., the roots of transmission trees).
Superspreading in Space and Time. Fig. 2A and B show a clear asymmetry in the average number of “offspring” at the individual level, quantifying the impact of superspreading. In particular, it was observed that most secondary cases generated less than one offspring on average. Thus, the epidemic growth appeared to be fueled mostly by only a few superspreaders (i.e., the outliers in the boxplot). A common empirical measure of degree-of-transmission heterogeneity and superspreading is the dispersion parameter \( k \), assuming that the offspring distribution is a negative binomial with variance \( \sigma^2 = \mu (1 + \mu/k) \), where \( \mu \) is the mean (9). Generally speaking, a lower \( k \) represents a higher degree of transmission heterogeneity and superspreading; and \( k < 1 \) implies substantial superspreading (compared with a geometric distribution, for which \( k = 1 \)). Our empirical estimate of \( k \) of our inferred mean offspring distribution (including index and secondary) was 0.37, and it is higher (i.e., implies less heterogeneity) than an estimate from an observational study in which \( k \) was estimated to be 0.16 (10, 11). This discrepancy in the estimate of \( k \) suggests that our estimate of the degree of superspreading may be conservative (Sensitivity Analysis), although it should be noted their estimate was made based on a study in a different geographical region and time frame. By sampling probabilistically consistent transmission networks among infected individuals (Materials and Methods), we were able to identify whether a case was a descendant of superspreaders by performing a backward search of sampled transmission tree from the case—for each case, we first identified its (most recent) direct infector (\( IF_{1} \)) from the sampled tree, from where we could subsequently identify the infector of \( IF_{1} \); We continued this backward searching until we reached an index case [i.e., the root of a (sub)tree]; a superspreader is an ancestor of this case if it happens to be one of the infectors during the backward searching. Fig. 2C shows that a few superspreaders (~3% of all of the cases) were responsible, either directly or indirectly, for a substantial proportion (with median 61%) of all of the cases generated, highlighting the key role of these superspreaders in driving the epidemic growth—had the superspreaders been identified and quarantined promptly, a majority of the infections could have been prevented.

In Fig. 3A, we show the time dependence of superspreading, illustrating that superspreading becomes relatively more important over time (i.e., within ~100 d after the epidemic peak). This figure suggests that, after the initial period of fast growth of the epidemic (i.e., time before peak), superspreaders may be crucial to sustaining and fueling epidemic growth and also prolonging the epidemic duration. Near the end of the epidemic (period 5 in Fig. 3A), most cases did not spread, and superspreading was non-significant, as reflected by \( k > 1 \). Fig. 3B shows that most of the transmission (including superspreading) occurred over relatively short distances (median 2.51 km), indicating that transmission tends to take place at the local community level.

Heterogeneity of Infectiousness by Age. Although superspreading in EBOV was evident and may be partly attributed to unsafe burial practice during the early stage of the outbreak (14), other drivers (e.g., social contact pattern) of this process remain unclear. In Fig. 4A, as expected, the infectious period had a clear positive relationship with mean offspring number. Despite the clear relationship between infectious period and the magnitude of superspreading, this covariate cannot be used as a predictor of superspreading, because it is not known a priori. More importantly, there is a significant difference in instantaneous infectious hazard exerted by different age groups (Fig. 4B)—cases <15 and >45 appear to have higher instantaneous transmissibility. Our

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Fig. 3. Spatial and temporal dependence of superspreading. (A) Reported weekly deaths and inferred mean offspring distributions and the corresponding empirical estimates of \( k \) at different time periods. The whole time period is divided into five periods—that is, period 1, from the time of first observation to the time of epidemic peak \( t_{peak} \); Period 2, \( t_{peak} + 20 \); period 3, \( t_{peak} + 20 \); period 4, \( t_{peak} + 50 \); and period 5, \( t_{peak} + 100 \), from \( t_{peak} + 100 \) to the time of last observation. Such a dividing was used so that we could use the peak time as a reference point and ensure a similar number of cases in most intervals. (B) Distribution of distance of transmission for all infected-infected pairs. Black dotted line represents the median (2.51 km) of the distribution. Red dotted line represents the median (2.61 km) of the subdistribution in which the infectors are superspreaders (defined as those who have mean offspring more than five here).
Missing Community Cases Missing Hospitalizations

Fig. 4. Heterogeneity of infectiousness in age. (A) Relation between mean offspring and infectious period. It is worth noting that here an infectious period is strictly referred to the mean of the posterior samples of imputed infectious period of an individual, rather than the assumed universal infectious period distribution. (B) Instantaneous risk exerted by different age groups.

results suggest that the combination of certain age groups (who have high instantaneous hazard) with a long infectious period (at the right tail of the infectious period distribution) constitutes a key driver of superspreading. The discrepancy of transmissibility in age may be rooted in social contact structure (15) or virological linkages (e.g., potential systematic variation among infected individuals) that cannot be established solely by using epidemiological data (16).

Sensitivity Analysis. Underreporting is a ubiquitous feature of epidemiological data (17, 18). In this section, we explore the effect of underreporting on our analysis under two probable scenarios: (i) All unreported cases were circulating in the community and not hospitalized; and (ii) all unreported cases were hospitalized and therefore not reported in our database. In both scenarios, we tested with constant underreporting rates, across the whole study period and region, ranging from a very low (10%) to a very high one (90%). Doing so allowed us to investigate the probable lower and upper bound of our estimates. We also tested with time-varying underreporting rates in both scenarios. Details of how to include underreported cases are provided in Materials and Methods.

We focused on investigating the effect on $k$, $R_0$ and transmission distance. Fig. 5A shows that, in general, superspread-

Fig. 5. Effect of constant underreporting rates on estimates of transmission dynamics. (A) Estimates of $k$. Bars represent the 95% C.L. and dots represent the median values. (B) Estimates of $R_0$. (C) Estimates of most probable distance of transmission. (D) Estimates of median transmission distance. Dotted lines represent the corresponding estimates using our data. At each underreporting rate, 10 independent simulations and corresponding inference were performed (Materials and Methods).
showing very similar estimates as the baseline case (Tables S1 and S2). We also tested alternative parameterization of priors in Table S3, giving virtually identical results compared with those obtained in the baseline case (see also Materials and Methods).

Discussion

Superspreading is a core process for the transmission of many infections (7, 8). However, the importance of superspreading in driving epidemics varies with context. For instance, its impact depends on how it persists over the course of an epidemic. Quantifying superspreading and identifying scenarios where it is more likely to occur can facilitate refining future epidemics predictions and help in devising targeted intervention strategies that may outperform population-level control measures (9). To date, a systematic understanding of how EBOV has been (super)spreading in the recent outbreak in Western Africa is lacking, particularly in terms of individual-level covariates, and across the spatiotemporal setting. The key contributions of this work are to highlight and quantify the importance of superspreading and to show that it is in some senses systematic.

Community-based surveillance data offer a valuable opportunity to study superspreading, by focusing on nonhospitalized cases that may have been involved in superspreading events and not detected by formal surveillance. Here, we introduce a continuous-time spatiotemporal model that integrates individual spatial information with other epidemiological information of community-based cases and deploy it to quantify superspreading and its drivers for EBOV. Our framework enabled us to sample likely realizations of the unobserved transmission network among cases from which the offspring distribution of each case could be inferred, providing explicitly a machinery for understanding superspreading in space and time.

Our analysis is broadly consistent with previous work, indicating values of $R_0$ of 2.39 [2.05, 2.84] for the outbreak in Sierra Leone (in particular, close to the 2.53 estimated in ref. 22). Our results show that EBOV exhibited a prominent superspreading pattern shared by SARS and MERS (7, 8, 23) [e.g., $k$ was estimated to be 0.16 for SARS (9)], which reinforces the finding that superspreading occurred during the recent EBOV outbreak (10).

We also extended previous analyses by showing that a substantial proportion of secondary cases were either direct or indirect descendants of a small number of superspreaders, underscoring the importance of superspreading in driving the epidemic—that is, had the superspreaders been identified and quarantined promptly, ~61% of the infections could have been prevented. Furthermore, we show that superspreaders may have particular importance in driving and sustaining the epidemic progression over the course of the outbreak. The increasing relative importance of superspreading over the later stages of the outbreak (Fig. 3A) is consistent with the rising availability of hospital beds (5)—that is, in the outbreak, most infected individuals were able to get a bed at an Ebola treatment center (ETC) and largely did not further transmit; as a result, those superspreaders in the community who did not make it to ETCs may have played an important role in (super)spreading EBOV in the local community.

Specifically, our results identify age groups that have higher instantaneous transmissibility and show that cases in the more infectious age groups tend to be superspreaders when combined with a relatively long infectious duration. One plausible explanation, from the social perspective, may be that the young and old are much more likely to have (and infect) lots of visitors, compared to other age groups; a parallel corollary is that the young and old might be more likely to have others caring for them. Also, our results highlight significant differences between community-based cases and cases notified in the clinical care system, with terminal community-based cases progressing significantly more rapidly. Our results stress the importance of characterizing superspreading of EBOV, enhance current understandings of its spatiotemporal dynamics, and highlight the potential importance of targeted control measures—for example, during the 2014–2015 EBOV epidemic, millions of dollars were spent implementing message strategies about Ebola prevention and control across entire countries; our results suggest that message strategies targeting individuals with higher risk may be useful to prevent superspreading events and the persistence of the outbreak.

There are limitations of our results. First of all, although community-based surveillance data complement formal surveillance by detecting cases that did not interface with clinical care, they contain only partial information about the epidemic, with hospitalized cases omitted. Also, it is possible that, by underreporting some community cases who generated subsequent cases, certain reported cases may be falsely attributed as sources of infection for those subsequent cases, overestimating the degree of superspreading. Accordingly, our sensitivity analysis evaluated the impact of these sources of underreporting, showing that our estimated degree of superspreading may in fact be conservative and represents a lower bound—superspreading in EBOV may be even more prominent in reality (Fig. 5). It is also worth noting that, by considering only safe burials, which tend to be less transmissible (relative to those who did not receive safe burials) among deaths (14), our estimate of superspreading may have been conservative. Conversely, because it was reported that individuals who eventually died might have a higher intrinsic transmissibility (24), our analysis might bias toward high transmitters by only using death data. Our methodology represents a transmission network-based approach that focused on constructing transmission trees among cases (25–28). Although such an approach captures contacts that caused infections, it does not account for “unsuccessful” contacts that correspond to escaped infections. Future theoretical work will need to include such contacts. Nevertheless, because unsuccessful contacts are not part of the transmission chain, ignoring them has limited effect on the transmission tree or on many overall topological characteristics (e.g., average number of offspring of an infected case) (25, 28, 29). Finally, although our analysis reveals the importance of age as demographic determinants of superspreading, future work in linking them with virological factors (e.g., age-specific viral loads) may shed further light (16).

Materials and Methods

Spatiotemporal Transmission Model. We developed a continuous-time spatiotemporal transmission model that allowed us to sample the transmission tree among cases, integrating observed spatial and temporal individual data. This approach allowed us to infer explicitly the mean offspring distribution of each case. Specifically, the total probability of becoming infected during time period $[t, t + dt]$ was given by

$$r(t, t + dt) = \left\{ \alpha + \sum_{j \in \xi(t)} \beta_j \times X(\mathbf{k} \cdot \mathbf{r}_j) \right\} dt + o(dt),$$

where $\xi(t)$ is the set of all infectious individuals at time $t$, $\alpha$ is the background rate of infection, and $\beta_j$ is the age-specific instantaneous infection hazard of a case in $\xi(t)$. We allowed five-level $\beta_j$ according to the age—that is, we had $\beta_0 = \beta_a$ for age between $[0, 15]$, $\beta_i$ for age between $[15, 30]$, $\beta_i$ for age between $[30, 45]$, $\beta_i$ for age between $[45, 60]$, and $\beta_i$ for age
Data Augmentation and Model Fit and Validation. We estimated θ (i.e., the parameter vector) in the Bayesian framework by sampling it from the posterior distribution \( P(\theta|x) \), where \( x \) is the observed data. Denoting the likelihood by \( L(\theta, x) \), the posterior distribution of \( \theta \) is \( P(\theta|x) \propto L(\theta; x) \), where \( P(\theta) \) is prior distribution for \( \theta \). Weak uniform priors for parameters in \( \theta \) were used (Table S4). Markov chain Monte Carlo (MCMC) techniques (30) were used to obtain the posterior distribution. The unobserved infection times and transmission network were imputed in the MCMC. Sampled transmission networks were recorded and used to infer the offspring distribution of each case. Details of the likelihood function and the MCMC algorithm are given in SI Text. Model fit was assessed by comparing the observed data with those simulated from the estimated model, suggesting a good fit (Fig. S4). Furthermore, for validating the implementation of our inference procedures, we generated multiple sets of pseudodata from the model and demonstrated that we could successfully reestimate the model parameters (Fig. S5).

Testing Underreporting. We divided the observational period into many 3-d-wide intervals. Within each time interval, we had the total number of unreported cases \( n_i^r = n_i - (1 - r)n_i \), where \( n_i \) and \( r \) were the observed cases in the interval and the assumed underreporting rate, respectively. Burial times and symptoms-onset time of these unreported cases were drawn from the empirical distribution of the observed cases. Finally, these \( n_i^r \) cases were distributed spatially by using the empirical distribution of (normalized) population densities across the study area. We also tested an underreporting rate that decreases with time (Fig. S1). For the scenario that considers unreported hospitalized cases, we drew the time from symptoms onset to hospitalization from the truncated above (at 7 d) empirical infectious period distribution of observed cases, effectively resulting in a shorter infectious period for unreported cases. These artificially generated data were combined with the observed data and fitted with our model.

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Supporting Information

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SI Text

Likelihood Function. Let \( E = (E_1, E_2, \ldots, E_n) \) be the vector of the exposure/infection times of the \( n = 200 \) cases, \( I = (I_1, I_2, \ldots, I_n) \) the times of becoming infectious, and \( R = (R_1, R_2, \ldots, R_n) \) the death times. The epidemic was observed up to time \( t_{\text{max}} \).

The incubation period was assumed to be a two-parameter density function \( f_{\omega}(t; a, b) \) characterized by parameters \( a \) and \( b \); similarly, for the infectious period (i.e., time from start of infectiousness to death), with density function \( f_{\alpha}(t; c) \). Finally, let \( \psi_j \) be the source of infection of case \( j \) and \( \psi \) be the collection set for \( n \) cases. The likelihood of the parameter vector \( \theta = (\alpha, \beta, \eta, a, b, c) \) given complete data can be expressed as

\[
L(\theta; E, I, R, \psi) = \prod_j P(j, \psi_j) \times Q(E_j)
\]

where

\[
P(j, \psi_j) = \begin{cases} 
\alpha, & \text{if } j \text{ is an index case,} \\
\beta \psi_j K(d_{\psi_j}; \eta), & \text{if } j \text{ infected by a case } \psi_j, 
\end{cases}
\]

is the (unnormalized) probability of case \( j \) to be an index case of infected by case \( \psi_j \), respectively, and

\[
Q(E_j) = \exp \left( -\int_0^{E_j} \{\alpha + \sum_{i \in \xi_j(t)} \beta K(d_{ij}; \eta)\} dt \right).
\]

MCMC Algorithm. Parameters in \( \theta \) were updated sequentially with a standard random-walk Metropolis–Hastings (M-H) algorithm (30, 31). For example, a new parameter value \( \alpha' \) was proposed from a normal distribution centered on the current value of \( \alpha \), that is,

\[
\alpha' = \alpha + N(0, \rho^2)
\]

where \( \rho \) controls the step size of the random-walk. Elements in infection times vector \( E \) were also treated as unobserved model parameters and were imputed in the same manner (30). Approximately 10% of the cases had invalid records of symptom onset time; hence, corresponding elements in \( I \) were also imputed similarly. We used (weak) uniform priors with upper bounds for all model parameters, and the maximum of the incubation period was assumed to be 21 d (32). Details of the priors and obtained posteriors are shown in Table S4.

Denote \( \omega_0 \) as the set of eligible candidates for a new source of infection \( \psi'_j \) for \( j \) (i.e., \( \omega_0 \) contains a set of cases whose are infectious at \( E_j \)). We propose a new infecting source \( i \in \omega_0 \) to be \( \psi'_j \) with probability

\[
p_{ij} \propto \beta K(d_{ij}; \eta).
\]

Note that the background infection can be accommodated by adding a permanent infectious source presenting an additional challenge of strength \( \alpha \) to individual \( j \). A newly proposed source is accepted or rejected depending on the M-H acceptance probability (29).

Fig. S1. Effect of time-varying underreporting on estimates of transmission dynamics. (A) Estimates of \( k \). Bars represent the 95% C.I., and dots represent the median values. (B) Estimates of \( R_0 \). (C) Estimates of most probable distance of transmission. (D) Estimates of median transmission distance. Dotted lines represent the corresponding estimates using our data. The underreporting rate is assumed to decrease with a step size 10%, from 90 to 10%, in the course of the epidemic: The study period is divided into nine equal intervals, and each interval takes an underreporting rate that is 10% lower than the previous one.

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Testing the assumption of an isotropic spatial dispersal. (A) The distribution of mean offsprings under different scenarios. (B) The distribution of $R_0$ under different scenarios. (C) The distribution of transmission distance under different scenarios. Here we considered three scenarios. In scenario 1 (base scenario), we assumed an isotropic dispersal and did not take into account the potential effect of population density. We considered in scenarios 2 and 3 that the dispersal kernel value was “moderated” by the relative population density of the 100 m × 100 m grid that a case resides in. Scenarios 2 and 3 differ in how the population density was normalized (to between [0, 1]) to obtain the discounting factor: In scenario 2, we normalized according to $\log(1 + \text{population density})$, and in scenario 3, we normalized according to the absolute scale of population density.
Fig. S3. Population density and spatial distribution of the cases in the study area. Other than the smaller clusters near the center of the study area, most cases were found in more populated regions. It was noted that the raw grid resolution is 100 m × 100 m (which is too fine to display), and here it is binned into 30 × 30 grids for better visualization. Lat, latitude; Lon, longitude.
Fig. S4. Assessing the model fit. We used the estimated model to simulate (500 times) forward the transmission path and timings of events (i.e., infection time, onset time, and death time). (A) Comparison of the observed weekly temporal distribution of the cases with that summarized from the simulated data. Gray area represents the 95% C.I., and the black dots and line are the observed data, with 5 of 500 random realizations (colored lines) of the simulated epidemics imposed. We compared the temporal autocorrelations (at lag = 1 and lag = 2) of the observed and simulated epidemics. We also compared the peak height, the growth rate before peak, and decay rate after peak between the observed and simulated (the growth and decay rates correspond to the slopes of best-fitted linear lines to the observed or simulated data). Dotted lines represent the values of the summary statistics corresponding to the observed data. (B) Comparison of the observed spatial autocorrelation and the simulated. Here we used two common measures, Moran’s I and Geary’s C indices (33, 34), which range from −1 to 1 (a value close 1 indicates strong clustering and close to −1 indicates strong dispersion). Dotted lines represent the values of the summary statistics corresponding to the observed data.
Fig. S5. Checking of the implementation of the inference procedures. We simulated 10 independent pseudodata from the model, with the model parameter values close to the posterior means obtained from fitting with the real dataset. The model is then fitted to each of the simulated datasets, and the resultant posterior distributions of the model parameters are shown. The true values of the model parameters are indicated by the red lines.
### Table S1. Testing alternative parameterizations of the incubation period

<table>
<thead>
<tr>
<th>Parameterization</th>
<th>Generation time, d</th>
<th>$R_0$</th>
<th>Dispersion parameter, k</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gamma (baseline)</td>
<td>10.9</td>
<td>2.39</td>
<td>0.37</td>
</tr>
<tr>
<td>Lognormal</td>
<td>10.9</td>
<td>2.46</td>
<td>0.35</td>
</tr>
<tr>
<td>Exponential</td>
<td>9.7</td>
<td>2.20</td>
<td>0.45</td>
</tr>
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The mean of generation time, $R_0$, and the dispersion parameter $k$ that quantifies superspreading are shown.

### Table S2. Testing alternative parameterizations of the infectious period

<table>
<thead>
<tr>
<th>Parameterization</th>
<th>Generation time, d</th>
<th>$R_0$</th>
<th>Dispersion parameter, k</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exponential (baseline)</td>
<td>10.9</td>
<td>2.39</td>
<td>0.37</td>
</tr>
<tr>
<td>Weibull</td>
<td>10.3</td>
<td>2.39</td>
<td>0.40</td>
</tr>
<tr>
<td>Gamma</td>
<td>10.43</td>
<td>2.40</td>
<td>0.38</td>
</tr>
</tbody>
</table>

The mean of generation time, $R_0$, and the dispersion parameter $k$ that quantifies superspreading are shown.

### Table S3. Testing alternative uninformative priors

<table>
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<tr>
<th>Priors</th>
<th>Generation time, d</th>
<th>$R_0$</th>
<th>Dispersion parameter, k</th>
</tr>
</thead>
<tbody>
<tr>
<td>U(0, 100) (baseline)</td>
<td>10.9</td>
<td>2.39</td>
<td>0.37</td>
</tr>
<tr>
<td>Exp(rate = 0.0001)</td>
<td>10.8</td>
<td>2.39</td>
<td>0.36</td>
</tr>
</tbody>
</table>

The mean of generation time, $R_0$, and the dispersion parameter $k$ that quantifies superspreading are shown.

### Table S4. Prior and posterior distributions of model parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Median [95% C.I.]</th>
<th>Prior</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\beta_a$, infectivity of first age group</td>
<td>0.76 [0.42, 1.39]</td>
<td>U(0,100)</td>
</tr>
<tr>
<td>$\beta_b$, infectivity of second age group</td>
<td>0.07 [0.002, 0.36]</td>
<td>U(0,100)</td>
</tr>
<tr>
<td>$\beta_c$, infectivity of third age group</td>
<td>0.4 [0.1, 0.66]</td>
<td>U(0,100)</td>
</tr>
<tr>
<td>$\beta_d$, infectivity of fourth age group</td>
<td>0.79 [0.27, 1.25]</td>
<td>U(0,100)</td>
</tr>
<tr>
<td>$\beta_e$, infectivity of fifth age group</td>
<td>0.56 [0.23, 0.92]</td>
<td>U(0,100)</td>
</tr>
<tr>
<td>$\psi$, spatial kernel parameter</td>
<td>0.42 [0.06, 0.87]</td>
<td>U(0,100)</td>
</tr>
<tr>
<td>$\alpha$ ($10^{-4}$), background hazard</td>
<td>4.6 [0.21, 12]</td>
<td>U(0,100)</td>
</tr>
<tr>
<td>$a$, mean of the incubation period</td>
<td>6.87 [5.34, 8.50]</td>
<td>U(0,100)</td>
</tr>
<tr>
<td>$b$, SD of the incubation period</td>
<td>4.02 [2.44, 5.44]</td>
<td>U(0,100)</td>
</tr>
<tr>
<td>$c$, mean (and SD) of the infectious period</td>
<td>3.96 [3.41, 4.60]</td>
<td>U(0,100)</td>
</tr>
</tbody>
</table>