Pandemic lockdown, isolation, and exit policies based on machine learning predictions

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Abstract
The widespread lockdowns imposed in many countries at the beginning of the COVID-19 pandemic elevated the importance of research on pandemic management when medical solutions such as vaccines are unavailable. We present a framework that combines a standard epidemiological SEIR (susceptible–exposed–infected–removed) model with an equally standard machine learning classification model for clinical severity risk, defined as an individual’s risk of needing intensive care unit (ICU) treatment if infected. Using COVID-19–related data and estimates for France as of spring 2020, we then simulate isolation and exit policies. Our simulations show that policies considering clinical risk predictions could relax isolation restrictions for millions of the lowest risk population months earlier while consistently abiding by ICU capacity restrictions. Exit policies without risk predictions, meanwhile, would considerably exceed ICU capacity or require the isolation of a substantial portion of population for over a year in order to not overwhelm the medical system. Sensitivity analyses further decompose the impact of various elements of our models on the observed effects. Our work indicates that predictive modeling based on machine learning and artificial intelligence could bring significant value to managing pandemics. Such a strategy, however, requires governments to develop policies and invest in infrastructure to operationalize personalized isolation and exit policies based on risk predictions at scale. This includes health data policies to train predictive models and apply them to all residents, as well as policies for targeted resource allocation to maintain strict isolation for high-risk individuals.

KEYWORDS
COVID-19, epidemic models, machine learning, personalized risk management, SIR

1 | INTRODUCTION

Many countries have adopted isolation restrictions, or “lockdowns,” to control the viral spread during the COVID-19 pandemic (Koo et al., 2020). Such nonpharmaceutical pandemic management strategies can complement pharmaceutical ones (e.g., treatments or vaccines) but also may be the only available tool if the latter either are not scientifically possible or take significant time to develop. While epidemic models have been used to inform such policies (Ferguson et al., 2020; Flaxman et al., 2020), how to best initiate, relax, and possibly reinitiate isolation restrictions is unclear. Some approaches rely on immunity tests (Petherick, 2020) or on testing and tracing technologies (Wang et al., 2020), but we study a different approach. Instead of using “ex post” immunity or diagnostic tests, utilizing “ex ante” predictive technologies, such as machine learning, which have been proved successful in other contexts. This approach can be used even in the absence of medical interventions because it relies solely on using data gathered in the pandemic’s early stages to identify factors that affect the severity of symptoms, not on developing and distributing tests, treatments, or vaccines. This approach can complement those relying on immunity and diagnostic tests or be considered independently.
The type of personalized isolation ("confinement") and exit ("deconfinement") policy that we study is as follows: First, the clinical risk score (risk of experiencing symptoms severe enough to require an intensive care unit [ICU] bed) for each individual is predicted. Second, those with predicted scores above a certain threshold are classified as "severe," or "high risk," and the remainder are classified as "mild" or "low risk." Third, those classified into the high-risk group are subject to stricter isolation and protection ("confined"), while those in the low-risk group are placed under softer restrictions or none at all ("released"). In practice, this can be achieved by targeted allocation of resources (e.g., providing masks and other personal protective equipment [PPE], dedicating health support, delivering groceries and other necessities for free) to the high-risk group, targeted government communication that differentiates between high- and low-risk individuals, and other targeted policies. Policy makers adjust the high versus low threshold over time to achieve the desired objectives while meeting required constraints. While we chose to minimize the time to complete exit while not exceeding ICU capacity at any point, other formulations of the underlying multiobjective problem are feasible.

The studied policies rely on two key assumptions: First, only a small percentage of the population is in the high-risk category. For example, estimates for France as of spring 2020 showed that the vast majority of the population, or >99% (Salje et al., 2020), will not experience severe symptoms needing an ICU if infected by the SARS-CoV-2 virus. Second, risk prediction models based, for example, on data from early infections can be developed and deployed if the necessary data are available. While the infrastructure required to achieve this during the COVID-19 pandemic appears limited despite the existence of risk prediction models (e.g., Bertsimas, 2020), appropriate policies may enable pandemic management based on the data-driven risk predictions in the future.

The intuition behind the studied policies is as follows: Using COVID-19 data as an example, were one to (i) determine who the ~ 1% of severe cases are and (ii) perfectly and temporarily isolate and protect them, the remainder of the population would be able to continue a more or less normal life. In such an ideal scenario, many low-risk people would get infected and would infect others but none would have severe symptoms because those high-risk individuals already have been correctly identified and perfectly protected. The medical system would not be overwhelmed, no one would die (though we note we do not consider long-term health effects for those infected), and both the society and economy would avoid a major shock from the indiscriminate lockdowns implemented in many countries.

In practice, the effectiveness of such a policy would depend on two critical imperfections. First, risk prediction models might occasionally make mistakes, for examples, false positive and false negative errors. Second, isolation would be imperfect as, for example, high-risk individuals who should be isolated may occasionally encounter those infected (e.g., due to PPE shortages, noncompliance, or family situation) and low-risk individuals who would be able to continue normal life may not do so (e.g., due to fear).

We study how these two imperfections impact the effectiveness of the aforementioned personalized pandemic isolation and exit policies. We extended a standard epidemic model, namely, a version of the susceptible–exposed–infected–removed (SEIR) model (Kucharski et al., 2020), to incorporate personalized predictions of severity risk (see Appendix A in the Supporting Information for alternative modeling frameworks). Using simulations, we investigated how prediction models for patient severity may inform policy in two scenarios: amid an ongoing outbreak, as was the case in France when lockdown began on March 17, 2020, and when the outbreak has been curbed and progressive loosening of isolation policies (exit or deconfinement) may take place, as was the case in France beginning May 11, 2020.

Our analysis is based on assuming hypothetical risk prediction models one may be able to develop for a pandemic. These can rely on factors known to affect the severity of symptoms if infected, when such factors exist. For example, existing research indicates differential impact of COVID-19 depending on age, body mass index, hypertension, diabetes, and other factors (Guan et al., 2020), which already have been used in emerging risk models, such as those reported in Bertsimas (2020).

To populate our simulation models, we used available COVID-19 estimates and data from France as of May 2020 (Di Domenico et al., 2020). At the end of the lockdown on May 11, there were about 2750 ICU beds occupied by people with COVID-19, down from a peak of 7148 against the French health system’s 6000-bed capacity. We used current estimates with a reproduction number value of $R_0 \approx 2.9$ prior to lockdown, and 1.5 million people who had been immune or infected when it started in France on March 17, 2020 (Salje et al., 2020). We analyzed uncertainty using approximate Bayesian computation (Marjoram et al., 2003).

Our simulations led to the following main observations and corresponding implications:

1. Isolation and exit policies when based on risk-model predictions could be substantially faster and safer. Utilizing realistic parameter values and a high-quality risk model at the upper end of Bertsimas (2020), simulations indicated that a complete exit from COVID-19 lockdown could be undertaken in three waves over 6 months, with only 10% of the population being under strict isolation for longer than 3 months—all without overwhelming the medical system and exceeding ICU capacity. Simulations indicated that without such a model, a complete exit would take 17 months and 40% of the population would be subject to strict isolation for over a year or ICU capacity would be exceeded four times over.

An alternative way to interpret our results is that even with a good risk model, ~30% of the population still must be strictly isolated for several months. In other words, the "herd immunity" approach some policy makers have endorsed is impossible without (i) a high-quality risk
prediction model and (ii) the ability to strictly isolate a substantial portion of the high-risk population.

**Implication:** Governments should invest in individual health data infrastructure to make such models implementable at scale. This entails infrastructure that not only collects data on the few thousand people who exhibited symptoms and went to hospitals but also collecting individual medical data on the entire population to obtain health risk predictions for all residents; see Evgeniou et al. (2020) for further discussion on the resultant data policies, privacy, and other related issues.

**Disclaimer:** because such data and policies do not exist in most countries, our policy was not suitable for managing COVID-19 in 2020. Rather, we study how personalized policies based on machine learning predictions could improve pandemic management in the future.

2. **Even moderate-quality risk models already could bring measurable improvements,** relaxing isolation for millions of people months sooner while adhering to existing constraints on medical resources. Further, and somewhat surprisingly, even with imperfect models, imperfect but targeted and optimally timed partial isolation policies can be more efficient than full nondiscriminatory lockdowns, as such policies allow safer immunity building while better protecting the high-risk population and keeping the average isolation percentage low.

**Implication:** For immediate action, focus on a “minimal viable product” data and models that can be used at scale. Even amid the COVID-19 pandemic, data on age, body mass index, and hypertension and diabetes—all of which can be assessed at a nearby pharmacy for all people within weeks—already can be used with a risk model such as in Bertsimas (2020) to inform policies that could be relevant for practice.

3. **Personalized policies based on risk-model predictions are highly sensitive to the protection level of confined people.** Interestingly, the impact of the protection level of deconfined people on simulated outcomes depends on risk-model quality. With a high-quality risk model, the optimal policy builds herd immunity, which can be done faster when deconfined people are less, not more, protected.

**Implication:** Personalize resource allocation to protect the confined predicted high-risk people: Distribute to them masks and other PPE, supply them with food and other necessities for free, prioritize testing those in contact with them, and so forth. Do not spread resources; practice targeted allocation.

4. **Lastly, whether an individual is classified as high- versus low-risk changes dynamically over time:** How one is classified depends not only on one’s individual characteristics but also on the state of the epidemic. Our proposed personalized policy combines the epidemic progression with data science principles and optimally adjusts the high-versus low-risk classification threshold so as to ensure safe and fast confinement and deconfinement over time.

**Implication:** A careful communication strategy, adjusted over time, is needed to convey such personalized policies to the public.

We now provide some comments on related work. As we do, Acemoglu et al. (2021) study the benefits of applying differential isolation restrictions within a multirisk SIR (susceptible–infectious–removed) model. Their risk groups, however, are static—young (20–44), middle (45–64), and old (65+)—so the resultant policies are rather limited. For instance, their optimal fully targeted policy keeps the “old” group (∼20% of the population) in isolation essentially indefinitely, waiting for a vaccine to arrive. In contrast, our high versus low risk assessment depends on the classification threshold in a machine learning model and changes dynamically as the epidemic progresses, allowing for much faster exits. Recall that with a high-quality risk model, the optimal exit takes 6 months and only 10% of the population is isolated for more than 3 months.

Our approaches also differ with regard to the multiobjective nature of managing a pandemic. Acemoglu et al. (2021) treat the problem as a weighted objective function and compute an efficient frontier between the economic and health metrics. We treat the problem as a constrained optimization. We optimize one objective (time to remove the isolation restrictions, which represents the socioeconomic goals) subject to a constraint on the availability of ICU beds (representing the health goal).

Gershon et al. (2020) and Duque et al. (2020) employ a similar constrained approach within their settings and objectives, which differ from ours. Similar to Acemoglu et al. (2021), Gershon et al. (2020) utilize static risk groups but they are not solely based just on age and include children, low-risk adults, high-risk adults, and nursing home occupants. The results are similar to Acemoglu et al. (2021) in the sense that the high-risk groups must be isolated indefinitely; under certain conditions, however, the low-risk may not be isolated at all in their model. Our results are similar in that up to 65% of the population should never be isolated. Duque et al. (2020) study the timing of nontargeted shelter-in-place isolation orders. We also study timing but our model isolates, that is, targets a different (smaller) fraction of the riskiest remaining population at each new epoch.

Targeting based on risk factors is, of course, not the only way: Birge et al. (2020) study spatial targeting and Camelo et al. (2021) study dual targeting based on risk groups and their activities. Such studies are complementary to ours.

We finally mention three companions to the present paper. First, given the large body of literature developed at the beginning of the COVID-19 pandemic, when multiple research articles were produced daily, we also present the detailed review in a “literature appendix”, Garin et al. (2021) in addition to the papers discussed above. Second, because our findings are based on numerical simulations, the code is available via GitHub at https://reine.cmla.ens-cachan.fr/boulant/seair, and the algorithmic details for how our model is implemented in the code are provided in Boulant et al. (2020).
This forms a “code appendix” to our paper, following the highest standards of reproducible research. Third, to facilitate dissemination of our results for the general public, we created a noncoding demo “simulator” for the differential policies that we study here: https://ipolcore.ipol.im/demo/clientApp/demo.html?id=305. This demo is prepopulated with the parameters for France used in our simulations but one can change the parameters and simulate the pandemic isolation and exit policies for other counties given their respective situations.

2 | MODEL

At a high level, our model is a combination of a rather standard SIR-like compartmentalized epidemiological model with an equally standard machine learning binary classification/risk model.

2.1 | Compartmentalized epidemiological model: Extended SEAIR

We start from the standard SIR model and extend it in two ways. First, we add three additional kinds of compartments:

- Exposed, to account for those who are infected but are neither infectious nor symptomatic;
- Asymptomatic, to account for those who have contracted the disease and are infectious but asymptomatic; and
- ICU, for people who experience severe enough symptoms and require intensive care (or, more generally, whatever one may define as severe cases for a pandemic).

Taken together, our model captures a pandemic where a proportion of infectious but initially asymptomatic individuals are assumed to develop mild symptoms and recover naturally, while the rest develop severe symptoms requiring hospitalization; see Figure 1. Such a separation of exposed individuals based on severity of symptoms has been a recurring aspect of most modeling approaches (whether deterministic or stochastic) by prominent epidemiologists; see, for instance, Di Domenico et al. (2020), Ferguson et al. (2020), Sonabend et al. (2021), and Salje et al. (2020). Including the “A” compartment captures the salient feature of a pandemic like COVID-19, where the disease spreads faster as many contagious individuals are unaware of being infectious. Technically, this also facilitates using the parameter estimates from these epidemiological studies.

Management studies that explicitly account for asymptomatic cases are rare. For instance, Kaplan (2020) considers the standard SIR model and Acemoglu et al. (2021) consider connected replicas of SIR, while both briefly mention SEIR. Birge et al. (2020) and Camelo et al. (2021) start with SEIR and, respectively, split infected individuals into clinical/subclinical and confirmed/unconfirmed through testing, the former being further divided based on symptoms. These approaches are similar to ours in spirit but suggest that eventually symptomatic individuals may initially be asymptomatic, which necessitates adding the “A”-type compartments.

Second, we split each S-E-A-I-R compartment into four categories, with the ICU compartment split in two. Together with a compartment for people who died from the disease, they add up to a total of 23 compartments. The four subcategories correspond to the so-called “confusion matrix” of the machine learning risk prediction model (see Table 2): (i) true positives, who would experience severe symptoms upon infection needing an ICU bed, and classified as high-risk and hence confined; (ii) false negatives, who would experience severe symptoms upon infection needing an ICU bed, but classified low-risk and hence released; (iii) false positives, who would experience only mild symptoms upon infection not needing an ICU bed, but classified high-risk and hence confined; and (iv) true negatives, who would experience only mild symptoms upon infection not needing an ICU bed, and classified low-risk and hence released. How each individual falls into one of these four groups is determined endogenously by our model, as we explain in the next section.

Because this notation is critically important going forward, we reiterate that the policy we study confines those who are predicted to be high risk and releases those who are predicted to be low risk. As with any model, our (assumed) prediction model makes mistakes, thus both the confined and released groups contain a mix of actually severe- and actually mild-symptom individuals. The sub- and superscripts \( j = \{s, m\} \), for (actually) “severe” versus “mild,” and \( i = \{c, r\} \), for “confined” (i.e., predicted severe) versus “released” (i.e., predicted mild), designate the subcategories in each compartment. For example, \( S^{(c)} \) refers to susceptible individuals who...
TABLE 1 Epidemiological parameters used

| Symbol | Description | Value(s) | Reference |
|--------|-------------|----------|-----------|
| $\beta$ | Transmission rate | Computed | Salje et al. (2020) |
| $R_0$ | Basic reproduction number | 2.9 | Salje et al. (2020) |
| $\varepsilon$ | Waiting rate to viral shedding | 1/3.7 day$^{-1}$ | Di Domenico et al. (2020) |
| $\sigma$ | Waiting rate to symptom onset | 1/1.5 day$^{-1}$ | Di Domenico et al. (2020) |
| $\eta$ | Waiting rate from symptom onset to ICU | 1/7 day$^{-1}$ | Salje et al. (2020) |
| $\gamma_m$ | Recovery rate from mild symptoms | 1/2.3 day$^{-1}$ | Di Domenico et al. (2020) |
| $\gamma_s$ | Recovery rate for people in ICU | 1/17 day$^{-1}$ | Salje et al. (2020) |
| $\alpha$ | Mortality rate for people in ICU | 1/11.7 day$^{-1}$ | Salje et al. (2020) |

are released but will get severe symptoms when infected, $S^{(r)}$ refers to all released, and so on.

We use $\rho \in [0, 1]$ as the control parameter of our policy, denoting the proportion of individuals who should be released, that is, classified in the low-risk group (correctly or incorrectly) and therefore subject to low isolation restrictions. We initially consider a single-release policy, optimizing over a scalar $\rho$, and then extend the analyses to the optimal control problem with multiple releases, optimizing over a vector $\{\rho_0, \rho_1, \ldots\}$ released at times $\{0, t_1, \ldots\}$.

We capture the impact of the differentiated isolation restrictions on people’s behavior using two behavioral parameters: $\delta_r$, for the group with low isolation restrictions, that is, released, and $\delta_c$, for the group with high isolation restrictions, that is, confined; $0 \leq \delta_r < \delta_c \leq 1$. These parameters capture a level of “protection” and aggregate several factors, such as respiratory and hand hygiene and how much a person has lowered the number of exits from home and social interactions.

Note that how individuals in group $i = \{r, c\}$ reduce their chances of contracting the disease depends not only on $\delta_r$ and $\delta_c$ but also on the proportion of people in each group, $\rho$ and $1 - \rho$. The so-called “contact rates,” $c_r$ and $c_c$, for the released and confined groups, important parameters in “standard” epidemiological models, satisfy $1 - c_r = (1 - \delta_r) \times ((1 - \delta_c) \rho + (1 - \delta_c)(1 - \rho))$, $i \in \{r, c\}$. These parameters have been used in the literature modeling COVID lockdowns, for example, Di Domenico et al. (2020) and Djidjou-Demasse et al. (2020). Our approach is fully aligned with those, as we endogenize contact rates per the preceding equation.

Figure 1 presents the simplified schematic of our SEAIR model, which corresponds to the following set of ordinary differential equations (ODEs), where all parameters are defined in Table 1:

\[
\dot{S}^{(r)} = -(1 - \delta_r)\beta^{(r)} S^{(r)}, \tag{ODE}
\]

\[
\dot{E}^{(r)} = (1 - \delta_r)\beta^{(r)} S^{(r)} - \varepsilon E^{(r)}, \tag{1}
\]

\[
\dot{S}^{(c)} = -(1 - \delta_c)\beta^{(c)} S^{(c)}, \tag{2}
\]

\[
\dot{E}^{(c)} = (1 - \delta_c)\beta^{(c)} S^{(c)} - \varepsilon E^{(c)}, \tag{3}
\]

\[
\dot{A} = \varepsilon E - \sigma A, \tag{4}
\]

\[
\dot{I}_m = \sigma A_m - \gamma_m I_m, \tag{5}
\]

\[
\dot{I}_s = \sigma A_s - \eta I_s, \tag{6}
\]

\[
\dot{U} = \eta I_s - (\gamma_s + \alpha) U, \tag{7}
\]

\[
\dot{R} = \gamma_m I_m + \gamma_s I_s, \tag{8}
\]

\[
\dot{D} = \alpha U. \tag{9}
\]

Here, the effective number of contagious people is $I^e = (1 - \delta_r)(A^{(r)} + I^{(r)}) + (1 - \delta_c)(A^{(c)} + I^{(c)})$. As is usual for such models, we consider the so-called basic reproduction number $R_0$ as measured prior to lockdown. In our model, the situation prior to lockdown corresponds to taking $\delta_r = \delta_c = 0$. Then, the transmission rate $\beta$ relates to $R_0$ and the other parameters through $\beta = R_0 \frac{1}{N_0} \times \frac{1}{p(\sigma^{-1} + \omega^{-1}) + (1 - p)(\sigma^{-1} + \eta^{-1})}$. This relation may be derived by linear stability analysis, as in Djidjou-Demasse et al. (2020), using the so-called next generation matrix method (Diekmann et al., 2010).

Remark 1. The set of ODEs and the schematic in Figure 1 present a simplified version of our model, reduced from the full set of 23 equations for ease of exposition; for example, even though the $A$ compartment is depicted as a single node in Figure 1 and is presented as a scalar in Equation (5), it is actually a four-vector $\{A^i\}$, where $i = \{c, r\}$, $j = \{s, m\}$. The full set of ODEs and the algorithmic details of their solution are presented in the “code appendix”, Boulant et al. (2020).

2.2 Risk prediction/classification model

Our model necessitates identifying individuals at the highest risk of severity and correspondingly advising them to remain in strict isolation, while relaxing isolation restrictions.
TABLE 2 Confusion matrix of the risk model. \( f_s, f_m \) denote class-conditional predictive distributions; \( T \) is the classification threshold, \( p \) is the proportion of people with mild symptoms in the population, and \( \rho \) is the proportion of the released population—the decision variable in our model.

| Actual, \( s \)          | Actual, \( m \)          | Total |
|-------------------------|-------------------------|-------|
| Model, \( s \rightarrow \text{confine} \) | False positives = \((1 - p) \times \int_0^{T^*} f_s(\omega) \, d\omega\) | \(1 - \rho\) |
| False negatives = \((1 - p) \times \int_0^{T^*} f_m(\omega) \, d\omega\) | \(p\) |
| Model, \( m \rightarrow \text{release} \) | True positives = \(p \times \int_0^{T^*} f_m(\omega) \, d\omega\) | \(\rho\) |
| Total:                   | 1 - \(p\)               | \(p\) |

for lower risk individuals. Such identification is done in two steps, following the common data science and machine learning approach. For step 1, a risk “score” is obtained for each individual, as in Bertsimas (2020), using a logistic regression, random forest, gradient boosting, or the like model. A standard metric to assess the discriminating power of such models is the area under the curve (AUC) of the receiver operating characteristic (ROC) curve (Fawcett, 2006). We chose this metric because it is widely used for biomedical applications involving screening populations with some score function (Lasko et al., 2005). The score orders the members of the population from low to high risk and the AUC is often referred to as the rate of concordant pairs (i.e., the fraction of pairs that are correctly compared by the score with respect to their actual risk status). For step 2, individuals with risk scores above a certain threshold, \( T \), are classified as high risk and are confined, while the rest are classified as low risk and are released. \( T \) is determined endogenously so the proportion of the released population equals \( \rho \), the decision variable in our model.

As mentioned in the introduction, we assume such a risk model would exist in practice. Training such models requires access to nontrivial personalized data and is outside the scope of this paper. Instead, our goal is to evaluate the efficacy of personalized pandemic management given such a model. Therefore, we utilize hypothetical risk models with AUCs that bracket existing COVID-19 models in the literature, for example, Bertsimas (2020).

To create a hypothetical risk model, let \( f_s \) and \( f_m \) denote the so-called “predictive distributions”—the PDFs of the risk scores for people with severe and mild symptoms, respectively, as predicted by the model. Together with the threshold, \( T \), and the proportion of the population with mild symptoms in the population, \( p \), these \( f_s, f_m \) define the model’s confusion matrix, per Table 2. See Clémençon and Vayatis (2007) for other performance measures derived in the context of a control parameter \( \rho \) applied to a personalized risk model.

As is evident from Table 2, given \( \rho, p \) and “the model,” that is, \( f_s, f_m \), and assuming the risk scores are between 0 and 1, the corresponding classification threshold \( T^* \) should be selected such that:

\[
(1 - p) \times \int_0^{T^*} f_s(\omega) \, d\omega + p \times \int_0^{T^*} f_m(\omega) \, d\omega = \rho. \tag{10}
\]

Let \( q^{FP} \) and \( q^{FN} \) denote the model’s false positive and false negative error rates, respectively; that is, \( q^{FP} = \int_0^{T^*} f_m(\omega) \, d\omega \) and \( q^{FN} = \int_0^{T^*} f_s(\omega) \, d\omega \). For notational convenience, we omit the dependency of \( q_s \) on \( T^* \) and, through that, on \( \rho \). Then (10) is equivalent to:

\[
(1 - p) \times q^{FN} + p \times (1 - q^{FP}) = \rho, \tag{11}
\]

which highlights the key relationship of (any) classification model that we exploit. Selecting a small \( \rho \) results in a small \( T^* \), thus a small \( q^{FN} \) as well. Relatively few people will be released but very few of the released would, by mistake, develop severe symptoms. Increasing \( \rho \) would not only increase \( T^* \), releasing more people, but it also would increase \( q^{FN} \), exerting a disproportionate impact on people who would require an ICU. An increase in \( q^{FN} \) will be smaller for a higher quality (higher AUC) model than for a lower quality one. This is because \( AUC = \int_0^1 (1 - q^{FN}) \, dq^{FP} \), thus a higher AUC implies, ceteris paribus, lower \( q^{FN} \) at a given \( T \).

Another important observation from Table 2 is that the released individuals consist of two groups: true negatives and false negatives. The latter will experience severe symptoms upon infection, requiring ICU beds. They are not, however, the only group to require ICU beds; because the confinement of the nonreleased is imperfect (\( \delta < 1 \)), some of the true positives will require ACU as well. Setting a smaller \( \rho \), decreases the number of false negatives but increases the number of true positives, leading to a nontrivial relationship between selecting \( \rho \), the resultant threshold \( T^* \), and ICU demand. This relationship depends on the model’s quality (AUC).

The goal is to select \( \rho \) so that, given the model’s quality, as few people as possible are confined but ICU capacity is not exceeded due to model errors or imperfect confinement.

2.3 Connecting risk and SEIR models

The risk model connects with the SEAIR model as follows: For \( Q \in \{S, E, A, I, U, R\} \) and a policy \( \rho \), we rescale the initial risk-independent epidemiologic conditions \( Q_0 \) to account for the distribution of people in the four groups:

\[
Q_s^{(0)}(0) \leftarrow (1 - \rho) q^{FN} Q_0, \quad Q_s^{(c)}(0) \leftarrow (1 - p)(1 - q^{FN}) Q_0, \tag{12}
\]

\[
Q_u^{(0)}(0) \leftarrow p (1 - q^{FP}) Q_0, \quad Q_u^{(c)}(0) \leftarrow p q^{FP} Q_0, \tag{13}
\]

This results in the starting “day 0” conditions for the 23 compartments in the extended SEAIR model, which all depend on \( \rho \).
As we investigate policies changing over time, we also update the number of people in each compartment when the decision maker increases $\rho$ from some value $\rho_{\text{old}}$ to $\rho_{\text{new}}$, with corresponding false positive and false negative rates $q_{\text{old}}^{FP}, q_{\text{old}}^{FN}$ and $q_{\text{new}}^{FP}, q_{\text{new}}^{FN}$, respectively:

$$Q_m^{(i)} \leftarrow Q_m^{(i)} + \frac{q_{\text{old}}^{FP} - q_{\text{new}}^{FP}}{q_{\text{old}}^{FP}} Q_m^{(c)},$$

$$Q_m^{(c)} \leftarrow Q_m^{(c)} - \frac{q_{\text{old}}^{FP} - q_{\text{new}}^{FP}}{q_{\text{old}}^{FP}} Q_m^{(c)},$$

$$Q_s^{(i)} \leftarrow Q_s^{(i)} + \frac{q_{\text{new}}^{FN} - q_{\text{old}}^{FN}}{1 - q_{\text{old}}^{FN}} Q_s^{(c)},$$

$$Q_s^{(c)} \leftarrow Q_s^{(c)} - \frac{q_{\text{new}}^{FN} - q_{\text{old}}^{FN}}{1 - q_{\text{old}}^{FN}} Q_s^{(c)}.$$  (UPD)

### 3 | ESTIMATION OF PARAMETERS AND OPERATIONALIZATION OF SIMULATIONS

We simulate the progression of the combined epidemic and risk model discussed in the previous section in two scenarios: setting “day 0” on March 17, 2020, France’s first day of national lockdown, or on May 11, 2020, the beginning of the lockdown exit. Doing so requires estimating several parameters, listed in Tables 1 and 3. Some of these parameters are inferred from the literature, while others are estimated from data, as we discuss next.

#### 3.1 | Risk-model parameters

The class-conditional predictive distributions are modeled as Beta distributions: $f_s \sim \text{Beta}(a_s, b_s)$ and $f_m \sim \text{Beta}(a_m, b_m)$. In simulated scenarios, the no model refers to $a_m = b_m = a_s = b_s = 1$. Otherwise, we fix $b_s = a_m = 2$ and vary $a_s = b_m$. The low AUC model refers to $a_s = b_m = 3$ (AUC $\sim 75\%$) and the high AUC model refers to $a_s = b_m = 5$ (AUC $\sim 96\%$). These parameters are selected so as to bracket the “low” (AUC $\sim 82\%$) and “high” (AUC $\sim 93\%$) models from Bertsimas (2020). For sensitivity analyses, we explore the range from $a_s = b_m = 2.5$ (AUC $\sim 65\%$) to $a_s = b_m = 6.5$ (AUC $\sim 99\%$), further bracketing the range of models that could possibly be available in practice for a disease like COVID-19.

#### 3.2 | Epidemic model parameters

We estimate the joint distribution of the model parameters by comparing the predictions from our model (at the given parameter values) to the actual data for ICU occupancy obtained from the official portal of the French government: https://dashboard.covid19.data.gouv.fr.

Two general approaches exist for doing so. With the “standard” statistical approach, one splits the data into training and testing sets (sequentially, given the time series nature of the data), learns the model parameters on the training data, and evaluates the predictive accuracy on the testing data. The error structure of the learned model, however, is likely highly nontrivial as the errors are not independent over time. For example, if one SIR-like curve is higher than another early in a time horizon, it must get lower at a later time as fewer susceptible individuals will remain. As a result, this approach could be used to learn the best point estimates but not their joint distribution.

A Bayesian approach can overcome this challenge but it has a noteworthy complication: The likelihood function for the resultant prediction errors is also unknown. To deal with this issue, we utilized an approximate Bayesian computation (ABC) method (Marjoram et al., 2003), which has been specifically designed for such situations. The ABC method was implemented with the root mean standard error as a distance function (Britton, 2010), with a maximum error set at 1000 ICU beds, which corresponds to the so-called “acceptance rate” of the ABC analyses of about 10%.

Parameters for the initial conditions $S_0$, $E_0$, $A_0$, $I_0$, $U_0$, and $R_0$ depend on the investigated scenario’s “day 0.” The initial number of susceptible individuals, $S_0$, is computed as

| Symbol | Description | Value(s) | Reference |
|--------|-------------|----------|-----------|
| $N_0$  | Total initial number of people in the population | $6.7 \times 10^7$ | population of France |
| $S_0$  | Total initial number of susceptible people in the population | | Computed |
| $E_0$  | Total initial number of exposed people in the population | | Case-dependent |
| $A_0$  | Total initial number of asymptomatic people in the population | | Case-dependent |
| $I_0$  | Total initial number of infected people in the population | | Case-dependent |
| $U_0$  | Total initial number of people in ICU | | Case-dependent |
| $R_0$  | Total initial number of immune people in the population | | Case-dependent |
| $I_{\text{max}}$ | Hospital capacity for COVID-19 ICU beds | 7250 | assumed |
| $p$    | Proportion with mild symptoms (prior with 90% confidence interval) | 0.9932 [0.9891 – 0.9961] | Salje et al. (2020)/estimated |

### Table 3 | Simulation parameters used

- $N_0$: Total initial number of people in the population
- $S_0$: Total initial number of susceptible people in the population
- $E_0$: Total initial number of exposed people in the population
- $A_0$: Total initial number of asymptomatic people in the population
- $I_0$: Total initial number of infected people in the population
- $U_0$: Total initial number of people in ICU
- $R_0$: Total initial number of immune people in the population
- $I_{\text{max}}$: Hospital capacity for COVID-19 ICU beds
- $p$: Proportion with mild symptoms (prior with 90% confidence interval)
- $N_0$: Total initial number of people in the population
- $S_0$: Total initial number of susceptible people in the population
- $E_0$: Total initial number of exposed people in the population
- $A_0$: Total initial number of asymptomatic people in the population
- $I_0$: Total initial number of infected people in the population
- $U_0$: Total initial number of people in ICU
- $R_0$: Total initial number of immune people in the population
- $I_{\text{max}}$: Hospital capacity for COVID-19 ICU beds
- $p$: Proportion with mild symptoms (prior with 90% confidence interval)
The number of samples from the prior distributions was set at \( n = 10,000 \) (resp. \( 100,000 \) for robustness when computing means). This led to around 1000 (resp. \( 10,000 \)) posterior samples as the acceptance rate was at \( \sim 10\% \). The mean posterior values were found to be \( p \sim 0.993 \) and \( E_0 + A_0 + I_0 \sim 1200000 \) (for March 17). The mean posterior value for \( c \) was found to be \( c \sim 69.2\% \), which is consistent with Salje et al. (2020). Reiterating prior discussion, the latter value is unused in our numerical experiments (because we investigate scenarios with differentiated isolation policies) but it is necessary to estimate the joint posterior of \( p \) and \( E_0 + A_0 + I_0 \) from the data about the nondifferentiated policy.

Remark 2. Although the obtained mean value of \( p \sim 0.993 \) may seem surprisingly high, leaving only \( \sim 0.7\% \) probability of dying from a COVID-19 infection, we emphasize that our work does not take into account deaths in nursing homes because those individuals are always confined. Also, in Salje et al. (2020), data show that about 15\% of deaths occur during the first day of hospitalization—in other words, those patients are never admitted to the ICU and are not included in an estimate for \( p \). That said, we acknowledge that this parameter is critical, hence we also performed sensitivity analyses with \( p = 0.98 \) in Appendix B in the Supporting Information.

3.3 Simulations with 95\% confidence intervals

In all figures showing the evolution of the number of people in the ICU (see Figure 2), the initial condition \( E_0 + A_0 + I_0 \) and the proportion of people \( p \) not requiring ICU admission were sampled according to their posterior distribution. The mean curve of Figure 2 was obtained by taking the average of all the sampled curves, and the 95\% confidence intervals were derived by removing the 2.5\% and 2.5\% upper and lower values for the computed number of ICU beds at each time.

3.4 Simulations with grid searches

As some numerical experiments (see Figures 3–5 and Table 4) require grid searches, we did not sample according to the posterior distribution for each scenario. Instead, we computed mean values in order to ease the computational burden. To account for uncertainty, we reduced the number of available ICU beds by the average width of the 95\% confidence interval from the corresponding simulations. We then operationalized the simulations as follows:

- For “day 0” of March 17, 2020: The initial number of utilized ICU beds is known, \( U_0 \sim 700 \), and the estimate for the total number \( E_0 + A_0 + I_0 \sim 1.5 \times 10^6 \) is available from Salje et al. (2020). Therefore, we took the average over the posterior and obtained \( E_0 + A_0 + I_0 \sim 12,000,000 \), resulting in \( R_0 \sim 3,000,000 \) people. Similarly, taking the mean along posterior samples resulted in the estimate of \( p \sim 0.993 \). The ODE system was then integrated up until 200 days beyond March 17, 2020.
- For “day 0” of May 11, 2020: Sampling according to the posterior for \( p \) and \( E_0 + A_0 + I_0 \) and integrating the ODE system from March 17 to May 11, we obtained a sample of initial conditions for May 11, of which we took the averages to obtain estimates for all parameters. The ODE system was then integrated up until 200 days past May 11, 2020.
RESULTS

We present the results of our simulations in four steps. First, we consider a partial exit problem with a single release. Second, we explore the sensitivity of the single release problem. Third, we consider the complete exit problem over multiple release epochs. Finally, we discuss its sensitivity, including the impact of “lockdown fatigue” (Goldstein et al., 2021).

4.1 Partial policies with a single release

Figure 2 displays the number of individuals requiring an ICU bed w.r.t. time \( t \). The March 17 scenario is in the left column and the May 11 scenario is in the right. Two risk models are considered: a “high” AUC \( \sim 95.99\% \) (top row) and a low AUC \( \sim 75.71\% \) (bottom row), bracketing the performance of initial risk models developed for COVID-19 (Bertsimas, 2020).

In each plot, \( \rho \) represents the maximal percentage of the population that can be released (i.e., subject to lighter restrictions), which is assumed to correspond to \( \delta_r = 0.1 \) in such a way that the 95% confidence interval of the number of individuals requiring an ICU bed when using the risk prediction model (green and orange curves) remains below the number of available ICU beds, 7250. In these first simulations, the rest of the population is confined with more restrictions, \( \delta_c = 0.9 \). Finally, the red curves show the number of individuals requiring an ICU bed w.r.t. time if the same \( \rho \) of population is released, but selected at random without any risk prediction model.

4.1.1 Initial lockdown

Figure 2a shows that on March 17, 2020, a high-AUC model (green curve) allows for \( \rho = 62\% \). That is, only 38% of the population should be ever confined. In France, which has a
FIGURE 3  Difference in maximal percentage of released people without exceeding ICU capacity (7250), compared to the case of not using a risk prediction model, plotted as a function of the AUC of a risk prediction model and the protection level $\delta_c$ for confined people. $p \sim 0.993$ for all figures. Appendix B in the Supporting Information presents similar figures with $p$ decreased to 0.98, and ICU capacity increased to 15,000 [Color figure can be viewed at wileyonlinelibrary.com]

FIGURE 4  Examples of gradual relaxation of isolation restrictions. High AUC model (green) and no model (red) with $\rho = [0.65, 0.7, 0.9, 1]$, $t = [60, 90, 180, 600]$, and no model (blue) with $\rho = [0.5, 0.55, 0.6, 1]$, $t = [30, 120, 510, 600]$. Vectors $t = [t_1, t_2, ...]$ give the release schedules $\rho = [\rho_1, \rho_2, ...]$ as follows: $\rho_1$, 100% of the population is released on day 0, then $(\rho_2 - \rho_1)$. 100% are released on day $t_1$, and so forth [Color figure can be viewed at wileyonlinelibrary.com]
population of 67 million, this corresponds to 25 million people. Critically, the remaining 42 million (62%) should never have been confined at all. Figure 2c shows the same for the low-AUC model (orange curve), which enables $\rho = 53\%$, or some $\sim 6$ million more people in isolation. Perhaps more importantly, without a model, for March 17, $\rho = 45\%$—a 17% and 8% difference, respectively, or 5 to 11 million people. In other words, the initial lockdown could have been managed much better if the government had the ability to first train and then utilize a severity risk model at scale.

4.1.2 Lockdown exit

Figures 2b,d also present the ICU occupancy over time, but for lockdown exit strategies as of May 11. Importantly, unlike what France did on May 11, it is not optimal to release the entire population. Even with a high-AUC model, some 30% should still remain in isolation (40% with the low-AUC model), otherwise ICU capacity would be exceeded in the second wave. Of course, our analyses assume that the released population is subject to $\delta_r = 0.1$ restrictions, while the policy actually implemented in France was aiming for a higher reduction in contact rates. That said, the French government in November 2020 implemented a second lockdown precisely to avoid overwhelming ICU admissions due to the second COVID wave, exactly as we predicted would happen if less than 30%–40% of the population remained in isolation.

These results, however, are for a single release, and a natural question is what one should do with those 30%–40%.
which we address in Section 4.3 after discussing the sensitivity analyses.

4.2 Sensitivity analyses of the single release policy

Figure 3 displays the results for the March 17 and May 11 scenarios showing the difference between the maximal percentage of people who may be released without exceeding ICU capacity with a risk model, relative to the same percentage, but without a risk model. Sensitivity is tested with respect to the discrimination performance of the risk prediction models (AUC) and the degree of isolation of the confined population ($\delta_c$). We also alter the degree of isolation for the released population ($\delta_r$) across different plots.

As expected, the higher the discrimination of the prediction model, the bigger the difference. However, the degree of isolation has a different impact depending on who is considered: For the confined population, the stricter the isolation (the higher $\delta_c$ is), the larger the impact of the risk prediction model. But for the released population ($\delta_r = 0.1$ or 0.2 in Figure 3), the results are more intricate. It is often better to isolate individuals more strictly, except when the risk prediction model is of very high quality and the confined people are in very strict isolation. In those situations, the optimal $\rho$ is large enough to achieve “herd immunity” ($1 - \frac{1}{R_0} \approx 65\%$), which can be achieved faster if the released population is less protected. The main implication of these analyses is that it is important to both assume in models and encourage in practice stricter isolation practices for the high-risk population—for example, by focusing distribution of PPE and other resources, strictly isolating nursing homes, and so forth. This not only better protects the high-risk group but also allows for a faster and more efficient exit from the pandemic for the rest.

4.3 Complete exit with multiple release epochs, a.k.a. gradual deconfinement

Next, we explore gradual exit strategies, where we optimize both over time and proportion of release with a given fixed number of policy updates $N$ while not exceeding a given ICU constraint $M > 0$. In other words, we set a time-optimal control problem, where the variables are the times $t_1 \leq \ldots \leq t_{N-1} \leq t_N$ and proportions of people released $\rho_1 \leq \ldots \leq \rho_N$ with $\rho_N = 1$, and we aim at minimizing $t_N$, that is, the moment at which all individuals are released and lockdown is over. The resulting time-optimal control problem is:

$$\begin{align*}
\min_{0 = t_1 \leq \ldots \leq t_{N-1} \leq t_N \leq 200} & \quad t_N \\
\text{s.t.} & \quad \text{(ODE) and (UPD)},
\end{align*}$$

where $U(t)$ is given by (8) and (ODE) and (UPD) are the systems of equations governing the evolution of the ODE system between the releases and the updating, per Section 2.

To consider practical and realistic scenarios, we solved the resultant optimization dynamic program, allowing releases every 30 days at multiples of 5% of the population, while ensuring the maximum number of utilized ICU beds did not exceed $M = 5250$. We found that implementing the full-blowed confidence interval analyses, as in Figure 2, was computationally intractable in the dynamic program setting. However, by observing that the maximum confidence range in Figure 2 was $\approx 2000$ beds, we reduced ICU capacity from the “base-case” of 7250 to $M = 5250$ to account for uncertainty in ICU demand. For a more realistic initialization, we first solved for the single-release $\rho_{max}$, as per Section 4.1 and Figure 2, then fixed $\rho_1 = \rho_{max}$. Similarly, for a realistic termination, we ran the ODE model for 200 days after $t_N$ to ensure the ICU constraint is not broken after the entire population is released.

Table 4 shows the minimal number of months, that is, the optimal solution to problem OCP, to release the entire population for different scenarios (no model, low-AUC model, high-AUC model), while keeping all other parameters constant for three different values of $\delta_c$. We considered only gradual releases in three or four epochs; the ICU system was overwhelmed when using only two epochs for most simulations. The main insight is that using a no-risk model would require more than a year in all scenarios, while an exit with risk-based models would lead to relaxing restrictions for the entire population in as quickly as 6 months.

Figure 4 shows similar optimal policies for May 11 corresponding to the four epochs from Table 4, and assuming $\delta_c = 0.9$ as in Figure 2. The insights complement those for single release policies: With risk-prediction models, a smaller percentage of the population may need to be confined. Consequently, one also could reach the moment when isolation measures could be lifted sooner.

For example, using the high-AUC model and without exceeding ICU capacity at any point, 65% of the lowest risk population could be released on May 11 (“day 0”), followed by another 5% on July 10 (“day 60”), another 20% on August 9 (“day 90”), and the final 10% on November 7 (“day 180”). Resultant ICU demand is shown as a green line in Figure 4a.

Implementing the same exit schedule without a risk model would lead to ICU demand of nearly 20,000 beds (red line). In contrast, a capacity-abiding exit strategy without a model (blue line) would require 17 months to reach full deconfinement. Only 50% of the population could be in low isolation on “day 0” (May 11), another 5% on “day 30” (June 10), additional 5% on “day 120” (September 8), and the last 40% only on “day 510” (October 3, 2021), or 11 months later than the similar risk-model–based strategy. Such an extended isolation also would apply to many more people: 10% with the model versus 40% without. For France, this means an additional $\approx 20$ million people in isolation for the additional 11 months.
For both scenarios, Figure 4b shows the percentage of the population that becomes immune over time. Because the model-based policy releases a larger portion of the low-risk population and does so faster, herd immunity is approached, allowing for the ultimate protection against the disease. In contrast, herd immunity is not achieved by a policy without the risk model: The disease is suppressed but could explode again.

4.4 | Sensitivity of multiple release policy to lockdown fatigue

The optimal exit policies reported above consist of several exit epochs at which a portion of the population is released from isolation, while the rest remain confined. Goldstein et al. (2021) argue that such prolonged restrictions represent a psychosociological burden on isolated individuals and could result in “isolation fatigue,” thus diminishing the degree of compliance. Our model can capture this by letting \( \delta_c \) depend on time, that is, by defining \( \delta_c(t) = [\delta_{c1}(t), \delta_{c2}(t), \delta_{c3}(t), \delta_{c4}(t)] \).

We explore two fatigue scenarios: “Low,” where the confined individuals increase their contact rates twofold by the final release epoch, and “high,” where the increase is threefold. For simplicity, we assume \( \delta_c(t) \) changes linearly over time. That is, the two scenarios correspond to \( \delta_c(t) = [0.9, 0.85, 0.8, 0.8] \) and \( \delta_c(t) = [0.9, 0.8, 0.7, 0.7] \), respectively.

Figure 5 illustrates two ways in which fatigue impacts our results. First, the black lines show the simulated ICU demand from applying the optimal policy without fatigue (per previous subsection) in a situation where fatigue is present. Clearly, ignoring fatigue could result in substantially exceeding ICU capacity. A policy that ignores fatigue is overly aggressive in relaxing the isolation restrictions as it anticipates the remaining population to be strictly confined. When that is not the case due to fatigue, the policy should be less aggressive early on, which will slow down the build-up of population immunity, also delaying subsequent release epochs.

Second, the green line shows ICU demand under the optimal policy that takes into account fatigue (and, critically, anticipated fatigue). Contrasting the waves in the two policies illustrates the differences between the optimal policies. For the benchmark, recall that a policy without fatigue releases 65% of population immediately, then releases another 5% on day 60, another 20% on day 90, and the remaining 10% on day 180, that is, completes the exit in 6 months. The optimal policy in the low-fatigue scenario also releases 65% immediately and completes the exit in 6 months but keeps an additional 10% of people—twice as many—in isolation after day 90, and 15% for days 120–180. The optimal policy in the high-fatigue scenario exacerbates the differences further, keeping an additional 20% of people—three times as many—in isolation beyond day 90, with release completed on day 210 as opposed to 180, and with 15% in isolation for days 120–210.

The driver of the differences is the same in both scenarios. The optimal policy aims at building enough population immunity so that when the highest risk individuals, many of whom would require ICU treatment upon infection, are released, the infection spreads slowly, allowing the last wave to stretch over an extended period of time, for example, see Figure 4a. Noncompliance due to fatigue restricts the ability to do so, shifting the release further in time, reducing the fraction of release individuals at each epoch, and enlarge the gaps between subsequent epochs.

Generally, the effect of noncompliance with isolation restrictions (due to fatigue or other reasons) is similar to that of lowering the quality of the risk model. Both restrict the ability to target individuals differently given their respective risk factors, thus emphasizing the two main messages of our paper: Successful targeted interventions require (i) identifying high-risk individuals and (ii) treating them differently.

5 | CONCLUSIONS

Data-driven prediction models, which made large impacts in many areas over the past decades, can help personalized policies for managing epidemic outbreaks. We detailed how prediction models for symptom severity upon infection could be used in epidemic simulations to study the effect of nonpharmaceutical policies, particularly isolation restrictions, during an outbreak. Our core goal is to explore the tension between the prediction model quality and epidemic containment performance. We used COVID-19 data from France as of spring 2020 as an example and provided sensitivity analyses to understand how different parameters could impact pandemic isolation and exit policies.

Simulations indicated that considering differential relaxation of isolation restrictions depending on predicted severity risk can decrease the immediate percentage of the population in France under stricter isolation by \( \sim 5\% - 20\% \) relative to not using such risk predictions. Doing so also would speed up a complete exit by several months, directly impacting the lives of millions of people.

We made our simulation engine available to a broad, non-technical audience via an interactive demo that is available at https://ipolcore.ipol.im/demo/clientApp/demo.html?id=305 and is illustrated in Figure 6. The demo is prepopulated with the model parameters for France that we used in this paper, but one can input parameters for other countries or regions and experiment with the envisioned policies. We presented our paper to the pandemic management task forces of multiple G7 countries and successfully used this tool to facilitate the dissemination of our results and raise awareness of the promise of personalized data-driven policies for pandemic management.

Sensitivity analyses showed that the qualitative insights from our simulations are robust to changes in risk prediction accuracy, percentage of severe-if-infected cases in the population, availability of resources (e.g., ICUs), and social distancing. Benefits increased when risk prediction
accuracy increased, percentage of severe-if-infected cases in the population decreased, availability of resources (e.g., ICUs) increased, and the isolation of high-risk individual increased.

5.1 Limitations and opportunities for future research

Our study is not without limitations, many of which provide opportunities for future research.

First and foremost, all results in this paper use hypothetical risk prediction models based on discrimination ranges in line with early indications from the literature, for example, Bertsimas (2020). To operationalize personalized isolation and exit policies based on risk predictions, governments need to develop policies and invest in infrastructure to enable building and using such models at scale. Critically, this involves both training models on early cases’ data pooled from the epicenter of the pandemic and using models to score entire populations. Currently, the latter presents a bigger problem because even countries with well-organized, single-payer healthcare systems mostly capture data about “sick” individuals (i.e., those who use the healthcare systems) and often lack accurate and recent data for the healthy population. As we discuss in the companion article, Evgeniou et al. (2020), addressing this challenge, involves policy matters such as standardization, privacy, and localization, as well as technical matters such as transfer and federated learning. All of the above offer opportunities for future research.

Second, epidemic models and their resultant conclusions rely on a number of parameters (e.g., virus incubation and recovery times, basic reproduction number $R_0$) that are uncertain and evolve dynamically, particularly as new virus variants emerge (Alban et al., 2020). The resultant policies are therefore contingent: Observing an ICU demand that is closer to an upper boundary of the confidence interval may require the next wave to be delayed or involve a smaller release percentage than our current simulations, built from day 0, suggest. Building models that specifically account for a potential shift in these parameters is an interesting future research direction.

Third, practical policy decisions for a given pandemic require careful context-specific robustness analysis of, for example, the benefit of combining isolation restrictions with other policies such as test-based ones, for example, Wang et al. (2020) and Petherick (2020), or vaccination-based ones, for example, Sonabend et al. (2021). Designing such combined policies is another fruitful research direction.

Finally, risk-predictions–based policies using epidemic simulations should be developed taking into account behavioral aspects that may help or hurt model predictions and policy actions: ethical issues, fear, widespread noncompliance to isolation measures, and the like. Behavioral operation studies could inform model building and guide lockdown policy implementation in isolation or in conjunction with other mechanisms. For example, governments in many countries have encouraged vaccination by implementing a de facto differential isolation policy for nonvaccinated individuals.

In conclusion, our simulations show that combining prediction models using data science and machine learning principles with epidemiological models may improve outbreak management policies. Governments, therefore, should make appropriate investments in the data infrastructure necessary to implement such models at scale and consider their predictions when developing pandemic isolation (confinement) and exit (deconfinement) policies.
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ENDNOTES
1 The individual prediction is not critical for our analyses; an alternative prediction target could, for example, be the risk that someone in a household develops severe symptoms. See Appendix A in the Supporting Information.
2 Given that the dependency of the COVID-19 clinical outcomes on age attracted enormous media attention, we note that policies based only on age may be unnecessarily simplistic. For example, ~10% (source: statista.com) of the French population is older than 75, yet only ~1% of the entire population will require an ICU bed if infected. It is thus not practical to implement differential isolation and exit policies based just on age. Instead one may need to only consider models that capture more nuanced and accurate patterns using multiple features, not just age.
3 Such long and massive lockdowns in the absence of a reasonable risk model are consistent with the literature. For example, Acemoglu et al. (2021) propose isolating all seniors for 18 months, and even that hinges on the assumption that a vaccine will be developed by then. Our analyses do not rely on the availability of vaccines.
4 Disclaimer: We do not advocate for herd immunity but building herd immunity may be optimal when high-risk individuals are well-protected and the risk model is of a sufficient quality to identify such individuals with a high-enough accuracy. Otherwise, a herd immunity strategy that does not overwhelm the medical system is impossible.
5 By definition, only severe cases go to the ICU but they can come from two categories: those classified by the model as high risk (true positives) and those classified as low risk (false negatives).
6 The symmetric ROC models we considered are not the only possible approach. While working on this paper, we explored multiple nonsymmetric settings and observed no qualitative impact on the simulation results.

REFERENCES
Acemoglu, D., Chernozhukov, V., Werning, I., & Whinston, M. D. (2021). Optimal targeted lockdowns in a multigroup SIR model. American Economic Review: Insights, 3(4), 487–502.
Alban, A., Chick, S., Dongelmann, D., Sluijs, A., Wiersinga, W., Vlaar, A., & Sent, D. (2020). ICU capacity management during the COVID-19 pandemic using a stochastic process simulation. Intensive Care Med., 18, 1624–1626.
Bertsimas, D. (2020). Mortality risk calculator. https://www.covidanalytics.io/calculator
Birge, J. R., Candogan, O., & Feng, Y. (2020). Controlling epidemic spread: Reducing economic losses with targeted closures. Working Paper 2020-57, University of Chicago, Becker Friedman Institute for Economics.
Boulant, O., Fekom, M., Pouchol, C., Evgeniou, T., Ovchinnikov, A., Porcher, R., & Vayatis, N. (2020). SEAIR framework accounting for a personalized risk prediction score: Application to the Covid-19 epidemic. Image Processing On Line, 10, 150–166.
Britton, T. (2010). Stochastic epidemic models: A survey. Mathematical Biosciences, 223(1), 24–35.
Camelo, S., Ciocan, D. F., Iancu, D. A., Wares, X. S., & Zoumpoulis, S. I. (2021). Quantifying the benefits of targeting for pandemic response. medRxiv. https://www.medrxiv.org/content/early/2021/04/02/2021.03.23.21254155
Clémençon, S., & Vayatis, N. (2007). Ranking the best instances. Journal of Machine Learning Research, 8(88), 2671–2699.
Di Domenico, L., Pullano, G., Sabbatini, C. E., Boëlle, P.-Y., & Colizza, V. (2020). Expected impact of lockdown in Île-de-France and possible exit strategies. BMC Medicine, 18(1), 240.
Diekmann, O., Heesterbeek, J., & Roberts, M. G. (2010). The construction of next-generation matrices for compartmental epidemic models. Journal of the Royal Society Interface, 7(47), 873–885.
Djidjou-Demasse, R., Michalakis, Y., Choisy, M., Sofonea, M. T., & Alizon, S. (2020). Optimal covid-19 epidemic control until vaccine deployment. https://doi.org/10.1101/2020.04.02.20049189
Duque, D., Morton, D. P., Singh, B., Du, Z., Pasco, R., & Meyers, L. A. (2020). Timing social distancing to avert unmanageable Covid-19 hospital surges. PNAS, 117(3), 19873–19878.
Evgeniou, T., Hardoon, R. D., & Ovchinnikov, A. (2020). Leveraging AI to battle this pandemic—And the next one. Harvard Business Review, April 2020.
Fawcett, T. (2006). An introduction to ROC analysis. Pattern Recognition Letters, 27(8), 861–874.
Ferguson, N., Laydon, D., Nedjati Gilani, G., Imai, N., Ainslie, K., Baguelin, M., Bhatia, S., Boonyasiri, A., Cucunubá, Z., Cuomo-Dannenburg, G., Dighe, A., Dorigatti, I., Hu, F., Gaythorpe, K., Green, W., Hamlet, A., Hinsley, W., Okell, L. C., van Elsland, S., & Ghani, A. C., et al. (2020). Report 9: Impact of non-pharmaceutical interventions (NPIS) to reduce Covid19 mortality and healthcare demand. https://www.imperial.ac.uk/mrc-global-infectious-disease-analysis/covid-19/report-9-impact-of-npis-on-covid-19/
Flaxman, S., Mishra, S., Gandy, A., Juliette, H., Unwin, T., Mellan, T. A., Coupland, H., Whittaker, C., Zhu, H., Berah, T., Eaton, J. W., Monod, M., Imperial College COVID-19 Response Team, Ghani, A. C., Donnelly, C. A., Riley, S., Vollmer, M. A. C., Ferguson, N. M., Okell, L. C., & Bhatt, S. (2020). Estimating the effects of non-pharmaceutical interventions on Covid-19 in Europe. Nature, 584, 257–261.
Garin, M., Limnios, M., Nicolai, A., Bargiottas, I., Boulant, O., Chick, S., Evgeniou, A. D. T., Fekom, M., Kalogeratos, A., Labourdette, C., Ovchinnikov, A., Porcher, R., Pouchol, C., & Vayatis, N. (2021). Epidemiological models for covid-19 during the first wave from February to May 2020: A methodological review. https://hal.archives-ouvertes.fr/hal-03332525
Gershon, D., Levy Yeyati, E., & Sartorio, L. (2021). Managing Covid-19 pandemic without destroying the economy. https://arxiv.org/abs/2004.10324
Goldstein, P., Levy Yeyati, E., & Sartorio, L. (2021). Lockdown fatigue: The diminishing effects of quarantines on the spread of Covid-19. https://doi.org/10.21203/rs.3.rs-621368/v1
Guan, W., Ni, Z., Hu, Y., Liang, W., Ou, C., He, J., Liu, L., Shan, H., Lei, C., Hui, D. S. C., Du, B., Li, L., Zeng, G., Yuen, K., Chen, R., Tang, C., Wang, T., Chen, P., Xiang, J., … Zhong, N. (2020). Clinical characteristics of coronavirus disease 2019 in China. The New England Journal of Medicine, 382(18), 1708–1720.
Kaplan, E. H. (2020). Covid-19 scratch models to support local decisions. Manufacturing & Services Operations Management, 4, 645–655.
Koo, J. R., Cook, A. R., Park, M., Sun, Y., Sun, H., Lim, J. T., Tam, C., & Dickens, B. L. (2020). Interventions to mitigate early spread of Sars-CoV-2 in Singapore: A modelling study. The Lancet Infectious Diseases, 20, 678–688.
Kucharski, A. J., Russell, T. W., Diamond, C., Liu, Y., Edmunds, J., Funk, S., Eggo, R. M., on behalf of the Centre for Mathematical Modelling of Infectious Diseases COVID-19 working group. (2020). Early dynamics of transmission and control of Covid-19: A mathematical modelling study. The Lancet Infectious Diseases, 5, 553–558.
Lasko, T. A., Bhagwat, J. G., Zou, K. H., & Ohno-Machado, L. (2005). The use of receiver operating characteristic curves in biomedical informatics. *Journal of Biomedical Informatics, 38*(5), 404–415.

Marjoram, P., Molitor, J., Plagnol, V., & Tavaré, S. (2003). Markov chain Monte Carlo without likelihoods. *Proceedings of the National Academy of Sciences, 100*(26), 15324–15328.

Petherick, A. (2020). Developing antibody tests for Sars-Cov-2. *The Lancet, World Report, 395*, 1101–1102.

Salje, H., Tran Kiern, C., Lefrancq, N., Courtejoie, N., Bosetti, P., Paireau, J., Andronico, A., Hozé, N., Richet, J., Dubost, C.-L., Le Strat, Y., Lessler, J., Levy-Bruhl, D., Fontanet, A., Opatowski, L., Boelle, P.-Y., & Cauchemez, S. (2020). Estimating the burden of Sars-Cov-2 in France. *Science, 369*(6500), 208–211.

Sonabend, R., Whittles, L. K., Imai, N., Perez-Guzman, P. N., Knock, E. S., Rawson, T., Gaythorpe, K. A. M., Djaafara, B. A., Hinsley, W., FitzJohn, R. G., Lees, J. A., Kanapram, D. T., Volz, E. M., Ghani, A. C., Ferguson, N. M., Baguelin, M., & Cori, A. (2021). Non-pharmaceutical interventions, vaccination, and the Sars-Cov-2 delta variant in England: A mathematical modelling study. *The Lancet, 398*(10313), 1825–1835.

Wang, C. J., Ng, C. Y., & Brook, R. H. (2020). Response to COVID-19 in Taiwan: Big data analytics, new technology, and proactive testing. *JAMA, 323*(14), 1341–1342.

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