

# Agent-based modeling for SARS-CoV-2 epidemic prediction and intervention assessment. A methodological appraisal

Mariusz Maziarz<sup>1</sup> and Martin Zach<sup>2</sup>

<sup>1</sup>Jagiellonian University

<sup>2</sup>Charles University

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## Abstract

Our purpose is to assess epidemiological agent-based models— or ABMs - of the SARS-CoV-2 pandemic methodologically. The rapid spread of the outbreak requires fast-paced decision-making regarding mitigation measures. However, the evidence for the efficacy of non-pharmaceutical interventions such as imposed social distancing and school or workplace closures is scarce: few observational studies use quasi-experimental research designs, and conducting randomized controlled trials seems infeasible. Additionally, evidence from the previous coronavirus outbreaks of SARS and MERS lacks external validity, given the significant differences in contagiousness of those pathogens relative to SARS-CoV-2. To address the pressing policy questions that have emerged as a result of COVID-19, epidemiologists have produced numerous models that range from simple compartmental models to highly advanced agent-based models. These models have been criticized for involving simplifications and lacking empirical support for their assumptions. In order to address these voices and methodologically appraise epidemiological ABMs, we consider AceMod (the model of the COVID-19 epidemic in Australia) as an example of the modeling practice. Our case study shows that, although epidemiological ABMs involve simplifications of various sorts, the key characteristics of social interactions and the spread of SARS-CoV-2 are represented sufficiently accurately. This is the case because these modelers treat empirical results as inputs for constructing modeling assumptions and rules that the agents follow; and they use calibration to assert the adequacy to benchmark variables. Given this, we claim that the best epidemiological ABMs are models of actual mechanisms and deliver both mechanistic and difference-making evidence. Furthermore, the efficacy claims are not only internally valid but also adequately describe the effects of interventions in the targets of the models. We also discuss the limitations of ABMs and put forward policy recommendations.

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Mariusz Maziarz & Martin Zach

Mariusz Maziarz

Interdisciplinary Centre for Ethics & Institute of Philosophy, Jagiellonian University, Kraków, Poland

*mariusz.maziarz@uj.edu.pl* Interdisciplinary Centre for Ethics, Jagiellonian University, Grodzka 52, 33-332 Krakow, Poland

Martin Zach

Department of Philosophy and Religious Studies, Faculty of Arts, Charles University in Prague, Prague, Czech Republic

Abstract

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Key-words: agent-based modeling, mechanism, causal inference, mechanistic evidence, difference-making evidence, SARS-CoV-2

## INTRODUCTION

In the aftermath of the outbreak of the novel coronavirus, governments around the globe have introduced non-pharmaceutical public health interventions aimed at slowing down the spread of the resultant pandemic. These measures range from relatively mild requirements like wearing face masks, washing hands, or avoiding close contacts to school closures and imposed isolation that are likely to have a detrimental and unpredictable influence on social and economic life.<sup>1</sup> Despite their significant impact, the introduction of many of these measures was not supported with high-quality evidence. First, conducting RCT would not be feasible for both ethical and practical constraints. Second, significant differences between the coronaviruses that caused the SARS and MERS outbreaks and SARS-CoV-2 (such as the likely airborne transmission<sup>2</sup> and asymptomatic infectiousness of the latter<sup>3,4</sup>) undermine extrapolation from the data gathered during these previous epidemics. Finally, the current pandemic has not lasted long enough to gather observational data in the amount and quality sufficient for the assessment of the efficacy of alternative public health interventions, since the first reports were published just weeks after the first measures were introduced.<sup>5</sup>

One of the many ways to address the issue concerning the impracticality of conducting RCTs and observational studies in the context of an ongoing pandemic is through scientific modeling, in particular epidemiological modeling. Here, we focus on the so-called agent-based modeling (ABM) approach, which differs from more traditional epidemiological modeling in several ways.

ABMs are a form of computational modeling strategy where agents are treated as entities interacting with each other and their environment in a locally-defined fashion described by a set of rules. The overall dynamics of the system are then computed, allowing for the simulation of complex patterns and an understanding of how these patterns arise.<sup>6,7</sup> ABMs are used in many scientific contexts, including modeling the spread of infectious diseases, and have proven successful in informing policy decisions before. For instance, Eisinger and Thulke<sup>8</sup> modified and then applied a previously-developed ABM of the spread of rabies, generating a rule-based model that represented specific spatial and behavioral characteristics of the fox population (e.g., with fox families represented as moving within home ranges and young foxes engaging in long-distance migratory behavior).<sup>6</sup> Whereas the classical differential equation models predicted that vaccinating at least

70% of the fox population would eliminate rabies, the ABM indicated that a successful vaccination strategy could do with much less than 70% of the population being immunized once the spatial arrangements of fox hosts were explicitly considered, saving millions of Euros as a result. Moreover, the ABM also suggested that the classical strategy would fail more often than not, and was successfully applied to deal with the rabies problem. However, despite the promising record of using ABMs in effective epidemiological interventions, its use in informing proposed measures against the novel coronavirus epidemic has raised criticism.<sup>9–11</sup>

Unfortunately for the assessment of healthcare interventions based on this type of epidemiological models, standard evidence hierarchies either exclude such studies all together or include theoretical or mechanistic inferences at the lowest level of the hierarchy. For example, the Oxford Centre for Evidence-Based Medicine lists mechanism-based reasoning at the lowest, fifth stage,<sup>12</sup> and National Institute for Health and Care Excellence (NICE) guidelines exclude both epidemiological mathematical models and mechanistic reasoning.<sup>13</sup> This can be explained by the novelty of agent-based modeling and the limited trust of EBMers in theoretical and mechanistic reasoning, which, despite being used implicitly to assess the possibility of confounding and the quality of results,<sup>14</sup> is downgraded or rejected as either subjective or fallacious.<sup>15</sup> However, such a view has been challenged by a group of philosophers advocating for improving the practices of evidence assessment in medicine by putting more weight on mechanistic reasoning in causal inference.<sup>16–18</sup> The position of the EBM+program<sup>16–18</sup> is encapsulated by the normative reading of the Russo-Williamson Thesis,<sup>19</sup> which states that causal claims should be based on both difference-making and mechanistic evidence.

The causal claims supported by agent-based models have been interpreted inconsonantly: either as being in line with the potential outcome approach (POA),<sup>20</sup> as delivering theory-driven understanding<sup>21</sup> or as providing mechanistic evidence.<sup>22</sup> Below, we show that all of these apparently inconsistent interpretations are correct, because the best contemporary ABMs bear a resemblance to the actual mechanisms and therefore allow for the counterfactual assessment of intervention efficacy in the target while also delivering an understanding of the phenomena of interest. Our argument proceeds by (1) discussing as a case study an ABM of SARS-CoV-2 epidemic in Australia, (2) showing that the best ABMs represent actual mechanisms despite the presence of various simplifications, and (3) considering the limitations of using ABMs as evidence for clinical and policy decisions.

## MODELING THE SARS-CoV-2 EPIDEMIC

Apart from the compartmental SIR (Susceptible, Infectious, Recovered) framework and its derivatives<sup>23–28</sup> or regression analysis,<sup>29,30</sup> most advanced models of the spread of the novel coronavirus are transformed versions of agent-based influenza pandemic models<sup>11,31</sup>. Such models have been used as evidence for introducing (sometimes severe) public health measures,<sup>32</sup> with the recent change in British policy being the prime example. In this section, we illustrate this approach to modeling the SARS-CoV-2 pandemic with an agent-based model of the epidemic in Australia<sup>31</sup> based on AceMod. Developed as a “framework for studying influenza pandemics in Australia,”<sup>33(p412)</sup> AceMod is an influenza spread model that addresses the need for simulating interventions responding to the outbreaks of future respiratory diseases. While the 2009 swine flu pandemic was the motivation for constructing AceMod, the model was not intended to accurately represent the outbreak of the H1N1 strain, but rather as a generalized framework for studying how an infectious disease spreads through the social interactions of Australians. AceMod utilizes census data to ascribe realistic spatial and social characteristics to almost 20 million agents inhabiting the model world. These agents are divided into different social groups of varying characteristics, with households differentiated proportionally according to statistical data on the prevalence of different types of families (singles, single parents, and couples with or without children). These features are ascribed to agents stochastically in a way that replicates the aggregate structure of statistical data. During the daytime, children and students meet in classrooms and at schools, adults go to work, and pensioners stay at home. During the nighttime, the agents encounter contacts at households and in their neighborhoods (e.g., at supermarkets, theaters).

The disease can be contracted by an agent in the event of meeting an infected individual in one of these settings. The probability that an agent  $i$  contracts the disease in a given step  $t$  depends on the number of sick individuals met in that step and the contagiousness of the disease, scaled by  $\beta$ . The modelers assume

that the infectivity of the disease decreases linearly over time. Asymptomatic cases are assumed to be 50% less infectious than symptomatic ones, and the flu lasts 5 days within the model. After this period, recovered agents cannot infect others. Additionally, those who experience symptoms do so after an incubation period lasting approximately three days. The influenza epidemic is started by agents coming to Australia via international airports and seeded into communities living near the airports at random.

In order to represent an epidemic of a particular strain of influenza with AceMod, the model requires calibration. Modelers can proceed with this step in two ways, depending on the accessibility of data. In the case of well-studied influenza strains, their infectivity and the ratios of transmission in different contexts are well-recognized, and parameter values can be chosen on the basis of empirical studies. However, if these data are missing, then parameter values have to be calibrated using statistical procedures such as simplex or genetic algorithms to maximize the fit of the model to a benchmark. After constructing and calibrating AceMod, modelers run simulations to obtain the estimates of prevalence, incidence, and attack rates, and choose the most common outcome (due to stochasticity, different runs of the model may lead to obtaining slightly different results).

Chang et al.<sup>31</sup> have used a significantly amended version of AceMod to address the question of the effectiveness of non-pharmaceutical interventions aimed at suppressing the SARS-CoV-2 epidemic in Australia. The selection of models constructed to control a novel and possibly deadly strain of the seasonal flu in this case is primarily the result of the rapid demand for evidence informing decisions regarding public health measures, which may raise doubts about the justification and soundness of their conclusions. For example, one can ask whether the efficacy claims assess healthcare interventions against the novel coronavirus epidemic or an artificial pathogen existing only within the model world that shares some features of influenza and others of SARS-CoV-2. To address this criticism (considered in depth below), we discuss the changes introduced to the model and argue that the process of model calibration and validation suggests that the model represents the actual mechanism of the SARS-CoV-2 epidemic.

ABMs such as AceMod can be seen as consisting of two parts: the rules specifying the behavior of agents and the creation of the model society, as well as the assumptions characterizing the infectivity of the pathogen causing the epidemic. Given that AceMod is based on 2016 census data and a major change in social behaviors is unlikely to have occurred since then, the model accurately represents the social interactions of present-day Australians. Hence, the former part of the model has been left mostly unchanged, beyond increasing the number of agents to over 24 million to adjust for the growing population. In addition to introducing a social structure sufficiently resembling the contact network of the present population, obtaining accurate predictions of epidemic development and policy assessment requires inputting data on transmission likelihoods that are true for the pathogen causing the modeled epidemic.<sup>34</sup> Most changes in the model are concerned with the assumptions specifying the infectivity of the disease. Even though several features of influenza epidemics are similar to the epidemic caused by the novel coronavirus, they differ with respect to infectivity and attack rates, mortality rates, the average duration of disease, the reproductive number  $R_0$ , and the distribution of asymptomatic cases. Therefore, these parameters in the model required recalibration.

The transmission probabilities remained mainly as specified in the influenza model. In order to account for the differences in the incubation period and disease length, Chang et al. set the time from contraction to the appearance of symptoms to 5 days on average and the duration of the disease to 12 days. Infectivity increases exponentially the day after an agent gets infected and then decreases linearly until the end of infection, so cases are most infectious at the start of symptoms. The length of the generation period was calibrated to 6.4 days in order to reflect this difference in the model. Additionally, the likelihood of contracting SARS-CoV-2 but staying asymptomatic was set to be age-dependent, and equaled 1/3 for adults while minors were set to be five times less likely to suffer from symptoms than adults. While this assumption is in agreement with the empirical findings that children represent a minor fraction of symptomatic cases, the calibration aimed at reproducing aggregate epidemic curves and may diverge from the actual chances of developing symptoms.

Within the AceMod framework, the reproductive number  $R_0$  is not one of the assumptions inputted into the model. Rather, its estimate results from a simulation of the scenario described by the rules and assump-

tions, some of which are stochastic. The assumptions considered and, particularly, the parameter denoting contagiousness of the disease ( ) have been calibrated such that  $R_0$  stays within the limit of (2.0-2.5), i.e., in agreement with empirical estimates of the reproductive number at the beginning of the SARS-CoV-2 outbreak.<sup>35,36</sup> The set of parameter values that result in the estimate of  $R_0 = 2.27$  create the epidemic dynamics reproducing the beginnings of the outbreak in a few countries experiencing the disease prior to Australia (China, Italy, Spain), where the growth rate of cumulative incidence equaled roughly 0.2. In addition to reproducing the empirical data for the beginning of the epidemic, the recalibrated AceMod allows for simulating what the future of the epidemic in Australia may look like. As the modelers admit, the *Baselines* scenario, which is based on the assumption that agents do not change their behavior in response to the epidemic, is unlikely given the widespread self-imposed isolation in other countries. However, it allows for counterfactual comparisons of the different possible (sets of) interventions relative to the Baseline scenario. In order to assess the efficacy of particular healthcare policies, Chang et al. modify relevant rules and assumptions to describe the spread of SARS-CoV-2 under either case isolation, school closure, along with three levels of compliance with social distancing, along with a few combinations of the three policies. For instance, in order to assess the effect of school closure (including primary and secondary schools, colleges, and universities), the parameter denoting the chance of meeting an infected agent in schools is set to zero, which describes the situation when both students and teachers stay at home (and hence cannot contract the virus). These counterfactual scenarios represent the effects of interventions on the model world. All interventions are modeled as taking place after the number of cases exceeds 1000. The comparison of most common outcomes (given the stochasticity of the assumptions and rules, they are indeterministic) including interventions with the baseline scenario allows for putting forward counterfactual causal claims that describe the effects of interventions on peak incidence and prevalence and the development of the epidemic in time. The conclusions are internally valid as long as no coding error occurs. However, the reliance of the model on simplifications generates a question as to whether the assessment of intervention efficacy holds for the novel coronavirus epidemic in Australia (i.e., of external validity).

## ABMS AS MODELS OF ACTUAL MECHANISMS

Before proceeding to our argument, let us first make several general remarks about modeling. These remarks should prove essential in clarifying the main issues that are often raised with regard to using simplified models, particularly in the context of policy decision-making. First of all, ABMs are instances of mechanistic models, for they clearly fit the characterization of what a mechanism is: a set of entities whose activities and interactions are organized such that they are responsible for the phenomenon.<sup>37</sup>

It should also be noted that much like any other kind of model, ABMs serve as simplified representations of their target phenomena. As the AceMod case clearly shows, modelers introduce various simplifications by which they purport to adequately capture the core dynamics of the modeled phenomenon. In this process, they first abstract away from the complexities of the real system by 'extracting' certain features that they believe to be of crucial importance and that will then be the focus of modeling, whereas other features that may or may not have a causal influence are disregarded in these early stages. Modeling is an iterative process during which the merits of the model's assumptions are continuously being evaluated, and if required, the assumptions are refined and additional assumptions added. More importantly, some of those extracted features are distorted to the extent that, if taken literally, they would misrepresent the actual state of things. However, introducing such distortions is often made in full awareness, with the ultimate goal of finding out whether the consequences they have for the behavior of the system make a difference and to what degree. Philosophers often refer to the former - i.e., the set of properties retained in a model - as an abstraction, while the latter case - i.e., the distortions of the system's features - is called an idealization.<sup>38</sup>

However, abstractions and idealizations do not exhaust the conceptual toolbox available to modelers. A popular way to attempt to model a given system realistically is to introduce various approximations. Although there are noteworthy differences between approximations and idealizations, we cannot afford to go into any detail here. In summary, models often effectively disregard, distort and otherwise simplify possibly important details. In light of this, many wonder whether we can gain insight into the modeled phenomenon

at all, and if so then how.

Although the SARS-CoV-2 ABM is fairly detailed and precise, it cannot do without some of the simplifications discussed above. Consider some of the following assumptions introduced in the model. On the one hand, the basic features of the social life of the majority of the population are extracted and considered in the model: e.g., the inclusion of day and night regimes with their respective differences in social behavior allows for modeling a more realistic scenario than in simpler models. On the other hand, the infectivity of symptomatic and asymptomatic cases is considered to be constant for all members of the two groups of agents, albeit it differs between the groups. In reality, we expect that infectivity varies, which is further supported by extreme cases of super-spreaders who infect a large number of people and thus may seed new local outbreaks, which could arguably impact the predictions.<sup>39–41</sup> Other parameter values also have a wide distribution but are treated as constant, often by calculating the mean value. The ABM also does not consider the potential impact of ethnic differences<sup>42–46</sup> in the population with respect to differing lifestyles, socioeconomic status and immune host responses, all of which could affect the dynamics of the spread.

Furthermore, some other assumptions exceed our current understanding of the epidemic and SARS-CoV-2's transmission mechanism. For example, one of the assumptions of the AceMod model is the linear reduction of infectivity over time. Unfortunately, empirical results<sup>47</sup> suggest only that infectivity reduces over time, but do not indicate the linearity of this process. Additionally, AceMod and its SARS-CoV-2 model put agents into working groups of 20 agents, despite the heterogeneity of their working conditions. Considering the differentiation of work duties (from healthcare workers and shop assistants to writers with virtually no social interaction), the chance of meeting an infected person at work is actually job-specific and therefore the model simplifies the reality.

Consequently, we concur with Andersen's claim that "no mechanism model can include all the actual, much less the potential, causal relationships in which such a mechanism may engage in a system."<sup>48(p995)</sup> This pessimistic view on simplified models has inspired the method known as exploratory modeling.<sup>49</sup> In cases when the values of parameters and assumptions inputted into the model cannot be established with certainty, researchers can simulate multiple possible worlds to discover the dependencies that are stable across the set of different models. In cases when only a fraction of assumptions are uncertain, researchers conduct sensitivity analyses to check if changes in the values of the parameters lead to changes in their conclusions.<sup>50</sup> The results that remain unchanged despite minor adjustments to assumptions are considered to be robust.<sup>51</sup> This, in turn, leads to choosing those interventions that are most effective across different sets of parameter values, known as robust decision-making.<sup>49</sup>

Others prefer to think in terms of the distinction between how-actually and how-possibly modeling, referring to models that describe an actual mechanism or a possible mechanism, respectively.<sup>52</sup> There are two general ways to unpack the concept of a how-possibly model. First, we may want to say that a model serves as a hypothesis to be confirmed or disconfirmed as new evidence emerges. In this sense, a how-possibly model will eventually either turn into a how-actually model, should the evidence confirm it, or be discarded if the evidence is contrary to the model's conclusions. The other general notion of a how-possibly model invites a different attitude. Rather than being in the position of having little data to establish whether or not the model does, in fact, represent the actual mechanism, we may interpret the model as representing something other than the potentially actual mechanism. On this view, claims about possible mechanisms do not attempt to pick out actual states, nor do they attempt to explain how a phenomenon actually occurs. Instead, they refer to conceivable states, and ask whether the hypothesized mechanism could, in principle, produce the phenomenon in question if certain assumptions are satisfied.

Here we argue that, notwithstanding the simplifications introduced in the discussed influenza and SARS-CoV-2 ABMs, the epidemiologists are, in fact, providing representations of actual mechanisms of the spread of the viruses. This can be supported by exploiting the relevant similarities<sup>53,54</sup> between the SARS-CoV-2 ABM and the actual outbreak. The respects in which an ABM can be judged similar to its target concern the features retained in that model, while the degree(s) of similarity concern the extent to which the model's features match those of the phenomenon. A good example is setting the parameter/assumption of incubation

period=5 days. This assumption was introduced based on empirical research: “We maintained the incubation period (the interval from the infection to the onset of disease in an individual) around the mean value of 5.0 days, as reported in several studies, e.g., the mean incubation period was reported as 5.2 days, 95% confidence interval (CI), 4.1 to 7.0, while being distributed around a mean of approximately 5 days within the range of 2–14 days with 95% CI.”<sup>31(p3)</sup>

To elaborate this further, Glennan<sup>55</sup> introduced a useful conceptual distinction between what he called behavioral adequacy and mechanical adequacy. According to Glennan, a model represents an actual mechanism if it reproduces the aggregate behavior of the phenomenon, and truthfully describes its parts and interactions. Concerning the behavioral adequacy, one should be asking if “the model predict[s] (quantitatively or qualitatively) the overall behavior of the mechanism?”<sup>55(p457)</sup>. By calibrating the model to data from the beginning of the epidemic, Chang et al.<sup>31</sup> showed that it reproduces the benchmark variables ( $R_0$  and attack rate).

Two remarks are in order here. First, one may oppose the claim that what is being represented is the actual mechanism by arguing that the mechanism underlying the beginning of the outbreak and the fully-fledged epidemic are distinct. Changes in social behavior or genetic mutations could undermine the behavioral adequacy of the model. Second, it is possible (at least in principle) that the model represents a false mechanism, but is calibrated to the relevant benchmark such that it reproduces it. For example, there is no data confirming (or disproving) the assumption that children are asymptomatic five times more often than adults. As the modelers admit, this assumption was made not only to account for the lower attack rate among minors, but also to make the model adequate to aggregate-level data. This approach to calibration resembles the estimation of statistical parameters (a.k.a. curve fitting) and is considered dubious. The main line of criticism highlights that it is in principle possible to construct a model that represents a possible mechanism and, using calibration, adjust parameter values so that it reproduces the represented phenomenon, i.e., obtains behavioral adequacy despite being false. However, while this criticism is indeed justified regarding models of mechanisms that are epistemically inaccessible in other ways (such as mechanisms in the social sciences<sup>56</sup>), it is not so in the case of epidemiological mechanisms whose transmission mechanism can be studied empirically and compared to the mechanism represented by the model.

This can establish that the mechanism represented by the model is similar (in relevant aspects and to relevant degrees) to the mechanism that generates the outbreak, i.e., achieves mechanical adequacy in Glennan’s terminology. Applying the list of Glennan’s<sup>55(p457)</sup> criteria for mechanical adequacy justifies the claim that the mechanism represented by Chang et al.<sup>31</sup> resembles the actual mechanism. First, according to our best contemporary understanding of the spread of the novel coronavirus, the model identifies all of the components of the mechanism. This would change if further studies identified other significant transmission routes, e.g., the fecal-oral route. Second, the model represents the entities of the mechanism in a localized way, given that it retains the spatial distribution of inhabitation in Australia. Additionally, the model simulates the development of an epidemic in time. This asserts that the “spatial and temporal organization of the mechanism” is accurately represented. Third, given that the number and place of social interactions are crucial for modeling the spread of contagious diseases, the model accurately captures relevant properties of the agents inhabiting the model world. Fourth, the calibration to census data asserts that the model provides “quantitatively accurate descriptions of the interactions and activities of each component,” at least on average for groups of agents. Finally, our background knowledge suggests that there is no other mechanism (different from the spread of the pathogen through human interactions) that could be responsible for the epidemic of SARS-CoV-2.

Given that AceMod fulfills Glennan’s criteria for behavioral and mechanical adequacy, considering our current understanding of the novel coronavirus, we can conclude that Chang’s et al.<sup>31</sup> model represents the actual mechanism of the spread of the disease in Australia. Given this, the claims assessing the efficacy of the mitigation measures under consideration are likely to be accurate not only within the model but also about its target. We claim this with several caveats in mind to be discussed in the next section.

## DISCUSSION AND RECOMMENDATIONS

Our study defends using ABMs for informing decisions regarding mitigation and suppression measures by arguing that its best epidemiological models represent actual mechanisms. Provided that the model's assumptions are calibrated and checked against the background empirical data - that is, the components, their activities, and spatiotemporal organization resemble (in relevant aspects and to a certain degree) the actual state of things - iterative runs of the simulations can indeed provide understanding and inform policy decisions. This is because the model delivers both difference-making and mechanistic evidence by satisfying the criteria of behavioral and mechanical adequacy, respectively.

In contrast to our claim, epidemiological SIR models and ABMs have been criticized for over-simplifying target phenomena and hence lacking relevance for policy decisions. For instance, Eubank et al. criticized the Imperial College London model<sup>11</sup> for its "reliance on a simplified picture of social interactions [that] limits its extensibility to counterfactuals. The general nature of conclusions based on such model can be expected to be similar to those of a simple compartmental model."<sup>57(pp5-6)</sup> Similarly, Squazzoni et al. suggested that even though AceMod is better calibrated than other epidemiological ABMs, "these [models] do not capture network effects nor people's reactive responses as the population states simply change via stochastic (randomized) processes determined by parameters (although the parameters derive from data)."<sup>9(p2.6)</sup> In our view, these highly-advanced epidemiological models, while being simplified representations of reality, account for relevant aspects of social interactions and crucial aspects of the novel coronavirus epidemic (e.g., contagiousness), therefore allowing them to be put forward as evidence for policy-relevant claims.

However, all models involve not only simplifications of the represented phenomena but also isolate certain aspects of interest. Epidemiological models usually do not account for the harms of non-pharmaceutical interventions. Severe mitigation measures such as imposed social distancing and business closures are likely to hamper economic and social life. All models are partial representations of reality and, given that the primary purpose of an epidemiological model is to address the efficacy of health care interventions, they isolate away certain factors and effects of interventions (economic and social) and are more accurate in predicting the spread of the disease. Other models<sup>58,59</sup> tradeoff epidemiological accuracy with accounting for social and economic effects, and may be more relevant for assessing the harms of mitigation measures.

Additionally, ABMs, much like the compartmental models, are dependent on the assumptions of the modelers.<sup>10</sup> Our claim that AceMod calibrated for SARS-CoV-2 bears similarity to the actual mechanism of the epidemic depends on the accuracy of the empirical results used as an input for this model. We need to acknowledge the provisional nature of these empirical results, given the novelty of the pathogen. If the parameter values in AceMod were miscalibrated, then the assessments of intervention efficacy could be wrong. This implies that neither the virus can mutate nor that people can significantly and unpredictably change their behavior since "the efficacy of implementation depends on people's reactions, [the stability of] pre-existing social norms, and structural societal constraints."<sup>9</sup> Furthermore, the effects of epidemiological agent-based modeling are highly dependent on social structure and carefully calibrated to social and economic characteristics. Therefore, the epidemiological ABMs are geographically-localized and their conclusions should not be extrapolated beyond their target systems,<sup>60</sup> as long as the models and their predictions are calibrated to particular settings. Finally, while AceMod is well-documented in the two publications discussed throughout our paper, neither its code nor detailed documentation regarding its use is published (this unfortunately also applies to some other ABMs of the SARS-CoV-2 epidemic). Given these limitations, the models should be carefully checked for coding errors and other possible flaws before applying their implications in the policy context.

In summary, we have argued that, despite the criticism raised against models being the appropriate vehicle for informing policies, the SARS-CoV ABM is suitable for this purpose because the mechanism described by the model sufficiently resembles the mechanism at work in the real world. Thus, our best contemporary epidemiological ABMs are representations of the actual mechanism of the spread of the virus. Unfortunately, such models have been left out from methodological discussions and are not explicitly listed by evidence hierarchies. While the need for appraising mechanistic reasoning in medicine is also voiced by EBMers,<sup>61</sup> there is no broadly-accepted view on how to amalgamate evidence of different types. Further research is



needed to assess the risk of bias in the epidemiological models that deliver both difference-making and mechanistic evidence. However, considering the current situation and pressing need for rapid and accurate decisions regarding mitigation measures, policymakers should take to heart the advice that “if no randomized trial has been carried out [...], we must follow the trail to the next best external evidence and work from there.”<sup>62(p74)</sup> In the current situation, accurately calibrated epidemiological ABMs are the best existing evidence.

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## CONFLICT OF INTERESTS

None of the authors reports conflict of interests.