Hawkes process modeling of COVID-19 with mobility leading indicators and spatial covariates

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Abstract

Hawkes processes are used in statistical modeling for event clustering and causal inference, while they also can be viewed as stochastic versions of popular compartmental models used in epidemiology. Here we show how to develop accurate models of COVID-19 transmission using Hawkes processes with spatial-temporal covariates. We model the conditional intensity of new COVID-19 cases and deaths in the U.S. at the county level, estimating the dynamic reproduction number of the virus within an EM algorithm through a regression on Google mobility indices and demographic covariates in the maximization step. We validate the approach on both short-term and long-term forecasting tasks, showing that the Hawkes process outperforms several models currently used to track the pandemic, including an ensemble approach and an SEIR-variant. We also investigate which covariates and mobility indices are most important for building forecasts of COVID-19 in the U.S.

Keywords: COVID-19 forecasting, Hawkes processes, Mobility indices, Spatial covariate

1. Introduction

Mathematical modeling and forecasting are playing a pivotal role in the ongoing SARS-CoV-2 (COVID-19) pandemic. In mid-March 2020, a report out of Imperial College London [1] forecasted severe consequences in the U.S. and ⁵ U.K. in the absence of significant public health interventions. In both nations, governments responded by closing schools, non-essential businesses and releasing general stay-at-home (shelter-in-place) orders. In the U.S., state and local policymakers are using mathematical models and projections to inform decisions about when and how to relax public health measures that have been put ¹⁰ in place. By and large, compartmental models that explicitly incorporate trans-

mission characteristics of infectious diseases have been favored over other statistical modeling approaches. High profile Susceptible-Exposed-Infected-Removed (SEIR) models include those out of the Institute for Health Metrics and Evaluation (IHME) [2], Columbia University [3], MIT [4], The Johns Hopkins University [5], and UCLA [6] (in the case of the UCLA model, an SEIR-variant

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Figure 1: The framework of the Hawkes process model for COVID-19 transmission. Demographic features at the county level impact the reproduction number of the Hawkes process. Lagged changes in mobility impact future secondary infections through a convolution with the inter-infection distribution w(t). The output of the model includes: (1) forecasts of future cases and mortality through simulation of the Hawkes process intensity, (2) an estimate of the dynamic reproduction number of the virus, and (3) regression results that allow for interpretation of the covariates that influence transmission differences across counties.

with an unreported compartment is fit using least-squares to reported infection and recovery data). Other strategies apart from SEIR models are CMU-TimeSeries ¹ and GT-DeepCOVID ². CMU-TimeSeries uses an auto-regressive time series model fit to case counts and deaths. GT-DeepCOVID is a purely data-driven approach using end-to-end deep learning models to predict mortality on a weekly basis. Our goal in this paper is to show that Hawkes processes, widely used in the statistical learning community to model contagion patterns in event data, are well suited for modeling and forecasting COVID-19 case and mortality data. They have several advantages that we will highlight, including being highly flexible in accommodating auxiliary spatio-temporal features such as county-level demographics and temporal mobility patterns, yet mathemat-

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¹https://delphi.cmu.edu/

²https://deepcovid.github.io/

ically they are connected to compartmental models [7] and allow for explicit incorporation of transmission dynamics (which we briefly review in the following section). Furthermore, extensive research has been conducted in the past several years on incorporating machine learning techniques into the point pro-

- ³⁰ several years on morporating machine learning techniques into the point process framework. Non-parametric Hawkes processes can be constructed where the triggering kernel is learned [8] and, more recently, fully neural network based point processes have been developed [9, 10]. Sparse linear combinations of Hawkes processes were a winning solution in the 2017 NIJ Crime Forecasting
- ³⁵ Challenge [11]. In certain cases a mixture of Hawkes processes may be needed to model more complex event contagion with high dimensional marks through Dirichlet processes [12, 13]. Hawkes processes can also be used for causal inference on networks [14] and recent efforts have also focused on training point processes through reinforcement learning [15, 16]. Hawkes processes also can
- take as input auxiliary covariates [17, 18, 19], including spatio-temporal features to model earthquake occurrences [20, 21, 22, 23] and environmental and demographic variables to model crime [18, 19]. We believe all of these methods have potential applications to modeling infectious diseases that are highly complex due to heterogeneity in the population, environment, and disparate public
- ⁴⁵ policies across regional and local jurisdictions. Despite these advantages, to our knowledge, the only U.S. state where a Hawkes process is being used to inform COVID-19 policy is in New Jersey (a collaboration with Facebook AI Research) ³.
- The outline of the paper is as follows. In Section 2, we introduce our Hawkes ⁵⁰ process model whose productivity (reproduction number) is dynamic and depends on spatio-temporal covariates. Unlike recently introduced models that incorporate covariates into the background rate of a Hawkes process [18, 19], our Hawkes process model may be viewed as a convolution of lagged mobility with an inter-infection time distribution to estimate the intensity of secondary
- ⁵⁵ infections in the future. This is important as phased reopening in the U.S. leads to mobility changes, the effects of which are not realized in the case and mortality data until days or weeks later. Hence the model can be used to forecast changes in transmission and new cases in real-time as mobility changes (see Figure 1). We estimate the intensity along with the dynamic reproduction number of the
- ⁶⁰ virus within an EM algorithm through a regression on Google mobility indices and demographic covariates in the maximization step. In Section 3, we validate the approach on both short-term and long-term forecasting tasks, showing that the Hawkes process outperforms several models from "The COVID-19 Forecast Hub Network ⁴." These models include SEIR models from Columbia University
- [3], Johns Hopkins University Applied Physics Lab [5], and an ensemble model from Berkeley that uses combined linear and exponential predictors with spatial covariates [24]. We also investigate which covariates and mobility indices are

 $^{{}^{3} \}rm https://ai.facebook.com/blog/using-ai-to-help-health-experts-address-the-covid-19-pandemic$

⁴https://covid19forecasthub.org/community/

most important for building forecasts of COVID-19 in the U.S. In Section 4, we discuss directions for future research and how the machine learning commu-⁷⁰ nity may be able to help improve Hawkes process models of COVID-19 as the pandemic continues to unfold.

2. Hawkes process model of COVID-19 transmission

In this section, we introduce a Hawkes process with spatio-temporal covariates for modeling COVID-19 case and death data. We then discuss the connection of the model to compartment models used in epidemiology and develop an expectation-maximization algorithm for inference.

2.1. Incorporating covariates into the Hawkes process

We propose a novel Hawkes process model that simultaneously estimates the intensity of events and tracks the dynamic reproduction number of the virus. Given the timestamps (or dates), $\mathcal{T} = \{t_1, t_2, \cdots, t_n\}$, of daily reported positive test cases or deaths, we model the rate of new cases (or deaths) in each country c as follows:

$$\lambda_c(t) = \mu_c + \sum_{\substack{t > t_j \\ t_j \in \mathcal{T}}} R(\boldsymbol{x}_c^{t_j - \Delta}, \theta) w(t - t_j), \text{ where } \boldsymbol{x}_c^{t_j - \Delta} = \begin{bmatrix} \boldsymbol{d}_c \\ \boldsymbol{m}_c^{t_j - \Delta} \end{bmatrix}, \quad (1)$$

where μ_c is the background rate modeling imported infections, w(t) is the interinfection time distribution, $\boldsymbol{m}_c^t = [\boldsymbol{m}_1^t, \boldsymbol{m}_2^t, \cdots]^{\mathsf{T}}$ are mobility indices on day t, and $\boldsymbol{d}_c = [d_1, d_2, \cdots]^{\mathsf{T}}$ are static demographic features. The time-varying reproduction number $R(\boldsymbol{x}_c^{t_j-\Delta}, \theta)$ is a function of mobility indices and demographic features. It can be interpreted as the average number of secondary infections caused by a primary infection. Because we are modeling reported infections rather than time of exposure, we introduce the parameter Δ to capture a po-

tential lag between a mobility change and the time t_j of a reported primary infection. Here, we combine the spatial and temporal covariates, and we model the dynamic reproduction number through a Poisson regression (Equation 2) where the coefficients θ are shared across the counties:

$$R(\boldsymbol{x}_{c}^{t_{j}-\Delta},\boldsymbol{\theta}) = \exp(\boldsymbol{\theta}^{\mathsf{T}} \; \boldsymbol{x}_{c}^{t_{j}-\Delta}).$$

$$\tag{2}$$

Our approach is related to those in recent preprints that incorporate mobility into compartment models [25, 5], however those approaches typically involve large-scale Monte Carlo simulations when performing inference. As we will show, the Hawkes process likelihood can be maximized without simulation via an efficient expectation-maximization algorithm.

2.2. Mathematical connection between Hawkes processes and compartmental models

Here we briefly review several variations of the Hawkes process in Equation 1 that can be connected to SEIR-type compartment models. The first variant is the SIR-Hawkes process. This model captures the long-term evolution of a pandemic by incorporating a pre-factor that accounts for the dynamic decrease in the number of susceptible individuals [7]:

$$\lambda^{\text{SIR}}(t) = (1 - \frac{I_c(t)}{N}) \left(\mu + \sum_{t_i < t} R_0 w(t - t_i) \right).$$
(3)

Here $I_c(t)$ is the cumulative number of infections that have occurred up to time t and N is the total population size. The point process governed by Equation 3 is a continuous time analog of a discrete stochastic SIR model when w(t) is specified to be exponential [7]. When w(t) is chosen to be gamma distributed, the Hawkes process also can approximate staged compartment models, like SEIR, if the average waiting time in each compartment is equal [26]. More complex parametric (or non-parametric) inter-infection time distributions w(t) may be employed within the Hawkes process framework in situations where disease dynamics cannot be captured by a SIR or SEIR model. In the early exponential growth stage of an epidemic, before finite population effects play a role, the Hawkes process in Equation 1 without the prefactor can be used to model new infections arising from SIR and SEIR models, as $\frac{I_c(t)}{N}$ will be small.

While a pre-factor in the Hawkes process involving the cumulative number of infections is necessary to model long-term disease dynamics, in the early stages of transmission a linear Hawkes process can be used (as the prefactor will be close to 1),

$$\lambda(t) \approx \mu + \sum_{t_i < t} R_0 w(t - t_i).$$
(4)

To illustrate this, we simulate a SEIR differential equation,

$$\frac{dS}{dt} = -\beta \frac{SI}{N}, \qquad \frac{dR}{dt} = \gamma I$$

$$\frac{dE}{dt} = \beta \frac{SI}{N} - \mu E, \quad \beta = \gamma R_0$$

$$\frac{dI}{dt} = \mu E - \gamma I,$$
(5)

where the parameters are chosen similar to those of COVID-19 estimates reported in [27, 28]. In particular we let $\gamma = .1$, $R_0 = 2$, $\mu = 1$, and $N = 5 \cdot 10^8$ and note that these parameters are not from any specific locations. We then fit the linear Hawkes process model in Equation 4 to new infections, μE , generated by the SEIR model. We use a non-parametric histogram estimator for w(t) and find a close fit between the Hawkes process and the SEIR model in Figure 2.

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Figure 2: (Main figure) The red plot shows new infections (μE) from the SEIR differential equation $\frac{dS}{dt} = -\beta \frac{SI}{N}$, $\frac{dE}{dt} = \beta \frac{SI}{N} - \mu E$, $\frac{dI}{dt} = \mu E - \gamma I$, $\frac{dR}{dt} = \gamma I$, where $\beta = \gamma R_0$, $\gamma = .1$, $R_0 = 2$, $\mu = 1$, and $N = 5 \cdot 10^8$. The blue squares show the linear Hawkes process $\lambda_t = \mu + \sum_{t>t_i} R_0 w(t-t_i)$ fit to the SEIR curve of new infections. Inset: Non-parametric histogram estimate for w(t).

In [7], the rate of events $\lambda(t)^{\text{SIR}}$ in a SIR-Hawkes process is established to be equal in expectation to new infections μE in the SEIR model after marginalizing out recovery events that are unobserved in a Hawkes process. In Figure 2, we show that in the early stage of spreading, the rate $\lambda(t)$ in a linear Hawkes process can also be used to approximate new infections μE .

2.3. EM algorithm for parameter inference

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We use an expectation-maximization (EM) algorithm to estimate the model in Equation 1, which has been widely used for Hawkes Process estimation [29, 30, 31]. First, we introduce latent random variables, $p_c(i, j)$, that represent the event that secondary infection *i* is caused by primary infection *j* in county *c*. We let $p_c(i, i)$ represent the event that case *i* is imported. The complete data log-likelihood is then given by,

$$\mathcal{L} = \sum_{c=1}^{|\mathcal{C}|} \left\{ \sum_{i=1}^{n} p_c(i,i) \log(\mu_c) - \int_0^T \mu_c \, dt + \sum_{j=1}^{n} \left\{ \sum_{i=j+1}^{n} p_c(i,j) \log[R(\boldsymbol{x}_c^{t_j - \Delta}, \theta) w(t_i - t_j | \alpha, \beta)] - \int_{t_j}^T R(\boldsymbol{x}_c^{t_j - \Delta}, \theta) w(t - t_j | \alpha, \beta) \, dt \right\} \right\}.$$
(6)

Here we use a Weibull distribution [32, 33, 34] with shape α and scale β to model inter-infection times, which we find accurately models the present data.

As the branching structure of the process is unobservable, we optimize the complete data log-likelihood in Equation 6 by iteratively alternating between an expectation step where the branching probabilities p_c are estimated and a

¹²⁵ maximization step where model parameters are updated by maximizing Equation 6. The EM-algorithm is equivalent to a projected gradient ascent on the likelihood of the Hawkes process [35].

2.3.1. Expectation step

During the expectation step, we estimate the latent variables $p_c(i, j)$ for each county. Given the parameters θ , α , β , and μ_c estimated from the last iteration, 130 the probabilities that case i was caused by case j (Equation 7a) or was imported (Equation 7b) are given by:

$$p_c(i,j) = \frac{R(\boldsymbol{x}_c^{t_j - \Delta}, \theta) w(t_i - t_j | \alpha, \beta)}{\lambda_c(t_i)},$$
(7a)

$$p_c(i,i) = \frac{\mu_c}{\lambda_c(t_i)}.$$
(7b)

Note that the rate $\lambda_c(t_i)$ in Equation 1 is considered to be an aggregation of triggering kernels from all previous historical events (i.e., all $t < t_i$) and the background rate μ_c . Therefore, we can consider the probability of case *i* caused 135 by case j, $p_c(i, j)$, as the contribution of primary infection j in the event rate at time t_i , i.e., $\lambda_c(t_i)$, and $p_c(i, i)$ can be seen as the contribution of the background rate.

2.3.2. Maximization step

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We then maximize the complete data log-likelihood with respect to the model parameters, conditioned on the estimated branching structure $p_c(i, j)$. During estimation we do not include event pairs (i, j) when j is within $\Psi =$ 14 days of the last day of the dataset, as the offspring events i have not yet been realized and the inclusion of these incomplete data biases parameter estimates. We choose $\Psi = 14$ as the incubation period for COVID-19 145 is thought to extend to 14 days given by the Clinical Care Guidance from the CDC: https://www.cdc.gov/coronavirus/2019-ncov/hcp/clinical-guidancemanagement-patients.html. For simplicity, the summation over n in the likelihood function, Equation 5, is replaced with \hat{n} in the description for the maximization step. Here, \hat{n} represents the number of events that are within $T - \Psi$ 150 $(\hat{n} = |t_i|t_i < T - \Psi|).$

Given the latent variable $p_c(i, j)$, the maximization of Equation 6 can be decoupled into three independent optimization problems. Starting with the coefficient θ from Poisson regression, the maximization of likelihood function can be rewritten as the following:

$$\hat{\theta} := \underset{\theta}{\operatorname{argmax}} \mathcal{L}_{\theta} = \sum_{c=1}^{|\mathcal{C}|} \left\{ \sum_{j=1}^{\hat{n}} \left\{ P_c(j) \log \left[R(\boldsymbol{x}_c^{t_j - \Delta}, \theta) w(t_i - t_j | \alpha, \beta) \right] - \int_{t_j}^T R(\boldsymbol{x}_c^{t_j - \Delta}, \theta) w(t - t_j | \alpha, \beta) \, dt \right\} \right\}, \text{where } P_c(j) = \sum_{i=j+1}^{\hat{n}} p_c(i, j).$$

$$\tag{8}$$

Because the last Ψ days are removed from the dataset and we assume that all possible offspring pairs (i, j) have been observed, we can therefore approximate the integrals for the inter-infection time w(t) in Equation 6 as is done in [36] by noting that $\int_{t_j}^T w(t-t_j|\alpha,\beta) \approx 1$. The optimization problem is therefore a Poisson regression, where we regress the observations $P_c(j) = \sum_{i=j+1}^{\hat{n}} p_c(i,j)$ against the covariates $\boldsymbol{x}_c^{t_j}$:

$$\hat{\theta} := \operatorname*{argmax}_{\theta} \mathcal{L}_{\theta} = \operatorname*{argmax}_{\theta} \sum_{c=1}^{|\mathcal{C}|} \left\{ \sum_{j=1}^{\hat{n}} P_c(j) \theta^{\mathsf{T}} \boldsymbol{x}_c^{t_j - \Delta} - \exp(\theta^{\mathsf{T}} \boldsymbol{x}_c^{t_j - \Delta}) \right\}.$$
(9)

The same optimization strategy can be applied on the shape and scale parameters, α and β . The optimization problem can then be solved as a weighted maximum likelihood estimation for the Weibull shape and scale parameters:

$$\hat{\alpha}, \hat{\beta} \coloneqq \operatorname*{argmax}_{\alpha,\beta} \mathcal{L}_{\alpha,\beta} = \operatorname*{argmax}_{\alpha,\beta} \sum_{c=1}^{|\mathcal{C}|} \left\{ \sum_{j=1}^{\hat{n}} \left\{ \sum_{i=j+1}^{\hat{n}} p_c(i,j) \log \left[w(t_i - t_j | \alpha, \beta) \right] \right\} \right\}.$$
(10)

where $p_c(i, j)$ is the weight of each inter-infection time observation $t_i - t_j$. Third, the background rate μ_c is determined analytically:

$$\hat{\mu_c} \coloneqq \operatorname*{argmax}_{\mu_c} \mathcal{L}_{\mu_c} = \operatorname*{argmax}_{\mu_c} \sum_{i=1}^{\hat{n}} p_c(i,i) \log(\mu_c) - \int_0^T \mu_c \, dt, \ \hat{\mu_c} = \sum_{i=1}^{\hat{n}} \frac{p_c(i,i)}{T}.$$
(11)

Pseudo code for the EM algorithm is presented in the Algorithm 1.

We note that the EM algorithm of the Hawkes process is also connected to ¹⁵⁵ the dynamic reproduction number estimator of Wallinga and Teunis [37], as the latter can be viewed as a 1-iteration EM algorithm where a histogram estimator is used for R_c^t with initial guess $R_c^t \equiv 1$. More details are discussed in the following section.

2.4. Connection of EM algorithm for Hawkes Process and dynamic R estimator of Wallinga and Teunis

Here we make the connection between the EM algorithm for the Hawkes process and the popular dynamic reproduction number estimator of Wallinga and Teunis [38, 37, 32]. The dynamic R estimator of Wallinga and Teunis is constructed as follows. The probability that individual i at time t_i was infected by individual j at time t_j is defined to be,

$$p_{ij} = \frac{w(t_i - t_j)}{\sum_{t_i > t_k} w(t_i - t_k)},$$
(12)

where the distribution of inter-infection times $w(t_i - t_j)$ is typically modeled as Weibull, Gamma, or log-normal [32]. The expected total number of individuals that j infects is then given by:

$$R_j = \sum_{i>j} p_{ij}.$$
 (13)

Algorithm 1 EM algorithm optimization

1: procedure $\mathbf{HkPR}_{M}^{+}(\mathcal{T}, \boldsymbol{x}, \Delta)$ $T \leftarrow \max \mathcal{T}, \ \alpha \leftarrow 2, \ \beta \leftarrow 2.$ ▷ Initialization 2: $\mu_c \leftarrow 0.5, \ R_c^t(t) \leftarrow 1, \ \forall c \in \mathcal{C} \ \text{and} \ 0 < t < T$. 3: while $\|\Delta\theta\|, |\Delta\alpha|, |\Delta\beta|, \|\Delta\mu\| > \text{tol } \mathbf{do}$ 4: Expectation step: 5: for $\forall i \geq j$ and 0 < i, j < T and $\forall c \in \mathcal{C}$ do 6: if i > j then 7: $p_c(i,j) \leftarrow \frac{R_c^{t_j}(\boldsymbol{x}_c^{t_j-\Delta},\theta)w(t_i-t_j|\alpha,\beta)}{\lambda_c(t_i)}.$ else if i = j then 8: 9: $p_c(i,i) \leftarrow \frac{\mu_c}{\lambda_c(t_i)}.$ 10: end if 11: end for 12: $\begin{aligned} \theta &\leftarrow \operatorname*{argmax}_{\theta} \sum_{c=1}^{|\mathcal{C}|} \left\{ \sum_{j=1}^{n} P_c(j) \theta^{\intercal} \boldsymbol{x}_c^{t_j - \Delta} - \exp(\theta^{\intercal} \boldsymbol{x}_c^{t_j - \Delta}) \right\}. \\ \alpha, \beta &\leftarrow \operatorname*{argmax}_{\alpha, \beta} \sum_{c=1}^{|\mathcal{C}|} \left\{ \sum_{j=1}^{n} \left\{ \sum_{i=j+1}^{n} p_c(i,j) \log \left[w(t_i - t_j | \alpha, \beta) \right] \right\} \right\}. \\ \mathbf{for} \ \forall c \in \mathcal{C} \ \mathbf{do} \\ \mu_c \leftarrow \sum_{i=1}^{n} \frac{p_c(i,i)}{T}. \\ \mathbf{end for} \end{aligned}$ 13:14:15: 16:17:18:end for 19:end while 20: 21: end procedure

Wallinga and Teunis then obtain an estimate of the dynamic reproduction number R(t) by averaging R_j over all observed cases j where the time of infection t_j occurred on day t:

$$R(t) = \frac{1}{N_t} \sum_{t \le t_j < t+1} R_j,$$
(14)

(here N_t is the number of observed infections on day t).

On the other hand, for the Hawkes process the intensity (rate) of infections is modeled as

$$\lambda(t) = \mu + \sum_{t>t_i} R(t_i)w(t-t_i), \qquad (15)$$

where w(t) and R(t) are the inter-infection time distribution and dynamic reproduction number respectively. Rather than modeling R(t) as dependent on mobility, we can instead model R(t) as a piece-wise constant function:

$$R(t) = \sum_{k=1}^{B} r_k 1\{t \in I_k\}.$$
(16)

Here the I_k are intervals discretizing time, B is the number of such intervals, and r_k is the estimated reproduction rate in interval k.

Given initial guesses for the model parameters and r_k , the EM algorithm for the Hawkes process iteratively updates the parameters and branching probabilities by alternating between the

E-step update:

$$p_{ij} = R(t_j)w(t_i - t_j)/\lambda(t_i)$$
(17)

$$p_{ii} = \mu / \lambda(t_i) \tag{18}$$

and M-step update:

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$$w(t) \sim MLE(\{t_i - t_j; p_{ij}\}) \tag{19}$$

$$\mu = \sum_{i} p_{ii}/T \tag{20}$$

$$r_k = \sum_{t_i > t_j} p_{ij} 1\{t_j \in I_k\} / N_k$$
(21)

where T is the total length of the observation period, N_k is the total number of events in interval k, and the w(t) is estimated via weighted MLE (for either a Gamma, Weibull or log-normal) using the inter-event times as observations and branching probabilities as weights. We also drop event pairs (i, j) when j is within $\Phi = 14$ days of the last day of the datasets in consideration of the incubation period.

Finally, we can compare Equation 17 to Equation 12. The dynamic R(t) estimator in Equation 12 is what you obtain with 1 step of the EM algorithm ¹⁷⁵ in Equation 17 with initial guess $R(t) \equiv 1$, $\mu = 0$ and 1 day chosen as the bin width for the histogram estimator.

2.5. Hawkes process forecasting

We forecast future events using the branching process representation of the Hawkes process. We first simulate immigrant events through the Poisson process based on the background rate. For each event in the history of the process, we then simulate a Poisson random variable with mean $R(\mathbf{x}_c^{t_j-\Delta}, \theta)$ representing the number of secondary infections caused by event j. For each of these infections we simulate the time of infection by drawing inter-event times from the estimated Weibull distribution. For example, for an event on day 4, it may cause a secondary infection on day 22 if we draw a sample as 18 from the Weibull distribution. Events falling in the future (past the forecasting date) are then used to update the forecasted intensity through Equation 1. We simulate multiple realizations of this process (100 times in our application) to estimate a mean intensity forecast along with confidence intervals.

¹⁹⁰ 3. Experiments and Results

In this section we first provide details on the datasets and baseline models used in our experiments. We then discuss the experimental results of several COVID-19 retrospective forecasting tasks at the U.S. county level. The source code and dataset are included in the supplemental material and are available online in a anonymous repository⁵.

3.1. Datasets

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3.1.1. Covid-19 daily cases and deaths reported by The New York Times

The New York Times (NYT) [39] ⁶ releases a daily report of the cumulative numbers of COVID-19 cases in the United States at the county level and over time. While NYT data closely tracks data aggregated by a project at Johns Hopkins University [40], NYT county level reporting started earlier and is therefore used in this study. In total, there are 3,217 counties with cases and/or deaths in the dataset. The time series data are compiled from state and local government health departments. In order to have sufficient data for statistical inference, we select the counties with confirmed cases greater than and equal to 10 (denoted by \mathcal{D}_{conf}) and the counties with at least 1 death (denoted by \mathcal{D}_{death}) by 11/10/2020 when the datasets. Parameter sharing may improve models in counties with less data through variance reduction, but can potentially bias estimates in more populated counties with more cases.

We therefore assess model performance over different subsets of counties grouped by case volume. We first rank counties by the number of confirmed cases and deaths by the cut-off date, 11/30/2020, and we then evaluate forecasting accuracy on the top-10% of counties (denoted by $Q_{10\%}^{\text{top}}$), the top-25% counties (denoted by $Q_{25\%}^{\text{top}}$), and counties between the top-25% and top-50% quantiles (denoted by $Q_{50\%}^{25\%}$).





Figure 3: Distribution of cumulative cases reported at 11/30/2020 at different quantiles.

 $^{^{5}} https://anonymous.4 open.science/r/d425dcf9-3cfb-4f82-a08c-ee583ab36291/$

⁶https://github.com/nytimes/covid-19-data



Figure 4: Example of the daily # of confirmed cases/deaths.

In Figure 3a and 3b, we present the distribution of the cumulative confirmed cases and deaths at three different quantiles up to the cute-off date 11/30/2020. As the counties at the top-50% have more than 1,000 confirmed cases and 10 deaths, some urban counties, mostly at the top-10%, had already surpassed 220 10,000 confirmed cases and accumulated more than 300 deaths. In Figure 4a and 4b, we show the daily reported confirmed cases and deaths of top-3 counties in $Q_{10\%}^{\text{top}}$ and $Q_{50\%}^{25\%}$ from $\mathcal{D}_{\text{conf}}$ and $\mathcal{D}_{\text{death}}$, respectively. Given different demographics and different COVID-19 regulations, each state went through different phases. For example, while Cook, IL seemed to contain the first spike after 225 May, the confirmed cases in Los Angeles, CA seem steadily increase and only slow down after July. The daily death toll of Maricopa, AZ only hit its record high only after August unlike Los Angeles, CA, which had already had their first wave in terms of deaths in April. Overall, the deaths are increasing as the U.S. heads into the winter months. Such differences in infection rates suggest 230 that different public health and social measures may need to be tailored county by county. Therefore, the proposed county-level forecasting model may aid local government policymakers in understanding the demographic and mobility factors that play a role in local reproduction of the virus.

235 3.1.2. Google mobility index reports

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We use Google daily mobility index reports at the county level [41] to estimate a dynamic reproduction number that tracks changes in movement patterns due to stay at home orders (and their staged removal). In total, there are 6 mobility types, including grocery & pharmacy, parks, transit stations, retail & recreation, residential and workplaces. Mobility indices for each category

and county are calculated with respect to a baseline value for that day of the week The baseline day of the week is the median value from the 5-week period from 01/03/2020 to 02/06/2020. That is, the values are the relative number of visitors for counties in each category. Note that during the model training, we

²⁴⁵ introduce the parameter Δ to capture a potential lag between a mobility change and the time t_j of a reported primary infection. As we make forecasts, we use the mobility in training data from the previous Δ days to infer the reproduction number. If the forecast target is more than Δ , we would use the most recent



Figure 5: Heat map of mobility indices across counties in \mathcal{D}_{conf} and over time.

value from the day of the week in the training data.

We drop "workplace" mobility from our analysis due to high collinearity with "residential" mobility. Some mobility data are missing when data is sparse for a given date. To deal with missing values, we adopt multivariate feature imputation ⁷, which estimates each missing mobility entry as a function of other mobility types on the same day in the same county. We show some heatmaps
of mobility patterns across counties and time in the Figure 5, where a major change can be observed coinciding with stay at home orders (the first statewide stay-at-home order was issued at 03/21/2020). Also, the reopening phase in most of the counties can be seen after May. For counties hit by COVID-19 the most (i.e., those in the top-10 %), we can also observe some strict regulations
in the "Retail and recreation" areas and better compliance with stay-at-home orders based on high mobility in "Residential" area.

3.1.3. County-level demographic covariates

We incorporate spatial demographic features that may be predictive of symptomatic cases of COVID-19 (which are more likely to result in testing and mortality). The dataset is available in a curated form [24] and is derived from CDC and census datasets. The data is at the county level and includes population, median age, number of hospitals and ICU beds, percentage of smokers and diabetes, and heart disease mortality.

In Figure 6, we present two examples of spatial demographic features at the county-level used to model variations in the reproduction number. In Figure 6a we observe that both the east and west coasts of the United States are more densely populated compared to midwestern and western regions. Diabetes percentage (shown in Figure 6b), on the other hand, is mostly higher in southern regions of the U.S.

 $^{^{7}}$ https://scikit-learn.org/stable/modules/impute.html#multivariate-feature-imputation



Figure 6: Examples of spatial demographic and health features at the county-level.

275 3.2. Baseline models and experimental setup for retrospective forecasting comparison

We compare the Hawkes process model in Equation 1 with several models including an SEIR model used in a pandemic tracking dashboard ⁸ out of Columbia University [3] (denoted by **PROJ**), an geospatial SEIR Model from

- the Johns Hopkins University Applied Physics Lab [5] (denoted by **BUCKY**), and an ensemble model with linear and exponential predictors from University of California, Berkeley [24] (denoted by **CLEP**). Note that all three competing models are tested directly from the released source code and we follow the same experimental protocol as for our proposed model. A simplified Hawkes process,
- denoted by Hawkes, where the reproduction number is held constant is used for comparison to demonstrate the effectiveness of tracking the reproduction number dynamically. We also compare our full Hawkes process model, denoted by HkPR⁺_m, to a Hawkes process, HkPR⁻_m, with only mobility features to determine the marginal improvement of adding demographics.
- ²⁹⁰ We backtest the six competing models on the \mathcal{D}_{conf} and \mathcal{D}_{death} datasets using the "walk-forward" validation approach. In particular, for 7-day forecasts we first train the models based on cases and deaths before the first cut-off date, 04/15/2020, and then forecast through 04/21/2020. We then slide the forecasting window, training on data before 4/22/2020 and forecasting from 04/22/2020
- to 04/28/2020. We repeat this process until the final date of 05/19/2020 (a similar approach is used for 14 and 28 day forecasts). The multivariate imputation models are also trained in the same walk forward fashion to avoid possible data leakage. The hyper-parameter of the lag parameter Δ ranges from 7, 14, 21, and 28 days in our experiments. For each of the forecasts, we simulate them 100 times and the point estimate is made through the average.

We evaluate the models according to mean absolute error, MAE, averaged across counties and forecasting windows of the same length, along with percentage error, PE. Mean absolute error (MAE) and the percentage error (PE) are

⁸https://covid19forecasthub.org/community/

calculated as follows:

$$\mathsf{MAE} = \frac{\sum_{c=1}^{|\mathcal{C}|} |n_c - \hat{n}_c|}{|\mathcal{C}|}, \ \mathsf{PE} = \frac{|\sum_{c=1}^{|\mathcal{C}|} n_c - \sum_{c=1}^{|\mathcal{C}|} \hat{n}_c|}{\sum_{c=1}^{|\mathcal{C}|} \hat{n}_c},$$
(22)

where \hat{n}_c , and n_c are the number of reported events and predicted events, respectively. We also compare the ranking quality of the competing models using Normalized Discounted Cumulative Gain (NDCG) [42], which can be used to evaluate the power of recommendations for counties with potential COVID-19 spikes in the near future.

3.3. Experimental results

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Table 1: MAE on predicted confirmed cases \mathcal{D}_{conf}

	7-days				14-days			28-days			
Model	$Q_{10\%}^{ m top}$	$Q_{25\%}^{\mathrm{top}}$	$Q^{25\%}_{50\%}$	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{\mathrm{top}}$	$Q^{25\%}_{50\%}$	$Q_{10\%}^{ m top}$	$Q_{25\%}^{ m top}$	$Q^{25\%}_{50\%}$		
PROJ	809.40	415.88	55.30	1664.90	857.93	117.86	3432.57	1779.56	252.43		
CLEP	238.30	134.83	33.86	585.81	324.32	88.87	1963.52	1090.36	207.81		
BUCKY	404.49	212.80	37.45	883.69	459.77	89.85	2085.88	1116.91	229.33		
Hawkes	224.35	120.61	24.02	569.49	300.06	55.45	1803.63	935.83	165.92		
$\mathbf{HkPR}_{\mathrm{m}}$	211.59	114.34	22.44	519.00	271.86	49.83	1573.58	835.59	136.60		
$\mathbf{HkPR}_{\mathrm{m}}^{+}$	210.72	114.69	22.38	522.92	276.28	49.86	1611.48	893.65	132.79		

The best performance is marked in **bold**.

Table 2:	MAE on	predicted	confirmed	cases	\mathcal{D}_{doubh}
10010 2.	TIAL OIL	predicted	communed	Cases	∠ death

	7-days			14-days			28-days		
Model	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{\mathrm{top}}$	$Q^{25\%}_{50\%}$	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{\rm top}$	$Q^{25\%}_{50\%}$	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{\rm top}$	$Q^{25\%}_{50\%}$
PROJ	15.56	7.89	1.12	30.66	15.55	2.20	56.85	29.37	4.41
CLEP	10.96	5.83	1.16	19.10	10.58	2.31	78.17	42.99	8.56
BUCKY	8.23	4.55	1.00	16.07	8.70	1.83	29.55	16.56	3.98
Hawkes	8.49	4.59	1.04	17.38	9.18	1.98	47.13	24.29	4.32
$\mathbf{HkPR}_{\mathrm{m}}$	7.19	4.07	1.01	13.40	7.55	1.78	33.30	18.23	3.74
$\mathbf{HkPR}_{\mathrm{m}}^{+}$	7.24	4.07	1.01	13.68	7.53	1.77	35.99	19.18	3.60

The best performance is marked in **bold**.

In Table 1 and Table 2, we present the experimental results for 7, 14, and 28 days window forecasts of MAE for all models applied to both confirmed cases (\mathcal{D}_{conf}) and deaths (\mathcal{D}_{death}) , and in Table 3 and Table 4, we report the results for PE. In terms of MAE and PE, both of our proposed models, $HkPR_m$ and $HkPR_m^+$, outperform the models, **PROJ** and **CLEP**, by a large margin in all three forecasting periods and across quantile subsets of the data. The improvements of MAE and PE can also be seen in the simplistic baseline Hawkes process, **Hawkes**. This suggests that the Hawkes process approach has a good potential on modeling infectious disease due to the self-exciting properties that lie in the COVID-19 cases.

		7-days		1	l4-days		28-days			
Model	$Q_{10\%}^{\rm top}$	$Q_{25\%}^{\rm top}$	$Q_{50\%}^{25\%}$	$Q_{10\%}^{ m top}$	$Q_{25\%}^{\rm top}$	$Q_{50\%}^{25\%}$	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{\rm top}$	$Q_{50\%}^{25\%}$	
PROJ	91.00	90.40	84.07	94.41	94.63	92.83	95.76	96.17	96.94	
CLEP	10.23	12.08	41.09	24.60	29.42	90.24	41.19	47.74	515.17	
BUCKY	19.61	20.12	35.58	28.05	27.90	69.61	47.91	49.24	97.86	
Hawkes	11.52	11.31	14.36	17.33	17.02	19.60	41.25	40.06	39.11	
$\mathbf{HkPR}_{\mathrm{m}}$	11.72	10.75	15.44	13.92	15.10	15.08	38.77	38.20	46.38	
$\mathbf{HkPR}_{\mathrm{m}}^{+}$	10.16	10.35	12.95	15.30	13.45	16.91	41.96	33.33	41.31	

Table 3: PE on predicted confirmed cases \mathcal{D}_{conf}

The best performance is marked in **bold**.

Table 4: PE on predicted confirmed cases \mathcal{D}_{death}

	7-days			14-days			28-days		
Model	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{\rm top}$	$Q^{25\%}_{50\%}$	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{\rm top}$	$Q^{25\%}_{50\%}$	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{\rm top}$	$Q^{25\%}_{50\%}$
PROJ	72.56	72.25	45.10	81.93	81.96	73.62	90.21	90.53	87.16
CLEP	23.39	26.35	18.05	19.23	19.27	24.74	73.77	78.90	173.88
BUCKY	17.64	16.08	14.71	15.63	13.93	20.22	15.36	13.20	36.19
Hawkes	17.97	16.99	15.81	20.03	20.71	16.17	48.79	44.15	28.17
$\mathbf{HkPR}_{\mathrm{m}}$	16.77	15.59	13.80	20.40	17.38	13.72	33.03	52.51	22.04
$HkPR_m^+$	17.53	16.92	14.05	18.18	15.23	16.93	38.31	44.78	17.66

The best performance is marked in **bold**.

We found that adding mobility indices improves **Hawkes**, where forecasting accuracy of **HkPR**_m also increases across the subsets and all forecasting window. For example, the improvements on MAE over **Hawkes** can go up to 13%, 11%, and 18% for 28 days forecast when **HkPR**_m is applied to $Q_{10\%}^{\text{top}}$, $Q_{25\%}^{\text{top}}$, and $Q_{50\%}^{25\%}$ in $\mathcal{D}_{\text{conf}}$, respectively. Similar decrease on MAE can be observed when **HkPR**_m is applied to three quantile subsets in $\mathcal{D}_{\text{death}}$, where **HkPR**_m outperforms **Hawkes** by 29%, 25%, and 13% in MAE, respectively. In terms of PE, **HkPR**_m stays ahead of **Hawkes** with only one exception at $Q_{50\%}^{25\%}$ of $\mathcal{D}_{\text{death}}$ in 28 days forecasting. This shows that by modeling the reproduction number through daily mobility indices we can enhance the forecasting accuracy and obtain more precise estimation on the spikes in the future.

By adding demographic features, we can marginally boost the MAE and PE of $\mathbf{HkPR_m^+}$ over $\mathbf{HkPR_m}$ in some cases. In general, the variation, $\mathbf{HkPR_m^+}$, also shows similar improvements over the competing models. In particular, $\mathbf{HkPR_m^+}$ has the best PE enhancement over $\mathbf{HkPR_m}$ at $Q_{50\%}^{25\%}$ in \mathcal{D}_{death} for 28 days forecast, which is 20%. This demonstrates that the major forecasting power comes from the joint modeling of mobility indices in the reproduction number while the choices of the background rate and inter-infection distribution may and only play a minor part.

Moreover, we notice that model **BUCKY** is a competitive baseline in $\mathcal{D}_{\text{death}}$ where it has better accuracy in a few cases, such as MAE and PE $Q_{10\%}^{\text{top}}$ and $Q_{25\%}^{\text{top}}$ for 28 days forecast. Possible explanation for its advantage could be the CDC-

recommended parameters that has been introduced to aid the model training especially for recovery and deaths compartments in its SEIR model. Those parameters include case fatality ratio, case hospitalization ratio, time between death and reporting, etc. However, introducing such pre-trained parameters from CDC may not be practical in real-time forecasting and may potentially bring in the data leakage issue.

Table 5: NDCG on predicted confirmed cases $\mathcal{D}_{\rm conf}$

		7-days			14-days		28-days			
Model	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{ m top}$	$Q^{25\%}_{50\%}$	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{ m top}$	$Q^{25\%}_{50\%}$	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{ m top}$	$Q_{50\%}^{25\%}$	
PROJ	0.7225	0.7039	0.8232	0.6946	0.6793	0.8412	0.6696	0.6614	0.8589	
CLEP	0.9626	0.9526	0.8620	0.9418	0.9402	0.8704	0.9015	0.8843	0.8739	
BUCKY	0.9283	0.9279	0.8600	0.9269	0.9216	0.8768	0.9013	0.8957	0.8813	
Hawkes	0.9738	0.9757	0.8680	0.9697	0.9704	0.8926	0.9414	0.9419	0.8879	
$\mathbf{HkPR}_{\mathrm{m}}$	0.9706	0.9728	0.8673	0.9715	0.9755	0.8956	0.9502	0.9521	0.8958	
$\mathbf{HkPR}_{\mathrm{m}}^{+}$	0.9734	0.9759	0.8672	0.9752	0.9758	0.8932	0.9493	0.9503	0.8918	

The best performance is marked in **bold**.

Table 6: NDCG on predicted confirmed cases \mathcal{D}_{death}

	7-days				14-days		28-days			
Model	$Q_{10\%}^{ m top}$	$Q_{25\%}^{\mathrm{top}}$	$Q_{50\%}^{25\%}$	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{\mathrm{top}}$	$Q^{25\%}_{50\%}$	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{\mathrm{top}}$	$Q_{50\%}^{25\%}$	
PROJ	0.6746	0.6607	0.7057	0.6823	0.6529	0.7666	0.7003	0.6837	0.8050	
CLEP	0.9126	0.9010	0.7451	0.9095	0.8945	0.7849	0.8696	0.8240	0.8043	
BUCKY	0.9161	0.9160	0.7301	0.9217	0.9222	0.7797	0.9074	0.9095	0.8278	
Hawkes	0.9506	0.9493	0.7548	0.9475	0.9469	0.8011	0.9293	0.9297	0.8212	
$\mathbf{HkPR}_{\mathrm{m}}$	0.9491	0.9476	0.7598	0.9446	0.9457	0.8007	0.9299	0.9315	0.8195	
$\mathbf{HkPR}_{\mathrm{m}}^{+}$	0.9504	0.9474	0.7597	0.9502	0.9514	0.7963	0.9368	0.9372	0.8176	

The best performance is marked in **bold**.

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uation. Generally, the proposed models **HawkPR** have a better NDCG performance when applied to confirmed cases for most of the quantile subsets. In terms of NDCG on the \mathcal{D}_{death} dataset, the baseline Hawkes process, **Hawkes**, performs better in some cases but proposed method consistently comes in second for most of the forecasting window. By generating rankings with good

In Table 5 and Table 6, we present the NDCG results for the ranking eval-

- qualities, **HawkPR** can serve as a recommender system for the hotspot counties and the public health policymakers can tailor strategies specifically for each region to contain the virus. We also note that in our model, we are estimating inter-event distributions of observed cases (ignoring asymptomatic cases) and
- ³⁵⁵ therefore these are observed or "effective" inter-event distributions, rather than true inter-infection distributions based on longitudinal data. We believe this approach is justified by the performance of the model in forecasting observed cases (and this approach is taken in other applications, like seismology where some earthquakes are not observed).

Table 7: Model coefficients (\mathcal{D}_{conf})

Covariate	coef	pValue
Retail/recreation	0.1303	0
Grocery/pharmacy	0.0029	8.46×10^{-09}
Transit stations	-0.0102	1.56×10^{-98}
Parks	-0.0355	0
Residential	-0.1063	0
Population density	0.0220	0
# ICU beds	0.0110	1.20×10^{-247}
# hospitals	0.0106	8.69×10^{-128}
Median age	-0.0049	3.85×10^{-38}
Population est.	-0.0214	0
Smokers %	-0.0361	0
Heart disease mort.	-0.0453	0
Diabetes %	-0.0589	0

The first 5 covariates are mobility indices, followed by static demographic covariates and two types of coefficients are sorted, respectively.

Table 8: Model coefficients (\mathcal{D}_{death})

Covariate	coef	pValue
Retail/recreation	0.1047	4.15×10^{-118}
Grocery/pharmacy	0.0746	8.88×10^{-111}
Transit stations	0.0276	2.48×10^{-11}
Residential	-0.0929	5.04×10^{-212}
Parks	-0.1294	0
# ICU beds	0.0423	2.17×10^{-64}
Population density	0.0409	0
Population est.	0.0062	5.18×10^{-2}
Median age	-0.0157	1.65×10^{-7}
Heart disease mort.	-0.0250	1.29×10^{-8}
# hospitals	-0.0423	6.54×10^{-28}
Diabetes %	-0.1041	1.33×10^{-86}
Smokers %	-0.1448	1.65×10^{-279}

The first 5 covariates are mobility indices, followed by static demographic covariates and two types of coefficients are sorted, respectively.

360 3.3.1. Importance of covariates

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In Table 7 and Table 8, we show the dynamic reproduction number coefficients of \mathbf{HkPR}_{m}^{+} estimated from the Poisson regression component (Equation 2) when applied to \mathcal{D}_{conf} and \mathcal{D}_{death} , respectively. The p-value is calculated from the Poisson regression analysis in the M-step after the EM algorithm reaches convergence. The absolute value of the coefficients indicates the magnitude of

the correlation between the reproduction number and the features. With the exception of population estimation in $\mathcal{D}_{\text{death}}$, the coefficients of all variables are statistically significant at the 10^{-7} level or below. The dynamic reproduction number is positively correlated with "Retail and recreation" while negatively correlated with "Residential", meaning that as mobility shifted away from com-

- mercial areas towards residences, the reproduction number decreased. In terms of spatial covariates, the reproduction number is positively correlated with "Population density" and "# of ICU beds." This suggests that the regions hit the hardest by COVID-19 are mostly urban areas, where most of intensive treat-
- ³⁷⁵ ment units are situated. The reproduction number is also negatively correlated with percent of the population with" Diabetes" and "Heart disease mortality rate." Several possible explanations for this observation include high-risk individuals are being more cautious or that they tend to live in areas with less cases, potentially with less population.

380 3.3.2. COVID-19 forecasting and reproduction number analysis

In Figure 7, we present an example of 28 days projection made through \mathbf{HkPR}_{m}^{+} from 10/28/2020 - 11/25/2020 for both \mathcal{D}_{conf} and \mathcal{D}_{death} . We can observe that \mathbf{HkPR}_{m}^{+} has very promising results in making projections, especially for the short term future, When the number of forecasting windows increases,

the forecasting error increase as the task also being more difficult. Moreover, the narrow confidence interval calculated through 100 Hawkes processes simulations



Figure 7: Forecasting for 28 days from 10/28/2020 - 11/25/2020

suggests that the proposed model can make relatively stable forecasting. Lastly, based on the projections, as the number of confirmed cases soon would hit over 500,000 in the top counties including Los Angeles, CA, Cook, IL, and etc. It is imperative to have a robust framework to help governments to design strategies to combat COVID-19 or even more, prioritize vaccine distribution.

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In Figure 8 and Figure 9, we find that the estimated dynamic reproduction number closely tracks lagged mobility, where the optimal lag parameter is determined as $\Delta = 14$ days for \mathcal{D}_{conf} and $\Delta = 21$ days for \mathcal{D}_{death} . The top-2 counties ³⁹⁵ in $Q_{10\%}^{top}$ have estimated reproduction number initially above 2.5. After stay-athome orders (around 04/11/2020), mobility in residential areas increased. On the other hand, mobility in retail and recreation decreased and the reproduction number fell to around 1, which explains why curves were relatively "flat" in many areas in the U.S. after the lockdown. However, as most of states reopened and lifted up the restrictions, the reproduction number increased after a large population resumed their daily routine, which can be also be observed by the increased mobility in retail and recreation after July. Lastly, to validate the reproduction numbers, we also compare our results to the ones estimated



Figure 8: Estimated R of confirmed cases \mathcal{D}_{conf} and lagged mobility changes ($\Delta = 14$ days)



Figure 9: Estimated R of deaths \mathcal{D}_{death} and lagged mobility changes ($\Delta = 21$ days)

by Stanford University ⁹ and our estimation match to their findings, which are around 1.5-2.5 initially and 0.5-1.5 up to the beginning of December in 2020.

3.3.3. Example of estimated event intensities and weekly forecasts

In Fig. 10, we present examples of the estimated intensities for the following four models: **Hawkes**, **HkPR**_d, **HkPR**_m and **HkPR**⁺_m, and we compare them with the number of cases/death in Cook, IL/Los Angles, CA, respectively. Note ⁴¹⁰ that we add **HkPR**_d, a Hawkes model in which only demographics is used. In these models, **HkPR**_m and **HkPR**⁺_m include mobility indices to estimate the reproduction number dynamically, and **Hawkes** and **HkPR**_d have a constant reproduction number for each county. Comparing **Hawkes** and **HkPR**_d, the

⁹https://web.stanford.edu/~chadj/Covid/Dashboard.html

marginal variance between the intensities suggests that demographic features ⁴¹⁵ may not significantly affect modeling the reproduction number in the present data. On the other hand, \mathbf{HkPR}_{m} and \mathbf{HkPR}_{m}^{+} show yield different fitted intensities compared to \mathbf{Hawkes} and \mathbf{HkPR}_{d} , indicating that mobility is playing an important role.

In Fig. 11, we present an example of weekly forecasts for all models except **PROJ**, which has relatively poor performance. In addition, we compare the forecasts against the true number of cases/deaths of Los Los Angeles, CA/Cook, IL respectively to provide a graphical presentation of the model fits. In general, all models can successfully capture the trend of the number of events, especially the valley around June and July and the spike in November, though the Hawkes process forecasts are more accurate than CLEP and BUCKY.



Figure 10: Example of the fitted effect from mobility and demographics



Figure 11: Example of the weekly forecasts

3.3.4. Dynamic background rate μ_c modeling

In this section, we investigate a potential improvement to the model with the incorporation of a dynamic background rate $\mu_c(t)$. For this purpose we again use mobility as a covariate and estimate the background rate through Poisson regression, where $\mu_c(t) = \exp(\theta_{\mu}^{\intercal} \boldsymbol{x}_c^{t-\Delta})$. The reasoning behind this choice is that imported case volume is correlated with mobility, especially in transit stations. Estimation for the corresponding parameter, θ_{μ} , is achieved through maximum likelihood estimation (MLE) corresponding to Poisson regression in the M-step of the EM algorithm. This new approach can be seen as a variation of our model that we denote as \mathbf{HkPR}_{μ} . We apply this variation of our **HawkPR** on the dataset from The New York Times (NYT) and we explore the improvements in the forecasting task for 7 and 14 days.

Table 9: Performance and improvements of \mathbf{HkPR}_{μ} on \mathcal{D}_{conf} and \mathcal{D}_{death}

data	ovl	7	'-days	14-	14-days				
	evi	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{ m top}$	$Q_{10\%}^{\mathrm{top}}$	$Q_{25\%}^{ m top}$				
	MAE	206.66 (1.93%	$\overline{(0.34\%)}$ 113.95 (0.34%)	509.88 (1.76%)	271.20 (0.24%)				
$\mathcal{D}_{\mathrm{conf}}$	PE	11.57 -	10.44 -	14.44 -	15.44 -				
	NDCG	0.9761 (0.24%	%) 0.9768 (0.09%)	0.9735 -	0.9739 -				
	MAE	7.12 (0.98%	6) 4.03 (0.98 %)	13.67 -	7.69 -				
$\mathcal{D}_{\mathrm{death}}$	PE	18.07 -	16.87 -	16.39 -	16.61 -				
	NDCG	0.9492 -	0.9475 -	$0.9515\ (\mathbf{0.14\%})$	$0.9520\ (\mathbf{0.06\%})$				

The performances which have an improvement over the best model from the previous experiments are marked in **bold** and the improvement (%) over the best models from previous performance is included. In this table, "evl" is the evaluation metrics and "dataset" is the set of COVID reports on which we apply the model.

In Table 9, we summarize the performance of the Hawkes model with dynamic background rate, where indicate improvements over the best model from the previous experiments in bold.

In Table 9, we can see some marginal improvements over the previous best models, especially for confirmed cases. Further improvements may be possible by monitoring cross-county and cross-state travel patterns, which would be a good direction for future investigation.

445 3.4. State-level Comparison to COVID-19 Forecast Hub

The COVID-19 Forecast Hub ¹⁰ is a repository that aggregates COVID-19 forecasts from a number of university and research groups following standardized data and forecasting formats. In specific, such ensemble was created by taking the arithmetic average of each prediction quantile for all eligible models for a given location. Recently, the COVID-19 Forecast Hub [43] has also intro-

duced an ensemble model that combines the various models submitted to the

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¹⁰https://covid19forecasthub.org/

hub into a single ensemble forecast. Comparing to other standalone models, it has demonstrated superior performance in forecasting deaths due to COVID-19 after May 2020 in the 50 states. To better validate the our Hawkes process framework, in this section we compare our model with several individual submissions and the ensemble model from the COVID-19 Forecast Hub.

Several differences between our county level experiments in the previous sections and the format of COVID-19 Forecast Hub submissions are worth noting. In particular, COVID-19 Forecast Hub forecasts are at the state level and some

- team contributions vary significantly in terms of the number of submissions, which locations are included, and whether cases and/or deaths are forecasted. In addition to the differences between the source of COVID-19 reports ¹¹, we also note that only a few teams have complete submissions at each county. Therefore, to fairly compare and contribute to the ensemble model, we adapt
- ⁴⁶⁵ our framework by training our models at the state-level using reports from the Johns Hopkins University dataset, from 02/15/2020 to 03/21/2021. Also, we incorporate Google mobility index for each state ¹² and with state-wise demographics ¹³ to model the event intensities, reproduction number, and the background rate in Equation 1.

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3.4.1. Weighted Interval Score WIS

The COVID-19 Forecast Hub uses a quantile based metric to evaluate forecasts, the weighted interval score (WIS), which considers the uncertainty in the predictive distribution [44]. Given a predictive distribution F in the format of quantiles, WIS can be seen as a measure of closeness between the entire distribution and the ground truth. To evaluate with WIS, we first calculate a single interval score IS_{α} .

$$IS_{\alpha}(F, \hat{n}_{c}) = (u - l) + \frac{2}{\alpha}(l - \hat{n}_{c}) \cdot 1(\hat{n}_{c} < l) + \frac{2}{\alpha}(\hat{n}_{c} - u) \cdot 1(\hat{n}_{c} > u),$$
(23)

where $1(\cdot)$ is the indicator function, l and u are the values at $\frac{2}{\alpha}$ and $1 - \frac{2}{\alpha}$ quantiles. We then calculate the weighted sum of interval scores by summarizing accuracy across the entire predictive distribution. The overall **WIS** is defined as a linear combination of K interval scores:

$$WIS_{\alpha,\mathbf{K}}(F,\hat{n}_{c}) = \frac{1}{K+0.5} \bigg\{ w_{0} \cdot |n_{c} - \hat{n}_{c}| + \sum_{k=1}^{K} w_{k} \cdot IS_{\alpha_{k}}(F,\hat{n}_{c}) \bigg\},$$
(24)

where $w_k = \frac{\alpha_k}{2}, k = 1, \dots, K$ and $w_0 = \frac{1}{2}$. In this manuscript, we use K = 11 and $\alpha = 0.02, 0.05, 0.1, 0.2, \dots, 0.9$ as in [44].

 $^{^{11}\}mathrm{COVID}\text{-}19$ Forecast Hub has used reports from Johns Hopkins University and we use reports from the New York Times in our previous application

¹²https://www.google.com/covid19/mobility/

¹³https://www.census.gov/

475 3.4.2. Selection Criteria and Comparison results

We first select the models which have passed the screening process in the COVID-19 Forecast Hub [45] based on the following criteria: (1) inclusion of forecasts for at least 25 states for deaths; (2) a complete set of quantiles; and (3) at least 19 eligible weeks. We further retain 9 models that have a better

- ⁴⁸⁰ WIS score than the baseline proposed by the COVID-19 Forecast Hub [45]. The forecasting evaluation starts with the week of 05/04/2020 - 05/10/2020 and then moves forward weekly until 03/15/2021 - 03/21/2021. In each weekly submission, each team makes forecasts for 1 to 4 weeks ahead. One challenge is that each model has a different number of states and forecasting weeks in the sub-
- ⁴⁸⁵ mission. Therefore, following a similar fashion in [45], we calculate the relative evaluation metric for a fair comparison. For each state-week combination, we first divide **HkPR**⁺_m's **WIS** and **MAE** by each models' evaluation results, respectively. We then report the geometric mean of relative scores from all state-week combinations in Table 10. Note that all
- ⁴⁹⁰ hyper-parameters are selected based on the best performance from the previous week for each state-week combination. Therefore, if the value is less than 1, we can suggest that there is an improvement in terms of the error measurement on average.

In terms of confirmed cases (denoted as \mathcal{D}_{conf}^{JHU}), $HkPR_m^+$ has consistently outperformed the selected models in MAE and WIS except for the COVIDhub-

- ⁴⁹⁵ performed the selected models in MAE and WIS except for the COVIDhubensemble in the 4th weeks window ahead. Overall, besides **HkPR**⁺_m, COVIDhubensemble has the most competitive results. COVIDhub-baseline and COVIDhubensemble model are designed by the COVID-19 Forecasts Hub [45]. Here, COVIDhub-baseline serves as a reference point and is generated by the me-
- dian number of cases from the most recent week, and COVIDhub-ensemble aggregates all the submissions to generate an ensemble forecast.

For the COVID-19 Forecast Hub to generate an accurate ensemble, diverse modeling perspectives can be beneficial. The promising results in $\mathcal{D}_{\text{conf}}^{\text{JHU}}$ indicate that forecasts for confirmed cases can benefit from modeling with a Hawkes pro-

- cess that incorporates dynamic covariates. Given that many forecasting groups are using compartmental models, we believe that \mathbf{HkPR}_{m}^{+} can potentially enhance the forecasting accuracy of ensemble forecasts through both its accuracy and diversity. Note that all the weekly forecasts were updated in the anonymous repository¹⁴.
- ⁵¹⁰ We note that, in the retrospective evaluation, our model may be slightlyfavored. Data available when prospective forecasting models were created mayhave been subject to reporting lags or have been updated at a later date. Tocompletely reconstruct such data is a non-trivial task, therefore we acknowledgethis as a limitation of the present work.

 $^{^{14} \}rm https://anonymous.4 open.science/r/d425 dcf9-3 cfb-4f82-a08 c-ee583 ab 36291/$

				$\mathcal{D}^{ m JH}_{ m con}$	IU nf					
Model -		Relative	e WIS			Relative MAE				
-	1 wk	2 wk	3 wk	4 wk	1 wk	2 wk	3 wk	4 wk		
$HkPR_m^+$	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00		
Covid19Sim-Simulator	0.51	0.53	0.60	0.69	0.72	0.70	0.79	0.88		
COVIDhub-baseline	0.80	0.74	0.78	0.87	0.91	0.80	0.84	0.91		
Karlen-pypm	0.68	0.70	0.73	0.77	0.77	0.73	0.74	0.78		
COVIDhub-ensemble	0.94	0.90	0.94	1.01	0.99	0.89	0.94	1.01		
	$\mathcal{D}_{ ext{death}}^{ ext{JHU}}$									
Model -		Relative	e WIS			Relative	e MAE			
-	1 wk	2 wk	3 wk	4 wk	1 wk	2 wk	3 wk	4 wk		
$HkPR_m^+$	1	1	1	1	1	1	1	1		
Covid19Sim-Simulator	0.77	0.89	1.05	1.39	0.88	1.07	1.22	1.61		
MOBS-GLEAM_COVID	0.81	0.87	0.97	1.25	0.88	0.99	1.08	1.32		
UA-EpiCovDA	0.85	0.83	0.92	1.15	1.19	1.17	1.15	1.49		
COVIDhub-baseline	0.86	0.8	0.87	1.11	1.36	1.36	1.35	1.67		
GT-DeepCOVID	0.9	0.95	1.01	1.21	0.92	1	1.05	1.18		
IHME-CurveFit	0.94	0.92	1.12	1.64	0.95	0.99	1.12	1.64		
CMU-TimeSeries	0.96	0.97	1.01	1.08	1.21	1.23	1.39	1.44		
YYG-ParamSearch	0.96	1.03	1.19	1.72	1.07	1.24	1.29	1.89		
COVIDhub-ensemble	1.05	1.08	1.2	1.52	1.5	1.42	1.56	1.94		

Table 10: Relative WIS and MAE on the JHU dataset

The best performance is marked in **bold** and the performances that \mathbf{HkPR}_{m}^{+} has outperformed are marked in **red**. In this table, "N wk" represents the forecasting for N weeks ahead, and " \mathcal{D}_{conf}^{JHU} " and " $\mathcal{D}_{death}^{JHU}$ " are the confirmed cases and deaths collected by Johns Hopkins University, respectively. All the metadata information for the competing models can be found in the following url: https://github.com/reichlab/covid19-forecast-hub/tree/master/data-processed.

515 4. Conclusion

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We showed how Hawkes processes can be combined with spatio-temporal covariates to accurately model COVID-19 transmission and forecast future cases and deaths. The model is competitive with several models used to forecast the pandemic, achieving improved MAE and NDCG scores on a majority of the experiments we conducted. Our hope is that this work will encourage more research into Hawkes process models of disease spreading that incorporate more advanced features and statistical learning principles. As vaccinations are rolled out across the U.S. (given recent FDA approval), local impacts on dynamic reproduction can be flexibly accommodated by our model and used to obtain more accurate and timely forecasts.

One potential direction for future research is investigating the combination of Hawkes process forecasts with compartmental models for improved ensembles. Our results using data from the COVID-19 Forecast Hub indicate this could be a promising direction. Another potential direction for future research is extending

the work here to neural network based point process models [9, 10]. These models may be able to capture more complicated relationships between mobility

patterns, demographics, and transmission. The challenges of such an approach include the potential for over-fitting with added parameters and determining how best to realistically model transmission in a neural point process (analogous to the SIR-Hawkes process), which will be important if neural point processes are to be used in long-term forecasting.

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Hawkes process modeling of COVID-19 with mobility leading indicators and spatial covariates

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Abstract

Hawkes processes are used in machine learning for event clustering and causal inference, while they also can be viewed as stochastic versions of popular compartmental models used in epidemiology. Here we show how to develop accurate models of COVID-19 transmission using Hawkes processes with spatial-temporal covariates. We model the conditional intensity of new COVID-19 cases and deaths in the U.S. at the county level, estimating the dynamic reproduction number of the virus within an EM algorithm through a regression on Google mobility indices and demographic covariates in the maximization step. We validate the approach on both short-term and long-term forecasting tasks, showing that the Hawkes process outperforms several benchmark models currently used to track the pandemic, including an ensemble approach and an SEIR-variant. We also investigate which covariates and mobility indices are most important for building forecasts of COVID-19 in the U.S.

Keywords: COVID-19 forecasting, Hawkes processes, Mobility indices, Spatial covariate

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