

# Nonlinear Modal Regression for Dependent Data with Application for Predicting COVID-19\*

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## Abstract

In this paper, under the stationary  $\alpha$ -mixing dependent samples, we develop a novel nonlinear modal regression for time series sequences and establish the consistency and asymptotic property of the proposed nonlinear modal estimator with a shrinking bandwidth  $h$  under certain regularity conditions. The asymptotic distribution is shown to be identical to the one derived from the independent observations, whereas the convergence rate ( $\sqrt{nh^3}$  in which  $n$  is the sample size) is slower than that in the nonlinear mean regression. We numerically estimate the proposed nonlinear modal regression model by the use of a modified modal-expectation-maximization (MEM) algorithm in conjunction with Taylor expansion. Monte Carlo simulations are presented to demonstrate the good finite sample (prediction) performance of the newly proposed model. We also construct a specified nonlinear modal regression to match the available daily new cases and new deaths data of the COVID-19 outbreak at the state/region level in the United States, and provide forward predictions up to 130 days ahead (from August 24, 2020 to December 31, 2020). In comparison to the traditional nonlinear regressions, the suggested model can fit the COVID-19 data better and produce more precise predictions. The prediction results indicate that there are systematic differences in spreading distributions among states/regions. For most western and eastern states, they have many serious COVID-19 burdens compared to Midwest. We hope that the built nonlinear modal regression can help policymakers to implement fast actions to curb the spread of the infection, avoid overburdening the health system, and understand the development of COVID-19 from some points.

**Keywords:** COVID-19, Dependent data, MEM algorithm, Modal regression, Nonlinear, Prediction.

**JEL Classification:** C01, C14, C22, C53.

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# 1 Introduction

COVID-19 is caused by a coronavirus called SARS-CoV-2 and was identified in Wuhan, the capital city of Hubei province, China, for the very first time in December of 2019. On January 30, 2020, the World Health Organization (WHO) declared the COVID-19 outbreak as a Public Health Emergency of International Concern (PHEIC). COVID-19 is a global threat spreading exponentially rather than linearly, i.e., the number of new cases is proportional to the existing number of cases, which has been dramatically affecting the health and safety of people all over the world. Based on the information from the Johns Hopkins Coronavirus Resource Center (Dong et al., 2020), due to the extensive spread of COVID-19, there are more than 23 million cases of COVID-19 and more than 800 thousand deaths worldwide as of August 23, 2020 (Figure 1 shows that compared to other countries, the United States (US) has suffered from COVID-19 in a more severe way (Yancy, 2020)). In the US alone, since the first US case of COVID-19 infection was identified in Washington state on January 20, 2020, more than 5.6 million COVID-19 cases and 170 thousand COVID-19 deaths have been identified across the US up to August 23, 2020 (Figure 2 indicates the urgency and necessity of providing reliable predictions to understand the growth behavior of COVID-19 in the US). WHO quotes 3.4% as the fatality rate (% people who contract the coronavirus and then die). The ongoing global outbreak of the COVID-19 pandemic, which was eventually classified as a pandemic on March 11, 2020 by WHO, poses serious challenges for countries/regions worldwide in designing tailored methods of epidemic control to provide effective and reliable health protection while allowing as much as possible societal and economic activity. It is unclear to anyone where this pandemic will lead us. In such an emergency situation without globally effective antiviral drugs for treating COVID-19 infections, a reliable prediction model for COVID-19 data is undoubtedly essential for policymakers to implement fast actions to curb the spread of the infection, avoid overburdening the health system, and understand the dynamics of the COVID-19 spread.

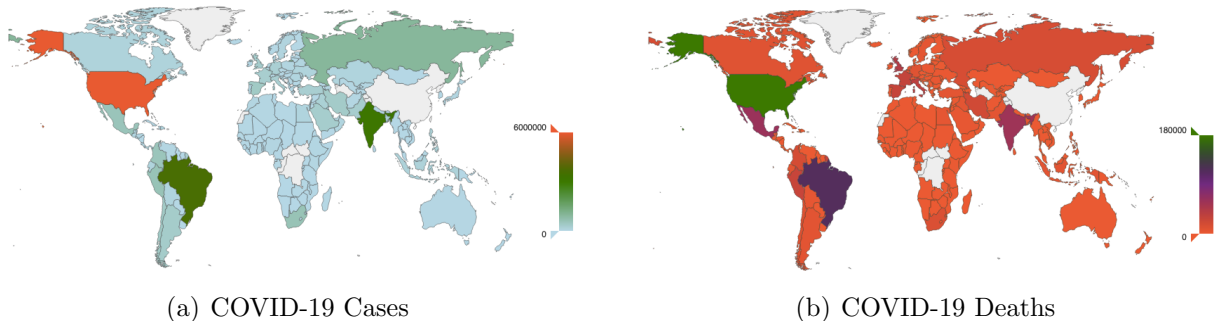


Figure 1: Visualization of the total number of cases and deaths in the world-August 23, 2020; data source: Tencent News <https://new.qq.com/ch/antip/>.

Most of the existing methods for predicting the incidence and prevalence of COVID-19

provided by researchers with backgrounds in epidemiology, biostatistics, and economics focus on some mechanistic models, such as the Susceptible-Exposed-Infectious-Recovered (SEIR) model (Grimm et al., 2020; Hauser et al., 2020; Maugeri et al., 2020), the Institute for Health Metrics and Evaluation (IHME) model (IHME, 2020; Jewell et al., 2020), and the Risk-Based model (Barda et al., 2020; Pueyo, 2020), or some statistical models/distributions (Deb and Majumdar, 2020; Fenga, 2020; Linton et al., 2020; Lu et al., 2020; Verity et al., 2020), such as the time series ARMA model and the machine learning model, for the number of cumulative deaths or cases. However, the accuracy of prediction largely depends on the reliability of data, and it is a widespread opinion in the scientific community that the current official COVID-19 data are often noisy with outliers, biased, skewed, and/or truncated (Linton et al., 2020; Rudnicki and Piliszek, 2020; Tuli et al., 2020). Therefore, the traditional statistical regression model built on mean might provide low accuracy and even misleading prediction results.

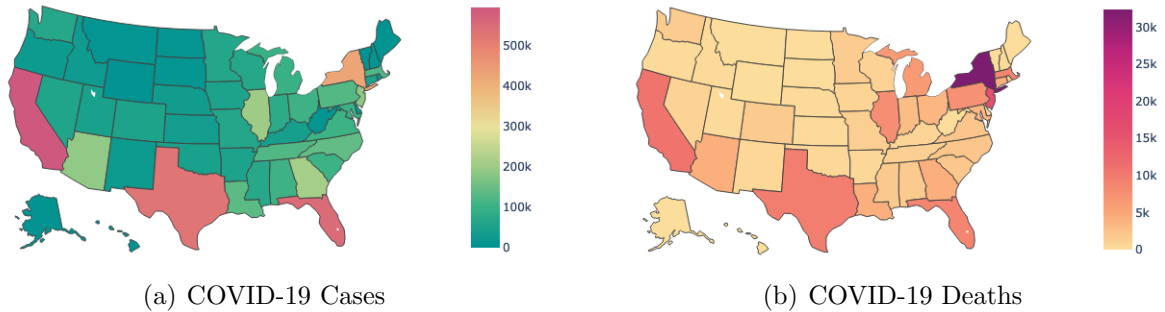


Figure 2: Visualization of the total number of cases and deaths in the US-August 23, 2020; data source: the GitHub repository managed by The New York Times <https://github.com/nytimes/covid-19-data>.

To meet the challenges of the noisy and skewed COVID-19 data, we propose a new statistical regression tool—nonlinear modal regression—that goes beyond the traditional regression models to investigate the dynamic of COVID-19 prevalence in different regions, where the dependent variable of our interest is the official number of daily new cases or new deaths of COVID-19 in a region that could be a state of US (we concentrate on the daily change value as it is a more representative indicator of epidemic severity and an important metric for assessing the effectiveness of COVID-19 regulation). Note that the built model can be applied to conduct predictions for some regions which are still in the early stage of the COVID-19 pandemic or when the COVID-19 pandemic happens again in the future (there is a growing belief among epidemiologists that COVID-19 will behave similarly to the seasonal flu and re-emerge annually in the winter).

It is well-known that the independence assumption for observations is not always valid in empirical applications. There are many statistical/economics analysis problems with high-dimensional data or information network data, where the data exhibit some sort of dependency,

such as Markovian chains, mixing sequences, long-range memory process, and so on. In these cases, the statistical properties of the estimator presented in the papers considering independent identically distributed (i.i.d.) samples may change. Because of this, there has been an extensive literature concerning the estimator for dependent data (Robinson, 1984; Härdle et al., 1997; Pagan and Ullah, 1999; Cai and Ould-Said, 2003; Fan and Yao, 2008; Bester et al., 2011). Nevertheless, nearly all of the existing models/methods regarding dependent samples were considered from the mean or quantile regression and are especially useful when there is no outlier in the data, or the density is not very skewed. When the time series dataset contains several outliers (or aberrant observations) or the data are skewed resulting in non-normally estimated standardized residuals (or heavy-tailed error distributions), which is a common feature of financial/macro-economics/panel time series data, the traditional mean or quantile regression may lose robustness/efficiency or have misspecification (Krief, 2017; Ullah et al., 2021). Thus, modal regression that focuses on the conditional mode, instead of the mean or quantile, of the response variable given the predictor may be more feasible for modeling processes in such cases. Furthermore, when the data are symmetrically distributed, where the modal regression line coincides with the mean regression line, modal regression can overcome the shortcoming of lack of robustness of mean regression to achieve robust and efficient estimators (Yao et al., 2012). To the best of our knowledge, besides Kemp et al. (2020) which considered the estimation of parametric vector autoregressive conditional mode models, there has not been any attempt to estimate modal regression for dependent samples. Substantially different to Kemp et al. (2020), in this paper, we fill the literature gap by focusing on the estimation of a nonlinear modal regression for stationary and weakly dependent samples under  $\alpha$ -mixing condition, which is indeed omnipresent in time series econometrics and is less restrictive than other mixing conditions available in the literature. Due to the space constraint, we leave the nonlinear modal-based robust regression for dependent data derived from mode value in another research, which is based on but significantly different from the proposed nonlinear modal regression in the current paper; see Remark 2.3.

This paper is primarily aimed at applying nonlinear modal regression to understand the characteristics of dependent samples from a mode perspective and settle theoretical properties rigorously. For the simplicity of notations, in what follows, we let  $\{(Y_t, X_t)_{t=1}^n\}$  be a stationary discrete-time random process, defined on the probability space  $(\Omega, \mathcal{F}, \mathcal{P})$ , where  $\Omega$  denotes the sample space,  $\mathcal{F}$  is the  $\sigma$ -algebra (the information) of events, and  $\mathcal{P}$  is a probability measure.  $\{(Y_t, X_t)_{t=1}^n\}$  has the same marginal distribution as  $(Y, X)$ , where  $Y_t$  is the dependent variable of the main interest and  $X_t \in \mathbb{R}^p$  denotes the covariates that may contain the lagged values of  $Y_t$  to reflect the dynamical characteristics of the underlying data generating mechanism. Let  $f(Y | X)$  be the conditional density function of  $Y$  given  $X$ . The conventional regression model usually employs the mean (or the median) of  $f(Y | X)$  to model the relationship between  $Y$  and  $X$ . For example, linear regression assumes that the mean or median of  $f(Y | X)$  is a



linear function of  $X$ . The main distinction of modal regression is to find the most probable value/scenario (i.e., mode) of a dependent variable  $Y$  given covariates  $X$ , which is defined as

$$\text{Mode}(Y | X) = \arg \max_Y f(Y | X). \quad (1.1)$$

When the dimension of  $X$  is not low, estimating (1.1) directly based on nonparametric kernel density estimation will raise many challenges due to the “curse of dimensionality”. We in this paper avoid directly estimating the conditional density by imposing certain model assumptions on the conditional mode of the response given the covariates (assuming that the global mode is uniquely defined), i.e.,  $\text{Mode}(Y | X)$ ; see Section 2 for more details. There is emerging literature on studying modal regression. Due to space limitations, we refer the interested readers to Yao and Li (2014), Chen (2018), Ullah et al. (2021), and the references therein for a comprehensive review of modal regression. Notice that Khardani and Yao (2017) extended the results in Kemp and Santos Silva (2012) to put forward a nonlinear modal regression for the independent samples. However, to the best of our knowledge, there is no existing literature investigating nonlinear modal regression under stationary  $\alpha$ -mixing dependent samples using a kernel smoother, which is one of the main contributions of the current paper. It is noteworthy that compared to mean or median regression, modal regression has the following noticeable advantages (Yao and Li, 2014; Ullah et al., 2021): i) better for reflecting the characteristics of skewed data; ii) better point prediction and narrower prediction intervals; iii) more robust to outliers and certain forms of measurement error; iv) consistent estimation even for truncated data. Therefore, the modal regression can overcome the limitations of the traditional existing regressions and naturally provide reliable (prediction) models for the noisy COVID-19 data, which is the main innovation of the present paper contributing to the rapidly growing literature on predicting the spread of the current COVID-19 pandemic. We also show a new and interesting theoretical result that the asymptotic theorem for the proposed nonlinear modal estimator under stationary  $\alpha$ -mixing dependent samples is the same as that for independent data under certain conditions, indicating the asymptotic negligence of the dependence. This remarkable result is intrinsic for nonparametric estimation for dependent samples and was already observed in the mean regression estimation (Cai and Ould-Said, 2003). Compared to the mean regression estimator, the modal regression estimator depends heavily on error term observations which are confined to the neighborhood of a given point (i.e., zero) and will unexpectedly have a much slower convergence rate (the modal estimation requires a shrinking bandwidth  $h$  due to the use of a small portion of data around the mode), which is the price to be paid in order to estimate mode (Parzen, 1962). The proposed nonlinear modal estimator is relatively simple to implement, where we develop a computationally efficient MEM algorithm in conjunction with Taylor expansion to numerically estimate it.

Generally, most new confirmed cases are infected via contact with the new confirmed cases

in recent days, indicating the necessity of incorporating lagged value for analyzing and predicting COVID-19 new cases. [Ho et al. \(2020\)](#) introduced a flexible statistical model for the infections and deaths caused by COVID-19 in New Zealand, in which the growth rate of the cumulative number of cases depends on the current cumulative number of cases. [Li and Linton \(2021\)](#) developed a quadratic time trend model that was applied to the log of new cases and obtained satisfying results for the trajectory of the epidemic in most countries. Based on these observations, we apply the proposed nonlinear modal regression to model the log of new cases/deaths as a function of time (to capture the trend or bell-shaped curve) and its own one-step lagged value (to capture the dynamics by autoregressive fluctuations) based on the general structure of the effects and process of infection from a mode perspective; see [Section 3](#) for more details on the model setting for COVID-19 data. Under the constraint imposed by a reasonable length of the paper, we compare the performance of nonlinear modal regression to that of nonlinear mean and median regressions for US COVID-19 data in the paper (for the sake of thoroughness, we also list the results associated with COVID-19 data obtained from the robust nonlinear mean regression with the bisquare weight in [Online Appendix B](#)). We emphasize that the outbreak spreads of COVID-19 are largely affected by the policies and social responsibilities of each state/region, it will be interesting in the future to compare the prediction results from the proposed model to some well-known predictions such as those from the IMHE model, SIR models in epidemiology, machine learning methods, or other models that can take policy effects into account).

The results indicate that the newly proposed model could have better fit performance in terms of  $R^2$  (the coefficient of determination) for most states/regions in the US. We also use Mean Square Error ( $MSE$ ) and Mean Absolute Percentage Error ( $MAPE$ ) to compare the out-of-sample prediction performance of the proposed nonlinear modal regression to that of nonlinear mean and median (and robust) regressions for the last 20 days of the samples, where we show that nonlinear modal regression can have considerably more precise predictions. We then apply the proposed nonlinear modal regression to predict COVID-19 new cases and new deaths in the US. Based on the prediction results up to the next 130 days (from August 24, 2020 to December 31, 2020), we can observe that there are systematic differences in spreading distributions across US states. Some states are showing a clear decreasing trend in the number of new cases and new deaths, such as Connecticut, Illinois, Maine, Maryland, Massachusetts, New Hampshire, New Jersey, New York, Pennsylvania, among others, while others, such as Alabama, Arkansas, California, Florida, Georgia, Mississippi, Montana, North Carolina, North Dakota, Oregon, Texas, Utah, Washington, and so on, are still in the first wave of the COVID-19 outbreak. Among the states, California, Florida, Texas, and Georgia are the worst affected ones in terms of the number of predicted new cases and new deaths for the next 130 days. For most western and eastern states, they have many serious COVID-19 burdens compared to the Midwest. It is interesting to note that the prediction results may reflect the effect of different

possible policy interventions, which can be interpreted as holding the current policies in place or under minimal interventions in each state/region. With the newly developed nonlinear modal regression, we hope that the prediction results can provide some timely information (i.e., turning point) to help policymakers to implement fast actions to curb the spread of the infection and avoid overburdening the health system.

The remainder of this paper is organized as follows. In Section 2, we introduce a nonlinear modal regression for dependent samples under the stationary  $\alpha$ -mixing condition and develop an efficient modal estimation algorithm. We also present the asymptotic distributional theory for the resulting estimator under mild conditions, which gives guidelines for practically selecting a reliable bandwidth. Monte Carlo simulations are conducted to show the good finite sample performance of the proposed model. A specified nonlinear modal regression is introduced in section 3. Based on the built model, we produce a modal multi-step-ahead point forecast framework for COVID-19 new cases and new deaths data, and present the out-of-sample prediction results of the behavior of COVID-19 at the state/region level in the US. The paper is concluded with some remarks in Section 4. We put additional numerical results, list all figures related to the prediction results, and outline the proofs for the main theorems in the online appendix.

## 2 Nonlinear Modal Regression

In order to streamline the discussion, we start in this section with the nonlinear modal estimator for dependent samples, where the numerical solutions are obtained via a modified MEM algorithm (Li et al., 2007; Yao, 2013) with the help of a first order Taylor expansion. Under the assumption of  $\alpha$ -mixing, we then present the asymptotic property and optimal bandwidth.

### 2.1 Nonlinear Modal Estimator

As mentioned previously, the traditional method of estimating (1.1) is to directly estimate the conditional density  $f(Y | X)$  nonparametrically based on the multivariate kernel density estimation; see the related discussions in Chen et al. (2016). However, due to the “curse of dimensionality”, such a method is practically infeasible when the dimension of covariates is moderate or high, which also contributes to the lack of enough research interest in modal regression. In this paper, similar to mean or quantile regression, we propose estimating the modal regression (1.1) by imposing some model assumptions on  $Mode(Y | X)$  directly (assuming that it is uniquely defined) to avoid the “curse of dimensionality” of the fully nonparametric kernel method. In particular, we assume the following baseline model

$$\begin{cases} Y_t = r(X_t, \beta) + \epsilon_t, \\ Mode(Y_t | X_t) = r(X_t, \beta), \quad t = 1, \dots, n, \end{cases} \quad (2.1)$$

where  $t$  represents calendar day that equals to one for the first date of the data,  $\beta \in \Theta$  is an unknown parameter vector with dimension  $p$ ,  $\Theta$  is the known compact parameter space,  $r(\cdot) : \mathbb{R}^p \times \Theta \rightarrow \mathbb{R}$  is a parametric nonlinear function measurable on  $\mathbb{R}^p$  for each  $\beta$  in  $\Theta$ , and  $\{\epsilon_t\}_{t=1}^n$  is a sequence of stochastic random variables with  $\text{Mode}(\epsilon_t | \mathcal{F}_t) = 0$  almost surely (a.s.) for every  $t$  in which  $\mathcal{F}_t$  is the  $\sigma$ -field generated by  $\{X_s, \epsilon_s\}_{s \leq t}$ . Different from the most existing regressions, we do not impose any second moment conditions on  $\epsilon_t$ , thus it can be conditional homoskedastic or conditional heteroskedastic. It is worth pointing out that in order to illustrate the applicability of nonlinear modal regression for time series data in a more general setting, we focus on dependent observations. However, (2.1) could also be an autoregressive time series model with finite order  $p$ , i.e.,  $\text{Mode}(Y_t | Y_{t-1}) = r(\{Y_{t-l}\}_{l=1}^p, \beta)$ , which characterizes the nonlinearity in terms of lags and could be considered as a special case of time series model in this section. The form of  $r(\cdot)$  for analyzing the COVID-19 data will be discussed in Section 3. Then, the modal parameter  $\beta$  can be estimated by maximizing the following kernel-based objective function given observations  $\{(Y_t, X_t)\}_{t=1}^n$  and knowledge of  $r(\cdot)$

$$Q_n(\beta) = \frac{1}{nh} \sum_{t=1}^n K\left(\frac{Y_t - r(X_t, \beta)}{h}\right), \quad (2.2)$$

where  $K(\cdot)$  is a nonnegatively symmetric kernel function such as the Gaussian kernel (i.e.,  $K(t) = (2\pi)^{-1/2} \exp[-(1/2)t^2]$ ) that we will use by default in this paper, and  $h := h(n)$  is a bandwidth that is assumed to go to 0 with  $n$  going to infinity (“ $:=$ ” denotes “equals by definition”). To keep the notation simple, we however suppress  $n$  throughout the paper. Notice that  $K(\cdot)$  is a function following the same rules as a probability density function, for example, it is positive and integrable. However, the role of bandwidth  $h$  (control mode) is different from that in nonparametric regression (control smoothness). According to Yao et al. (2012), the choice of kernel function is not very important in modal regression compared to the choice of bandwidth. We choose the Gaussian kernel in this paper for the sake of simplicity. In particular, we can obtain an explicit expression for the M-Step in Algorithm 1.

**Remark 2.1.** When  $r(X_t, \beta) = \beta^*$ , only an intercept term,  $Q_n(\beta^*)$  is a kernel density estimate of  $Y$ , and thus the maximizer of (2.2) is the estimated mode of  $f(Y)$ . Here, we extend this kernel-type objective function to estimate the modal regression parameter  $\beta$  in the regression setting. When  $r(X_t, \beta) = X_t^T \beta$  in which  $T$  represents the transpose of a matrix or a vector, the modal regression (2.1) is simplified to the linear modal regression (Kemp and Santos Silva, 2012; Yao and Li, 2014). Note that if  $K(t) = 2^{-1}I(|t| \leq h)$ , a uniform kernel, then (2.2) tries to find the curve  $r(X_t, \hat{\beta})$  such that the band  $r(X_t, \hat{\beta}) \pm h$  contains the largest number of response  $Y_t$ , where  $\hat{\beta}$  is the modal estimator. Therefore, the modal regression provides more meaningful point predictions and shorter prediction intervals than the mean regression.

It is well-known that the estimation of nonlinear models is a notoriously difficult problem,

especially for modal regression, as maximizing (2.2) does not have an explicit solution. We thus develop a modified MEM Algorithm 1 originally proposed by Li et al. (2007) and Yao (2013) to simplify the computations, which decomposes the optimization (2.2) into E-Step and M-Step. Given the initial value  $\beta^{(0)}$  (e.g., nonlinear least squares (NLS) estimate), we shall repeat the two steps in the algorithm until it converges. Note that if  $r(X_t, \beta)$  is a linear function of  $X_t$ , then M-Step is just a weighted LS estimation and has an explicit solution. To simplify the computation of M-Step for a general nonlinear function  $r(\cdot)$ , we approximate  $r(X_t, \beta)$  by a first order Taylor expansion around the current parameter estimate. It can be proved that each iteration of the above algorithm monotonically nondecreases the objective function (2.2) following the procedures in Yao and Li (2014), i.e., at each iteration  $Q_n(\beta^{(g+1)}) \geq Q_n(\beta^{(g)})$  and the equality holds if and only if  $\beta^{(g+1)} = \beta^{(g)}$ . Therefore, the algorithm is very stable and converges. However, for the bandwidth  $h$  with a small value, the objective function may have multiple maxima, and there is no guarantee that the MEM algorithm will converge to the global maximizer. Accordingly, it is important to try different starting points on each occasion to compare the values of the target function to choose the best optimal one (Yao and Li, 2014; Ullah et al., 2021).

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**Algorithm 1** MEM Algorithm for Nonlinear Modal Regression

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**E-Step.** Calculate the weight  $\pi(t | \beta^{(g)})$

$$\pi(t | \beta^{(g)}) = \frac{K\left(\frac{Y_t - r(X_t, \beta^{(g)})}{h}\right)}{\sum_{t=1}^n K\left(\frac{Y_t - r(X_t, \beta^{(g)})}{h}\right)} \propto K\left(\frac{Y_t - r(X_t, \beta^{(g)})}{h}\right),$$

where  $g$  is the iteration indicator.

**Expansion.** Approximate  $r(X_t, \beta)$  by a first order Taylor expansion around  $\beta^{(g)}$

$$r(X_t, \beta) \approx r(X_t, \beta^{(g)}) + \frac{\partial r(X_t, \beta)}{\partial \beta^T} \Big|_{\beta=\beta^{(g)}} (\beta - \beta^{(g)}).$$

**M-Step.** Update  $\beta^{(g+1)}$  by

$$\begin{aligned} \beta^{(g+1)} &= \arg \max_{\beta} \sum_{t=1}^n \left\{ \pi(t | \beta^{(g)}) \log K\left(\frac{Y_t - r(X_t, \beta)}{h}\right) \right\} \\ &= \left[ \sum_{t=1}^n \pi(t | \beta^{(g)}) \frac{\partial r(X_t, \beta^{(g)})}{\partial \beta} \frac{\partial r(X_t, \beta^{(g)})}{\partial \beta^T} \right]^{-1} \left[ \sum_{t=1}^n \pi(t | \beta^{(g)}) \frac{\partial r(X_t, \beta^{(g)})}{\partial \beta} Y_t^{(g)} \right], \end{aligned}$$

where  $Y_t^{(g)} = Y_t - r(X_t, \beta^{(g)}) + \frac{\partial r(X_t, \beta^{(g)})}{\partial \beta^T} \beta^{(g)}$ .

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Based on the above algorithm, it can be seen that the major difference between the mean regression by the LS estimation and the modal regression lies in the weight  $\pi(t | \beta^{(g)})$  used in

E-Step. For the LS estimation, each observation has an equal weight  $1/n$ . On the other hand, for modal regression estimate, the weight  $\pi(t | \beta^{(g)})$  calculated in E-step depends on how close  $Y_t$  is to the modal regression curve  $r(X_t, \beta)$ . This weighting scheme allows modal regression to reduce the effect of observations far away from the modal regression curve to achieve robustness, which is one of the advantages of modal regression over mean regression.

## 2.2 Asymptotic Property

Before proceeding to the asymptotic theorem for the estimator under the  $\alpha$ -mixing assumption, it is convenient to introduce some notations that will be used in the remaining part of this section. We define  $T_n(x) = T(x) + o_p(s_n)$  (or  $O_p(s_n)$ ) uniformly for  $x \in \mathcal{X}$  if  $\sup_{x \in \mathcal{X}} |T_n(x) - T(x)| = o_p(s_n)$  (or  $O_p(s_n)$ ), and use “ $\xrightarrow{d}$ ” to represent convergence in distribution. We say that  $f(n) = o(g(n))$  if for all  $c > 0$ , there exists some  $k > 0$  (not depend on  $n$ ) such that  $0 \leq f(n) < cg(n)$  for all  $n \geq k$ . Let  $\|\cdot\|$  denote the Euclidean norm, i.e.,  $\|A\| = [tr(AA^T)]^{1/2}$  in which  $tr(A)$  is the trace of the matrix or vector  $A$ . For positive sequences  $\{a_n\}$  and  $\{b_n\}$ , we write  $a_n \asymp b_n$  if  $a_n/b_n + b_n/a_n$  is bounded for all large  $n$ . To facilitate the derivation of the consistency and asymptotic theorem for the estimator from (2.2) in a general framework, we impose the following regularity conditions.

- C1 The true value of parameter  $\beta_0$  defined in (2.1) is in the interior of the known compact parameter space  $\Theta$ , which is a subset of  $\mathbb{R}^p$ .
- C2 The kernel function  $K(\cdot)$  is a nonnegatively symmetric density function with bounded support and integrates to one. It is four times continuous differentiable with all derivatives bounded in absolute value. Furthermore,  $\int t^{2+\delta} K^{2+\delta}(t) dt < \infty$  with probability one in which  $\delta \in [0, 1)$  is a constant.
- C3 The regression function  $r(\cdot)$  has at least a continuous first derivative on an open set that contains the true parameter point  $\beta_0$ . In addition,  $n^{-1} \sum_{t=1}^n \{\partial r(X_t, \beta)/\partial \beta\} \{\partial r(X_t, \beta)/\partial \beta\}^T$  converges to a finite positive definite matrix at  $\beta = \beta_0$ .
- C4 The conditional density of  $\epsilon$  given  $X$  denoted by  $q(\epsilon | X) : \mathbb{R} \rightarrow \mathbb{R}$  is bounded away from zero and infinity, and has the third continuous derivative.  $q^{(c)}(\cdot | X)$  denotes the  $c$ th derivative of  $q(\cdot | X)$ . Furthermore,  $q(\epsilon | X) < q(0 | X)$  for all  $\epsilon \neq 0$  and  $X$ , and the first derivative  $q^{(1)}(\epsilon | X) = 0$ .
- C5  $\{(Y_t, X_t)\}$  is a stationary  $\alpha$ -mixing process, and the mixing coefficient  $\rho(n) = \sup_{A \in \mathcal{F}_{-\infty}^0, B \in \mathcal{F}_t^\infty} |P(A \cap B) - P(A)P(B)|$  tending to zero for  $n \rightarrow \infty$  satisfies  $\sum_{n \geq 1} n^\gamma (\rho(n))^{\delta/(2+\delta)} < \infty$  for some  $\gamma > \delta/(2+\delta)$ , where  $\delta$  is a constant given in C2 and  $\mathcal{F}$  is the  $\sigma$ -algebra of events generated by the random variables  $\{(Y_t, X_t)\}$ . Moreover, there is a sequence of positive integers  $d_n$  such that  $d_n \rightarrow \infty$ ,  $d_n h \rightarrow 0$ , and  $h^4 \sum_{k=d_n}^n [\rho(k)]^{\delta/(2+\delta)} = o(nh^{-3})$ .



C6 As  $n \rightarrow \infty$ ,  $n^{-1} \sum_{t=1}^n q^{(2)}(0 \mid X_t) \left\{ \frac{\partial r(X_t, \beta)}{\partial \beta} \right\} \left\{ \frac{\partial r(X_t, \beta)}{\partial \beta} \right\}^T$  converges in probability to a negative definite matrix.

Most of the above conditions have been used in [Kemp and Santos Silva \(2012\)](#), [Yao and Li \(2014\)](#), and [Ullah et al. \(2021\)](#). *Condition C1* is a common condition and can be easily satisfied in practice, as there are no constraints on  $\beta$ . The bounded support in *Condition C2* imposed on the kernel function  $K(\cdot)$  is for the brevity of proofs, and may be relaxed somewhat if we impose certain restrictions on the tail of the kernel function; for example, the Gaussian kernel is allowed ([Ullah et al., 2021](#)), which is the default kernel used in this paper. *Condition C3* is a commonly used condition on the smoothness of the nonlinear function and the information matrix to ensure the existence of the asymptotic mean and variance for the proposed nonlinear modal estimator, as the modal estimator  $\hat{\beta}$  must satisfy  $-\frac{1}{nh^2} \sum_{t=1}^n K^{(1)}\left(\frac{Y_t - r(X_t, \hat{\beta})}{h}\right) r^{(1)}(X_t, \hat{\beta}) = 0$  where  $K^{(1)}(\cdot)$  and  $r^{(1)}(\cdot)$  are the first derivatives of  $K(\cdot)$  and  $r(\cdot)$ , respectively. *Condition C4* implies a certain smoothness of  $q(\epsilon_t \mid X_t)$  in the neighborhood of zero, which is necessary for identification. It imposes that the conditional density of  $\epsilon$  has a well-defined global mode at zero ([Kemp and Santos Silva, 2012](#); [Ullah et al., 2021](#)). It is to be conceded that this assumption is used for simple illustration; when the population is not homogeneous, the proposed method could also be applied to the multimode setting to capture different modal regression lines, under which the newly developed nonlinear modal regression can reveal the possible heterogeneity of COVID-19 development patterns across different states/regions. *Condition C5* is a condition on the data generating process that permits, and is the standard requirement for the  $\alpha$ -mixing process, which is used to control the dependence between two random variables as the time distance increases. It is reasonably weak and is known to be satisfied by many stochastic processes, such as the stationary Markov process and the stationary autoregressive-moving average process. A sufficient condition for the mixing coefficient  $\rho(n)$  to satisfy *Condition C5* is to set  $\rho(n) = O(n^{-d})$  for some  $d > 2(\gamma + 1)/\gamma$  ([Cai and Ould-Said, 2003](#)). When  $\{(Y_t, X_t)\}_{t=1}^n$  are independent in which  $\delta = 0$ , the results in this paper also hold. *Condition C6* is the classic rank condition placing restrictions on the moments of covariates, which is necessary for deriving the asymptotic property of the proposed nonlinear modal estimator. All conditions related to bandwidth  $h$  are specified for each of the theorems stated below.

We are now in a position where we can state the main asymptotic results for the proposed nonlinear modal estimator. The results are presented in the following Theorems [2.1](#) and [2.2](#), where the modal convergence rate  $\sqrt{nh^3}$  can be considered as a new one in the literature of nonlinear regression models for dependent samples.

**Theorem 2.1.** *Under the regularity conditions C1-C6, with probability approaching one, as  $n \rightarrow \infty$ ,  $h \rightarrow 0$ , and  $nh^5 \rightarrow \infty$ , there exists a consistent maximizer  $\hat{\beta}$  of [\(2.2\)](#) such that*

$$\|\hat{\beta} - \beta_0\| = O_p\left((nh^3)^{-1/2} + h^2\right).$$



**Theorem 2.2.** *With  $nh^5 = O(1)$ , under the same conditions as Theorem 2.1, the parameter satisfying the consistency result in Theorem 2.1 has the following asymptotic result*

$$\sqrt{nh^3} \left[ \hat{\beta} - \beta_0 - \frac{h^2}{2} J^{-1} M \{1 + o_p(1)\} \right] \xrightarrow{d} N \left\{ 0, \int t^2 K^2(t) dt J^{-1} L J^{-1} \right\}.$$

*If we allow  $nh^5 \rightarrow 0$ , the asymptotic theorem becomes*

$$\sqrt{nh^3} (\hat{\beta} - \beta_0) \xrightarrow{d} N \left\{ 0, \int t^2 K^2(t) dt J^{-1} L J^{-1} \right\},$$

where  $J = \mathbb{E} \left[ q^{(2)}(0 | X_t) \left\{ \frac{\partial r(X_t, \beta)}{\partial \beta} \right\} \left\{ \frac{\partial r(X_t, \beta)}{\partial \beta} \right\}^T \right]$ ,  $M = \mathbb{E} \left[ q^{(3)}(0 | X_t) \left\{ \frac{\partial r(X_t, \beta)}{\partial \beta} \right\} \right]$ , and  $L = \mathbb{E} \left[ q(0 | X_t) \left\{ \frac{\partial r(X_t, \beta)}{\partial \beta} \right\} \left\{ \frac{\partial r(X_t, \beta)}{\partial \beta} \right\}^T \right]$  at  $\beta = \beta_0$ .

The proofs of the above two theorems are outlined in Online Appendix C. For Theorem 2.1, the first term  $(nh^3)^{-1/2}$  in the convergence rates characterizes the magnitude of the estimation variance, while the second term  $h^2$  characterizes the magnitude of the estimation bias. It is necessary to emphasize that these results are consistent with those in Yao and Li (2014) and Ullah et al. (2021) for the i.i.d. data. Theorem 2.2 shows that the asymptotic bias term is mainly determined by the bandwidth and can be successfully removed under certain undersmoothing conditions. However, the asymptotic mean squared error (AMSE) optimal bandwidth  $h$  satisfies  $h \asymp n^{-1/7}$ , which does not meet the condition that  $nh^5 \rightarrow 0$ . Hence, undersmoothing is required, i.e.,  $\lim_{n \rightarrow \infty} \sqrt{nh^7} \rightarrow 0$ , which will be incorporated into this paper when selecting bandwidth in practice. We remark that the asymptotic results hold for both i.i.d. data and dependent samples under mild conditions including strongly mixing ( $\alpha$ -mixing). The asymptotic negligence of dependence with a large sample size is intrinsic to nonparametric estimation for dependent samples and it was already observed in the mean regression estimator; see Cai and Ould-Said (2003). This should be expected as a heuristic principle for nonlinear modal regression as well due to the fact that under the  $\alpha$ -mixing process, the covariance between random variables  $\epsilon_t$  and  $\epsilon_j$  such that  $\epsilon_t, \epsilon_j \in (\epsilon - h, \epsilon + h)$  is dominated by the variance of  $\epsilon_t$  through the conditions imposed on the smoothing parameter; see Lemma 1 in Online Appendix C. Thus, the dependence between the random variables  $\epsilon_t$  and  $\epsilon_j$  in a short interval is of “short memory” which makes them behave as if they were independent (Härdle et al., 1997).

**Remark 2.2.** *The convergence rate of the proposed nonlinear modal estimator  $\hat{\beta}$ ,  $n^{2/7}$  with the MSE-optimal bandwidth, is slower than the root- $n$  convergence rate of the traditional NLS estimator, which is the cost we need to pay in order to estimate the conditional mode (Parzen, 1962). How to improve the convergence rate of the nonlinear modal estimator needs to be researched further in the future. For example, we may assume a certain analytical relationship*

among mean, median, and mode to help estimate the nonlinear modal regression line. Nevertheless, for skewed data with moderate sample size, the modal regression usually provides better prediction performance than the mean and median regressions, as the mode is trying to capture the most likely data points; see the Monte Carlo simulation results in Yao and Li (2014) for cross sectional data and Ullah et al. (2021) for fixed effects panel data. Our analysis of COVID-19 time series data in Section 3 also shows the superior prediction performance of the proposed nonlinear modal regression over the nonlinear mean and median (and robust) regressions.

**Remark 2.3.** *It is observed that the proposed nonlinear modal regression focuses on asymmetric data to reveal the characteristics of the data that have been neglected by mean or quantile regression. In practice, it is also common to encounter symmetric data with outliers/aberrant observations or heavy tails. In such a case, we might still be interested in estimating the mean regression, while the proposed modal regression may not be directly applicable owing to the slower convergence rate and the traditional LS estimator is not robust to outliers or heavy tailed data. One way in the literature to handle this kind of data is to utilize robust regression models, like M-estimation, which will lose efficiency for normal errors. We can then supplement the robust regression literature by demonstrating that with symmetric data having only one mode at the center and a heavy-tailed distribution, the nonlinear modal regression can be used alternatively as a robust regression to achieve robustness and efficiency. Compared to the proposed nonlinear modal regression, the main feature of the nonlinear modal-based robust regression is that we treat bandwidth  $h$  as a constant, which does not depend on sample size. Under suitable conditions, we can establish the asymptotic normality for the proposed modal-based robust estimator with  $\sqrt{n}$  consistency, and demonstrate that the modal-based robust estimator could be more efficient than the NLS estimator with a heavy-tailed distribution, or as efficient as the NLS estimator with a normal distribution. Due to the space constraint, we leave the detail of the nonlinear modal-based robust regression for dependent data derived from mode value in another research.*

## 2.3 Optimal Bandwidth

Compared to the bandwidth selection method for density estimation in order to estimate modes, it is more challenging to calculate the optimal bandwidth for modal regression, as the value of bandwidth can strongly affect the regression estimates. Particularly, if bandwidth is large enough, the modal regression will instead capture the mean estimate; see Remark 2.3. In addition, bandwidth plays an important role in estimating dependent observations, as the dependency can be controlled with the observations in a small window. There exist some methods for selecting the optimal bandwidths for nonparametric estimation of conditional modes based on kernel density estimation; see Chen (2018) and Zhou and Huang (2019). However, the methods for bandwidth selection in modal regression by directly imposing structural assumptions on  $\text{Mode}(Y | X)$  are rather limited. One of them is related to the plug-in bandwidth selection

method for linear modal regression, which was presented in Yao and Li (2014) and Ullah et al. (2021) by replacing the unknown quantities with the corresponding estimates. Nevertheless, such a plug-in method places a heavy burden on calculation. In this part, we discuss the asymptotic optimal bandwidth for  $h$  and suggest a simple data adaptive method to obtain the bandwidth.

To derive the asymptotically optimal bandwidth, we minimize the *AMSE* of the proposed nonlinear modal estimator, i.e.,

$$\mathbb{E} \left\{ (\hat{\beta} - \beta_0)^T W (\hat{\beta} - \beta_0) \right\} \approx M^T J^{-1} W J^{-1} M h^4 / 4 + (n h^3)^{-1} \text{tr} (J^{-1} L J^{-1} W) \int t^2 K^2(t) dt, \quad (2.3)$$

where the symbol “ $c_n \approx d_n$ ” indicates that  $c_n/d_n \rightarrow 1$  as  $n \rightarrow \infty$  and  $W$  is a weight function, such as an identity matrix, reflecting which coefficient is more important in inference. Therefore, the asymptotically optimal bandwidth  $h$  is

$$\hat{h}_{opt} = \left[ \frac{3 \int t^2 K^2(t) dt \text{tr} (J^{-1} L J^{-1} W)}{M^T J^{-1} W J^{-1} M} \right]^{1/7} n^{-1/7}. \quad (2.4)$$

If  $W = (J^{-1} L J^{-1})^{-1}$ , which is proportional to the inverse of the asymptotic variance of  $\hat{\beta}$ , then  $\text{tr} (J^{-1} L J^{-1} W) = p$ , and we can have

$$\hat{h}_{opt} = \left[ \frac{3p \int t^2 K^2(t) dt}{M^T L^{-1} M} \right]^{1/7} n^{-1/7}. \quad (2.5)$$

The optimal bandwidth in the above equation depends on the unknown density  $q(\cdot)$  in a complicated manner, which is not available in practice. However, the expression can give some guidelines on how to select the optimal data-driven bandwidth in practice. To simplify the calculations, we can follow Kemp and Santos Silva (2012) to choose the bandwidth, and let  $\hat{h} = 1.6 \text{MAD} n^{-0.143}$  (-0.13 comes from the rate -1/7 and undersmoothing requirement) be a normalized median absolute deviation (MAD) estimate, where

$$\text{MAD} = \text{med}_j \{ |(Y_j - r(X_j, \hat{\beta}_m)) - \text{med}_t(Y_t - r(X_t, \hat{\beta}_m))| \}, \quad (2.6)$$

$\hat{\beta}_m$  represents the corresponding mean estimate, and *med* representing the median value. Besides the above procedure, researchers could also follow the cross-validation method or the weighted integrated squared error method developed in Zhou and Huang (2019) to select the bandwidth.

## 2.4 Monte Carlo Experiments

To illustrate that the asymptotic result investigated in the above subsection provides a good approximation of the finite sample behavior of the proposed nonlinear modal estimator, we

present two numerical examples based on Monte Carlo experiments (one is shown in Online Appendix A). We mainly focus on asymmetric data and use DGP to represent the data generating process in what follows. For comparison, both nonlinear modal regression and mean regression are considered to estimate parameters with  $M = 200$  replications and sample size  $n \in \{200, 400, 600, 1000\}$ . We examine how estimators behave in finite samples in terms of bias, standard error, and MSE,

$$MSE(\hat{\beta}) = \frac{1}{M} \sum_{j=1}^M \|\hat{\beta}^{(j)} - \beta\|^2$$

in which  $\hat{\beta}^{(j)}$  is the estimate in the  $j$ th replication and  $\beta$  is the true value. In order to validate the asymptotic normality property, we present the shape of the empirical density of the standardized (recentered and rescaled) modal estimate. The coverage probabilities to assess the prediction performance of the proposed nonlinear modal regression are reported as well.

**DGP 1** We generate the dependent data from the following model

$$Y_t = X_{1,t} + \exp(2X_{2,t}) + X_{1,t}\epsilon_t,$$

where  $X_{1,t} = -0.3X_{1,t-1} + u_{1,t}$ ,  $u_{1,t} \stackrel{i.i.d.}{\sim} \mathcal{N}(0, 0.8^2)$ ,  $X_{2,t} = 0.4X_{2,t-1} + u_{2,t}$ ,  $u_{2,t} \stackrel{i.i.d.}{\sim} \mathcal{N}(0, 0.5^2)$ , and  $\epsilon_t \stackrel{i.i.d.}{\sim} 0.5\mathcal{N}(-1, 2.5^2) + 0.5\mathcal{N}(1, 0.5^2)$  with  $\mathbb{E}(\epsilon_t) = 0$  and  $Mode(\epsilon_t) = 1$  (Yao and Li, 2014; Ullah et al., 2021). We then have

$$\begin{cases} \text{Mean Regression: } \mathbb{E}(Y_t | X_{1,t}, X_{2,t}) = X_{1,t} + \exp(2X_{2,t}), \\ \text{Modal Regression: } Mode(Y_t | X_{1,t}, X_{2,t}) = 2X_{1,t} + \exp(2X_{2,t}). \end{cases}$$

Notice that the median value of  $\epsilon_t$  is around 0.67, which indicates that the nonlinear median regression line is  $Median(Y_t | X_{1,t}, X_{2,t}) = 1.67X_{1,t} + \exp(2X_{2,t})$ . For space considerations, we do not present results for median estimates, but they are available upon request.

Table 1: Results of Simulations—DGP 1

Modal Estimation					Mean Estimation			
Sample Size	$\beta_1$ (SE)	MSE( $\beta_1$ )	$\beta_2$ (SE)	MSE( $\beta_2$ )	$\beta_{m,1}$ (SE)	MSE( $\beta_{m,1}$ )	$\beta_{m,2}$ (SE)	MSE( $\beta_{m,2}$ )
$n=200$	1.9329 (0.2288)	0.0566	2.0078 (0.0574)	0.0033	1.0073 (0.2371)	0.0560	1.9974 (0.0402)	0.0016
$n=400$	1.9604 (0.0924)	0.0101	1.9995 (0.0313)	0.0010	1.0000 (0.1816)	0.0328	2.0008 (0.0241)	0.0006
$n=600$	1.9620 (0.0817)	0.0081	2.0003 (0.0237)	0.0006	0.9956 (0.1440)	0.0207	1.9999 (0.0193)	0.0004
$n=1000$	1.9620 (0.0603)	0.0051	1.9985 (0.0178)	0.0003	0.9886 (0.1043)	0.0110	1.9990 (0.0153)	0.0002
True Value	$\beta_1 = 2$		$\beta_2 = 2$		$\beta_{m,1} = 1$		$\beta_{m,2} = 2$	

The simulation results are summarized in Table 1 ( $\beta_m$  represents the coefficients of mean regression), from which we can see that the proposed nonlinear estimation procedure can recover

modal coefficients well with finite samples. Also, when the sample size increases, the performance of all estimators improves as expected, both in terms of biases and standard errors. With skewed data where the mean, median, and mode differ by a location shift, it is necessary to perform the nonlinear modal regression to complement the nonlinear mean or quantile regression and capture the most likely effect that the existing regressions cannot directly reveal.

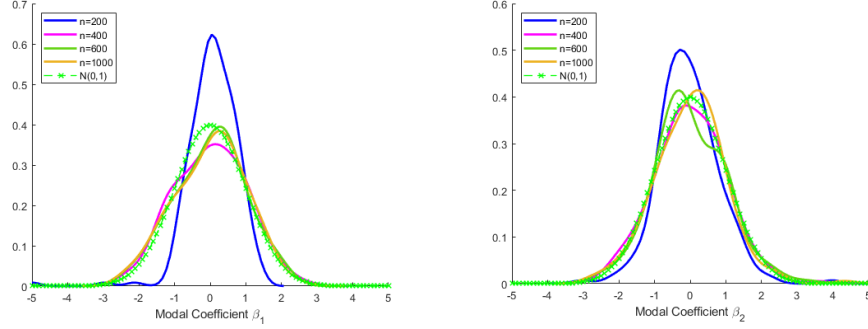


Figure 3: Empirical Density of the Standardized Estimate

We present the shape of the empirical density of the standardized modal estimate in Figure 3 to examine the asymptotic normality property of the nonlinear modal estimator. In accordance with the theoretical findings, most of the results manifest asymptotic normality as the sample size  $n$  increases. It is noticed that the performance of the asymptotic normality approximation is not perfectly good. We attribute it to the value of the bandwidth selected, which has a substantial effect on the estimation of parameters. How to develop a more efficient way to select the optimal bandwidth needs to be carefully researched in the future.

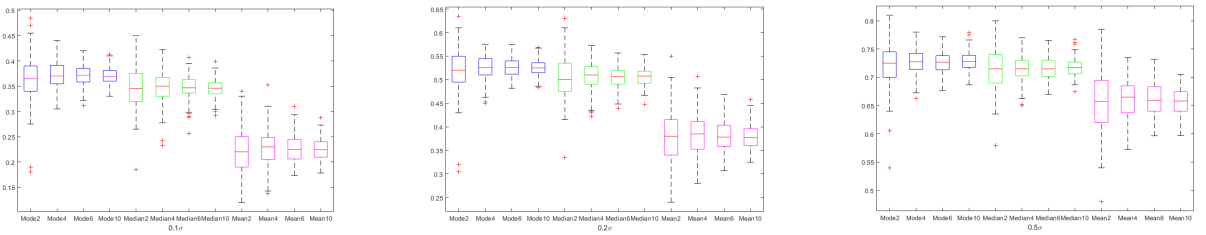


Figure 4: Boxplots of average of coverage probabilities: the numbers 2, 4, 6, and 10 represent the values of  $n = 200, 400, 600$ , and  $1000$ , respectively.

To show the advantage of the proposed nonlinear modal regression in prediction, we follow Yao and Li (2014) and Ullah et al. (2021) to report the coverage probabilities of prediction intervals of three different lengths ( $0.1\sigma, 0.2\sigma, 0.5\sigma, \sigma = \sqrt{\text{Var}(\epsilon_t)} \approx 2$ ). We use the same DGP procedure as before but implement the out-of-sample prediction with 200 repetitions for the additional  $n$  data points. The representative results of the coverage probabilities of the proposed nonlinear modal regression model and the nonlinear mean and median regression models

are reported in Figure 4, which shows that in comparison to the nonlinear mean and median regressions, the nonlinear modal regression tends to have superior predictive performance by providing the highest coverage probabilities. Although median regression outperforms mean regression due to the skewness of the error distribution, its performance is worse than that of modal regression. As expected, the nonlinear modal regression and mean and median regressions would have closer coverage probabilities with the increase of the interval length. These simulation findings encourage the use of nonlinear modal regression in prediction.

### 3 Nonlinear Modal Regression for COVID-19 Data

The prediction advantage of modal regression illustrated in the above section provides underlying support for building a nonlinear modal regression to predict COVID-19. We in this section develop a nonlinear modal regression based on the general structure of the effects and process of infection from a mode view and use it to predict COVID-19 new cases and new deaths in the US, which are the key quantities that determine the epidemic peak. We aim to investigate how well the proposed model could be used to guide the modeling of the dynamic of the spread.

#### 3.1 Model Framework

We first discuss the choice of a nonlinear modal function  $r(X_t, \beta)$  according to the transmission characteristics of COVID-19. It has been shown that the COVID-19 spread follows an exponential distribution, and the number of new cases/deaths does not follow a standard distribution like Gaussian or Exponential due to the large number of outliers and noise; see the related literature summarized in Tuli et al. (2020). In addition, Tuli et al. (2020) showed that the COVID-19 cases/deaths data follow the Generalized Inverse Weibull (GIW) Distribution better than the Gaussian, which has the following probability density function (de Gusmão et al., 2011)

$$f(y) = abc^b y^{-(b+1)} \exp \left[ -a \left( \frac{c}{y} \right)^b \right], \quad y > 0 \quad (3.1)$$

with three parameters  $a \in \mathbb{R} > 0$ ,  $b \in \mathbb{R} > 0$ , and  $c \in \mathbb{R} > 0$ . It can be easily proven that (3.1) is a probability density function by substituting  $u = -ac^b y^{-b}$ . Instead of considering probability distribution, Tuli et al. (2020) treated (3.1) as a regression function and used it to establish a mean regression model of cross countries COVID-19 prediction between a dependent variable  $Y_t$  and time trend  $t$ , which is expressed as follows

$$Y_t = abc^b t^{-(b+1)} \exp \left[ -a \left( \frac{c}{t} \right)^b \right], \quad (3.2)$$

where  $t > 0$  is the time in the number of days from the first case. Tuli et al. (2020) introduced a machine-learning-based iterative weighting strategy to fit (3.2) with the number of cases data and compared the prediction performance with the Gaussian fitting by  $MSE$ ,  $MAPE$ , and  $R^2$ , where they showed that the proposed GIW model performs significantly better.

By coincidence, the same phenomenon, i.e., the data with a large number of noise follow a GIW-type shape, appears when we plot the new cases/deaths data against time for most states in the US, which motivates us to develop a regression model for the COVID-19 data in the US based on (3.2). This paper however does not use this mean regression model directly as it only depends on time  $t$  and cannot capture the dynamics of COVID-19. Since previous studies have suggested that the log of new cases/deaths is more suitable to be the dependent variable (Deb and Majumdar, 2020; Schüttler et al., 2020; Wang et al., 2020; Li and Linton, 2021), as the logarithm value can weight more evenly values close to the maximum of the objective function and disregard other values, we then instead take the logarithm on both sides of (3.2), from which we can see that  $\log(Y_t)$  is linearly associated with  $\log(t)$  and  $t^\delta$  ( $\delta$  is a constant number). We emphasize that it is reasonable to use log value due to the increase in the number of new cases/deaths rose by multiple orders of magnitude in a short period of time and sensible to include  $t^\delta$  to capture the fact that most states in the US experience a decreasing trend after approaching the peak number of cases/deaths per day (in the early stage of the COVID-19 epidemic, the data usually show an exponential growth trend. After a period of time, as the number of uninfected people decreases, the growth rate starts to decelerate and the number of cases keeps rising until reaching a peak. Subsequently, the number of new infections begins to decline). Furthermore, to incorporate the time series structure of the data and the fact that each infected person will create a chain of new infections, we include the lag variable  $\log(Y_{t-1})$  in the model. Along with the above arguments, we propose the following nonlinear modal regression for modeling COVID-19 data by taking into consideration the effect of progress evolving over time

$$\log(Y_t) = \alpha + \beta \log(t) + \eta \log(Y_{t-1}) + \gamma t^\delta + \epsilon_t, \quad t = 2, \dots, n, \quad (3.3)$$

where error term  $\{\epsilon_t\}_{t=2}^n$  is a sequence of stochastic random variables with  $Mode(\epsilon_t | \mathcal{F}_{t-1}) = 0$  almost surely (a.s.) for model identification in which  $\mathcal{F}_{t-1}$  is the  $\sigma$ -field generated by  $\{Y_{t-1-s}\}_{s=0}^\infty$ . Therefore, the nonlinear modal regression line is defined as

$$r(X_t, \theta) = \alpha + \beta \log(t) + \eta \log(Y_{t-1}) + \gamma t^\delta, \quad (3.4)$$

where  $X_t = (1, \log(t), \log(Y_{t-1}), t)$  and  $\theta = (\alpha, \beta, \eta, \gamma, \delta)^T$ . Compared to (3.2), the proposed nonlinear modal regression model can better incorporate other covariates into the mode structure, such as the lag variable or social distance variables. In addition, the new model uses the conditional mode instead of mean or quantile to model the nonlinear relationship among variables. We also note that, although not presented here, the model developed in (3.3) performs



better in terms of  $MSE$  and  $MAPE$  than using a polynomial regression for  $t$ , the model (3.4) without the lag variable ( $Y_{t-1}$ ), and the model (3.4) with two lag variables ( $Y_{t-1}$  and  $Y_{t-2}$ ).

Different from the mean or median regression, we propose estimating the modal regression (3.3) using the following kernel-based objective function

$$Q_n(\theta) = \frac{1}{(n-1)h} \sum_{t=2}^n K \left( \frac{\log(Y_t) - \alpha - \beta \log(t) - \eta \log(Y_{t-1}) - \gamma t^\delta}{h} \right), \quad (3.5)$$

whose estimation relies on the choice of the regularization parameter—the bandwidth  $h$ . We propose to choose the bandwidth according to Kemp and Santos Silva (2012), where we minimize  $MSE$  and  $MAPE$  for a grid of 50 values of  $h$  between  $50MAD$  and  $0.5MAD(n-1)^{-0.143}$  with  $MAD = med_t\{|\log(Y_t) - r_m(X_t, \hat{\theta}_m) - med_t(\log(Y_t) - r_m(X_t, \hat{\theta}_m))|\}$  in which  $\hat{\theta}_m(\cdot)$  representing the corresponding NLS estimate.

With the available parameter estimate  $\hat{\theta} = (\hat{\alpha}, \hat{\beta}, \hat{\eta}, \hat{\gamma}, \hat{\delta})^T$  obtained from Algorithm 1, we can formulate a  $k$ -step ahead prediction to capture the dynamic behavior of COVID-19 by fitting the nonlinear modal regression (3.3) recursively for the entire horizon

$$Mode(\log(\hat{Y}_{t+k|t}) \mid t+k, \log(\hat{Y}_{t+k-1})) \approx \hat{\alpha} + \hat{\beta} \log(t+k) + \hat{\eta} \log(\hat{Y}_{t+k-1}) + \hat{\gamma}(t+k)^{\hat{\delta}}, \quad (3.6)$$

where  $\log(\hat{Y}_{t+k|t})$  represents the estimate of  $\log(Y_{t+k})$  based on the data  $\log(Y_1), \dots, \log(Y_t), \log(\hat{Y}_{t+1}), \dots, \log(\hat{Y}_{t+k-1})$ . Particularly, we pretend the pre-step estimate was the true value of  $Y_t$  at the corresponding step and use it as part of the input variable for predicting the next step. To graphically present the prediction procedure, we have the following roadmap

$$Y_t \xrightarrow{(Y_t, t+1)} \hat{Y}_{t+1} \xrightarrow{(\hat{Y}_{t+1}, t+2)} \hat{Y}_{t+2} \xrightarrow{(\hat{Y}_{t+2}, t+3)} \hat{Y}_{t+3} \quad \dots \quad \xrightarrow{(\hat{Y}_{t+k-1}, t+k)} \hat{Y}_{t+k}.$$

**Remark 3.1.** To reduce the computation time, we apply the same modal estimates with the bandwidth  $h$ , which are constructed using samples  $\{Y_t\}_{t=1}^n$  and the corresponding time sequence for all predictions. However, the prediction performance can be improved if we dynamically reestimate modal parameters each time to incorporate the substantial information contained in the intermediate variables  $Y_{t+1}, \dots, Y_{t+k-1}$  about the conditional mode when the pre-stage estimated forecast is added to the samples (for example, we estimate  $\theta$  with the data  $\{Y_t, t\}_{t=1}^n$  and use the corresponding estimate to predict the value of  $Y_{n+1}$ . After that, we use the data  $(\{Y_t, t\}_{t=1}^n, \hat{Y}_{n+1})$  to reestimate  $\theta$  and use the corresponding estimate to predict  $Y_{n+2}$ . Iterative this procedure until we achieve all predictions). Although the suggested recursive prediction procedure performs well for COVID-19 data in this paper, we notice that the accuracy of the predictions may deteriorate when  $k$  is too large, which is due to the accumulation of errors with the predicting horizon. Therefore, compared to the long-term prediction, the proposed model is better to be used for the short-term prediction.

**Remark 3.2.** It is noticed that there is a basic assumption for (3.6) such that the predicted value  $\hat{Y}_{t+k-1}$  performs almost the same as the true value  $Y_{t+k-1}$  with  $\text{Mode}(\epsilon_{t+1} \mid \hat{Y}_{t+k-1}) = 0$ , which is the main reason we use “ $\approx$ ” sign in (3.6). How to release this assumption to provide a more reliable prediction for modal regression needs to be researched further. However, compared to the prediction procedure of the  $k$ -step-ahead predictions based only on the observed data, as is standard in macro settings, our procedure should be more reliable. For instance, as mode does not have the additive property, it is difficult to guarantee that  $\text{Mode}(\eta\epsilon_t + \epsilon_{t+1} \mid Y_{t-1}) = 0$  with equation  $\log(Y_{t+1}) = \alpha + \beta \log(t+1) + \eta(\alpha + \beta \log(t) + \eta \log(Y_{t-1}) + \gamma t^\delta) + \gamma(t+1)^\delta + \eta\epsilon_t + \epsilon_{t+1}$ .

**Remark 3.3.** We in this paper model the new cases and new deaths datasets with equation (3.3), separately, which indicates that the predicted new deaths and new cases do not appear to be linked to each other. Such a univariate model may ignore possible comovements with other available time series. In practice, it is extremely likely that new cases and new deaths are collectively impactful on observable trends, i.e., there is a dependency nature in the series. Thus, it is possible to improve predictions and the explanatory power of the model by jointly predicting these two through a nonlinear vector autoregressive modal regression by extending the results in [Kemp et al. \(2020\)](#), i.e.,  $Y_{jt} = r(Y_{-jt}, \{Y_{jt-l}\}_{l=1}^L, X_{jt}, \gamma) + e_{jt}$  with finite order  $L$  for  $j = 1, 2$  in which  $Y_{-jt}$  collects all but the  $j$ th observation at time  $t$  and  $X_{jt}$  includes all possible factors that affect both cases and deaths. With the stationary condition and  $\text{Mode}(e_{jt} \mid \mathcal{F}_{t-1}) = 0$  in which  $\mathcal{F}_{t-1}$  is the  $\sigma$ -field generated by  $\{Y_{-jt}, \{Y_{jt-l}\}_{l=1}^L\}$ , it can be shown that the estimator of  $\gamma$  is identified and asymptotically normally distributed. In addition, due to the computation burden, we do not compute the confidence interval for predictions. This should be easily carried out based on the bootstrapped modal regression method introduced in [Ullah et al. \(2021\)](#), where we independently draw bootstrapped pseudo samples of residuals from the estimated regression, use the pseudo residual to minus the corresponding mode value to ensure the mode of residual is zero, and then follow the standard procedure as in mean regression to get the modal confidence interval. Future studies could fruitfully explore these issues further.

## 3.2 Modal Prediction Results

We use publicly available COVID-19 data on the daily number of reported cases and deaths to fit the proposed model (we use the case and death data from each state/region to fit the model (3.4) and fully expect that the parameters vary across the states/regions, as different states/regions are at different stages of the epidemic cycle and have taken different approaches to managing it), and perform an out-of-sample prediction analysis for all states/regions in the US (including the District of Columbia and Puerto Rico) to predict the number of daily new cases and deaths. We remark that the daily data are superior for short-term/medium tactical predicting and are more informative than weekly or monthly data, as they can reflect the turning point of the curve timely and encourage policymakers and people to take flexible actions at any moment. The data

of aggregated US COVID-19 cases/deaths we use are from the GitHub repository managed by The New York Times (<https://github.com/nytimes/covid-19-data>), which was accessed on August 24, 2020 and used to calculate the daily new cases and new deaths data through the differencing transformation (the last date for the data in this paper is August 23, 2020), i.e.,  $\text{New Cases} = \text{Cases}_t - \text{Cases}_{t-1}$  and  $\text{New Deaths} = \text{Deaths}_t - \text{Deaths}_{t-1}$ . We set all negative values in the new dataset to be zero for calculation. Due to space limitation, we do not put the results of the descriptive statistics of data here, but they are available upon request. Note that this dataset automatically updates every day with new information.

The accuracy and reliability of a model can be tested by comparing the actual values with the predicted values. Following Tuli et al. (2020), we compare the performance (model accuracy) of the newly developed nonlinear modal regression to those of nonlinear mean and median (and robust) regressions using  $R^2$  (higher value indicates better fit). We also use performance metrics— $MSE$  and  $MAPE$  (lower values indicate better fit)—to determine the residuals between predictions and actual values in order to compare the out-of-sample prediction validity of the proposed nonlinear modal regression and mean and median (and robust) regressions for the last 20 days of the samples (they are treated as validation data, while the other data are used for training)

$$MSE = \frac{1}{20} \sum_t (\log(Y_t) - \log(\hat{Y}_t))^2, \quad (3.7)$$

$$MAPE = \frac{1}{20} \sum_t \frac{|\log(Y_t) - \log(\hat{Y}_t)|}{\log(Y_t)} \times 100, \quad t \in \text{last 20 days}. \quad (3.8)$$

The model comparison results are summarized in Table 2 (and Table 7 in Online Appendix B), with the best performing model highlighted in bold font. As we can see from Table 2 (and Table 7 in Online Appendix B), the proposed nonlinear modal regression succeeds in predicting the new cases/deaths for 20 days ahead with better accuracy compared to the nonlinear mean and median (and robust) regressions for most states/regions. It can fit the data significantly better with higher  $R^2$ , and also has more precise predictions with lower  $MSEs$  and  $MAPEs$  for most states with the observed data. Overall, we can see that the proposed nonlinear modal regression model outperforms other competing models in terms of prediction accuracy and can give reliable guidance on the trend of the epidemic in the future. There is no special reason for comparing model predictions in terms of  $MSE$  and  $MAPE$  over 20 days, which was chosen arbitrarily. To show the results robust to choosing alternative time horizons, we also compared the prediction performance for the last 30 days of the samples, which does not reveal the large difference in prediction or comparison results.

We then apply the proposed nonlinear modal regression to predict the number of new cases and new deaths for up to 130 days (August 24, 2020–December 31, 2020) to show how the epidemic has evolved over time, which has some differences from many other papers focusing on

the long-term trajectory of COVID-19 using mean regression. To conduct the prediction for the latest 130 days, we use the same bandwidth obtained from the training data (when comparing the model prediction performance) to reestimate nonlinear modal regression with a full sample

Table 2: Model Comparison Results

State/ Region	New Cases-Mode			New Deaths-Mode			New Cases-Mean			New Deaths-Mean			New Cases-Median			New Deaths-Median		
	MSE	MAPE	R <sup>2</sup>	MSE	MAPE	R <sup>2</sup>	MSE	MAPE	R <sup>2</sup>	MSE	MAPE	R <sup>2</sup>	MSE	MAPE	R <sup>2</sup>	MSE	MAPE	R <sup>2</sup>
AL	0.1712	4.5563	0.9338	0.6943	35.5300	0.3363	0.1734	5.0252	0.8930	0.6953	36.3657	0.3110	0.1384	4.4649	0.7963	0.9213	50.8288	0.4650
AK	0.1569	7.5698	0.7537	0.1095	62.6997	0.1844	0.7199	17.8641	0.7350	0.1198	79.8023	0.0130	2.3017	33.9916	0.3400	0.1682	89.3547	0.1818
AZ	0.3095	6.4026	0.9989	0.7599	24.2938	0.7294	0.9339	12.2357	0.9740	0.8226	24.8236	0.7220	1.5375	17.1057	0.8314	3.2440	32.2442	0.7978
AR	0.1280	4.4861	0.7671	0.4956	23.8808	0.4168	0.1595	4.9279	0.7350	0.6668	28.4478	0.4080	0.0868	3.7454	0.6070	0.9910	34.4291	0.3265
CA	0.2040	4.3817	0.9919	0.5272	13.5211	0.9555	0.2576	4.6241	0.9780	0.5313	13.4917	0.9170	12.0594	36.3353	0.9434	0.6386	13.9525	0.9928
CO	0.1226	4.7919	0.9989	1.2861	63.6700	0.5278	0.1576	5.7275	0.9120	1.5153	72.3817	0.4740	0.4542	10.7173	0.9936	2.0659	66.9498	0.3467
CT	5.0400	44.6875	0.5327	1.1784	133.0601	0.7671	5.4543	51.1498	0.4930	2.1439	179.9120	0.7150	5.2566	48.3522	0.5686	6.5791	171.9621	0.3973
DE	0.2388	7.7782	0.6795	0.4570	85.1702	0.5880	0.2454	8.3739	0.5870	0.8193	125.5970	0.3850	0.2779	5.6415	0.6342	0.7644	36.6193	0.4012
DC	0.4210	14.2005	0.9999	0.4204	82.0909	0.7721	0.7385	19.4940	0.7550	1.6516	175.4791	0.6370	0.4884	15.3523	0.9598	0.8240	55.8350	0.5495
FL	0.1081	3.1617	0.8969	0.2720	8.9638	0.8726	0.2255	4.4887	0.9330	0.4822	11.2948	0.7900	0.1124	3.4112	0.8697	0.3507	9.4744	0.8612
GA	0.0933	3.1374	0.8604	1.1771	22.4311	0.5551	0.0953	3.2290	0.8210	1.3582	24.5763	0.5160	0.2448	5.4379	0.7784	0.6647	16.6072	0.5868
HI	0.8501	15.0967	0.7482	0.3821	73.1096	0.5342	4.5141	36.6774	0.7120	0.5695	95.8169	0.0353	8.1706	51.3927	0.6466	0.6140	97.5388	0.1575
ID	0.1737	6.1939	0.7379	0.9100	43.6542	0.2868	0.4780	10.1412	0.7080	1.1558	49.4825	0.1880	1.5833	20.1322	0.5941	2.6503	91.1905	0.4947
IL	0.2880	5.9163	0.9999	2.0747	45.7357	0.9838	0.3712	7.1152	0.9870	2.8030	523.2636	0.9220	0.5619	9.2363	0.9297	1.6185	48.3631	0.8174
IN	0.1591	5.0592	0.9331	0.5063	29.2463	0.8413	0.3958	8.5265	0.9530	1.5668	40.3907	0.6930	0.1087	4.2200	0.9162	1.1215	51.2706	0.6255
KS	1.3407	19.9173	0.5557	0.6090	42.7460	0.2992	1.3489	19.4022	0.5230	0.6648	44.1607	0.2040	1.5040	18.6129	0.5247	0.6261	42.8812	0.2032
KY	0.2305	5.9564	0.7583	0.4734	21.5414	0.4976	0.3371	8.3026	0.6530	0.8958	36.7502	0.3060	0.2675	7.3576	0.6038	0.4541	23.6336	0.3849
LA	6.5341	14.2872	0.3179	2.1630	29.0675	0.5084	6.5771	11.4450	0.1560	2.9178	42.0480	0.3880	6.5429	19.8462	0.9944	2.6146	37.7073	0.4861
IA	0.1406	5.1942	0.7727	1.0872	39.7959	0.4968	0.1445	5.5619	0.8730	1.2304	44.0175	0.4830	0.2799	7.1513	0.9453	0.4754	23.9231	0.6961
ME	0.7938	23.4699	0.4109	0.1436	43.9784	0.3659	0.8567	25.3532	0.4030	0.1676	91.5552	0.1460	0.5190	31.6516	0.3985	0.3118	124.5080	0.2344
MD	0.0368	2.5438	0.9435	0.8368	33.7025	0.9999	0.1471	5.1483	0.9390	1.1255	41.9215	0.8370	0.0374	2.5885	0.9050	1.6134	62.2135	0.7214
MA	0.1779	6.1324	0.9999	0.4949	27.5922	0.9999	0.4097	9.5495	0.9700	1.6480	42.7498	0.9070	4.0086	84.2134	0.9524	2.1522	61.7372	0.7854
MI	0.4083	7.6109	0.7484	2.0608	50.8842	0.7454	0.86664	13.7508	0.6960	3.3902	76.3392	0.6810	1.1114	15.5525	0.9989	2.4700	59.1702	0.7377
MN	0.0797	3.0202	0.8551	2.1624	62.5585	0.8691	0.1180	4.0501	0.8040	2.8030	73.1131	0.7680	0.7755	12.7442	0.8128	0.2617	23.5088	0.9505
MS	0.2104	5.5623	0.6057	0.4832	21.6170	0.5995	0.2119	5.9219	0.5680	0.5972	24.6271	0.4960	0.3400	7.6840	0.5632	0.4983	21.1192	0.5929
MO	0.1533	4.8664	0.9425	0.4962	17.8628	0.5882	0.2882	7.1869	0.9170	0.9254	30.6017	0.4330	2.0424	19.4914	0.9423	0.7243	26.3912	0.4522
MT	1.0004	19.3589	0.8605	0.4924	60.1072	0.2565	1.6448	26.1567	0.7550	0.4806	59.7919	0.1240	5.3211	48.6101	0.6491	0.7768	95.9944	0.2628
NE	0.1410	5.4894	0.9529	0.4809	37.3174	0.2980	0.4029	9.7425	0.9360	0.4927	36.9213	0.2960	0.5193	11.0812	0.9999	0.5364	38.3479	0.9596
NV	0.0630	3.0931	0.9096	0.8165	32.1520	0.3850	0.0603	3.0970	0.8540	1.2392	36.1752	0.2290	0.2832	7.4281	0.7900	1.3796	36.8643	0.2530
NH	0.2892	15.0186	0.9191	0.6415	125.4310	0.4923	0.5377	20.4409	0.8020	0.7696	142.2721	0.3570	0.4894	19.4310	0.8084	0.3534	22.4967	0.8373
NJ	1.0870	17.8764	0.8583	0.7045	25.4733	0.8089	2.0952	25.1485	0.8170	1.0478	40.9881	0.7440	1.3815	20.2057	0.8490	7.0072	153.6538	0.6002
NM	0.2643	8.5568	0.9046	0.1674	21.8560	0.7441	0.3769	10.7114	0.8780	0.3389	28.0368	0.5900	0.3004	9.3521	0.9010	0.2964	32.2350	0.8484
NY	0.0449	3.0657	0.9968	0.6965	17.1554	0.9841	0.1438	5.3248	0.9600	1.2468	38.8174	0.8900	1.1538	15.5820	0.9992	2.7453	64.9087	0.9858
NC	0.3222	7.0517	0.9310	0.5711	27.0946	0.8095	0.4167	8.2013	0.9430	0.5678	26.9968	0.6890	0.2933	6.6726	0.9529	0.6438	28.1160	0.9207
ND	0.3865	10.4769	0.8256	0.2729	26.3197	0.5332	0.4928	12.2427	0.7420	0.5751	72.0739	0.0923	0.8129	15.8381	0.6120	0.3398	42.7843	0.2809
OH	0.0573	2.8443	0.9999	0.8720	46.1271	0.8514	0.0643	2.8045	0.9540	1.2982	46.2033	0.5820	0.3008	7.2960	0.9999	2.1708	73.6845	0.6820
OK	0.1821	5.2804	0.9135	1.0089	39.1482	0.4061	0.2220	5.9597	0.8970	1.5633	48.5069	0.2630	1.0923	15.2377	0.7034	1.4307	46.3320	0.3782
OR	0.0407	2.7310	0.8674	0.6189	48.0773	0.3032	0.0434	2.8604	0.8620	0.6585	46.4323	0.1930	1.2971	16.7406	0.8478	0.9339	38.1471	0.3004
PA	0.0485	2.6820	0.9332	2.0029	44.4196	0.8239	0.1419	4.8088	0.9430	3.6654	60.8070	0.7110	0.2093	5.9761	0.9074	1.7553	42.5175	0.6950
PR	1.1170	13.2338	0.5786	1.3052	48.8159	0.4122	1.4482	15.9853	0.6820	2.7094	75.2085	0.1060	2.1473	21.9538	0.6611	3.3145	83.7110	0.1182
RI	5.0243	41.8085	0.5909	0.3811	51.8988	0.8547	7.0779	61.0051	0.4840	1.9419	164.4378	0.5520	9.9050	70.2124	0.8195	0.9570	69.1517	0.9258
SC	0.0835	3.5217	0.9999	0.4014	16.4416	0.6935	0.2432	6.1175	0.9400	0.6080	18.1253	0.5910	0.1153	4.0323	0.8632	0.7424	20.3135	0.4535
SD	0.1802	6.7710	0.8605	0.2033	21.1258	0.1757	0.3197	9.9167	0.6880	0.2088	23.0946	0.1040	1.0279	18.9569	0.6765	0.7436	100.0031	0.3583
TN	0.1058	3.6718	0.9785	0.3449	16.2219	0.6200	0.1475	4.2999	0.8990	0.6678	21.8297	0.4620	0.2820	6.3490	0.7760	0.3492	16.3966	0.5805
TX	0.0595	2.1483	0.9928	0.2397	8.5471	0.9690	0.3448	5.6411	0.9760	0.7228	14.0341	0.8860	0.1099	2.6563	0.9530	0.2809	9.1692	0.9474
UT	0.2180	6.6142	0.9601	0.5060	33.4387	0.3575	0.5707	11.4507	0.9510	0.5043	33.1763	0.3140	0.2272	6.3151	0.9275	0.7189	45.1290	0.5394
VA	0.0619	2.7185	0.7473	0.9535	33.2575	0.3727	0.0958	3.7875	0.7250	1.0224	38.1446	0.4330	0.1015	3.4359	0.7826	1.6647	55.0330	0.6826
WA	0.1298	4.8612	0.9781	0.7005	26.9027	0.5764	0.3073	7.4457	0.9380	0.7100	27.0861	0.5520	0.1055	4.0141	0.8863	1.0449	31.2122	0.3787
WV	0.0965	5.6020	0.6729	1.4237	84.2442	0.3243	0.3392	10.2841	0.5900	1.5321	87.9995	0.1940	1.0661	20.3992	0.6033	2.3734	112.7576	0.3147
WI	0.0695	3.3494	0.9246	0.6409	33.9752	0.7993	0.											

for each state/region (training data+validation data), and then apply the suggested recursive prediction procedure with the new parameter estimates. We remark that there exist large variations in the parameter fittings, which indicates that long-term predictions are complicated. However, the long-term prediction in comparison with the short-term prediction can provide the pattern of the epidemic. Also, there is an underlying assumption for the prediction results, which is that the data used are reliable and the outbreak will continue to follow the past pattern in the future (Petropoulos and Makridakis, 2020). We acknowledge that this assumption is actually the key issue for predicting the transmission of COVID-19, and it is necessary to update predictions by the suggested model when new information/data is available. However, one advantage of modal regression is that it can cope with some forms of measurement errors. Thus, applying modal regression to predict COVID-19 still has an advantage compared to traditional regressions.

Table 3: Modal Prediction Results

State/ Region	Predictions of Modal Regression 08/24/2020-12/31/2020				State/ Region	Predictions of Modal Regression 08/24/2020-12/31/2020			
	09/30	10/31	11/30	12/31		09/30	10/31	11/30	12/31
	Total New Cases					Total New Deaths			
AL	45570	46151	52293	61847	AK	2759	2735	3025	3503
AZ	2772	712	737	945	AR	38586	45110	56629	73506
CA	296260	380190	446670	530800	CO	10461	73826	63510	58704
CT	83	0	0	0	DE	2360	1704	1450	1329
DC	1569	1107	946	881	FL	270170	323480	384240	473570
GA	101620	108200	125330	151120	HI	8046	7494	8178	9367
ID	8692	9657	11487	14158	IL	45842	14954	4759	1301
IN	21824	16657	14641	13683	KS	7895	6849	6749	7039
KY	17470	17932	20270	23934	LA	18705	15715	15186	15651
IA	19712	16834	16233	16550	ME	508	300	206	148
MD	15219	8544	5695	4027	MA	1050	0	0	0
MI	17657	13613	12608	12549	MN	32478	32085	35103	40228
MS	34407	35406	40023	47283	MO	35374	35535	39682	46351
MT	12332	15629	17901	21389	NE	4193	2000	1059	636
NV	22775	24441	28663	34930	NH	295	85	19	0
NJ	3104	1123	523	264	NM	8305	7397	7328	7681
NY	12006	6420	4858	4076	NC	82554	81715	88581	100270
ND	5588	5533	6095	7036	OH	45069	47630	52908	61855
OK	24683	26744	31587	38723	OR	13887	14217	15953	18724
PA	15625	10309	8261	7179	PR	16332	18541	22792	29017
RI	82	0	0	0	SC	70597	91851	120560	163670
SD	2663	2052	1859	1799	TN	73717	77510	89298	107170
TX	994910	1679200	2255200	3082600	UT	19034	19507	20285	22113
VA	35771	30929	30338	31505	WA	44721	63229	81864	108910
WV	4456	4308	4623	5217	WI	35719	36597	40896	47835
VT	89	53	37	28	WY	1509	1448	1555	1756

**Note:** The results represent the total number of modal predicted new cases and new deaths between 08/24 and 09/30, between 10/01 and 10/31, between 11/01 and 11/30, and between 12/01 and 12/31, separately.

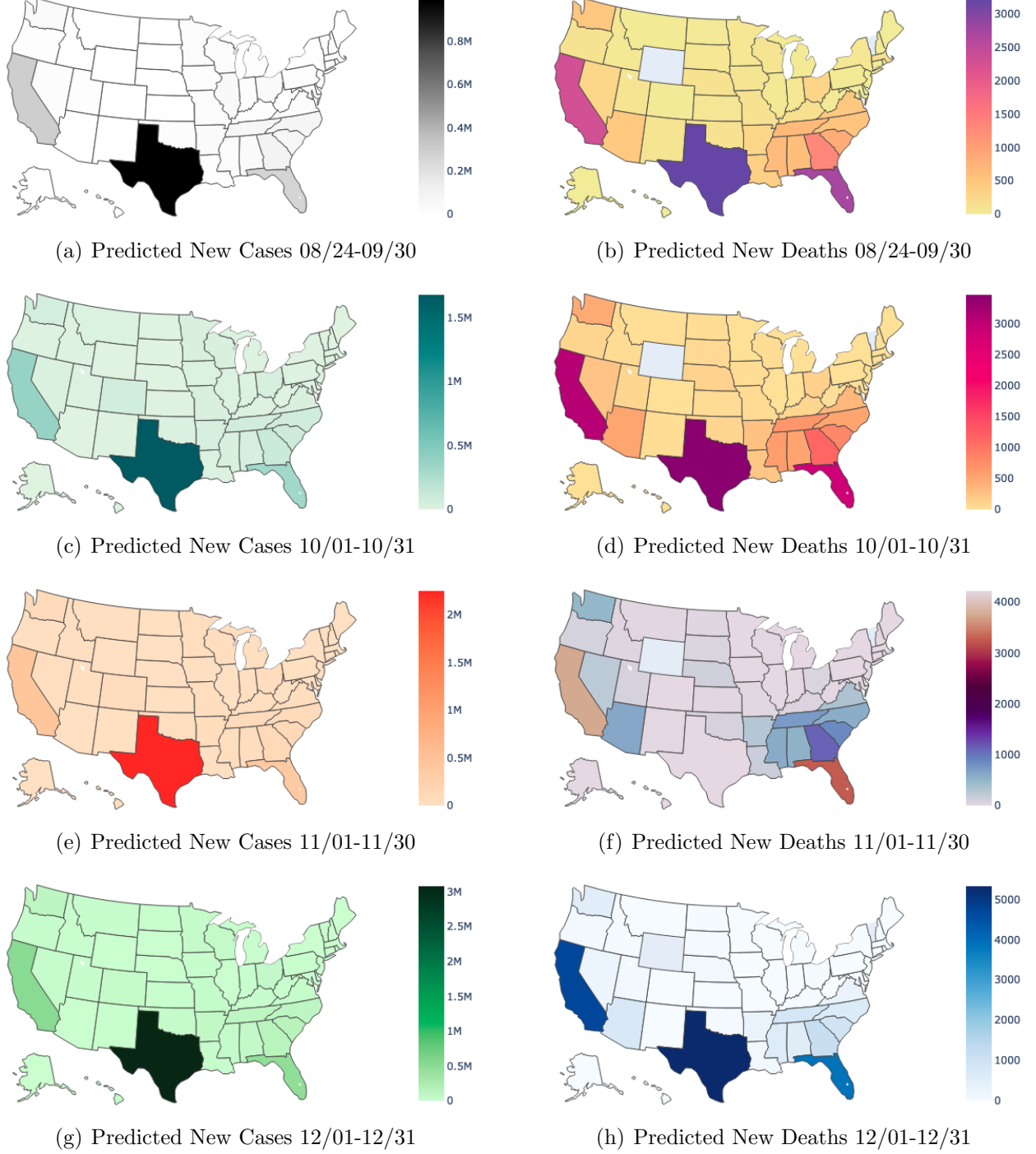


Figure 5: Visualization of the Total Number of Modal Predicted New Cases and New Deaths across the US

To clearly show the dynamic of the COVID-19 spread in the US, we divide the prediction period into four stages, which are 08/24-09/30, 10/01-10/31, 11/01-11/30, and 12/01-12/31. The prediction results are presented in Table 3, from which we can observe that the COVID-19 outbreak in the US is dynamic both in time and across different states/regions. Some



states/regions are showing a clear decreasing trend in the number of new cases and new deaths (it is tempting to speculate that this result is due to the rapid imposition of alert levels and ever tighter lockdowns for these states. The detailed analysis of the effect of lockdowns and social distancing policies on the transmission of COVID-19 is beyond the scope of this paper; see the related discussions in Section 4), e.g., Connecticut, Illinois, Maine, Maryland, Massachusetts, New Hampshire, New Jersey, New York, Pennsylvania, Rhode Island, among others, while some other states/regions are still in the first wave of the COVID-19 outbreak with an increase in the number of new cases and new deaths, e.g., Alabama, Arkansas, California, Florida, Georgia, Mississippi, Montana, North Carolina, North Dakota, Oregon, Texas, Utah, Washington, and so on. Furthermore, as Figure 5 shows (the darker the color, the more severe the infection), it is clear that there are systematic differences in spreading distributions among states/regions (heterogeneous across the states/regions). In particular, for the next 130 days, California, Florida, Texas, and Georgia are the most severe states in terms of the number of predicted new cases and new deaths, which indicates the urgency for these states to take actions to keep social distancing and necessary precautions.

It should be noted that the number of predicted new cases and new deaths across different states/regions has orders of magnitude differences, resulting in the almost uniform color in Figure 5 for other states/regions having small numbers. To better reveal the situations of other states/regions from visualization, we remove the first three states with the largest numbers of predicted new cases and new deaths from Figure 5. The new visualizing results are presented in Figure 6 which shows a stark heterogeneity across states/regions. We find that for most western and eastern states, the total numbers of new cases and new deaths are incredibly large for the next 130 days based on the prediction results, and these states are in fact experiencing significantly more serious COVID-19 burdens compared to the Midwest (under the stress of economic stagnation, many states/regions have reopened their economies. However, based on the analysis of modal prediction results, it is clear that the outbreak has not been sufficiently controlled in many states up to the date of this paper).

Last but not the least, Online Appendix B contains the nonlinear modal prediction figures for each state/region (including the District of Columbia and Puerto Rico) in terms of new cases and new deaths, which further demonstrates that the trend of daily confirmed new cases and new deaths is being nicely captured (except for some noisy fluctuations) and the significant new trend is detected by the proposed nonlinear modal regression. Based on these figures, we can also observe that for some states/regions, they have already arrived at a saturation stage and show a decreasing trend for the number of new cases and new deaths, e.g., Colorado, Connecticut, Delaware, Maine, Massachusetts, New Hampshire, and Pennsylvania, while for some other states/regions, such as Alabama, Arkansas, California, Florida, Idaho, Nebraska, Tennessee, and Texas, they will still be at the initial phase of the epidemics and show an increase of the trend for the number of new cases and new deaths if the control and intervention policy



is not implemented more effectively. We also list the prediction results for the nonlinear mean regression, median regression, and robust regression (including  $R^2$  and performance metrics) in Online Appendix B, although we have shown that nonlinear modal regression is of higher prediction quality than nonlinear mean and median regressions. The results indicate that there are systemic prediction differences among these models.

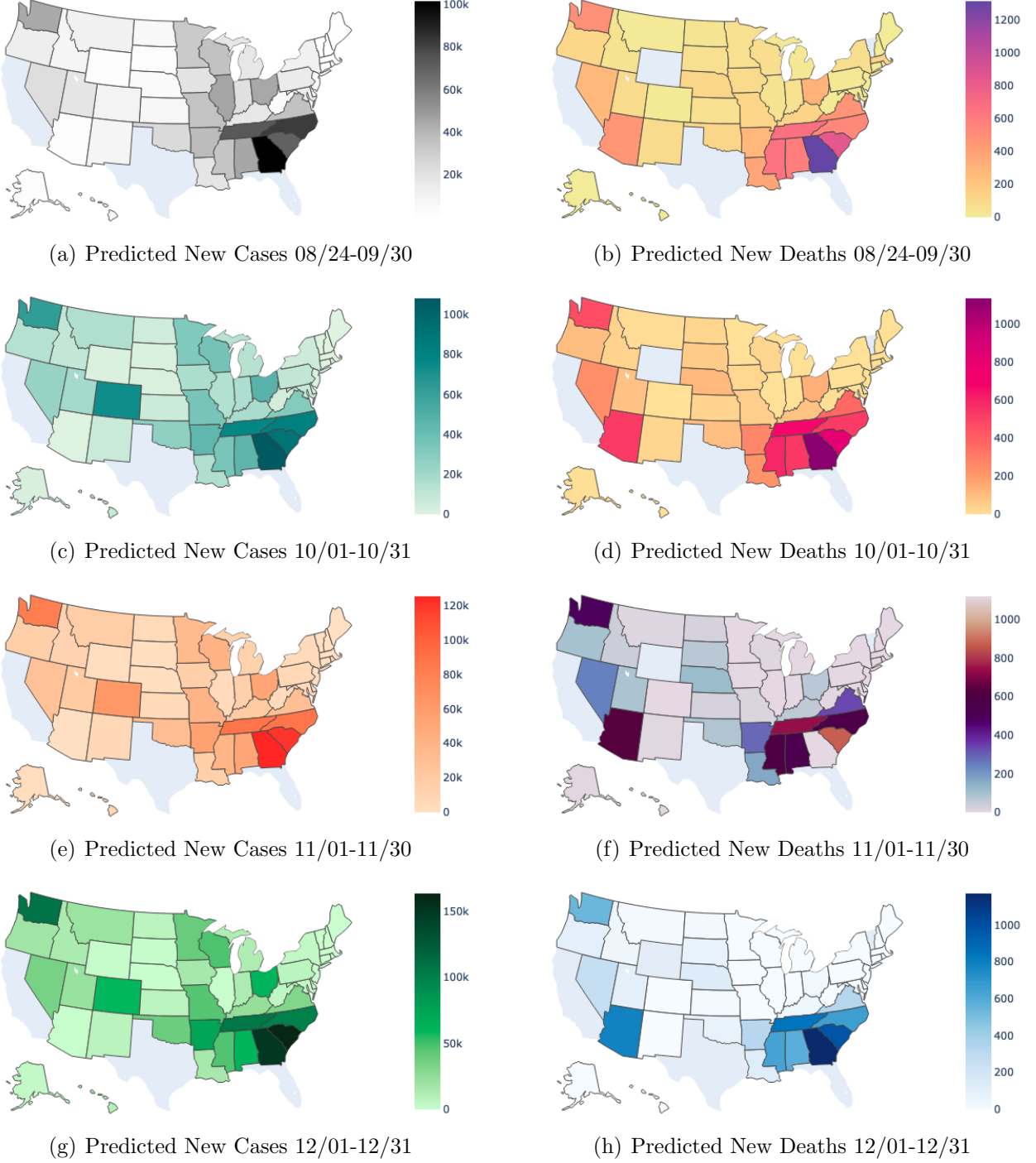


Figure 6: Visualization of the Total Number of Modal Predicted New cases and New Deaths across the US after Removing CA, TX, and FL

## 4 Concluding Remarks

The outbreak of COVID-19 has been unprecedentedly affecting the health and safety of people all over the world, which implies the urgency and importance of accurate prediction. In this paper, we propose a new model, namely parametric nonlinear modal regression for dependent samples, which is particularly useful for handling noisy, skewed, or truncated data (such as the COVID-19 data) and can complement the existing mean or quantile regression. The new model uses the conditional mode instead of mean or quantile to model the nonlinear relationship among variables. We employ a kernel-based objective function to simplify the computation and numerically estimate the proposed model by virtue of a modified MEM algorithm. The asymptotic theorem and the optimal bandwidth are investigated under mild conditions. We then use the proposed nonlinear modal regression to predict the COVID-19 outbreak in the US at the state/region level. We compare the predictions for this novel model with the predictions for nonlinear mean and median (and robust) regressions, and show that the proposed modal regression model can quantify the observed dynamics and provide more precise predictions. Although the outbreak spreads of COVID-19 are largely affected by the policies and social responsibilities of each state/region, we hope that the newly proposed model can be applied to analyze and classify the characteristics of COVID-19 in the US to provide more timely information to help policymakers to implement fast actions to curb the spread of the infection, avoid overburdening the health system, and understand the development of COVID-19 from some points.

This work paves the way for a number of exciting research directions in the analysis of modal regression and COVID-19. In this paper, we focus on parametric nonlinear modal regression. As pointed out by a referee, the results could be extended to the nonparametric modal regression for dependent samples under  $\alpha$ -mixing without imposing any kind of structural assumptions on the data generating process. Specifically, we can employ a kernel-based objective function with the local linear approximation. For our case, as we have both discrete (time variable) and continuous regressors in the model, we need to smooth the discrete variable using discrete kernels such that  $\Lambda_\lambda(Z_i, z_0) = \prod_{j=1}^q \lambda_j^{I\{Z_{i,j} \neq z_{0,j}\}}$ , where  $I(\cdot)$  denotes the usual indicator function,  $Z_i$  is a  $q$ -dimensional discrete random vector, and  $\lambda = (\lambda_1, \dots, \lambda_q)^T$ ,  $\lambda_j \in [0, 1]$  is the bandwidth for the  $j$ th discrete covariate  $Z_{i,j}$ . We can then make the bandwidths in the discrete kernel be a vector of zeros, and the model will be reduced to the local linear modal regression, which splits the full sample into several subsamples according to different values of the discrete variables. Nevertheless, such a naive sample-splitting method may increase the estimation variance (Li and Racine, 2004). How to derive asymptotic properties and provide asymptotic analysis on the selection of optimal bandwidths for the nonparametric modal regression with mixed discrete and continuous data would be an interesting but challenging future research topic.

Furthermore, in the current paper, we focus on the new cases and new deaths in the

US. However, the proposed model could be easily generalized to other countries, say country-level data, and other quantities of interest, e.g., cumulative recorded cases and deaths, or the number of people needing hospitalization in an Intensive Care Unit (ICU) each day for a set of regions. Different from the existing research about COVID-19 data, we can also use the proposed model to predict the unconditionally most likely (mode) value of new cases/deaths, which is one of the most important variables/factors when fighting the COVID-19 pandemic. When new cases/deaths reach their mode value, the healthcare system may have the biggest pressure and the largest chance of being overwhelmed, which could in turn affect the death rate. The importance of the mode value can also be seen by noticing that even if the total number of cases is fixed, if we could spread the cases over time and reduce the mode value, the health care system can function much better and thus reduce the fatality rate. In addition, it is important to note that the model for COVID-19 presented in this paper has certain limitations (we have to take such predictions reticently, as the prediction error will accumulate over time), as it does not account for any mitigation measures and policy changes. To understand the factors that contribute to the spread of COVID-19, in the future, we could include many other covariates into the model, i.e., the factors that might affect new cases/deaths such as social distancing measures as well as the timing of their implementation, the demographics and health condition of the population, the state of the epidemic, the capacity of the healthcare system, the population density, and so on. We can then model how the number of cases/deaths depends on the above collected covariates to find out whether there are some clusters of countries having a similar relationship between dependent variable and covariates. Also, it will be interesting to study the spatiotemporal pattern in the spread of COVID-19 by incorporating the spatial correlation into the modal regression.

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## Online Appendix A: Simulation Results

**DGP 2** To further illustrate the proposed nonlinear modal regression, we generate the data from the following model

$$Y_t = \beta_1 + \beta_2 X_t + \exp(\beta_3 X_t^2) + X_t \epsilon_t,$$

where  $\beta_1 = 1.5$ ,  $\beta_2 = 2$ ,  $\beta_3 = 2$ ,  $X_t = 0.4X_{t-1} + u_t$ ,  $u_t \sim \text{Uniform}[0, 1]$ , and the error  $\epsilon_t$  follows  $0.5Ga(k_1, \theta) + 0.5Ga(k_2, \theta)$  in which  $Ga$  represents the gamma distribution,  $k_j \in \mathbb{N}_{>0}$ ,  $j = 1, 2$ , is the shape parameter that can adjust the skewness of  $v_{it}$  (coefficient of skewness =  $\sqrt{4/k}$ ), and  $\theta \in \mathbb{N}_{>0}$  is the scale parameter; see [Ullah et al. \(2021\)](#). Note that  $\mathbb{E}(\epsilon_t) = 0.5(k_1 + k_2)\theta$  and  $\text{Mode}(\epsilon_t) = 0.5(k_1 + k_2 - 1)\theta$ . In this simulation, we set  $k_1 = 1$ ,  $k_2 = 2$ , and  $\theta = 0.5$ . We then have  $\mathbb{E}(\epsilon_t) = 0.5(k_1 + k_2)\theta = 0.75$ ,  $\text{Mode}(\epsilon_t) = 0.5(k_1 + k_2 - 1)\theta = 0.5$ , and

$$\begin{cases} \text{Mean Regression: } \mathbb{E}(Y_t | X_t) = 1.5 + 2.75X_t + \exp(2X_t^2), \\ \text{Modal Regression: } \text{Mode}(Y_t | X_{1,t}, X_{2,t}) = 1.5 + 2.5X_t + \exp(2X_t^2). \end{cases}$$

Notice that if  $X \sim Ga(\alpha, \theta)$  and  $Y \sim Ga(\beta, \theta)$  are independently distributed with the same scale parameter, then  $X + Y$  follows  $Ga(\alpha + \beta, \theta)$  with variance  $(\alpha + \beta)\theta^2$ . [Ullah et al. \(2021\)](#) considered two different settings with different values of  $k_j$  to formulate distributions with different magnitudes of skewness. They concluded that when the dataset is from the less skewed setting, modal regression offers close but less accurate regression estimates than mean regression. We thus focus on the more skewed case here.

Table 4: Results of Simulations—DGP 3

Sample Size	$\beta_1$ (SE)	MSE( $\beta_1$ )	$\beta_2$ (SE)	MSE( $\beta_2$ )	$\beta_3$ (SE)	MSE( $\beta_3$ )
<u>Modal Regression</u>						
$n=200$	1.5622 (0.1381)	0.0228	2.4782 (0.4692)	0.2195	1.8776 (0.3344)	0.1263
$n=400$	1.5637 (0.0907)	0.0122	2.4892 (0.3130)	0.0976	1.8699 (0.2295)	0.0693
$n=600$	1.5515 (0.0683)	0.0073	2.5207 (0.2325)	0.0542	1.8789 (0.1683)	0.0428
$n=1000$	1.5478 (0.0507)	0.0049	2.5195 (0.1692)	0.0289	1.8819 (0.1232)	0.0290
<u>Mean Regression</u>						
$n=200$	1.4949 (0.0996)	0.0099	2.7607 (0.1593)	0.0254	1.9999 (0.0031)	0.0955
$n=400$	1.5062 (0.0741)	0.0055	2.7376 (0.1207)	0.0146	2.0002 (0.0018)	0.0344
$n=600$	1.4983 (0.0629)	0.0039	2.7562 (0.1003)	0.0100	2.0000 (0.0015)	0.0233
$n=1000$	1.5029 (0.0478)	0.0023	2.7424 (0.0756)	0.0057	2.0001 (0.0011)	0.0129
True Value	$\beta_1 = 1.5$	$\beta_{2,mode} = 2.5$	$\beta_{2,mean} = 2.75$	$\beta_3 = 2$		

From the above equations, we can see that the nonlinear modal regression shares the same coefficients with the nonlinear mean regression, but the intercepts are different. Following DGP 1, we estimate parameters with  $M=200$  replications and consider sample size



$n \in \{200, 400, 600, 1000\}$ . The results are summarized in Table 4, where we report the bias, standard error, and MSE. The advantage of the proposed nonlinear modal regression observed from Table 4 shows that employing nonlinear modal estimation can reveal some characteristics of the data that have been ignored by the nonlinear mean or quantile regression. Although the nonlinear modal estimation is biased slightly regarding the coefficients, overall it estimates all coefficients reasonably well.

To confirm the asymptotic property of the proposed nonlinear modal estimator, we report the empirical density of the standardized modal estimate in Figure 7, which shows that the distributions are well approximated by standard normal asymptotics for all cases, coinciding with our expectation. Also, we find that due to the slow convergence rate, the empirical density converges to the density of the standard normal distribution slowly.

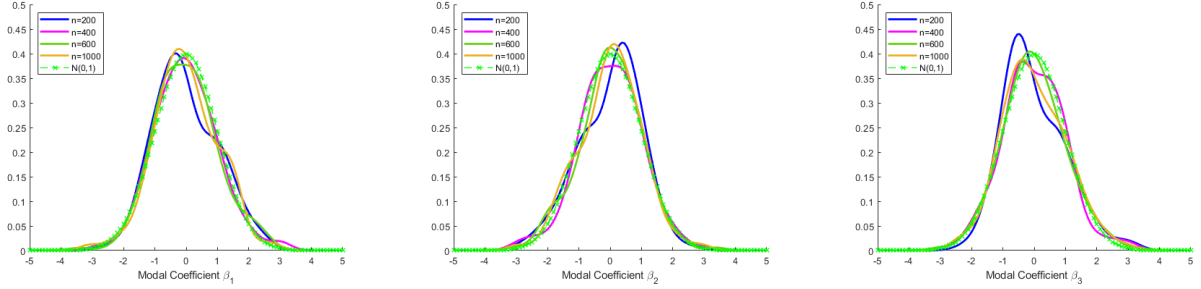


Figure 7: Empirical Density of the Standardized Estimate

Finally, we compare the prediction performance of the proposed nonlinear modal regression to those of nonlinear mean and median regressions by calculating the coverage probabilities of prediction intervals of three different lengths ( $0.1\sigma, 0.2\sigma, 0.5\sigma, \sigma = \sqrt{Var(\epsilon_t)} \approx 0.4330$ ). We use the same DGP procedure as before, except that we implement the out-of-sample prediction with 200 repetitions for the extra  $n$  data points. The results are shown in Figure 8, from which we observe the similar patterns as those in DGP 1. As expected, the nonlinear modal regression performs better than the other two regressions, and the average of coverage probabilities obtained from the nonlinear median regression is close to that of the nonlinear modal regression.

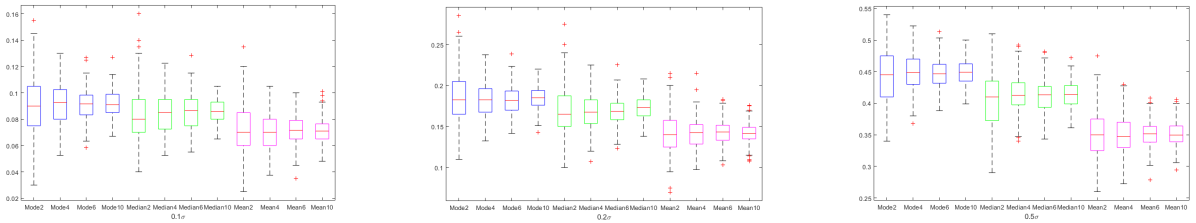
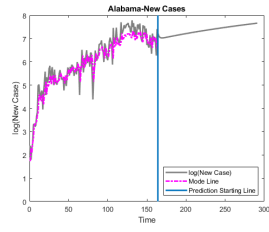
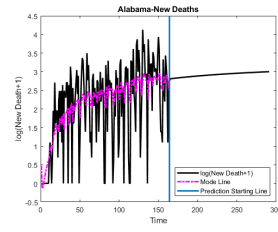


Figure 8: Boxplots of average of coverage probabilities: the numbers 2, 4, 6, and 10 represent the values of  $n = 200, 400, 600$ , and  $1000$ , respectively.

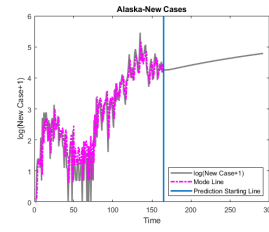
# Online Appendix B: Prediction Results



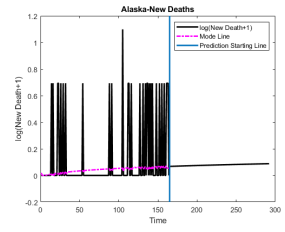
(a) AL-New Cases



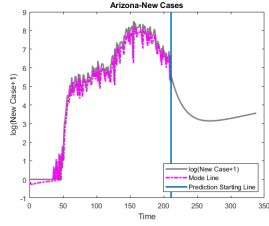
(b) AL-New Deaths



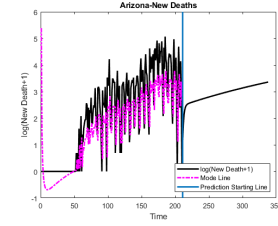
(c) AK-New Cases



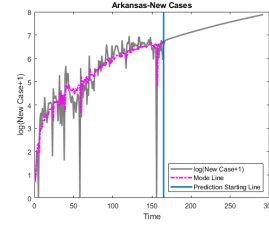
(d) AK-New Deaths



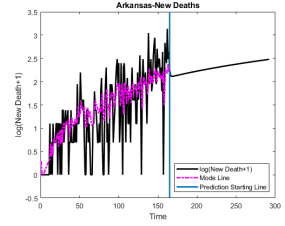
(e) AZ-New Cases



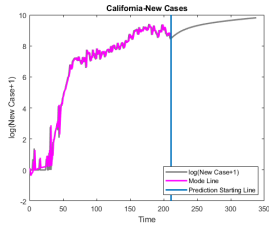
(f) AZ-New Deaths



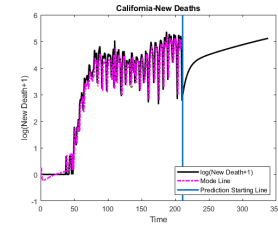
(g) AR-New Cases



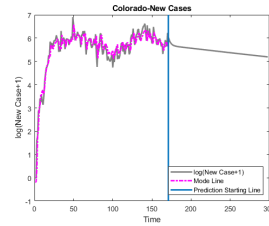
(h) AR-New Deaths



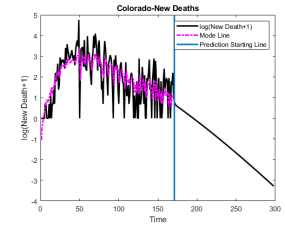
(i) CA-New Cases



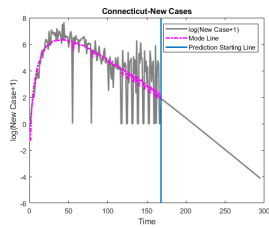
(j) CA-New Deaths



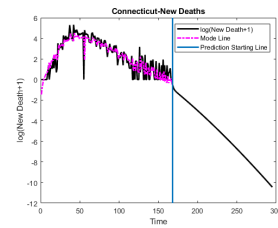
(k) CO-New Cases



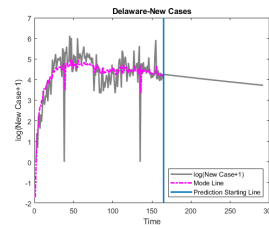
(l) CO-New Deaths



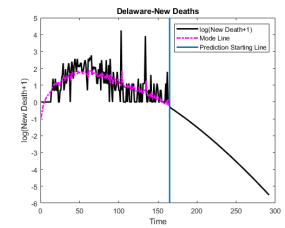
(m) CT-New Cases



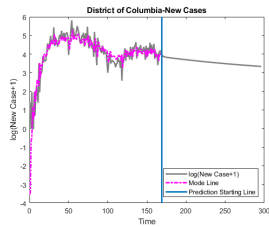
(n) CT-New Deaths



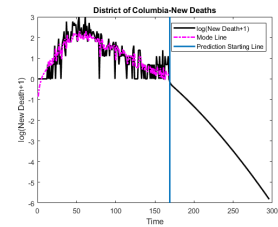
(o) DE-New Cases



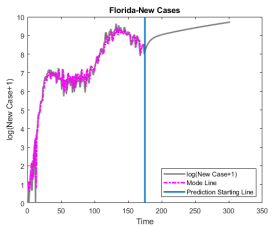
(p) DE-New Deaths



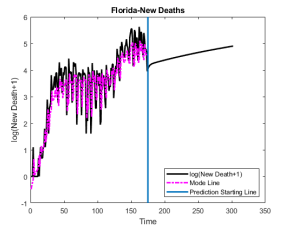
(q) DC-New Cases



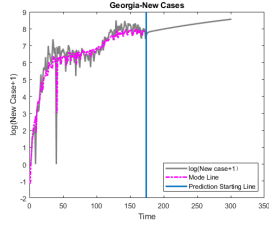
(r) DC-New Deaths



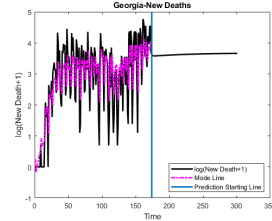
(s) FL-New Cases



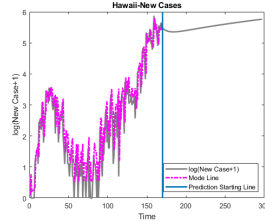
(t) FL-New Deaths



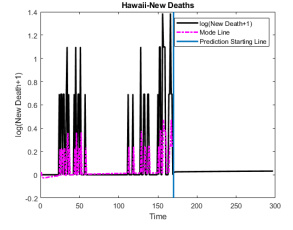
(a) GA-New Cases



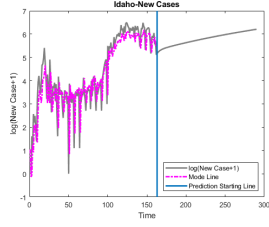
(b) GA-New Deaths



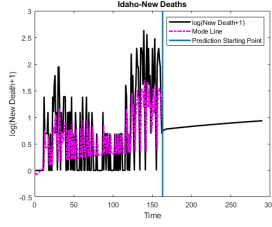
(c) HI-New Cases



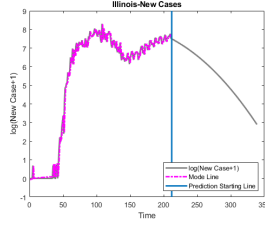
(d) HI-New Deaths



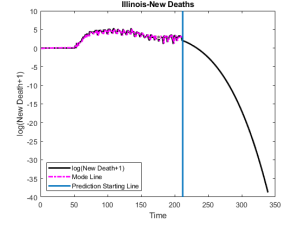
(e) ID-New Cases



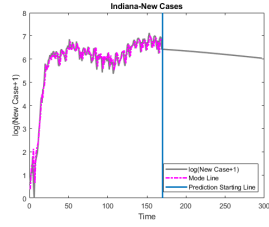
(f) ID-New Deaths



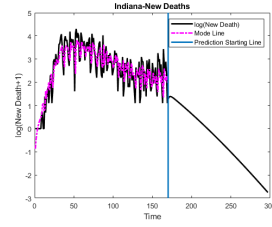
(g) IL-New Cases



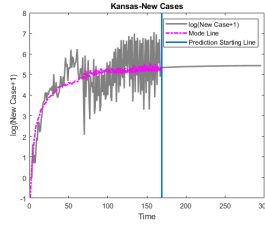
(h) IL-New Deaths



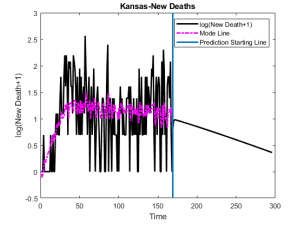
(i) IN-New Cases



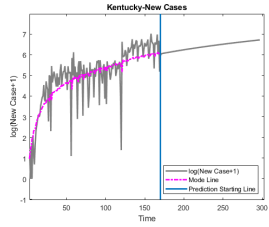
(j) IN-New Deaths



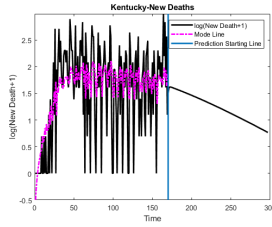
(k) KS-New Cases



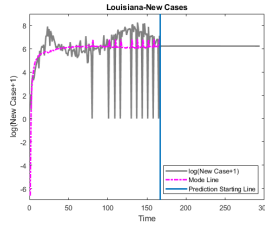
(l) KS-New Deaths



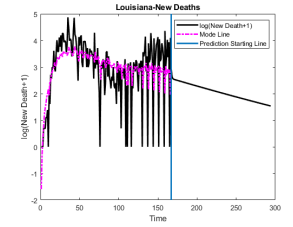
(m) KY-New Cases



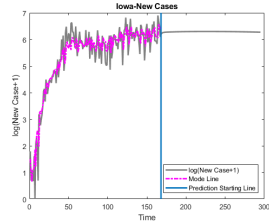
(n) KY-New Deaths



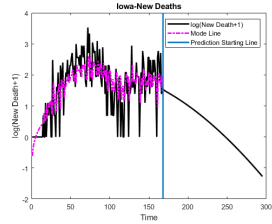
(o) LA-New Cases



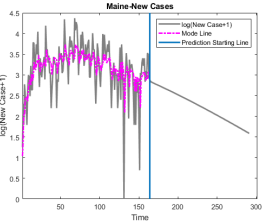
(p) LA-New Deaths



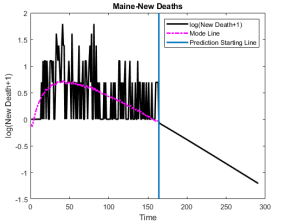
(q) IA-New Cases



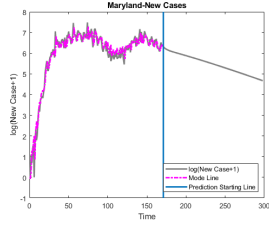
(r) IA-New Deaths



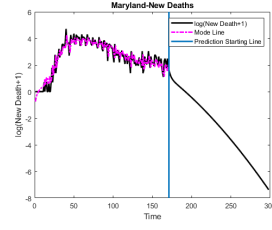
(s) ME-New Cases



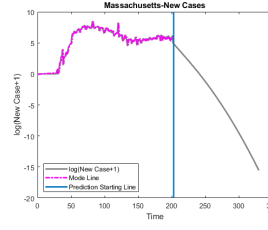
(t) ME-New Deaths



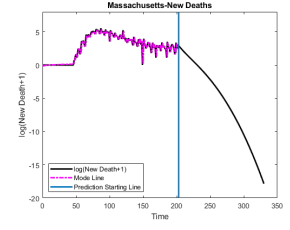
(a) MD-New Cases



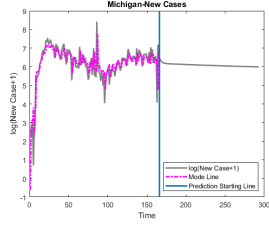
(b) MD-New Deaths



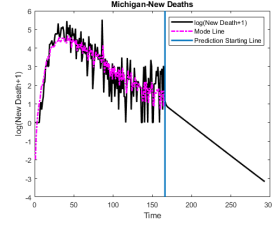
(c) MA-New Cases



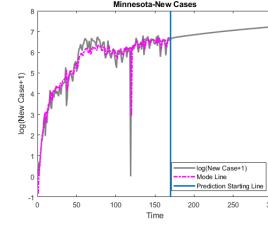
(d) MA-New Deaths



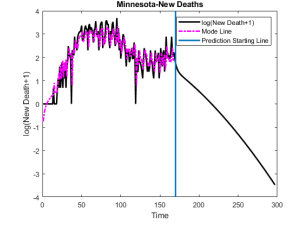
(e) MI-New Cases



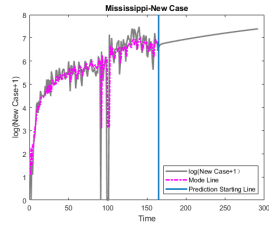
(f) MI-New Deaths



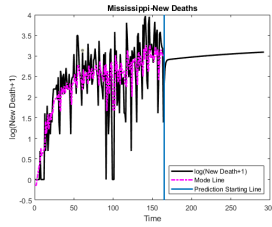
(g) MN-New Cases



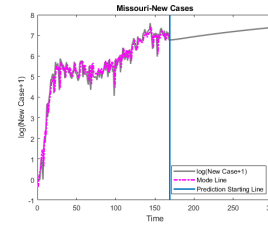
(h) MN-New Deaths



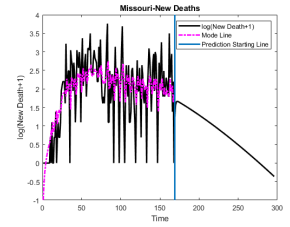
(i) MS-New Cases



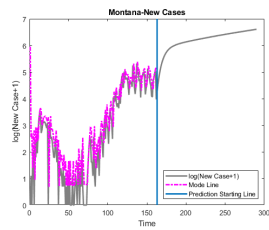
(j) MS-New Deaths



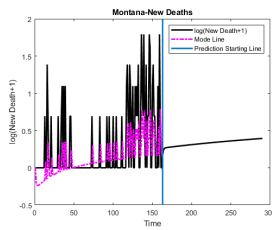
(k) MO-New Cases



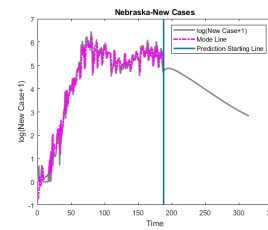
(l) MO-New Deaths



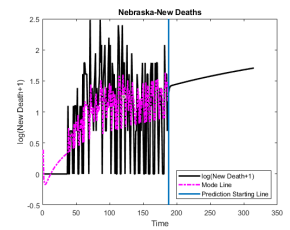
(m) MT-New Cases



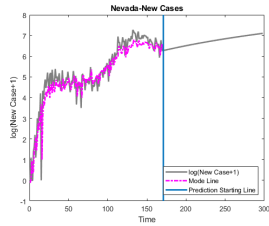
(n) MT-New Deaths



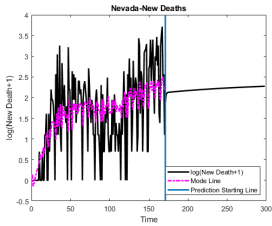
(o) NE-New Cases



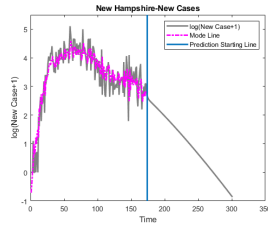
(p) NE-New Deaths



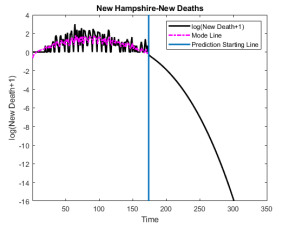
(q) NV-New Cases



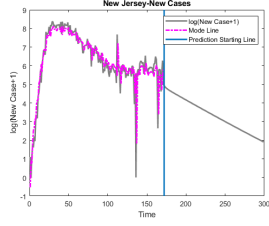
(r) NV-New Deaths



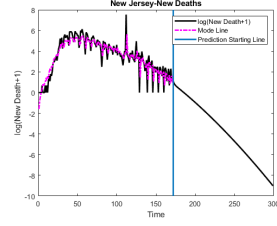
(s) NH-New Cases



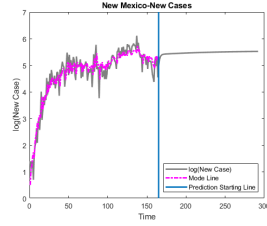
(t) NH-New Deaths



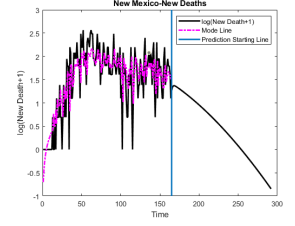
(a) NJ-New Cases



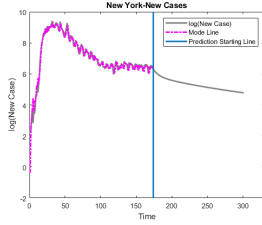
(b) NJ-New Deaths



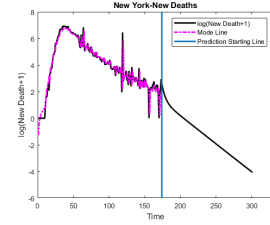
(c) NM-New Cases



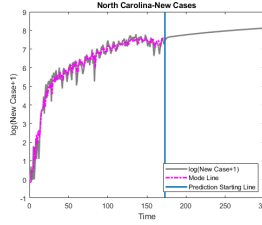
(d) NM-New Deaths



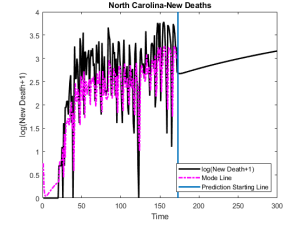
(e) NY-New Cases



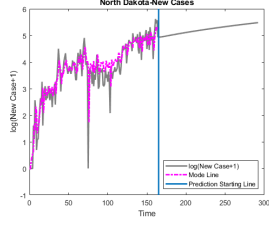
(f) NY-New Deaths



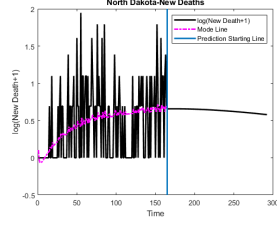
(g) NC-New Cases



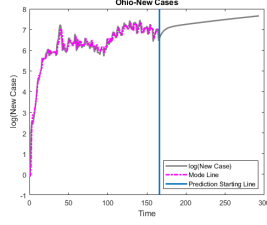
(h) NC-New Deaths



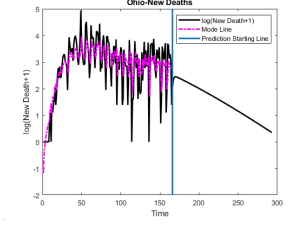
(i) ND-New Cases



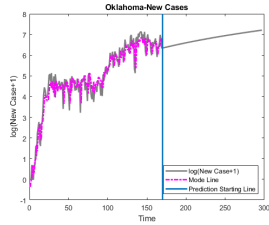
(j) ND-New Deaths



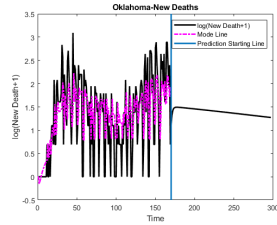
(k) OH-New Cases



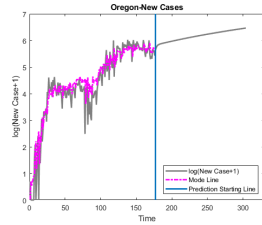
(l) OH-New Deaths



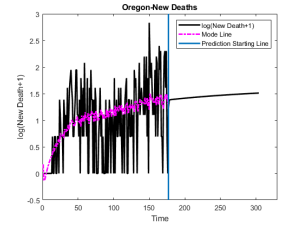
(m) OK-New Cases



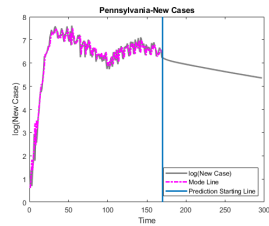
(n) OK-New Deaths



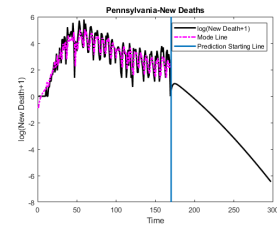
(o) OR-New Cases



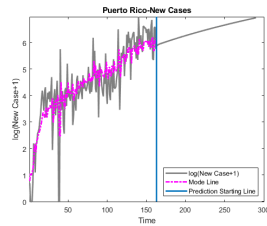
(p) OR-New Deaths



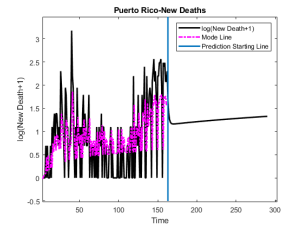
(q) PA-New Cases



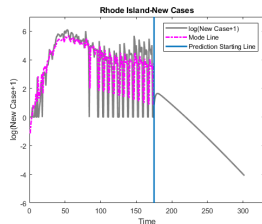
(r) PA-New Deaths



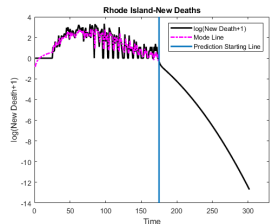
(s) PR-New Cases



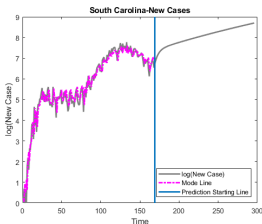
(t) PR-New Deaths



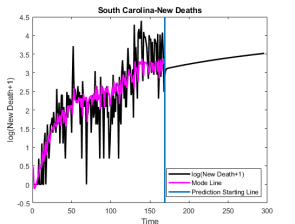
(u) RI-New Cases



(v) RI-New Deaths



(w) SC-New Cases



(x) SC-New Deaths

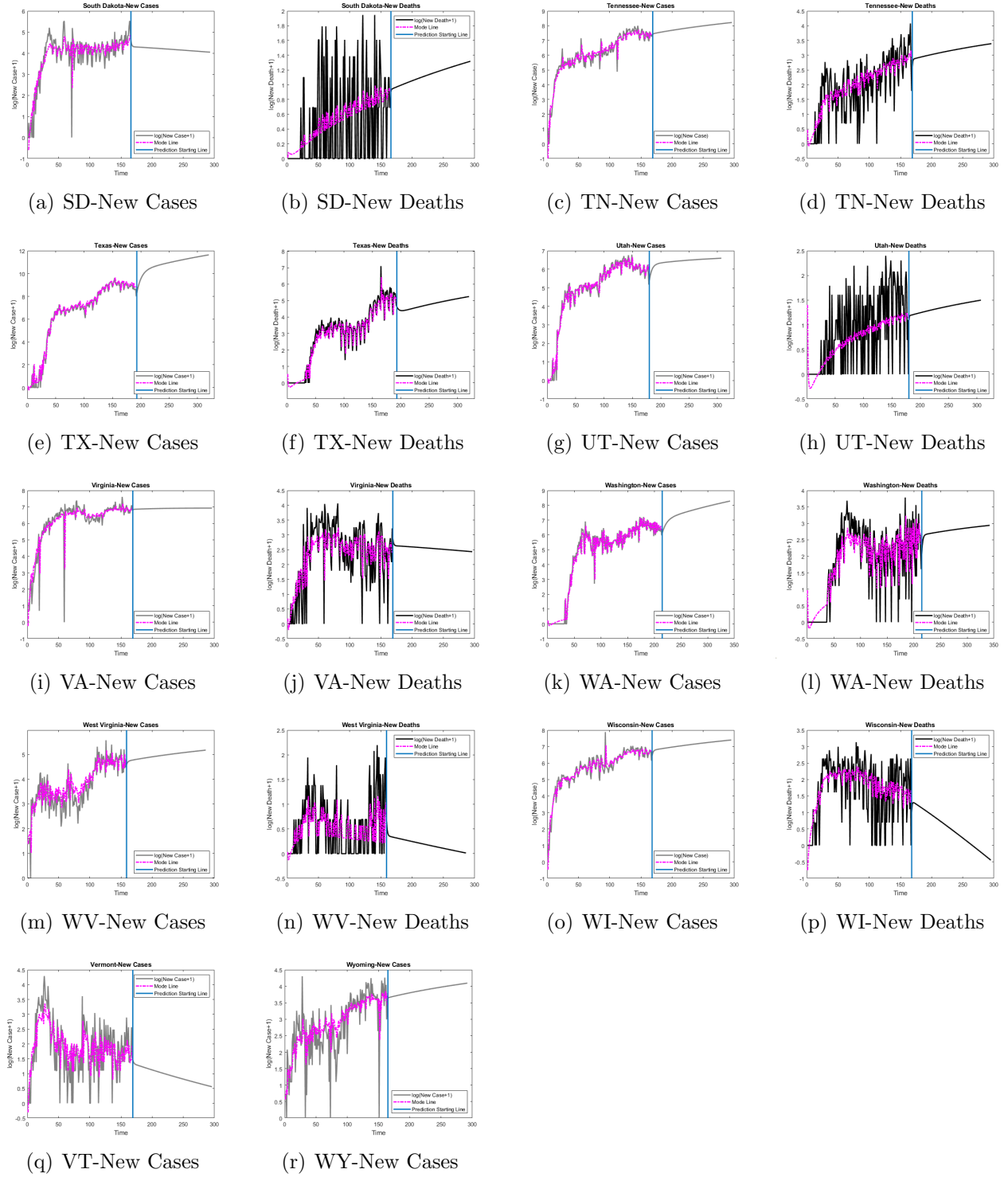


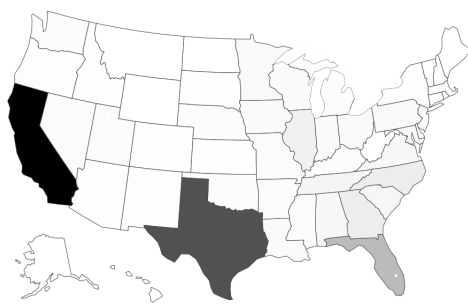
Figure 9: Modal Predicted Curves for COVID-19 Data (with 102 Subfigures)

Table 5: Mean Prediction Results

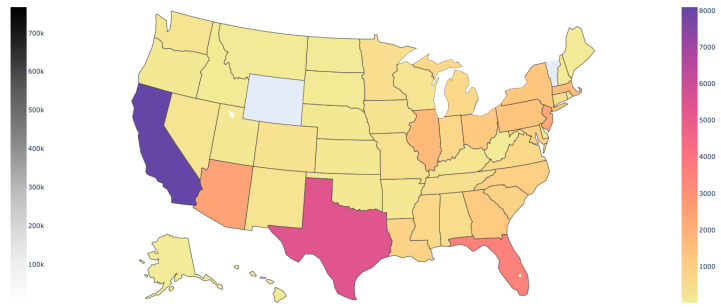
State/ Region	Predictions of Mean Regression 08/24/2020-12/31/2020				State/ Region	Predictions of Mean Regression 08/24/2020-12/31/2020			
	09/30	10/31	11/30	12/31		09/30	10/31	11/30	12/31
Region	Total New Cases				Region	Total New Cases			
AL	48870/50820/57789/68443				AK	1194/1050/1128/1272			
AZ	15024/20065/27064/37230				AR	24261/25938/30280/36676			
CA	766530/1373100/2227000/3624300				CO	14752/13408/13540/14506			
CT	4072/3177/2977/2993				DE	1669/851/513/327			
DC	3222/3074/3132/3809				FL	292230/345800/428590/546660			
GA	107570/116780/137450/167750				HI	927/446/456/495			
ID	7119/6703/7395/8535				IL	92107/106490/120150/140530			
IN	29051/29013/30678/34128				KS	7363/6300/6076/6188			
KY	17889/19020/22091/26655				LA	29173/27480/29072/32398			
IA	30160/33773/39667/48306				ME	1017/891/881/926			
MD	43610/44329/48377/55328				MA	15324/16555/16320/16952			
MI	21141/17795/17438/18211				MN	36291/39496/46507/56771			
MS	25951/26253/29474/34424				MO	27022/25925/28757/22249			
MT	1724/1983/2578/3449				NE	17731/21128/25909/32780			
NV	27695/29232/34130/41366				NH	2164/2159/2284/2546			
NJ	21756/20664/20015/20690				NM	10935/11152/12340/14282			
NY	15157/8463/6178/5053				NC	108280/127690/159380/204510			
ND	3057/2947/3206/3659				OH	40331/38591/40959/45758			
OK	19182/20454/23635/28380				OR	11209/11941/13942/16907			
PA	36024/32024/31966/33884				PR	9894/10666/12575/15378			
RI	2968/2572/2653/2895				SC	58550/66348/80307/100330			
SD	4299/4348/4811/5569				TN	61660/66311/77172/93257			
TX	576360/843030/1235800/1825800				UT	27866/30558/35982/43925			
VA	44744/43204/46378/52139				WA	21885/19643/20076/21652			
WV	3148/3167/3577/4223				WI	32805/34060/38521/45430			
VT	156/128/119/119				WY	1247/1254/1402/1642			

**Note:** The results represent the total number of mean predicted new cases and new deaths between 08/24 and 09/30, between 10/01 and 10/31, between 11/01 and 11/30, and between 12/01 and 12/31, separately.

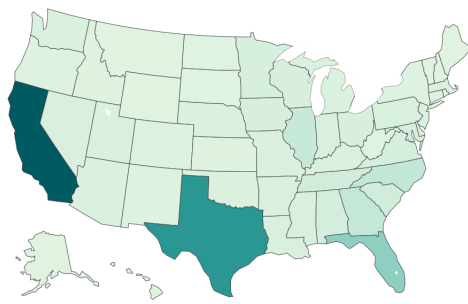




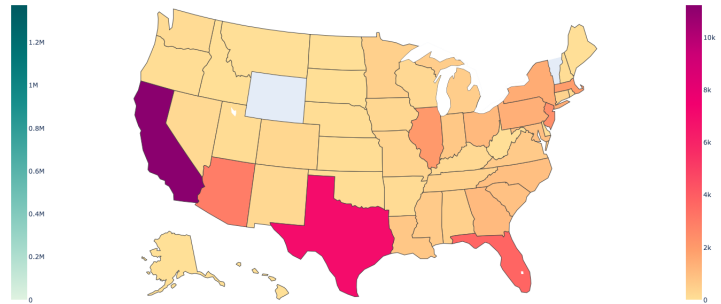
(a) Predicted New Cases 08/24-09/30



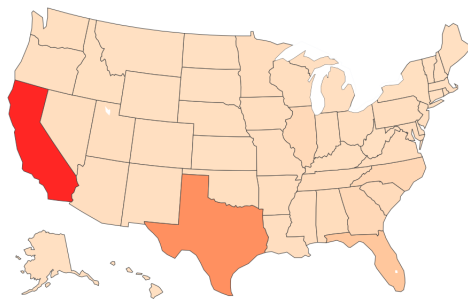
(b) Predicted New Deaths 08/24-09/30



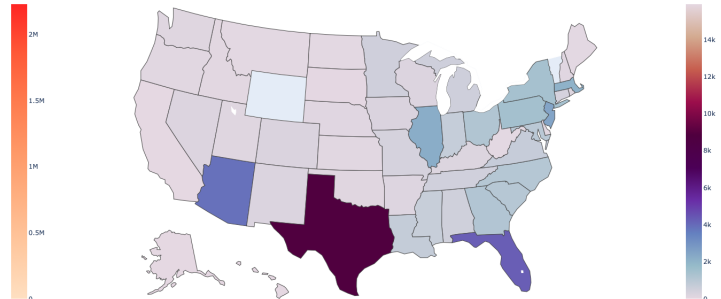
(c) Predicted New Cases 10/01-10/31



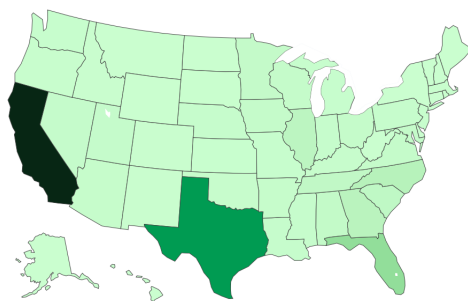
(d) Predicted New Deaths 10/01-10/31



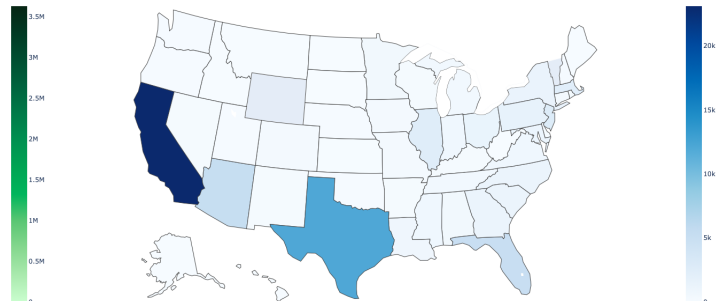
(e) Predicted New Cases 11/01-11/30



(f) Predicted New Deaths 11/01-11/30

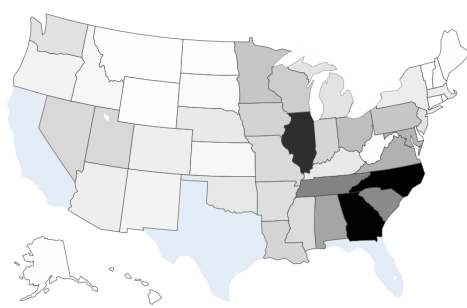


(g) Predicted New Cases 12/01-12/31

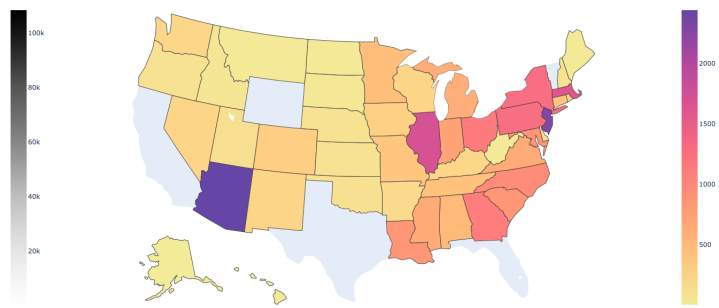


(h) Predicted New Deaths 12/01-12/31

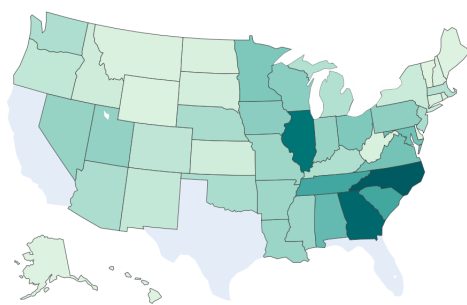
Figure 10: Visualization of the Total Number of Mean Predicted New Cases and New Deaths across the US



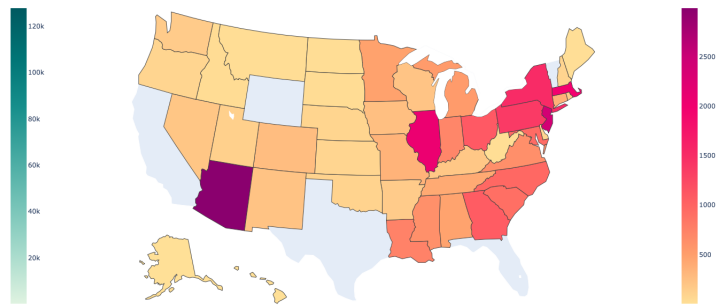
(a) Predicted New Cases 08/24-09/30



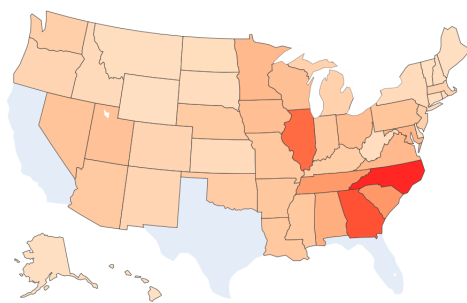
(b) Predicted New Deaths 08/24-09/30



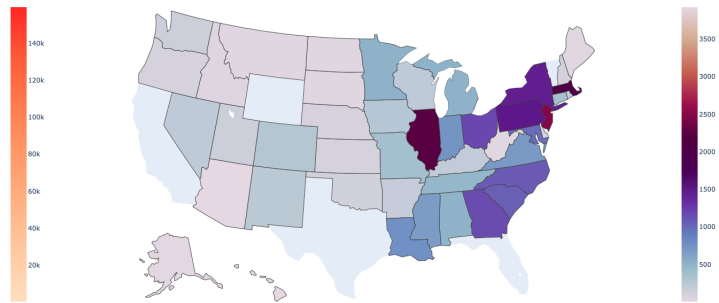
(c) Predicted New Cases 10/01-10/31



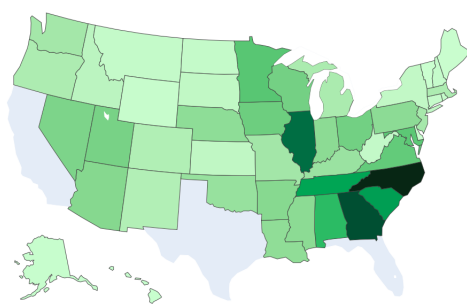
(d) Predicted New Deaths 10/01-10/31



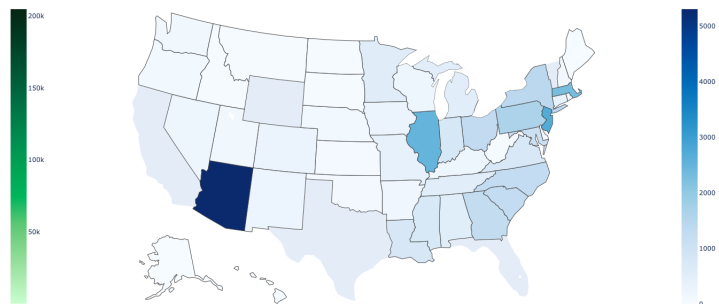
(e) Predicted New Cases 11/01-11/30



(f) Predicted New Deaths 11/01-11/30



(g) Predicted New Cases 12/01-12/31



(h) Predicted New Deaths 12/01-12/31

Figure 11: Visualization of the Total Number of Mean Predicted New cases and New Deaths across the US after Removing CA, TX, and FL

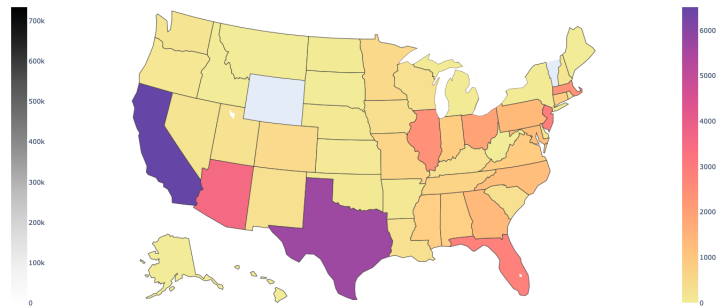
Table 6: Median Prediction Results

Predictions of Median Regression 08/24/2020-12/31/2020					Predictions of Median Regression 08/24/2020-12/31/2020												
State/	09/30	10/31	11/30	12/31	09/30	10/31	11/30	12/31	State/	09/30	10/31	11/30	12/31	09/30	10/31	11/30	12/31
Region	Total New Cases				Total New Deaths				Region	Total New Cases				Total New Deaths			
AL	43268/36844/36099/37303				749/679/688/731				AK	475/223/132/83				1/0/0/0			
AZ	96349/104610/119830/142240				3541/5354/8463/13644				AR	25176/23421/24351/26575				92/81/82/86			
CA	14308/525/560/712				6510/12405/23738/45784				CO	5869/2371/1245/722				463/465/510/589			
CT	191/0/0/0				772/815/922/1090				DE	2094/1140/738/509				78/87/105/132			
DC	1026/467/259/154				114/143/187/253				FL	171590/100470/69169/51631				2852/2293/2036/1878			
GA	65768/48212/40964/37139				1343/1255/1255/1312				HI	280/36/1/0				0/0/0/0			
ID	3656/1733/1063/717				6/0/0/0				IL	30027/16429/11529/8701				2423/4198/7188/12498			
IN	25731/22931/22638/23685				832/830/892/1004				KS	5696/4340/3774/3484				60/36/25/17			
KY	17300/16992/18468/20974				297/315/365/443				LA	8605/3188/1535/826				281/125/65/32			
IA	29469/32280/37435/44906				326/425/587/833				ME	537/262/137/65				0/0/0/0			
MD	23548/15083/11461/9302				1202/1678/2420/3590				MA	4304/1379/567/249				2393/3960/6374/10460			
MI	8786/3015/1431/764				28/0/0/0				MN	33305/42029/56323/77542				509/782/1225/1954			
MS	26843/30351/37510/48298				762/756/825/942				MO	8769/3497/1906/1152				610/600/650/739			
MT	294/70/16/0				1/0/0/0				NE	18871/39136/80801/166650				95/131/185/265			
NV	14496/8446/6044/4696				177/136/122/115				NH	320/60/0/0				157/190/234/298			
NJ	3563/955/360/143				2951/3435/4158/5222				NM	9164/8242/8142/8429				269/345/463/640			
NY	9993/2968/1304/655				35/0/0/0				NC	109250/131800/168280/220850				1224/1898/3053/5003			
ND	2267/1673/1478/1396				27/29/34/41				OH	33541/27396/26027/26434				1944/2155/2550/3138			
OK	8657/5115/3685/2881				74/44/28/17				OR	7796/5594/4716/4246				127/121/131/149			
PA	16090/7599/4916/3497				1371/2155/3367/5361				PR	4351/2828/2188/1823				14/6/1/0			
RI	203/0/0/0				115/189/309/505				SC	49197/48444/52641/59809				314/281/285/306			
SD	1378/724/448/291				0/0/0/0				TN	39326/30302/26894/25423				665/758/946/1230			
TX	734130/1615100/3953800/11162000				5684/9608/17281/31416				UT	19514/17215/16870/17393				236/281/362/485			
VA	28189/19544/15676/13191				864/1290/1960/3046				WA	17706/11465/8894/7425				149/67/31/9			
WV	1379/800/573/447				0/0/0/0				WI	22313/17704/16013/15417				317/286/287/304			
VT	20/0/0/0				-				WY	766/516/413/356				-			

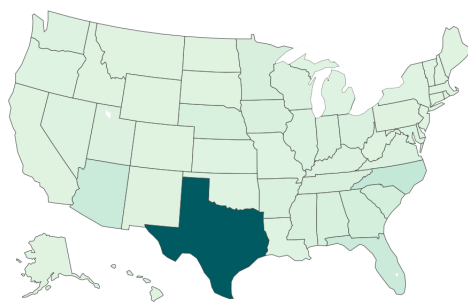
**Note:** The results represent the total number of median predicted new cases and new deaths between 08/24 and 09/30, between 10/01 and 10/31, between 11/01 and 11/30, and between 12/01 and 12/31, separately.



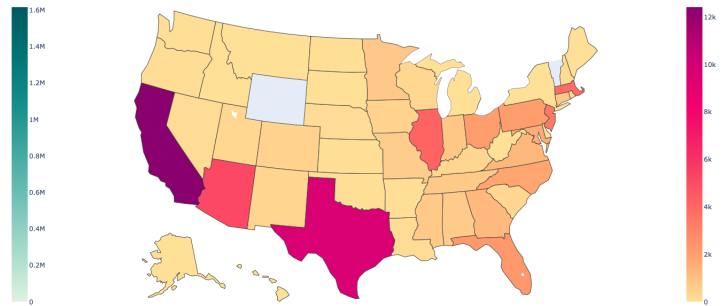
(a) Predicted New Cases 08/24-09/30



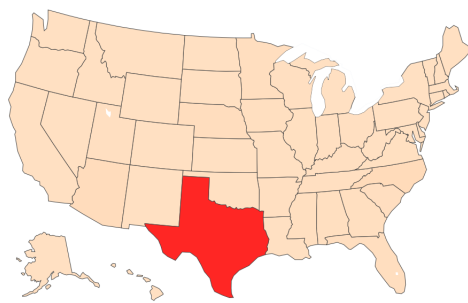
(b) Predicted New Deaths 08/24-09/30



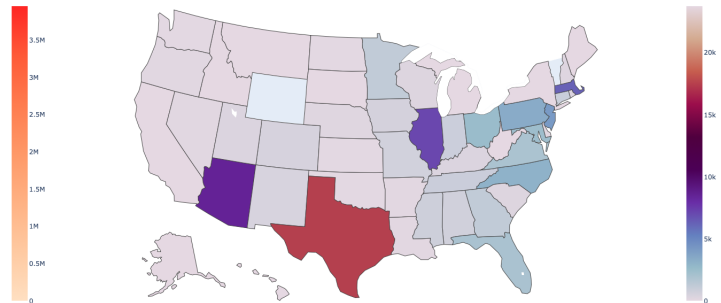
(c) Predicted New Cases 10/01-10/31



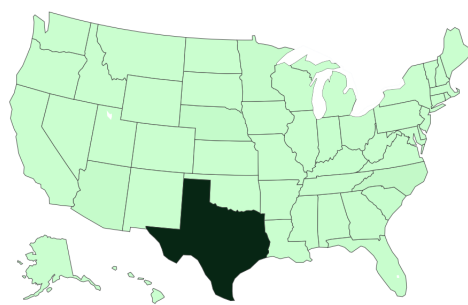
(d) Predicted New Deaths 10/01-10/31



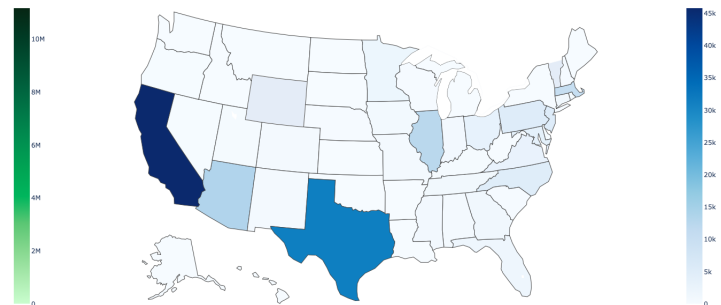
(e) Predicted New Cases 11/01-11/30



(f) Predicted New Deaths 11/01-11/30

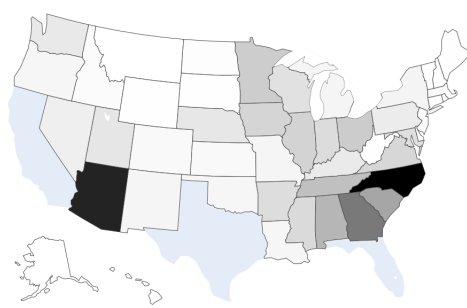


(g) Predicted New Cases 12/01-12/31

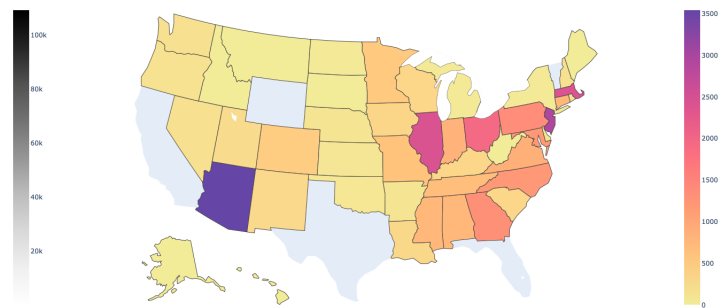


(h) Predicted New Deaths 12/01-12/31

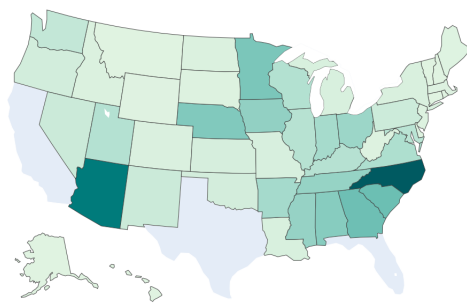
Figure 12: Visualization of the Total Number of Median Predicted New Cases and New Deaths across the US



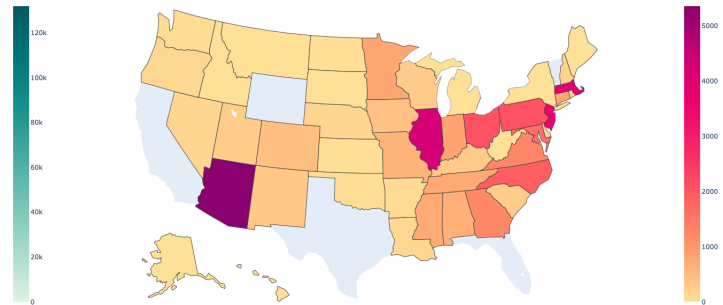
(a) Predicted New Cases 08/24-09/30



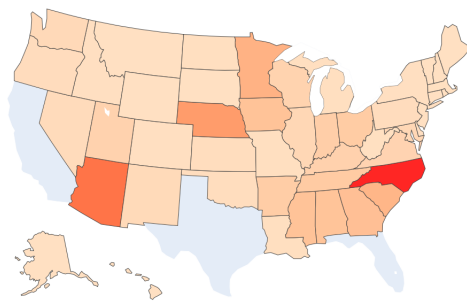
(b) Predicted New Deaths 08/24-09/30



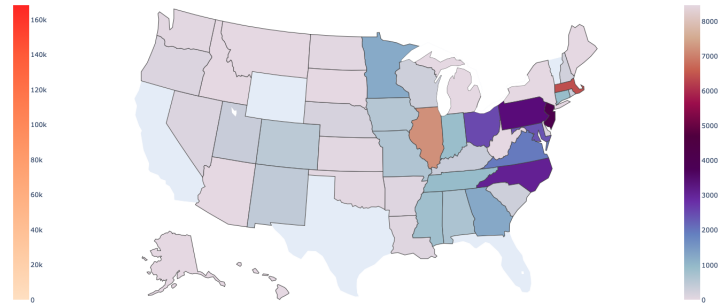
(c) Predicted New Cases 10/01-10/31



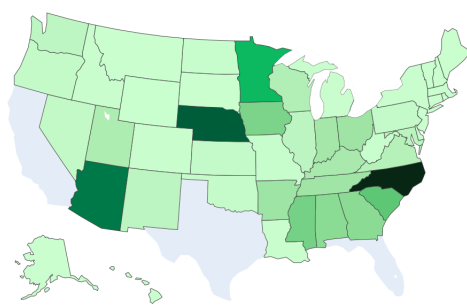
(d) Predicted New Deaths 10/01-10/31



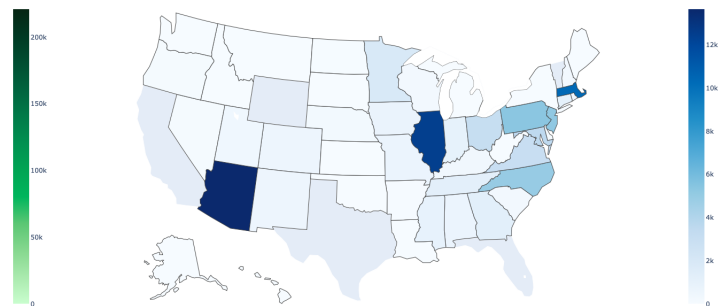
(e) Predicted New Cases 11/01-11/30



(f) Predicted New Deaths 11/01-11/30



(g) Predicted New Cases 12/01-12/31



(h) Predicted New Deaths 12/01-12/31

Figure 13: Visualization of the Total Number of Median Predicted New cases and New Deaths across the US after Removing CA, TX, and FL

Table 7: Performance of Robust Nonlinear Regression

State/ Region	New Cases-Robust			New Deaths-Robust			State/ Region	New Cases-Robust			New Deaths-Robust		
	<i>MSE</i>	<i>MAPE</i>	<i>R</i> <sup>2</sup>	<i>MSE</i>	<i>MAPE</i>	<i>R</i> <sup>2</sup>		<i>MSE</i>	<i>MAPE</i>	<i>R</i> <sup>2</sup>	<i>MSE</i>	<i>MAPE</i>	<i>R</i> <sup>2</sup>
AL	0.2007	5.4940	0.9161	0.7066	35.3855	0.1122	AK	0.3885	12.7831	0.7275	0.1595	97.3091	0.1463
AZ	2.5575	22.5294	0.9460	2.5937	26.9784	0.6549	AR	2.2192	3.7485	0.7661	0.5898	26.5847	0.4290
CA	0.0906	2.8343	0.9777	0.8732	16.4897	0.8931	CO	0.3207	8.7383	0.8942	1.8599	82.806	0.5567
CT	4.8553	26.5439	0.5000	4.9880	260.7927	0.8783	DE	0.2780	5.5840	0.6285	1.0676	148.8698	0.4284
DC	0.6002	17.3181	0.9171	0.7145	54.4436	0.6071	FL	0.1135	3.4220	0.9260	0.4740	11.2328	0.8565
GA	0.1183	3.6592	0.8044	2.4378	34.6543	0.5156	HI	3.3976	31.5063	0.7168	0.6253	101.0654	0.1578
ID	0.2622	7.7879	0.8689	1.4770	59.1572	0.1742	IL	0.8745	11.5423	0.9596	1.6400	40.5490	0.8292
IN	0.3373	7.7480	0.8994	1.5867	40.6630	0.7309	KS	1.3625	18.5649	0.5903	0.6607	44.1437	0.2176
KY	0.3196	8.0646	0.6920	0.7973	34.2550	0.2377	LA	6.5711	11.3220	0.1565	2.1475	28.6599	0.5012
IA	0.3185	7.0682	0.9028	0.9446	35.3794	0.3372	ME	0.5315	32.0621	0.3515	0.1434	82.6993	0.1461
MD	0.0739	3.4723	0.8834	0.8983	37.1860	0.8073	MA	0.6194	12.2318	0.9179	0.8348	30.0713	0.8133
MI	0.6607	11.6073	0.8139	3.2961	74.8745	0.8800	MN	0.0831	3.6945	0.8830	1.9138	59.7755	0.6962
MS	0.2668	6.7109	0.5675	0.6071	24.9590	0.4331	MO	0.3487	7.9736	0.8765	0.5841	23.2154	0.3250
MT	1.2728	22.7149	0.8096	0.7873	99.9689	0.3158	NE	0.2141	7.0617	0.9322	0.9591	70.7803	0.1873
NV	0.0582	3.1247	0.8741	1.5149	37.9698	0.1333	NH	0.6283	22.143	0.8065	0.4255	107.9180	0.2807
NJ	1.5676	21.6682	0.9248	0.9672	37.9099	0.7485	NM	0.2635	8.6004	0.9100	0.2207	23.2218	0.4272
NY	0.5781	11.0520	0.9675	0.8059	26.2690	0.9534	NC	0.2796	6.4587	0.9072	0.6852	29.3216	0.3412
ND	0.4570	11.6518	0.6758	0.5641	71.2429	0.0470	OH	0.0538	2.8937	0.9547	0.8634	45.9123	0.4345
OK	0.2534	6.5370	0.8761	1.6995	51.0717	0.1887	OR	0.0430	2.8567	0.8400	0.7043	45.5382	0.0751
PA	0.1930	5.7194	0.8966	2.6139	49.9138	0.6553	PR	0.8039	12.2884	0.6582	2.8252	76.9209	0.1113
RI	7.0776	55.1564	0.8410	1.9236	171.2987	0.5591	SC	0.3297	7.1750	0.9707	0.5805	17.6734	0.5032
SD	0.9844	18.4277	0.7414	0.2325	34.3005	0.0426	TN	0.0811	3.2367	0.8982	0.6446	21.3525	0.3131
TX	0.3289	5.4990	0.9555	0.9205	16.1135	0.6834	UT	0.2908	7.8331	0.9353	0.4397	33.3464	0.2023
VA	0.2202	5.7012	0.7957	1.9174	50.2956	0.5348	WA	0.0928	3.6163	0.8321	1.1136	30.4684	0.3710
WV	0.3458	10.4524	0.6618	1.5842	89.7621	0.2224	WI	0.0753	3.4719	0.9029	0.9968	40.6244	0.2209
VT	0.7406	35.9913	0.3202	-	-	-	WY	0.7861	10.6908	0.5088	-	-	-

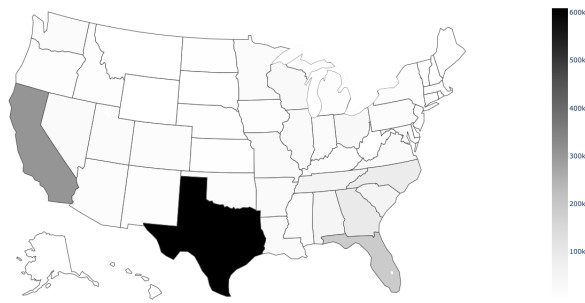
**Note:** When the dataset has zero values, we instead use  $\log(Y_t + 1)$  transformation for the whole data. When calculating *MAPE*, we eliminate all  $\log(1)=0$  values. For VT and WY, the existing death data are not sufficient for predicting (most values are zero). Thus, we do not have the predicted new deaths results for these two states.

Table 8: Robust Nonlinear Prediction Results

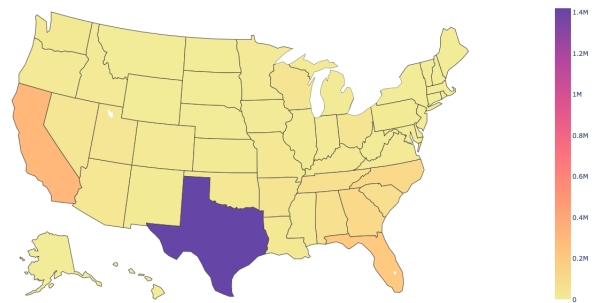
Predictions of Median Regression 08/24/2020-12/31/2020			Predictions of Median Regression 08/24/2020-12/31/2020		
State/ Region	09/30 10/31 11/30 12/31	09/30 10/31 11/30 12/31	State/ Region	09/30 10/31 11/30 12/31	09/30 10/31 11/30 12/31
	Total New Cases	Total New Deaths		Total New Cases	Total New Deaths
AL	52931/55706/63849/76134	552/510/532/586	AK	1536/1437/1569/1795	0/0/0/0
AZ	15206/20274/27283/37434	1865/1949/2241/2682	AR	26990/28443/32930/39603	183/171/180/201
CA	306260/308790/306840/304910	6897/7681/9077/11127	CO	14451/13052/13165/14089	419/395/402/434
CT	3255/1603/1335/1218	454/482/479/504	DE	2118/1233/865/652	69/64/63/66
DC	818/300/127/49	0/0/0/0	FL	189400/186470/205470/237050	4150/4407/5062/6051
GA	95096/98265/110910/130400	1249/1250/1338/1506	HI	1003/455/458/490	0/0/0/0
ID	10573/10666/12514/15211	33/27/27/28	IL	25964/12102/7621/5195	84/0/0/0
IN	18089/13205/11413/10561	73/0/0/0	KS	6870/5620/5205/5104	54/34/25/19
KY	16815/16205/17302/19313	80/27/3/0	LA	23041/18606/17290/17112	429/240/162/117
IA	11305/6250/3692/2184	74/4/0/0	ME	497/216/96/32	0/0/0/0
MD	17625/10218/7245/5476	49/0/0/0	MA	5306/1726/822/428	97/0/0/0
MI	10870/6402/4764/3865	15/0/0/0	MN	20185/13920/10458/7955	7/0/0/0
MS	20143/18454/19001/20586	756/737/796/901	MO	23143/20811/22070/24579	148/58/18/0
MT	1204/972/1039/1167	1/0/0/0	NE	5283/1703/502/102	95/90/95/105
NV	24047/23057/25641/29808	183/149/139/138	NH	285/52/0/0	91/81/81/85
NJ	3366/989/404/179	130/0/0/0	NM	8883/7962/7913/8276	71/9/0/0
NY	9184/3145/1621/942	160/0/0/0	NC	88199/94810/109980/132090	574/450/408/393
ND	3088/2880/3055/3407	9/0/0/0	OH	35499/31629/31941/34096	431/175/72/21
OK	17795/18535/21066/24923	66/43/31/24	OR	10236/10619/12164/14505	93/83/84/90
PA	15969/8497/6114/4801	87/0/0/0	PR	9143/9436/10764/12782	24/16/12/9
RI	328/0/0/0	75/56/54/55	SC	54864/60776/72346/89090	836/858/968/1139
SD	1464/827/555/397	38/36/37/41	TN	56251/58664/66665/78847	411/383/401/444
TX	608440/932910/1419800/2176000	2981/2836/3102/3555	UT	21981/21948/23970/27318	106/97/101/110
VA	24413/15732/11791/9318	627/627/657/726	WA	19508/16521/16055/16500	159/79/43/21
WV	2516/2298/2430/2706	5/0/0/0	WI	28588/28124/30467/34550	72/29/10/0
VT	66/34/20/11	-	WY	1171/1157/1275/1475	-

**Note:** The results represent the total number of median predicted new cases and new deaths between 08/24 and 09/30, between 10/01 and 10/31, between 11/01 and 11/30, and between 12/01 and 12/31, separately.

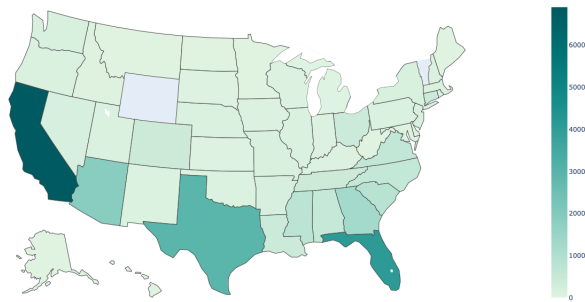




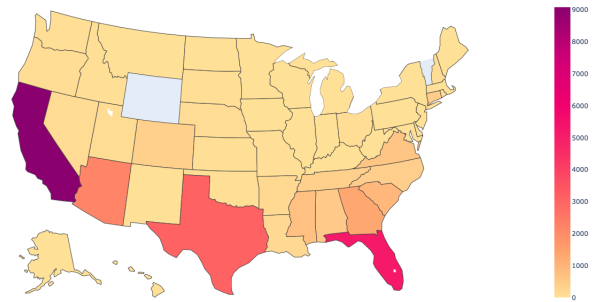
(a) Predicted New Cases 08/24-09/30



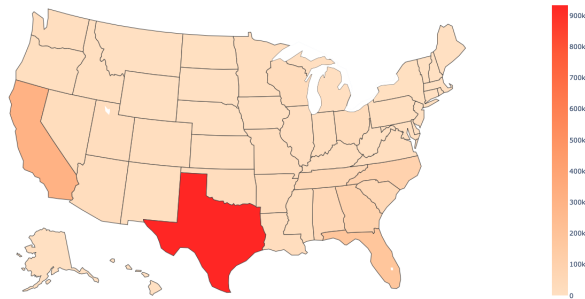
(b) Predicted New Deaths 08/24-09/30



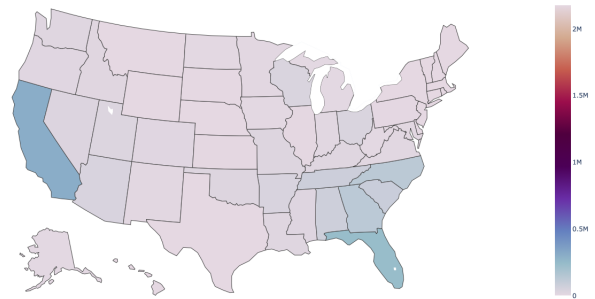
(c) Predicted New Cases 10/01-10/31



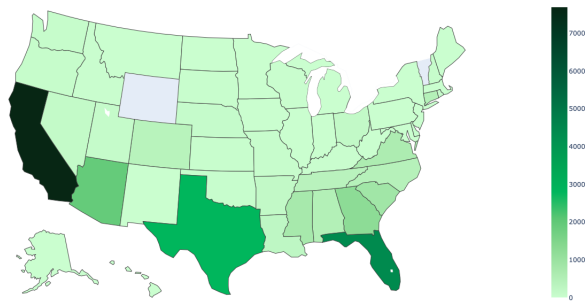
(d) Predicted New Deaths 10/01-10/31



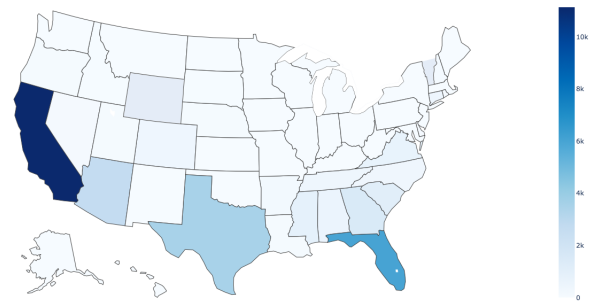
(e) Predicted New Cases 11/01-11/30



(f) Predicted New Deaths 11/01-11/30

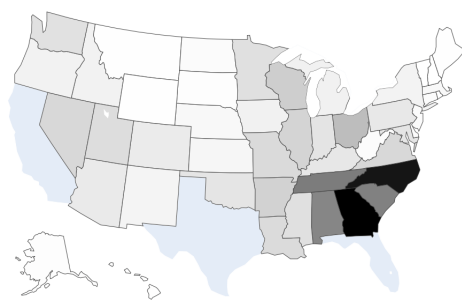


(g) Predicted New Cases 12/01-12/31

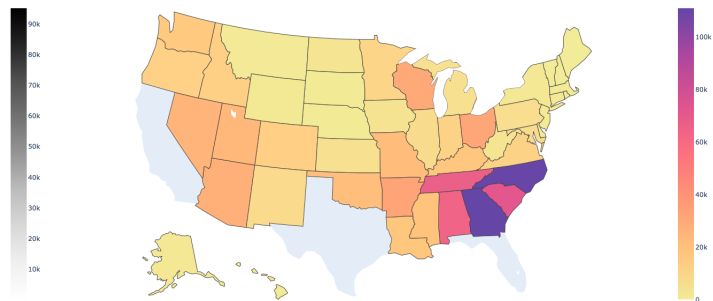


(h) Predicted New Deaths 12/01-12/31

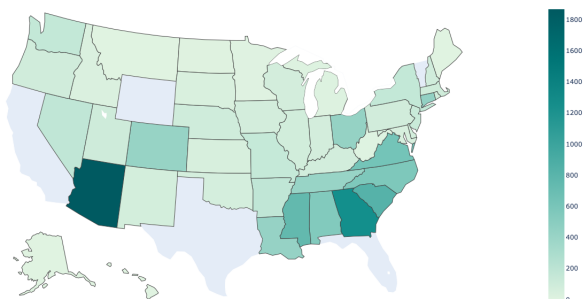
Figure 14: Visualization of the Total Number of Robust Predicted New Cases and New Deaths across the US



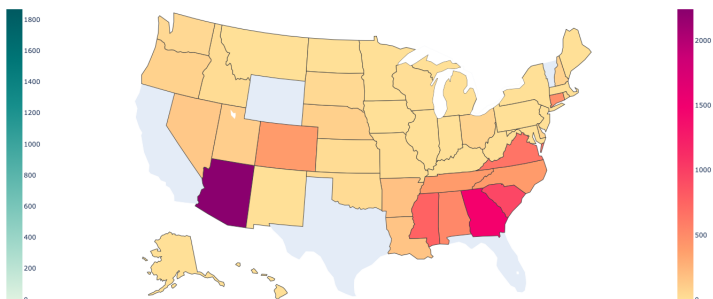
(a) Predicted New Cases 08/24-09/30



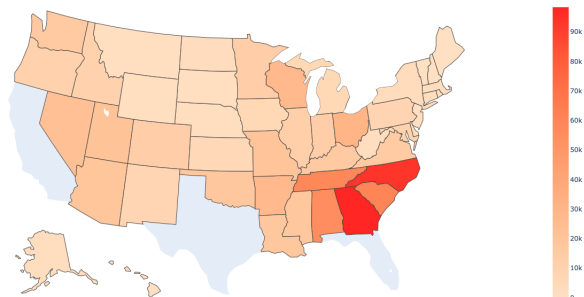
(b) Predicted New Deaths 08/24-09/30



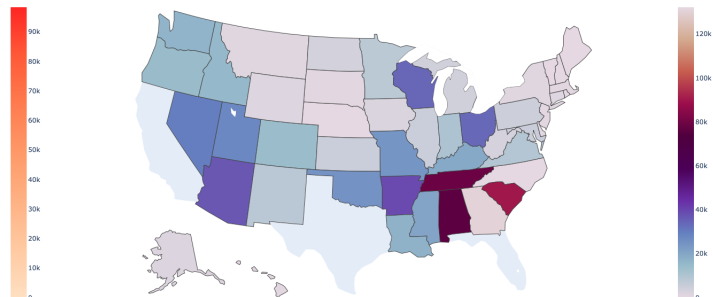
(c) Predicted New Cases 10/01-10/31



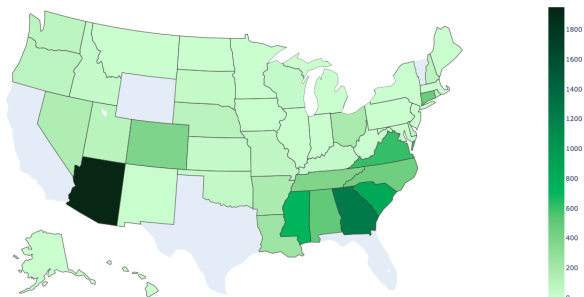
(d) Predicted New Deaths 10/01-10/31



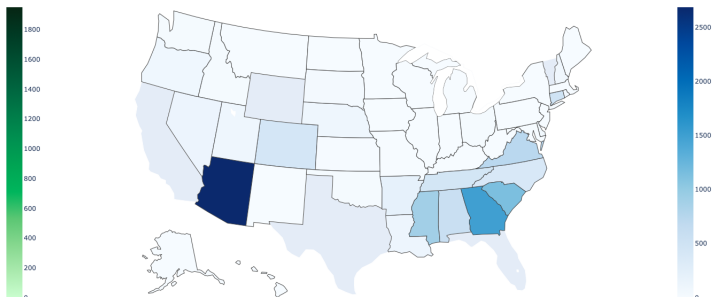
(e) Predicted New Cases 11/01-11/30



(f) Predicted New Deaths 11/01-11/30



(g) Predicted New Cases 12/01-12/31



(h) Predicted New Deaths 12/01-12/31

Figure 15: Visualization of the Total Number of Robust Predicted New cases and New Deaths across the US after Removing CA, TX, and FL

## Online Appendix C: Outline Proofs

To show the asymptotic negligence of dependence with a large sample size for the nonlinear modal regression proposed in this paper, we establish the following lemma, where  $K^{(j)}(\cdot)$  represents the  $j$ th derivative of  $K(\cdot)$ .

**Lemma 1.** *Under suitable conditions, with  $nh^3 \rightarrow \infty$  held, we have*

$$-\frac{1}{nh} \sum_{t=1}^n K^{(1)}\left(\frac{\epsilon_t}{h}\right) = \frac{h^2}{2} q^{(3)}(0 | X)(1 + o(1)).$$

*Proof.* Define  $Z_{n,t} = -\frac{1}{h} K^{(1)}\left(\frac{\epsilon_t}{h}\right)$ , we obtain

$$\mathbb{E}(Z_{n,1} | X) = \int \frac{\epsilon}{h^3} K\left(\frac{\epsilon}{h}\right) q(\epsilon | X) d\epsilon = \frac{1}{h} \int t K(t) q(th | X) dt = \frac{h^2}{2} q^{(3)}(0 | X)(1 + o(1)).$$

Therefore, we have

$$\mathbb{E}\left\{-\frac{1}{nh} \sum_{t=1}^n K^{(1)}\left(\frac{\epsilon_t}{h}\right)\right\} = \frac{h^2}{2} q^{(3)}(0 | X)(1 + o(1)).$$

Note that

$$\sum_{t=1}^n Z_{n,t} = \mathbb{E}\left(\sum_{t=1}^n Z_{n,t}\right) + O_p\left(\sqrt{\text{Var}\left(\sum_{t=1}^n Z_{n,t}\right)}\right),$$

and the stationary of  $\{\epsilon_t\}_{t=1}^n$  gives

$$\text{Var}\left(\sum_{t=1}^n Z_{n,t}\right) = n\mathbb{E}Z_{n,1}^2 + 2\sum_{j=2}^n (n-j+1) \text{Cov}(Z_{n,t}, Z_{n,j}).$$

Conditioning on  $X$ , we can have

$$\begin{aligned} \mathbb{E}(Z_{n,1}^2 | X) &= \frac{1}{h^2} \int \frac{\epsilon^2}{h^4} K^2\left(\frac{\epsilon}{h}\right) q(\epsilon | X) d\epsilon = \frac{1}{h^3} \int t^2 K^2(t) q(th | X) dt \\ &= \frac{1}{h^3} \int q(0 | X) t^2 K^2(t) dt (1 + o(1)). \end{aligned}$$

To obtain an upper bound for the second term on the right-hand side of  $\text{Var}(\sum_{t=1}^n Z_{n,t})$ , without loss of the generality, let  $d_n$  be a sequence of positive integers satisfying  $d_n \rightarrow \infty$  and  $d_n h \rightarrow 0$  as  $n \rightarrow \infty$ , we can split it into two terms

$$\sum_{j=2}^n |\text{Cov}(Z_{n,1}, Z_{n,j})| = \sum_{j=2}^{d_n} |\text{Cov}(Z_{n,1}, Z_{n,j})| + \sum_{j=d_n+1}^n |\text{Cov}(Z_{n,1}, Z_{n,j})|.$$

Conditioning on  $X_t$  and  $X_j$ , we can have the following result, where

$$\begin{aligned}
|\mathbb{E}(Z_{n,t}Z_{n,j} \mid X_t, X_j)| &\leq \mathbb{E}|(Z_{n,t}Z_{n,j} \mid X_t, X_j)| = \mathbb{E}\left|\mathbb{E}\left[\frac{1}{h^2}K^{(1)}\left(\frac{\epsilon_t}{h}\right)K^{(1)}\left(\frac{\epsilon_j}{h}\right) \mid X_t, X_j\right]\right| \\
&= \mathbb{E}\left|\iint \frac{1}{h^2}\frac{\epsilon}{h^2}K\left(\frac{\epsilon}{h}\right)\frac{\epsilon^*}{h^2}K\left(\frac{\epsilon^*}{h}\right)q(\epsilon, \epsilon^* \mid X_t, X_j)d\epsilon d\epsilon^*\right| \\
&\leq C_1\frac{1}{h^2}
\end{aligned}$$

in which  $C_1$  is a constant number. Therefore, we have

$$\sum_{j=2}^{d_n} |\text{Cov}(Z_{n,t}, Z_{n,j})| \leq C_1 \frac{1}{h^2} \sum_{j=2}^{d_n} 1 = o(nh^{-3}).$$

By applying Davydov's inequality, we have

$$|\text{Cov}(Z_{n,1}, Z_{n,j})| \leq C_2[\rho(j-1)]^{\delta/(2+\delta)} (\mathbb{E}|Z_{n,1}|^{2+\delta})^{2/(2+\delta)}$$

and obtain

$$\mathbb{E}(|Z_{n,t}|^{2+\delta} \mid X_t) = \mathbb{E}\left|\mathbb{E}\left[\frac{1}{h}K^{(1)}\left(\frac{\epsilon_t}{h}\right) \mid X_t\right]\right|^{2+\delta} \leq C_3h^{(2+\delta)^2},$$

where  $C_2$  and  $C_3$  are constants. By choosing  $d_n$  such that  $d_n^\gamma h^{-7} = O(1)$ , we have

$$\begin{aligned}
\sum_{j=d_n+1}^n |\text{Cov}(Z_{n,1}, Z_{n,j})| &\leq C_3 \sum_{j=d_n+1}^n C_2[\rho(j-1)]^{\delta/(2+\delta)} (h^{(2+\delta)^2})^{2/(2+\delta)} = C_4h^4 \sum_{k=d_n}^n [\rho(k)]^{\delta/(2+\delta)} \\
&\leq C_4d_n^{-\gamma}h^4 \sum_{k=d_n}^n [\rho(k)]^{\delta/(2+\delta)} = o(nh^{-3}),
\end{aligned}$$

where  $C_4$  is a constant. Then, we obtain

$$\text{Var}\left(\sum_{t=1}^n Z_{n,t}\right) = O(nh^{-3}).$$

With the assumption that  $nh^3 \rightarrow \infty$ , we obtain the result of Lemma 1.

□

*Outline the Proof of Theorem 2.1* Based on the result from Lemma 1, we can observe that under suitable conditions, the covariance of two different error terms can be dominated by the expectation of the squared error, which is the underlying result to prove the consistency and

asymptotic normality of modal estimator. Let  $r^{(j)}(\cdot)$  represent the  $j$ th derivative  $r(\cdot)$ . Define  $\delta_n = h^2 + \sqrt{(nh^3)^{-1}}$ , then it is sufficient to show that for any given  $\eta$ , there exists a large number constant  $c$  such that

$$P \left\{ \sup_{\|\mu\|=c} Q_n(\beta_0 + \delta_n \mu) < Q_n(\beta_0) \right\} \geq 1 - \eta,$$

where  $\|\cdot\|$  represents the Euclidean distance. It implies that with probability tending to 1, there is a local maximum in the ball  $\{\beta_0 + \delta_n \mu : \|\mu\| \leq c\}$ . Using the Taylor expansion, it follows that

$$\begin{aligned} & Q_n(\beta_0 + \delta_n \mu) - Q_n(\beta_0) \\ &= \frac{1}{nh} \sum_{t=1}^n \left[ K \left( \frac{Y_t - r(X_t, \beta_0 + \delta_n \mu)}{h} \right) - K \left( \frac{Y_t - r(X_t, \beta_0)}{h} \right) \right] \\ &\approx \frac{1}{nh} \sum_{t=1}^n \left[ K \left( \frac{\epsilon_t - r^{(1)}(X_t, \beta_0) \delta_n \mu}{h} \right) - K \left( \frac{\epsilon_t}{h} \right) \right] \\ &= \frac{1}{nh} \sum_{t=1}^n \left\{ -K^{(1)} \left( \frac{\epsilon_t}{h} \right) \left( \frac{r^{(1)}(X_t, \beta_0) \delta_n \mu}{h} \right) + \frac{1}{2} K^{(2)} \left( \frac{\epsilon_t}{h} \right) \left( \frac{r^{(1)}(X_t, \beta_0) \delta_n \mu}{h} \right)^2 \right. \\ &\quad \left. - \frac{1}{6} K^{(3)} \left( \frac{\epsilon_t^*}{h} \right) \left( \frac{r^{(1)}(X_t, \beta_0) \delta_n \mu}{h} \right)^3 \right\} \\ &=: I_1 + I_2 + I_3, \end{aligned}$$

where  $\epsilon_t^*$  is between  $\epsilon_t$  and  $\epsilon_t - r^{(1)}(X_t, \beta_0) \delta_n \mu$ . Based on the result  $T_n = \mathbb{E}(T_n) + O_p(\sqrt{\text{Var}(T_n)})$ , we could consider each part of above Taylor expansion following [Yao and Li \(2014\)](#) and [Ullah et al. \(2021\)](#), where we have

$$\begin{aligned} I_1 &= O_p(\delta_n c h^2) + O_p(\sqrt{\delta_n^2 c^2 (nh^3)^{-1}}) = O_p(\delta_n^2 c), \\ I_2 &= O_p(\delta_n^2 c^2 h^2) + O_p((\delta_n c)^4 h^{-5}), \\ I_3 &= O_p(\delta_n^3 c^3) + O_p((\delta_n c)^6 h^{-7}). \end{aligned}$$

Based on these, we can choose  $c$  bigger enough such that  $I_2$  dominates both  $I_1$  and  $I_3$  with probability  $1 - \eta$  under certain conditions. Because the second term is negative, thus  $P \left\{ \sup_{\|\mu\|=c} Q_n(\beta_0 + \delta_n \mu) < Q_n(\beta_0) \right\} \geq 1 - \eta$  holds. □

*Outline the Proof of Theorem 2.2* Following the same steps as proving Theorem 2.1, recall that

$$-\frac{1}{nh^2} \sum_{t=1}^n K^{(1)} \left( \frac{Y_t - r(X_t, \hat{\beta})}{h} \right) r^{(1)}(X_t, \hat{\beta}) = 0.$$

By taking Taylor expansion, we could obtain

$$\begin{aligned}
& -\frac{1}{nh^2} \sum_{t=1}^n K^{(1)}\left(\frac{\epsilon_t}{h}\right) r^{(1)}(X_t, \beta) \\
& + \frac{1}{nh^3} \sum_{t=1}^n K^{(2)}\left(\frac{\epsilon_t}{h}\right) r^{(1)}(X_t, \beta) (-\{r^{(1)}(X_t, \beta)\}^T (\hat{\beta} - \beta_0)) \\
& - \frac{1}{nh^4} \sum_{t=1}^n K^{(3)}\left(\frac{\tilde{\epsilon}_t^*}{h}\right) r^{(1)}(X_t, \beta) \left(-\{r^{(1)}(X_t, \beta)\}^T (\hat{\beta} - \beta_0)\right)^2 = 0
\end{aligned}$$

at  $\beta = \beta_0$ , where  $\tilde{\epsilon}_t^*$  is between  $\epsilon_t$  and  $\epsilon_t - r^{(1)}(X_t, \beta)(\hat{\beta} - \beta_0)$ . From the Proof of Theorem 2.1, we can see that the third part of the above equation is dominated by the second part. We then mainly focus on the first two parts of the above equation. By some direct calculations, we can obtain

$$\begin{aligned}
& \mathbb{E} \left( -\frac{1}{nh^2} \sum_{t=1}^n K^{(1)}\left(\frac{\epsilon_t}{h}\right) r^{(1)}(X_t, \beta) \right) = \frac{h^2}{2} q^{(3)}(0 | X) r^{(1)}(X_t, \beta), \\
& \mathbb{E} \left( \frac{1}{nh^3} \sum_{t=1}^n K^{(2)}\left(\frac{\epsilon_t}{h}\right) r^{(1)}(X_t, \beta) \{r^{(1)}(X_t, \beta)\}^T \right) = q^{(2)}(0 | X) r^{(1)}(X_t, \beta) \{r^{(1)}(X_t, \beta)\}^T.
\end{aligned}$$

Based on the above two equations, we can achieve

$$\hat{\beta} - \beta_0 = \frac{h^2}{2} [q^{(2)}(0 | X) r^{(1)}(X_t, \beta) \{r^{(1)}(X_t, \beta)\}^T]^{-1} [q^{(3)}(0 | X) r^{(1)}(X_t, \beta)].$$

Combining the results obtained from Lemma 1, we could obtain

$$\text{Var} \left( -\frac{1}{nh^2} \sum_{t=1}^n K^{(1)}\left(\frac{\epsilon_t}{h}\right) r^{(1)}(X_t, \beta) \right) = \frac{\int t^2 K^2(t) dt}{nh^3} q(0 | X) r^{(1)}(X_t, \beta) \{r^{(1)}(X_t, \beta)\}^T (1 + o_p(1)).$$

Define  $W_n = -\sqrt{\frac{h}{n}} \sum_{t=1}^n K^{(1)}\left(\frac{\epsilon_t}{h}\right) r^{(1)}(X_t, \beta)$ . To show Theorem 2.2, it is sufficient to show that

$$T_n = \sqrt{nh^3} W_n \xrightarrow{d} \mathcal{N}(0, T),$$

where  $T = \int t^2 K^2(t) dt q(0 | X) r^{(1)}(X_t, \beta) \{r^{(1)}(X_t, \beta)\}^T$ . By Slutsky's theorem and the above two equations, we can obtain Theorem 2.2. To show the above equation, we prove that for any unit vector  $d \in \mathbb{R}^p$ ,

$$\{d^T \text{Cov}(T_n) d\}^{-1/2} \{d^T T_n - d^T \mathbb{E}(T_n)\} \xrightarrow{d} N(0, 1).$$

Then, we can check Lyapunov's condition by following the arguments in Yao and Li (2014) directly, where we can show  $n\mathbb{E}|\xi_1|^3 \rightarrow 0$  and  $\xi_t = -\frac{1}{\sqrt{nh}} K^{(1)}\left(\frac{\epsilon_t}{h}\right) d^T r^{(1)}(X_t, \beta)$ .  $\square$