



Mathematical Models for COVID-19 Pandemic: A Comparative Analysis

Downloaded from: <https://research.chalmers.se>, 2024-07-17 23:10 UTC

Citation for the original published paper (version of record):

Adiga, A., Dubhashi, D., Lewis, B. et al (2020). Mathematical Models for COVID-19 Pandemic: A Comparative Analysis. *Journal of the Indian Institute of Science*, 100(4): 793-807.

<http://dx.doi.org/10.1007/s41745-020-00200-6>

N.B. When citing this work, cite the original published paper.



Mathematical Models for COVID-19 Pandemic: A Comparative Analysis

Aniruddha Adiga^{1*}, Devdatt Dubhashi², Bryan Lewis¹, Madhav Marathe^{1,3*},
Srinivasan Venkatramanan¹ and Anil Vullikanti^{1,3}

Abstract | COVID-19 pandemic represents an unprecedented global health crisis in the last 100 years. Its economic, social and health impact continues to grow and is likely to end up as one of the worst global disasters since the 1918 pandemic and the World Wars. Mathematical models have played an important role in the ongoing crisis; they have been used to inform public policies and have been instrumental in many of the social distancing measures that were instituted worldwide. In this article, we review some of the important mathematical models used to support the ongoing planning and response efforts. These models differ in their use, their mathematical form and their scope.

1 Introduction

The ongoing COVID-19 pandemic is the most significant pandemic since the 1918 Influenza pandemic. It has already caused over 21 Million confirmed cases and 758,000 deaths.¹ The economic impact is already in trillions of dollars. As in other pandemics, researchers and public health policy makers are interested in questions such as,² (i) How did it start? (ii) How is it likely to progress and how can we control it? (iii) How can we intervene while balancing public health and economic impact? (iv) Why did some countries do better than other countries thus far into the pandemic? In particular, models and their projections/forecasts have received unprecedented attention. With a multitude of modeling frameworks, underlying assumptions, available datasets and the region/timeframe being modeled, these projections have varied widely, causing confusion among end-users and consumers. We believe an overview (non-exhaustive) of the current modeling landscape will benefit the readers and also serve as a historical record for future efforts.

1.1 Role of Models

Models have been used by mathematical epidemiologists to support a broad range of policy questions. Their use during COVID-19 has been widespread. In general, the type and form of models used in epidemiology depend on the phase of the epidemic. Before an epidemic, models are used for planning and identifying critical gaps and prepare plans to detect and respond in the event of a pandemic. At the start of a pandemic, policy makers are interested in asking questions such as: (i) where and how did the pandemic start, (ii) risk of its spread in the region, (iii) risk of importation in other regions of the world, (iv) basic understanding of the pathogen and its epidemiological characteristics. As the pandemic takes hold, researchers begin investigating: (i) various intervention and control strategies; usually pharmaceutical interventions do not work in the event of a pandemic and thus non-pharmaceutical interventions are most appropriate, (ii) forecasting the epidemic incidence rate, hospitalization rate and mortality rate, (iii) efficiently allocating scarce medical resources to treat the patients and (iv) understanding the change in individual and collective behavior and adherence to public policies. After the pandemic starts to slow down, modelers are interested in developing models related to recovery and long-term impacts caused by the pandemic.

¹ The numbers reported are as of August 14, 2020. See <https://coronavirus.jhu.edu/map.html> and <https://nssac.bii.virginia.edu/covid-19/dashboard/> for most up to date surveillance information.

² see <https://www.nytimes.com/news-event/coronavirus>.

¹ Biocomplexity Institute and Initiative, University of Virginia, Charlottesville, USA.

² Department of Computer Science and Engineering, Chalmers University, Gothenburg, Sweden.

³ Department of Computer Science, University of Virginia, Charlottesville, USA.

*aniruddha@virginia.edu
marathe@virginia.edu

As a result comparing models needs to be done with care. When comparing models: one needs to specify: (a) the purpose of the model, (b) the end user to whom the model is targeted, (c) the spatial and temporal resolution of the model, (d) and the underlying assumptions and limitations. We illustrate these issues by summarizing a few key methods for *projection and forecasting* of disease outcomes in the US and Sweden.

Organization. The paper is organized as follows. In Sect. 2 we give preliminary definitions. Section 3 discusses US and UK centric models developed by researchers at the Imperial College. Section 4 discusses metapopulation models focused on the US that were developed by our group at UVA and the models developed by researchers at Northeastern University. Section 5 describes models developed Swedish researchers for studying the outbreak in Sweden. In Sect. 6 we discuss methods developed for forecasting. Section 8 contains discussion, model limitations and concluding remarks. In a companion paper that appears in this special issue, we address certain complementary issues related to pandemic planning and response, including role of data and analytics.

Important note. The primary purpose of the paper is to highlight some of the salient computational models that are currently being used to support COVID-19 pandemic response. These models, like all models, have their strengths and weaknesses—they have all faced challenges arising from the lack of timely data. Our goal is **not** to pick winners and losers among these model; each model has been used by policy makers and continues to be used to advice various agencies. Rather, our goal is to introduce to the reader a range of models that can be used in such situations. A simple model is no better or worse than a complicated model. The suitability of a specific model for a given question needs to be evaluated by the decision maker and the modeler.

2 Background: Computational Methods for Epidemiology

Epidemiological models fall in two broad classes: statistical models that are largely data driven and mechanistic models that are based on underlying theoretical principles developed by scientists on how the disease spreads.

Data-driven models use statistical and machine learning methods to forecast outcomes, such as case counts, mortality and hospital demands. This is a very active area of research, and a broad class of techniques have been

developed, including auto-regressive time series methods, Bayesian techniques and deep learning^{1, 2, 3, 4, 5, 6}. Mechanistic models of disease spread within a population^{7, 8, 9, 10} use mechanistic (also referred to as procedural or algorithmic) methods to describe the evolution of an epidemic through a population. The most common of these is the SIR type models. Hybrid models that combine mechanistic models with data driven machine learning approaches are also starting to become popular, e.g.,¹¹.

2.1 Mass Action Compartmental Models

There are a number of models, which are referred to as SIR class of models. These partition a population of N agents into three sets, each corresponding to a disease state, which is one of: susceptible (S), infective (I) and removed or recovered (R). The specific model then specifies how susceptible individuals become infectious, and then recover. In its simplest form (referred to as the basic compartmental model)^{7, 9, 10}, the population is assumed to be completely mixed. Let $S(t)$, $I(t)$ and $R(t)$ denote the number of people who are susceptible, infected and recovered states at time t , respectively. Let $s(t) = S(t)/N$, $i(t) = I(t)/N$ and $r(t) = R(t)/N$; then, $s(t) + i(t) + r(t) = 1$. Then, the SIR model can be described by the following system of ordinary differential equations

$$\frac{ds}{dt} = \beta si, \quad \frac{di}{dt} = \beta si - \gamma i, \quad \frac{dr}{dt} = \gamma i,$$

where β is referred to as the transmission rate, and γ is the recovery rate. A key parameter in such a model is the “reproductive number”, denoted by $R_0 = \beta/\gamma$. At the start of an epidemic, much of the public health effort is focused on estimating R_0 from observed infections¹².

Mass action compartmental models have been the workhorse for epidemiologists and have been widely used for over 100 years. Their strength comes from their simplicity, both analytically and from the standpoint of understanding the outcomes. Software systems have been developed to solve such models and a number of associated tools have been built to support analysis using such models.

2.2 Structured Metapopulation Models

Although simple and powerful, mass action compartmental models do not capture the inherent heterogeneity of the underlying populations. Significant amount of research has been conducted

to extend the model, usually in two broad ways. The first involves structured metapopulation models—these construct an abstraction of the mixing patterns in the population into m different sub-populations, e.g., age groups and small geographical regions, and attempt to capture the heterogeneity in mixing patterns across sub-populations. In other words, the model has states $S_j(t), I_j(t), R_j(t)$ for each subpopulation j . The evolution of a compartment $X_j(t)$ is determined by mixing within and across compartments. For instance, survey data on mixing across age groups¹³ have been used to construct age structured metapopulation models¹⁴. More relevant for our paper are spatial metapopulation models, in which the subpopulations are connected through airline and commuter flow networks^{15, 16, 17, 18, 19}.

Main steps in constructing structured metapopulation models. This depends on the disease, population and the type of question being studied. The key steps in the development of such models for the spread of diseases over large populations include

- Constructing subpopulations and compartments: the entire population V is partitioned into subpopulations V_j , within which the mixing is assumed to be complete. Depending on the disease model, there are S_j, E_j, I_j, R_j compartments corresponding to the subpopulation V_j (and more, depending on the disease)—these represent the number of individuals in V_j in the corresponding state
- Mixing patterns among compartments: state transitions between compartments might depend on the states of individuals within the subpopulations associated with those compartments, as well as those who they come in contact with. For instance, the $S_j \rightarrow E_j$ transition rate might depend on I_k for all the subpopulations who come in contact with individuals in V_j . Mobility and behavioral datasets are needed to model such interactions.

Such models are very useful at the early days of the outbreak, when the disease dynamics are driven to a large extent by mobility—these can be captured more easily within such models, and there is significant uncertainty in the disease model parameters. They can also model coarser interventions such as reduced mobility between spatial units and reduced mixing rates. However, these models become less useful to model the effect of detailed interventions (e.g., voluntary

home isolation, school closures) on disease spread in and across communities.

2.3 Agent-Based Network Models

Agent-based networked models (sometimes just called as agent-based models) extend metapopulation models further by explicitly capturing the interaction structure of the underlying populations. Often such models are also resolved at the level of single individual entities (animals, humans, etc.). In this class of models, the epidemic dynamics can be modeled as a diffusion process on a specific undirected contact network $G(V, E)$ on a population V —each edge $e = (u, v) \in E$ implies that individuals (also referred to as nodes) $u, v \in V$ come into contact³. Let $N(v)$ denote the set of neighbors of v . For instance, in the graph in Fig. 1, we have $V = \{a, b, c, d\}$ and $E = \{(a, b), (a, c), (b, d), (cd)\}$. Node a has b and c as neighbors, so $N(a) = \{b, c\}$. The SIR model on the graph G is a dynamical process in which each node is in one of the S, I or R states. Infection can potentially spread from u to v along edge $e = (u, v)$ with a probability of $\beta(e, t)$ at time instant t after u becomes infected, conditional on node v remaining uninfected until time t —this is a discrete version of the rate of infection for the ODE model discussed earlier. We let $I(t)$ denote the set of nodes that become infected at time t . The (random) subset of edges on which the infections spread represents a disease outcome, and is referred to as a *dendogram*. This dynamical system starts with a configuration in which there are one or more nodes in state I and reaches a fixed point in which all nodes are in states S or R . Figure 1 shows an example of the SIR model on a network.

Main steps in setting up an agent-based model. While the specific steps depend on the disease, the population, and the type of question being studied, the general process involves the following steps:

- Construct a network representation G : the set V is the population in a region, and is available from different sources, such as Census and Landscan. However, the contact patterns are more difficult to model, as no real data are available on contacts between people at a large scale. Instead, researchers have tried to model

³ Note that though edge e is represented as a tuple (u, v) , it actually denotes the set $\{u, v\}$, as is common in graph theory.

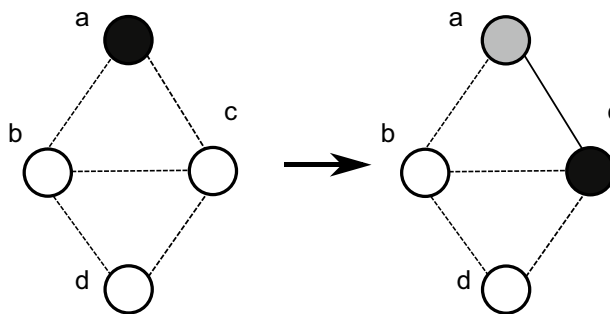


Figure 1: The SIR process on a graph. The contact graph $G = (V, E)$ is defined on a population $V = \{a, b, c, d\}$. The node colors white, black and gray represent the Susceptible, Infected and Recovered states, respectively. Initially, only node a is infected, and all other nodes are susceptible. A possible outcome at $t = 1$ is shown, in which node c becomes infected, while node a recovers. Node a tries to independently infect both its neighbors b and c , but only node c gets infected—this is indicated by the solid edge (a, c) . The probability of getting this outcome is $(1 - p(a, b))p(a, c)$.

activities and mobility, from which contacts can be inferred, based on co-location. Multiple approaches have been developed for this, including random mobility based on statistical models, and very detailed models based on activities in urban regions, which have been estimated through surveys, transportation data, and other sources, e.g.,^{20, 21, 8, 22, 23}.

- Develop models of within-host disease progression: such models can be represented as finite state probabilistic timed transition models, which are designed in close coordination with biologists, epidemiologists, and parameterized using detailed incidence data (see⁹ for discussion and additional pointers).
- Develop high-performance computer (HPC) simulations to study epidemic dynamics in such models, e.g.,^{24, 25, 26, 27}. Typical public health analyses involve large experimental designs, and the models are stochastic; this necessitates the use of such HPC simulations on large computing clusters.
- Incorporate interventions and behavioral changes: interventions include closure of schools and workplaces^{22, 28} and vaccinations²¹; whereas, behavioral changes include individual level social distancing, changes in mobility, and use of protective measures.

Such a network model captures the interplay between the three components of computational epidemiology: (i) individual behaviors of agents, (ii) unstructured, heterogeneous multi-scale networks, and (iii) the dynamical processes on these networks. It is based on the hypothesis that a better understanding of the

characteristics of the underlying network and individual behavioral adaptation can give better insights into contagion dynamics and response strategies. Although computationally expensive and data intensive, network-based epidemiology alters the types of questions that can be posed, providing qualitatively different insights into disease dynamics and public health policies. It also allows policy makers to formulate and investigate potentially novel and context-specific interventions.

2.4 Models for Epidemic Forecasting

Like projection approaches, models for epidemic forecasting can be broadly classified into two broad groups: (i) statistical and machine learning-based data-driven models, (ii) causal or mechanistic models—see^{29, 30, 2, 31, 32, 6, 33} and the references therein for the current state of the art in this rapidly evolving field.

Statistical methods employ statistical and time series-based methodologies to learn patterns in historical epidemic data and leverage those patterns for forecasting. Of course, the simplest yet useful class is called *method of analogs*. One simply compares the current epidemic with one of the earlier outbreaks and then uses the best match to forecast the current epidemic. Popular statistical methods for forecasting influenza-like illnesses (that includes COVID-19) include, e.g., generalized linear models (GLM), autoregressive integrated moving average (ARIMA), and generalized autoregressive moving average (GARMA)^{34, 31, 35}. Statistical methods are fast, but they crucially depend on the availability of training data. Furthermore, since they are purely

data driven, they do not capture the underlying causal mechanisms. As a result, epidemic dynamics affected by behavioral adaptations are usually hard to capture. Artificial neural networks (ANN) have gained increased prominence in epidemic forecasting due to their self-learning ability without prior knowledge (see^{1,11,36} and the references therein). Such models have used a wide variety of data as surrogates for producing forecasts. This includes: (i) social media data, (ii) weather data, (iii) incidence curves and (iv) demographic data.

Causal models can be used for epidemic forecasting in a natural manner^{30, 3, 37, 32, 38, 39}. These models calibrate the internal model parameters using the disease incidence data seen until a given day and then execute the model forward in time to produce the future time series. Compartmental as well as agent-based models can be used to produce such forecasts. The choice of the models depends on the specific question at hand and the computational and data resource constraints. One of the key ideas in forecasting is to develop ensemble models—models that combine forecasts from multiple models^{40, 6, 38, 39}. The idea which originated in the domain of weather forecasting has found methodological advances in the machine learning literature. Ensemble models typically show better performance than the individual models.

3 Models from the Imperial College Modeling Group (UK Model)

Background. The modeling group led by Neil Ferguson was to our knowledge the first model to study the impact of COVID-19 across two large countries: US and UK, see²². The basic model was first developed in 2005—it was used to inform policy pertaining to H5N1 pandemic and was one of the three models used to inform the federal pandemic influenza plan and led to the now well-accepted targeted layered containment (TLC) strategy. It was adapted to COVID-19 as discussed below. The model was widely discussed and covered in the scientific as well as popular press⁴¹. We will refer to this as the IC model.

Model structure. The basic model structure consists of developing a set of households based on census information for a given country. The structure of the model is largely borrowed from their earlier work, see^{42, 28}. Landscan data were used to spatially distribute the population. Individual members of the household interact with other members of the household. The data to produce these households are obtained using

Census information for these countries. Census data are used to assign age and household sizes. Details on the resolution of census data and the dates were not clear. Schools, workplaces and random meeting points are then added. The school data for US were obtained from the National Centre of Educational Statistics, while for UK schools were assigned randomly based on population density. Data on average class sizes and staff-student ratios were used to generate a synthetic population of schools distributed proportional to local population density. Data on the distribution of workplace size were used to generate workplaces with commuting distance data used to locate workplaces appropriately across the population. Individuals are assigned to each of these locations at the start of the simulation. The gravity-style kernel is used to decide how far a person can go in terms of attending work, school or community interaction place. The number of contacts between individuals at school, work and community meeting points are calibrated to produce a given attack rate.

Each individual has an associated disease transmission model. The disease transmission model parameters are based on the data collected when the pandemic was evolving in Wuhan; see page 4 of²².

Finally, the model also has rich set of interventions. These include: (i) case isolation, (ii) voluntary home quarantine, (iii) Social distancing of those over 70 years, (iv) social distancing of the entire population, (v) closure of schools and universities; see page 6²². The code was recently released and is being analyzed. This is important as the interpretation of these interventions can have substantial impact on the outcome.

Model predictions. The Imperial college (IC Model) model was one of the first models to evaluate the COVID-19 pandemic using detailed agent-based model. The predictions made by the model were quite dire. The results show that to be able to reduce R to close to 1 or below, a combination of case isolation, social distancing of the entire population and either household quarantine or school and university closure is required. The model had tremendous impact—UK and US both decide to start considering complete lock downs—a policy that was practically impossible to even talk about earlier in the Western world. The paper came out around the same time that Wuhan epidemic was raging and the epidemic in Italy had taken a turn for the worse. This made the model results even more critical.

Strengths and limitations. IC model was one of the first models by a reputed group to report the

potential impact of COVID-19 with and without interventions. The model was far more detailed than other models that were published until then. The authors also took great care parameterizing the model with the best disease transmission data that was available until then. The model also considered a very rich set of interventions and was one of the first to analyze pulsing intervention. On the flip side, the representation of the underlying social contact network was relatively simple. Second, often the details of how interventions were represented were not clear. Since the publication of their article, the modelers have made their code open and the research community has witnessed an intense debate on the pros and cons of various modeling assumptions and the resulting software system, see⁴³. We believe that despite certain valid criticisms, overall, the results represented a significant advance in terms of the when the results were put out and the level of details incorporated in the models.

4 Spatial Metapopulation Models: Northeastern and UVA Models (US Models)

Background. This approach is an alternative to detailed agent-based models, and has been used in modeling the spread of multiple diseases, including Influenza^{15, 18}, Ebola¹⁷ and Zika¹⁹. It has been adapted for studying the importation risk of COVID-19 across the world¹⁶. Structured metapopulation models construct a simple abstraction of the mixing patterns in the population, in which the entire region under study is decomposed into fully connected geographical regions, representing subpopulations, which are connected through airline and commuter flow networks. Thus, they lack the rich detail of agent-based models, but have fewer parameters, and are, therefore, easy to set up and scale to large regions.

Model structure. Here, we summarize GLEaM¹⁵ (Northeastern model) and PatchSim¹⁸ (UVA model). GLEaM uses two classes of datasets—population estimates and mobility. Population data are used from the “Gridded Population of the World”⁴⁴, which gives an estimated population value at a 15×15 minutes of arc (referred to as a “cell”) over the entire planet. Two different kinds of mobility processes are considered—airline travel and commuter flow. The former captures long-distance travel; whereas, the latter captures localized mobility. Airline data are obtained from the International Air Transport Association (IATA)⁴⁵, and the

Official Airline Guide (OAG)⁴⁶. There are about 3300 airports world wide; these are aggregated at the level of urban regions served by multiple airport (e.g., as in London). A Voronoi tessellation is constructed with the resulting airport locations as centers, and the population cells are assigned to these cells, with a 200 mile cutoff from the center. The commuter flows connect cells at a much smaller spatial scale. We represent this mobility pattern as a directed graph on the cells, and refer to it as the mobility network.

In the basic SEIR model, the subpopulation in each cell j is partitioned into compartments S_j, E_j, I_j and R_j , corresponding to the disease states. For each cell j , we define the force of infection λ_j as the rate at which a susceptible individual in the subpopulation in cell j becomes infected—this is determined by the interactions the person has with infectious individuals in cell j or any cell j' connected in the mobility network. An individual in the susceptible compartment S_j becomes infected with probability $\lambda_j \Delta t$ and enters the compartment E_j , in a time interval Δt . From this compartment, the individual moves to the I_j and then the R_j compartments, with appropriate probabilities, corresponding to the disease model parameters.

The PatchSim¹⁸ model has a similar structure, except that it uses administrative boundaries (e.g., counties), instead of a Voronoi tessellation, which are connected using a mobility network. The mobility network is derived by combining commuter and airline networks, to model time spent per day by individuals of region (patch) i in region (patch) j . Since it explicitly captures the level of connectivity through a commuter-like mixing, it is capable of incorporating week-to-week and month-to-month variations in mobility and connectivity. In addition to its capability to run in deterministic or stochastic mode, the open source implementation⁴⁷ allows fine-grained control of disease parameters across space and time. Although the model has a more generic force of infection mode of operation (where patches can be more general than spatial regions), we will mainly summarize the results from the mobility model, which was used for COVID-19 response.

What did the models suggest? GLEaM model is being used in a number of COVID-19-related studies and analysis. In⁴⁸, the Northeastern University team used the model to understand the spread of COVID-19 within China and relative risk of importation of the disease internationally. Their analysis suggested that the spread of COVID-19 out of Wuhan into other parts of mainland China was not contained well due

to the delays induced by detection and official reporting. It is hard to interpret the results. The paper suggested that international importation could be contained substantially by strong travel ban. While it might have delayed the onset of cases, the subsequent spread across the world suggest that we were not able to arrest the spread effectively. The model is also used to provide weekly projections (see <https://covid19.gleamproject.org/>); this site does not appear to be maintained for the most current forecasts (likely because the team is participating in the CDC forecasting group).

The PatchSim model is being used to support federal agencies as well as the state of Virginia. Due to our past experience, we have refrained from providing longer term forecasts, instead of focusing on short-term projections. The model is used within a *Forecasting via Projection Selection* approach, where a set of counterfactual scenarios are generated based on on-the-ground response efforts and surveillance data, and the best fits are selected based on historical performance. While allowing for future scenarios to be described, they also help to provide a reasonable narrative of past trajectories, and retrospective comparisons are used for metrics such as ‘cases averted by doing X’. These projections are revised weekly based on stakeholder feedback and surveillance update. Further discussion of how the model is used by the Virginia Department of Health each week can be found at <https://www.vdh.virginia.gov/coronavirus/covid-19-data-insights/#model>.

Strength and limitations. Structured meta-population models provide a good tradeoff between the realism/compute of detailed agent-based models and simplicity/speed of mass action compartmental models and need far fewer inputs for modeling, and scalability. This is especially true in the early days of the outbreak, when the disease dynamics are driven to a large extent by mobility, which can be captured more easily within such models, and there is significant uncertainty in the disease model parameters. However, once the outbreak has spread, it is harder to model detailed interventions (e.g., social distancing), which are much more localized. Further, these are hard to model using a single parameter. Both GLEaM and PatchSim models also faced their share of challenges in projecting case counts due to rapidly evolving pandemic, inadequate testing, a lack of understanding of the number of asymptomatic cases and assessing the compliance levels of the population at large.

5 Models by KTH, Umea and Uppsala Researchers (Swedish Models)

Sweden was an outlier amongst countries in that it decided to implement public health interventions without a lockdown. Schools and universities were not closed, and restaurants and bars remained open. Swedish citizens implemented “work from home” policies where possible. Moderate social distancing based on individual responsibility and without police enforcement was employed but emphasis was attempted to be placed on shielding the 65+ age group.

5.1 Simple Model

Background. Statistician Tom Britton developed a very simple model with a focus on predicting the number of infected over time in Stockholm.

Model structure. Britton⁴⁹ used a very simple SIR general epidemic model. It is used to make a coarse grain prediction of the behavior of the outbreak based on knowing the basic reproduction number R_0 and the doubling time d in the initial phase of the epidemic. Calibration to calendar time was done using the observed number of case fatalities, together with estimates of the time between infection to death, and the infection fatality risk. Predictions were made assuming no change of behavior, as well as for the situation where preventive measures are put in place at one specific time-point.

Model predictions. One of the controversial predictions from this model was that the number of infections in the Stockholm area would quickly rise towards attaining *herd immunity* within a short period. However, mass testing carried out in Stockholm during June indicated a far smaller percentage of infections.

Strength and limitations. Britton’s model was intended as a quick and simple method to estimate and predict an on-going epidemic outbreak both with and without preventive measures put in place. It was intended as a complement to more realistic and detailed modeling. The estimation-prediction methodology is much simpler and straight-forward to implement for this simple model. It is more transparent to see how the few model assumptions affect the results, and it is easy to vary the few parameters to see their effect on predictions so that one could see which parameter uncertainties have biggest impact on predictions, and which parameter uncertainties are less influential.

5.2 Compartmentalized Models I: FHM Model

Background. The Public Health Authority (FHM) of Sweden produced a model to study the spread of COVID-19 in four regions in Sweden: Dalarna, Skåne, Stockholm, and Västra Götaland.⁵⁰

Model structure. It is a standard compartmentalized SEIR model and within each compartment, it is homogeneous; so, individuals are assumed to have the same characteristics and act in the same way. Data used in the fitting of the model include point prevalences found by PCR-testing in Stockholm at two different time points.

Model predictions. The model estimated the number of infected individuals at different time points and the date with the largest number of infectious individuals. It predicted that by July 1, 8.5% (5.9–12.9%) of the population in Dalarna will have been infected, 4% (2.4–9.9%) of the population in Skåne will have been infected, 19% (17.7–20.2%) of the population in Stockholm will have been infected, and 9% (6.3–12.2%) of the population in Västra Götaland will have been infected. It was hard to test these predictions because of the great uncertainty in immune response to SARS-CoV-2—prevalence of antibodies was surprisingly low but recent studies show that mild cases never seem to develop antibodies against SARS-CoV-2, but only T-cell-mediated immunity⁵¹.

The model also investigated the effect of increased contacts during the summer that stabilizes in autumn. It found that if the contacts in Stockholm and Dalarna increase by less than 60% in comparison to the contact rate in the beginning of June, the second wave will not exceed the observed first wave.

Strength and limitations. The simplicity of the model is a strength in ease of calibration and understanding but it is also a major limitation in view of the well-known characteristics of COVID-19: since it is primarily transmitted through droplet infection, the social contact structure in the population is of primary importance for the dynamics of infection. The compartmental model used in this analysis does not account for variation in contacts, where few individuals may have many contacts, while the majority have fewer. The model is also not age stratified, but COVID-19 strikingly affects different age groups differently; e.g., young people seem to get milder infections. In this model, each infected individual has the same infectivity and the same risk of becoming a reported case, regardless of age. Different age groups normally

have varied degrees of contacts and have changed their behavior differently during the COVID-19 pandemic. This is not captured in the model.

5.3 Compartmentalized Models II

Background. A group around statistician Joacim Rocklöv developed a model to estimate the impact of COVID-19 on the Swedish population at the municipality level, considering demography and human mobility under various scenarios of mitigation and suppression. They attempted to estimate the time course of infections, health care needs, and the mortality in relation to the Swedish ICU capacity, as well as the costs of care, and compared alternative policies and counterfactual scenarios.

Model structure.⁵² used a SEIR compartmentalized model with age structured compartments (0–59, 60–79, 80+) susceptibles, infected, in-patient care, ICU and recovered populations based on Swedish population data at the municipal level. It also incorporated inter-municipality travel using a radiation model. Parameters were calibrated based on a combination of values available from international literature and fitting to available outbreak data. The effect of a number of different intervention strategies was considered ranging from no intervention to modest social distancing and finally to imposed isolation of various groups.

Model predictions. The model predicted an estimated death toll of around 40,000 for the strategies based only on social distancing and between 5000 and 8000 for policies imposing stricter isolation. It predicted ICU cases of up to 10,000 without much intervention and up to 6000 with modest social distancing, way above the available capacity of about 500 ICU beds.

Strength and limitations. The model showed a good fit against the reported COVID-19-related deaths in Sweden up to 20th of April, 2020. However, the predictions of the total deaths and ICU demand turned out to be way off the mark.

5.4 Agent-Based Microsimulations

Background. Finally,^{53, 54} used an individual-based model parameterized on Swedish demographics to assess the anticipated spread of COVID-19.

Model structure.⁵³ employed the individual agent-based model based on work by Ferguson et al.²². Individuals are randomly assigned an age based on Swedish demographic data and they are also assigned a household. Household

size is normally distributed around the average household size in Sweden in 2018, 2.2 people per household. Households were placed on a lattice using high-resolution population data from Landscan and census data from the Statistics Sweden and each household is additionally allocated to a city based on the closest city center by distance and to a county based on city designation. Each individual is placed in a school or workplace at a rate similar to the current participation in Sweden.

Transmission between individuals occurs through contact at each individual's workplace or school, within their household, and in their communities. Infectiousness is, thus, a property dependent on contacts from household members, school/workplace members and community members with a probability based on household distances. Transmissibility was calibrated against data for the period 21 March–6 April to reproduce either the doubling time reported using pan-European data or the growth in reported Swedish deaths for that period. Various types of interventions were studied including the policy implemented in Sweden by the public health authorities as well as more aggressive interventions approaching full lockdown.

Model predictions. Their prediction was that “under conservative epidemiological parameter estimates, the current Swedish public-health strategy will result in a peak intensive-care load in May that exceeds pre-pandemic capacity by over 40-fold, with a median mortality of 96,000 (95% CI 52,000 to 183,000)”.

Strength and limitations. This model was based on adapting the well-known Imperial model discussed in Sect. 3 to Sweden and considered a wide range of intervention strategies. Unfortunately the predictions of the model were woefully off the mark on both counts: the deaths by June 18 are under 5000 and at the peak the ICU infrastructure had at least 20% unutilized capacity.

6 Forecasting Models

Forecasting is of particular interest to policy makers as they attempt to provide actual counts. Since the surveillance systems have relatively stabilized in recent weeks, the development of forecasting models has gained traction and several models are available in the literature. In the US, the Centers for Disease Control and Prevention (CDC) has provided a platform for modelers to share their forecasts which are analyzed and combined in a suitable manner to produce ensemble

multi-week forecasts for cumulative/incident deaths, hospitalizations and more recently cases at the national, state, and county level. Probabilistic forecasts are provided by 36 teams as of July 28, 2020 (there were 21 models as of June 24, 2020) and the CDC with the help of⁵⁵ has developed uniform ensemble model for multi-step forecasts⁵⁶.

6.1 COVID-19 Forecast Hub Ensemble Model

It has been observed previously for other infectious diseases that an ensemble of forecasts from multiple models perform better than any individual contributing model³⁹. In the context of COVID-19 case count modeling and forecasting, a multitude of models have been developed based on different assumptions that capture specific aspects of the disease dynamics (reproduction number evolution, contact network construction, etc.). The models employed in the CDC Forecast Hub can be broadly classified into three categories, data-driven, hybrid models, and mechanistic models with some of the models being open source.

Data-driven models. They do not model the disease dynamics but attempt to find patterns in the available data and combine them appropriately to make short-term forecasts. In such data-driven models, it is hard to incorporate interventions directly; hence, the machine is presented with a variety of exogenous data sources such as mobility data, hospital records, etc. with the hope that its effects are captured implicitly. Early iterations of Institute of Health Metrics and Evaluation (IHME) model³⁴ for death forecasting at state level employed a statistical model that fits a time-varying Gaussian error function to the cumulative death counts and is parameterized to control for maximum death rate, maximum death rate epoch, and growth parameter (with many parameters learnt using data from outbreak in China). The IHME models are undergoing revisions (moving towards the hybrid models) and updated implementable versions are available at⁵⁷. The University of Texas at Austin COVID-19 Modeling Consortium model⁵⁸ uses a very similar statistical model as³⁴ but employs real-time mobility data as additional predictors and also differ in the fitting process. The Carnegie Mellon Delphi Group employs the well known auto-regressive (AR) model that employs lagged version of the case counts and deaths as predictors and determines a sparse set that best describes the observations from it by using

LASSO regression^{59,60} is a deep learning model which has been developed along the lines of⁴ and attempts to learn the dependence between death rate and other available syndromic, demographic, mobility and clinical data.

Hybrid models. These methods typically employ statistical techniques to model disease parameters which are then used in epidemiological models to forecast cases. Most statistical models^{34,58} are evolving to become hybrid models. A model that gained significant interest is the Youyang Gu (YYG) model and uses a machine learning layer over an SEIR model to learn the set of parameters (mortality rate, initial R_0 , post-lockdown R) specific to a region that best fits the region's observed data. The authors (YYG) share the optimal parameters, the SEIR model and the evaluation scripts with general public for experimentation⁶¹. Los Alamos National Lab (LANL) model³⁵ uses a statistical model to determine how the number of COVID-19 infections changes over time. The second process maps the number of infections to the reported data. The number of deaths is a fraction of the number of new cases obtained and is computed using the observed mortality data.

Mechanistic models. GLEaM and JHU models are county-level stochastic SEIR model dynamics. The JHU model incorporates the effectiveness of state-wide intervention policies on social distancing through the R_0 parameter. More recently, model outputs from UVA's PatchSim model were included as part of a multi-model ensemble (including autoregressive and LSTM components) to forecast weekly confirmed cases.

7 Comparative Analysis Across Modeling Types

We end the discussion of the models above by qualitatively comparing model types. As discussed in the preliminaries, at one end of the spectrum are models that are largely data driven: these models range from simple statistical models (various forms of regression models) to the more complicated deep learning models. The difference in such model lies in the amount of training data needed, the computational resources needed and how complicated the mathematical function one is trying to fit to the observed data. These models are strictly data driven and, hence, unable to capture the constant behavioral adaptation at an individual and collective level. On the other end of the spectrum SEIR, meta-population and agent-based network models are based on the underlying procedural representation of the

dynamics—in theory, they are able to represent behavioral adaptation endogenously. But both class of models face immense challenges due to the availability of data as discussed below.

- (1) Agent-based and SEIR models were used in all the three countries in the early part of the outbreak and continue to be used for counter-factual analysis. The primary reason is the lack of surveillance and disease specific data and hence, purely data-driven models were not easy to use. SEIR models lacked heterogeneity but were simple to program and analyze. Agent-based models were more computationally intensive, required a fair bit of data to instantiate the model but captured the heterogeneity of the underlying countries. By now it has become clear that use of such models for long term forecasting is challenging and likely to lead to mis-leading results. The fundamental reason is adaptive human behavior and lack of data about it.
- (2) Forecasting, on the other hand, has seen use of data-driven methods as well as causal methods. Short-term forecasts have been generally reasonable. Given the intense interest in the pandemic, a lot of data are also becoming available for researchers to use. This helps in validating some of the models further. Even so, real-time data on behavioral adaptation and compliance remain very hard to get and is one of the central modeling challenges.

8 Models and Policy Making

Were some of the models wrong? In a recent opinion piece,⁴ Professor Vikram Patel of the Harvard School of Public Health makes a stinging criticism of modeling:

Crowning these scientific disciplines is the field of modeling, for it was its estimates of mountains of dead bodies which fuelled the panic and led to the unprecedented restrictions on public life around the world. None of these early models, however, explicitly acknowledged the huge assumptions that were made,

A similar article in NY Times recounted the *mistakes* in COVID-19 response in Europe⁵; also see⁶².

Our point of view. It is indeed important to ensure that assumptions underlying mathematical models be made transparent and explicit. But we respectfully disagree with Professor Patel's statement: most of the *good* models tried to be very explicit about their assumptions. The mountains of deaths that are being referred to are explicitly calculated when **no** interventions are put in place and are often used as a worst case scenario. Now, one might argue that the authors be explicit and state that this worst case scenario will never occur in practice. Forecasting dynamics in social systems is inherently challenging: individual behavior, predictions and epidemic dynamics *co-evolve*; this coevolution immediately implies that a dire prediction can lead to extreme change in individual and collective behavior leading to reduction in the incidence numbers. Would one say forecasts were wrong in such a case or they were influential in ensuring the worst case never happens? None of this implies that one should not explicitly state the assumption underlying their model. Of course our experience is that policy makers, news reporters and common public are looking exactly for such a forecast—we have been constantly asked “when will peak occur” or “how many people are likely to die”. A few possible ways to overcome this tension between the unsatiable appetite for forecasts and the inherent challenges that lie in doing this accurately, include:

- We believe that, in general, it might not be prudent to provide long term forecasts for such systems.
- State the assumptions underlying the models as clearly as possible. Modelers need to be much more disciplined about this. They also need to ensure that the models are transparent and can be reviewed broadly (and expeditiously).
- Accept that the forecasts are provisional and that they will be revised as new data comes in, society adapts, the virus adapts and we understand the biological impact of the pandemic.
- Improve surveillance systems that would produce data that the models can use more effectively. Even with data, it is very hard to estimate the prevalence of COVID-19 in society.

Communicating scientific findings and risks is an important topical area in this context, see^{41, 63, 64, 65}.

Use of models for evidence-based policy making. In a new book,⁶⁶ *Radical Uncertainty*, economists John Kay and Mervyn King (formerly Governor of the Bank of England) urge caution when using complex models. They argue that models should be valued for the insights they provide but not relied upon to provide accurate forecasts. The so-called “evidence-based policy” comes in for criticism where it relies on models but also supplies a false sense of certainty where none exists, or seeks out the evidence that is desired *ex ante*—or “cover”—to justify a policy decision. “*Evidence-based policy has become policy-based evidence*”.

Our point of view. The authors make a good point here. But again, everyone, from public to citizens and reporters clamor for a forecast. We argue that this can be addressed in two ways: (i) viewing the problem from the lens of control theory so that we forecast only to control the deviation from the path we want to follow and (ii) not insisting on exact numbers but general trends. As Kay and King opine, the value of models, especially in the face of *radical uncertainty*, is more in exploring alternative scenarios resulting from different policies:

a model is useful only if the person using it understands that it does not represent the “the world as it really is” but is a tool for exploring ways in which a decision might or might not go wrong.

Supporting science beyond the pandemic.

In his new book *The Rules of Contagion*, Adam Kucharski⁶⁷ draws on lessons from the past. In 2015 and 2016, during the Zika outbreak, researchers planned large-scale clinical studies and vaccine trials. But these were discontinued as soon as the infection ebbed.

This is a common frustration in outbreak research; by the time, the infections end, fundamental questions about the contagion can remain unanswered. That is why building long-term research capacity is essential.

Our point of view. The author makes an important point. We hope that today, after witnessing the devastating impacts of the pandemic on the economy and society, the correct lessons will be learnt: sustained investments need to be

⁴ *Indian Express*, July 30, 2020.

⁵ NY Times July 20, 2020: <https://www.nytimes.com/2020/07/20/world/europe/coronavirus-mistakes-france-uk-italy.html>.

made in the field to be ready for the impact of the next pandemic.

9 Concluding Remarks

The paper discusses a few important computational models developed by researchers in the US, UK and Sweden for COVID-19 pandemic planning and response. The models have been used by policy makers and public health officials in their respective countries to assess the evolution of the pandemic, design and analyze control measures and study various what-if scenarios. As noted, all models faced challenges due to availability of data, rapidly evolving pandemic and unprecedented control measures put in place. Despite these challenges, we believe that mathematical models can provide useful and timely information to the policy makers. On one hand the modelers need to be transparent in the description of their models, clearly state the limitations and carry out detailed sensitivity and uncertainty quantification. Having these models reviewed independently is certainly very helpful. On the other hand, policy makers should be aware of the fact that using mathematical models for pandemic planning, forecast response rely on a number of assumptions and lack data to overcome these assumptions.

Publisher's Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Acknowledgements

The authors would like to thank members of the Biocomplexity COVID-19 Response Team and Network Systems Science and Advanced Computing (NSSAC) Division for their thoughtful comments and suggestions related to epidemic modeling and response support. We thank members of the Biocomplexity Institute and Initiative, University of Virginia for useful discussion and suggestions. This work was partially supported by National Institutes of Health (NIH) Grant 1R01GM109718, NSF BIG DATA Grant IIS-1633028, NSF DIBBS Grant ACI-1443054, NSF Grant No.: OAC-1916805, NSF Expeditions in Computing Grant CCF-1918656, CCF-1917819, NSF RAPID CNS-2028004, NSF RAPID OAC-2027541, US Centers for Disease Control and Prevention 75D30119C05935, DTRA subcontract/ARA S-D00189-15-TO-01-UVA. Any opinions, findings, and conclusions

or recommendations expressed in this material are those of the author(s) and do not necessarily reflect the views of the funding agencies.

Received: 27 August 2020 Accepted: 14 September 2020
Published online: 30 October 2020

References

- Adhikari B, Xu X, Ramakrishnan N, Prakash BA (2019) Epiddeep: exploiting embeddings for epidemic forecasting. In: Proceedings of the 25th ACM SIGKDD international conference on knowledge discovery and data mining, KDD '19, pp 577–586, New York, NY, USA. Association for Computing Machinery
- Desai A, Kraemer M, Bhatia S, Cori A, Nouvellet P, Herring M, Cohn E, Carrion M, Brownstein J, Madoff L, Lassmann B (2019) Real-time epidemic forecasting: challenges and opportunities. *Health Secur* 17(08):268–275
- Funk S, Camacho A, Kucharski AJ, Eggo RM, Edmunds WJ (2018) Real-time forecasting of infectious disease dynamics with a stochastic semi-mechanistic model. *Epidemics* 22:56–61 (The RAPIDD Ebola Forecasting Challenge)
- Murray C (2020) Forecasting the impact of the first wave of the Covid-19 pandemic on hospital demand and deaths for the USA and European economic area countries. 04. <https://doi.org/10.1101/2020.04.21.20074732>
- Perone G (2020) An ARIMA model to forecast the spread and the final size of Covid-2019 epidemic in Italy (first version on SSRN 31 march). SSRN Electron J. <https://doi.org/10.2139/ssrn.3564865>
- Reich NG, McGowan CJ, Yamana TK, Tushar A, Ray EL, Osthus D, Kandula S, Brooks LC, Crawford-Crudell W, Gibson GC, Moore E, Silva R, Biggerstaff M, Johansson MA, Rosenfeld R, Shaman JL (2019) Accuracy of real-time multi-model ensemble forecasts for seasonal influenza in the US. *PLoS Comput Biol* 15:e1007486
- Allen Linda JS, Brauer F, Van den Driessche P, Wu J (2008) *Mathematical epidemiology*, vol 1945. Springer, Berlin
- Eubank S, Kumar VSA, Marathe MV, Srinivasan A, Wang N (2006) Structure of social contact networks and their impact on epidemics. *DIMACS Ser Discrete Math Theor Comput Sci* 70:181
- Marathe M, Vullikanti A (2013) Computational epidemiology. *Commun ACM* 56(7):88–96
- Newman MEJ (2003) The structure and function of complex networks. *SIAM Rev* 45(2):167–256
- Wang L, Chen J, Marathe M (2019) Deep learning based epidemic forecasting with synthetic information. In: AAAI, DEFSI
- Lipsitch M, Cohen T, Cooper B, Robins JM, Ma S, James L, Gopalakrishna G, Chew SK, Tan CC, Samore MH (2003) Transmission dynamics and

- control of severe acute respiratory syndrome. *Science* 300(5627):1966–1970
13. Mossong J, Hens N, Jit M, Beutels P, Auranen K, Mikolajczyk R, Massari M, Salmaso S, Tomba GS, Wallinga J, Heijne J (2008) Social contacts and mixing patterns relevant to the spread of infectious diseases. *PLoS Med* 5:e74
 14. Medlock J, Galvani AP (2009) Optimizing influenza vaccine distribution. *Science* 325(5948):1705–1708
 15. Balcan D, Colizza V, Gonçalves B, Hao H, Ramasco JJ, Vespignani A (2009) Multiscale mobility networks and the spatial spreading of infectious diseases. *Proc Natl Acad Sci* 106:21484–21489
 16. Chinazzi M, Davis JT, Ajelli M, Gioannini C, Litvinova M, Merler S, Piontti AP, Mu K, Rossi L, Sun K, Viboud C, Xiong X, Yu H, Halloran ME, Longini IM, Vespignani A (2020) The effect of travel restrictions on the spread of the 2019 novel coronavirus (Covid-19) outbreak. *Science* 368(6489):395–400
 17. Gomes MF, Piontti AP, Rossi L, Chao D, Longini I, Halloran ME, Vespignani A (2014) Assessing the international spreading risk associated with the 2014 West African Ebola outbreak. *PLoS Curr* 6
 18. Venkatramanan S, Chen J, Fadikar A, Gupta S, Higdon D, Lewis B, Marathe M, Mortveit H, Vullikanti A (2019) Optimizing spatial allocation of seasonal influenza vaccine under temporal constraints. *PLoS Comput Biol* 15(9):e1007111
 19. Zhang Q, Sun K, Chinazzi M, Piontti AP, Dean NE, Rojas DP, Merler S, Mistry D, Poletti P, Rossi L, Bray M, Halloran ME, Longini IM, Vespignani A (2017) Spread of Zika virus in the Americas. *PLoS One* 12(2):E4334–E4343
 20. Barrett CL, Beckman RJ, Khan M, Kumar VA, Marathe MV, Stretz PE, Dutta T, Lewis B (2009) Generation and analysis of large synthetic social contact networks. In: Winter simulation conference, pp 1003–1014. Winter Simulation Conference
 21. Eubank S, Guclu H, Kumar VSA, Marathe MV, Srinivasan A, Toroczkai Z, Wang N (2004) Modelling disease outbreaks in realistic urban social networks. *Nature* 429(6988):180–184
 22. Ferguson N, Laydon D, Nedjati Gilani G, Imai N, Ainslie K, Baguelin M, Bhatia S, Boonyasiri A, Cucunuba Perez ZU, Cuomo-Dannenburg G et al (2020) Report 9: impact of non-pharmaceutical interventions (NPIs) to reduce covid19 mortality and healthcare demand
 23. Longini IM, Nizam A, Shufu X, Ungchusak K, Hanshaworakul W, Cummings DA, Halloran EM (2005) Containing pandemic influenza at the source. *Science* 309(5737):1083–1087
 24. Barrett CL, Bisset KR, Eubank SG, Feng X, Marathe MV (2008) Episimdemics: an efficient algorithm for simulating the spread of infectious disease over large realistic social networks. In: Proceedings of the 2008 ACM/IEEE conference on supercomputing, p 37. IEEE Press
 25. Bisset KR, Chen J, Feng X, Kumar VA, Marathe MV (2009) Epifast: a fast algorithm for large scale realistic epidemic simulations on distributed memory systems. In: Proceedings of the 23rd international conference on supercomputing, pp 430–439. ACM
 26. Deodhar S, Bisset K, Chen J, Ma Y, Marathe M (2012) Enhancing user-productivity and capability through integration of distinct software in epidemiological systems. In: Proceedings of the 2nd ACM SIGHIT international health informatics symposium, pp 171–180. ACM
 27. Grefenstette JJ, Brown ST, Rosenfeld R, DePasse J, Stone NTB, Cooley PC, Wheaton WD, Fyshe A, Galloway DD, Sriram A (2013) Fred (a framework for reconstructing epidemic dynamics): an open-source software system for modeling infectious diseases and control strategies using census-based populations. *BMC Public Health* 13(1):1
 28. Halloran ME, Ferguson NM, Eubank S, Longini IM, Cummings DAT, Lewis B, Xu S, Fraser C, Vullikanti A, Germann TC (2008) Modeling targeted layered containment of an influenza pandemic in the united states. *Proc Natl Acad Sci* 105(12):4639–4644
 29. Brooks L (2020) Pancasting: forecasting epidemics from provisional data. PhD thesis, Centers for Disease Control and Prevention
 30. Chretien J-P, George D, Shaman J, Chitale RA, McKenzie FE (2014) Influenza forecasting in human populations: a scoping review. *PLoS ONE* 9(4):e94130
 31. Kandula S, Shaman J (2019) Near-term forecasts of influenza-like illness: an evaluation of autoregressive time series approaches. *Epidemics* 27:41–51
 32. Nsoesie EO, Brownstein JS, Ramakrishnan N, Marathe MV (2014) A systematic review of studies on forecasting the dynamics of influenza outbreaks. *Influenza Other Respir Viruses* 8(3):309–316
 33. Tabataba FS, Chakraborty P, Ramakrishnan N, Venkatramanan S, Chen J, Lewis B, Marathe M (2017) A framework for evaluating epidemic forecasts. *BMC Infect Dis* 17(1):345
 34. IHME COVID, Murray Christopher JL et al (2020) Forecasting covid-19 impact on hospital bed-days, ICU-days, ventilator-days and deaths by US state in the next 4 months. *MedRxiv*
 35. LANL. LANL Covid-19 cases and deaths forecasts. <https://covid-19.bsvgateway.org/>
 36. Wang L, Chen J, Marathe M (2020) TDEFISI: theory-guided deep learning-based epidemic forecasting with synthetic information. *ACM Trans Spatial Algorithms Syst (TSAS)* 6(3):1–39
 37. Fadikar A, Higdon D, Chen J, Lewis B, Venkatramanan S, Marathe M (2018) Calibrating a stochastic, agent-based model using quantile-based emulation. *SIAM/ASA J Uncertain Quant* 6(4):1685–1706
 38. Tabataba FS, Lewis B, Hosseinipour M, Tabataba FS, Venkatramanan S, Chen J, Higdon D, Marathe M (2017) Epidemic forecasting framework combining agent-based models and smart beam particle filtering. In: 2017 IEEE

- international conference on data mining (ICDM), pp 1099–1104. IEEE
39. Yamana TK, Kandula S, Shaman J (2017) Individual versus superensemble forecasts of seasonal influenza outbreaks in the United States. *PLoS Comput Biol* 13(11):e1005801
 40. Chakraborty P, Khadivi P, Lewis B, Mahendiran A, Chen J, Butler P, Nsoesie EO, Mekaru SR, Brownstein JS, Marathe MV et al (2014) Forecasting a moving target: ensemble models for ILI case count predictions. In: Proceedings of the 2014 SIAM international conference on data mining, pp 262–270. SIAM
 41. Adam D (2020) Modelling the pandemic the simulations driving the world's response to Covid-19. *Nature* 580(7803):316–318
 42. Ferguson NM, Cummings DAT, Fraser C, Cajka JC, Cooley PC, Burke DS (2006) Strategies for mitigating an influenza pandemic. *Nature* 442(7101):448–452
 43. Dalmeeth Singh Chawla (2020) Critiqued coronavirus simulation gets thumbs up from code-checking efforts. *Nature* 582(7812):323–324
 44. Lloyd CT, Sorichetta A, Tatem AJ (2017) High resolution global gridded data for use in population studies. *Sci Data* 4(1):1–17
 45. IATA Air traffic statistics. <https://www.iata.org/en/services/statistics/air-transport-stats/>. Last accessed Apr 2020
 46. OAG Official airline guide. <https://www.oag.com/>. Last accessed Apr 2020
 47. Venkatramanan S Nssac/patchsim: code for simulating the metapopulation SEIR model. <https://github.com/NSSAC/PatchSim>. Accessed on 08/14/2020
 48. Kraemer MUG, Yang C-H, Gutierrez B, Wu C-H, Klein B, Pigott DM, Plessis LD, Faria NR, Li R, Hanage WP (2020) The effect of human mobility and control measures on the Covid-19 epidemic in china. *Science* 368(6490):493–497
 49. Britton T (2020) Basic prediction methodology for Covid-19: estimation and sensitivity considerations. medRxiv
 50. Public Health Agency of Sweden (2020) Estimates of the number of infected individuals during the Covid-19 outbreak in the Dalarna Region, Skåne Region, Stockholm Region, and Västra Götaland Region, Sweden. <https://www.folkhalsomyndigheten.se/publicerat-material/publikationsarkiv/e/estimates-of-the-number-of-infected-individuals-during-the-covid-19-outbreak/>
 51. Karolinska COVID-19 Study Group (2020) Robust T cell immunity in convalescent individuals with asymptomatic or mild Covid-19. *Cell* 183(1):158–168
 52. Rocklov J (2020) Covid-19 healthcare demand and mortality in Sweden in response to non-pharmaceutical (NPIs) mitigation and suppression scenarios. ARBO-PREVENT: climate change, human mobility and emerging arboviral outbreaks: new models for risk characterization, resilience and prevention. <https://doi.org/10.1101/2020.03.20.20039594>
 53. Gardner JM, Willem L, van der Wijngaart W, Kamerlin SC, Brusselaers N, Kasson P (2020) Intervention strategies against Covid-19 and their estimated impact on Swedish healthcare capacity. *Medrxiv*. <https://doi.org/10.1101/2020.04.11.20062133>
 54. Kamerlin Shina CL, Kasson Peter M (2020) Managing Covid-19 spread with voluntary public-health measures: Sweden as a case study for pandemic control. *Clin Infect Dis*. <https://doi.org/10.1093/cid/ciaa864>
 55. Reich N Reich lab. <https://reichlab.io/>
 56. CDC. Covid-19 forecasthub. <https://viz.covid19forecasthub.org/>
 57. IHME. <https://github.com/ihmeuw/covid-model-seiir-pipeline>
 58. Woody S, Tec MG, Dahan M, Gaither K, Lachmann M, Fox S, Meyers LA, Scott JG (2020) Projections for first-wave Covid-19 deaths across the us using social-distancing measures derived from mobile phones. medRxiv
 59. Carnegie Mellon Delphi Group. <https://delphi.cmu.edu>
 60. DeepCOVID. <https://deepcovid.github.io/>
 61. Gu Youyang (YYG). https://github.com/youyanggu/covid_19_projections
 62. Avery C, Bossert W, Clark A, Ellison G, Ellison SF (2020) Policy implications of models of the spread of coronavirus: perspectives and opportunities for economists. Technical report, National Bureau of Economic Research
 63. Fischhoff B (2019) Evaluating science communication. *Proc Natl Acad Sci* 116(16):7670–7675
 64. Metcalf CJE, Morris DH, Park SW (2020) Mathematical models to guide pandemic response. *Science* 369(6502):368–369
 65. Vaezi A, Javanmard SH (2020) Infodemic and risk communication in the era of CoV-19. *Adv Biomed Res* 9:10. https://doi.org/10.4103/abr.abr_47_20
 66. Kay J, King M (2020) Radical uncertainty: decision-making beyond the numbers. W. W. Norton & Company, New York
 67. Kucharski A (2020) The rules of contagion: why things spread-and why they stop. Basic Books, New York



Aniruddha Adiga is a Postdoctoral Research Associate at the NSSAC Division of the Biocomplexity Institute and Initiative. He completed his PhD from the Department of Electrical Engineering, Indian Institute of Science (IISc), Bangalore, India and

has held the position of Postdoctoral fellow at IISc and North Carolina State University, Raleigh, USA. His research areas include signal processing, machine learning, data mining, forecasting, big data analysis etc. At NSSAC, his primary focus has been the analysis and development of forecasting systems for epidemiological signals such as influenza-like illness and COVID-19 using auxiliary data sources.



Devdatt Dubhashi is a Professor in the Data Science and AI Division in the Department of computer Science and Engineering, Chalmers University Sweden. His research interests are in machine learning, algorithms and cognitive science. He received

his Ph.D. in Computer Science at Cornell University USA and has held positions at the Max Planck Institute for Computer Science, Saarbrücken Germany, BRICS (Basic Research in computer Science) a center of the Danish National Science Foundation at the University of Aarhus Denmark and at the Indian Institute of Technology, Delhi India.



Bryan Lewis is a research associate professor in the Network Systems Science and Advanced Computing division. His research has focused on understanding the transmission dynamics of infectious diseases within specific populations through both analysis

and simulation. Lewis is a computational epidemiologist with more than 15 years of experience in crafting, analyzing, and interpreting the results of models in the context of real public health problems. As a computational epidemiologist, for more than a decade, Lewis has been heavily involved in a series of projects forecasting the spread of infectious disease as well as evaluating the response to them in support of the federal government. These projects have tackled diseases from Ebola to pandemic influenza and melioidosis to cholera.



Madhav Marathe is a Distinguished Professor in Biocomplexity, the division director of the Networks, Simulation Science and Advanced Computing (NSSAC) Division at the Biocomplexity Institute and Initiative, and a Professor in the Department of Computer

Science at the University of Virginia (UVA). His research interests are in network science, computational epidemiology, AI, foundations of computing, socially coupled system science and high-performance computing. Before joining UVA, he held positions at Virginia Tech and the Los Alamos National Laboratory. He is a Fellow of the IEEE, ACM, SIAM and AAAS.



Srinivasan Venkatramanan is a Research Scientist at the Biocomplexity Institute & Initiative, University of Virginia and his research focuses on developing, analyzing and optimizing computational models in the field of network epidemiology. He

received his PhD from the Department of Electrical and Communication Engineering, Indian Institute of Science (IISc), and did his postdoctoral research at Virginia Tech. His areas of interest include network science, stochastic modeling and big data analytics. He has used in-silico models of society to study the spread of infectious diseases and invasive species. Recent research includes modeling and forecasting emerging infectious disease outbreaks (e.g., Ebola, COVID-19), impact of human mobility on disease spread and resource allocation problems in the context of epidemic control (e.g., seasonal influenza vaccines).



Anil Vullikanti is a Professor in the Biocomplexity Institute and the Department of Computer Science at the University of Virginia. He got his PhD at the Indian Institute of Science, and was a postdoctoral associate at the Max Planck Institute for Computer

Science, Saarbrücken, Germany, and the Los Alamos National Lab. His research interests are in the broad areas of network science, dynamical systems, combinatorial optimization, and distributed computing, and their applications to computational epidemiology and social networks.