Modeling the Spatiotemporal Epidemic Spreading of COVID-19 and the Impact of Mobility and Social Distancing Interventions

Alex Arenas,1,* Wesley Cota,2,3,4 Jesús Gómez-Gardeñes,2,4,5 Sergio Gómez,1 Clara Granell,1,2,4 Joan T. Matamalas,6 David Soriano-Paños,2,4 and Benjamin Steinegger1

1Departament d’Enginyeria Informàtica i Matemàtiques, Universitat Rovira i Virgili, E-43007 Tarragona, Spain
2Department of Condensed Matter Physics, University of Zaragoza, E-50009 Zaragoza, Spain
3Departamento de Fisica, Universidade Federal de Viçosa, 36570-900 Viçosa, Minas Gerais, Brazil
4GOTHAM Lab-BIFI, University of Zaragoza, E-50018 Zaragoza, Spain
5Center for Computational Social Science (CCSS), Kobe University, Kobe 657-8501, Japan
6Center for Interdisciplinary Cardiovascular Sciences, Cardiovascular Division, Department of Medicine, Brigham and Women’s Hospital, Harvard Medical School, Boston, Massachusetts 02115, USA

(Received 11 August 2020; revised 1 October 2020; accepted 12 November 2020; published 18 December 2020)

On 31 December, 2019, an outbreak of a novel coronavirus, SARS-CoV-2, that causes the COVID-19 disease, was first reported in Hubei, mainland China. This epidemic’s health threat is probably one of the biggest challenges faced by our interconnected modern societies. According to the epidemiological reports, the large basic reproduction number $R_0 \sim 3.0$, together with a huge fraction of asymptomatic infections, paved the way for a major crisis of the national health capacity systems. Here, we develop an age-stratified mobility-based metapopulation model that encapsulates the main particularities of the spreading of COVID-19 regarding (i) its transmission among individuals, (ii) the specificities of certain demographic groups with respect to the impact of COVID-19, and (iii) the human mobility patterns inside and among regions. The full dynamics of the epidemic is formalized in terms of a microscopic Markov chain approach that incorporates the former elements and the possibility of implementing containment measures based on social distancing and confinement. With this model, we study the evolution of the effective reproduction number $R(t)$, the key epidemiological parameter to track the evolution of the transmissibility and the effects of containment measures, as it quantifies the number of secondary infections generated by an infected individual. The suppression of the epidemic is directly related to this value and is attained when $R < 1$. We find an analytical expression connecting $R$ with nonpharmacological interventions, and its phase diagram is presented. We apply this model at the municipality level in Spain, successfully forecasting the observed incidence and the number of fatalities in the country at each of its regions. The expression for $R$ should assist policymakers to evaluate the epidemic’s response to actions, such as enforcing or relaxing confinement and social distancing.

DOI: 10.1103/PhysRevX.10.041055

I. INTRODUCTION

As of August 2020, the novel coronavirus, SARS-CoV-2, has infected more than 20,000,000 persons worldwide with COVID-19, causing more than 700,000 deaths. From its onset, the scientific community has embarked on an unprecedented search for efficient ways to stop the advance of the epidemic wave. This extraordinary effort has involved different disciplines: epidemiological studies to characterize the transmission patterns of the new coronavirus, large-scale clinical trials to test treatments that improve the course of the most severe cases, and the design of vaccines that safely provide immunity to the world’s population.

Once the epidemiological parameters were found [1–10], the mathematics and physics community made use of the most advanced epidemic models [11–18] to track and anticipate the spread of the epidemics (see Ref. [19] for an exhaustive review on these efforts). Nevertheless, the particularities of both SARS-CoV-2 transmission and the heterogeneous clinical evolution of patients with COVID-19 call for a rethinking of conventional models toward adapted models that explicitly take into account

*alexandre.arenas@urv.cat
†gardenes@unizar.es

Published by the American Physical Society under the terms of the Creative Commons Attribution 4.0 International license. Further distribution of this work must maintain attribution to the author(s) and the published article’s title, journal citation, and DOI.
these details and the characteristics of the populations affected by the pandemic. Among these characteristics, capturing the specific demographic distribution, the complex patterns of social contacts, and the geographic mobility networks among populations is essential to understand and forecast the impact that nonpharmaceutical containment measures, such as social distancing or confinement, have on the spread of the pathogen.

Here, we propose a mathematical (mechanistic) model particularly tailored to capture the main features of the propagation of SARS-CoV-2. This model relies on previous metapopulation models by the authors [20–24] to incorporate the specific characteristics of SARS-CoV-2 transmission and the clinical features of COVID-19 patients. Concerning transmission, we consider the important effect that the large fraction of asymptomatic infections (around 40% [25]) have on the covert spread of the disease in the early stages, causing a delayed response of containment measures based on the observed incidence of symptomatic patients. For the clinical aspect, we consider the long hospitalization times required by critical cases admitted to the intensive care unit (ICU), since the saturation of these units constitutes one of the main political and health threats of COVID-19 epidemics.

To incorporate the former knowledge into a single framework, we build an epidemiological model with ten compartments characterizing the epidemiological and clinical status of individuals in each patch that make up the metapopulation. In addition, we split the former partition into three age strata (young, adults, and older adults), thus allowing us to capture in a stylized way the main epidemiological, clinical, and behavioral differences between these groups. On the one hand, SARS-CoV-2 importation and exportation events between patches are mostly due to the mobility of the active population. On the other hand, the medical evolution of COVID-19 displays significant differences across age strata [13,26,27]. In this regard, infections in the young group lead to null or mild symptoms [28], whereas, for older individuals, the infection evolves toward more severe symptoms and often requires hospitalization.

The detailed structure of the metapopulation framework offers the possibility of designing and evaluating the impact of contention policies to stop the propagation of SARS-CoV-2. In particular, we focus on those policies relying on global or targeted quarantine measures due to the extreme relevance of these interventions for the course of the pandemic. This way, our formalism allows us to incorporate the temporal evolution of the fraction of the confined population and to assess the performance of lockdown policies to mitigate the pandemic. Taking advantage of this possibility, we explore several epidemic scenarios characterized by different contention measures and evaluate their impact on the decrease of the epidemic prevalence and the saturation of the Spanish health system during March, April, and May 2020. As a very relevant application, our model enables us to derive the selection of the minimum degree of confinement needed to avoid the health system crisis while not disrupting the economic fabric of the country.

In a nutshell, the proposed model takes into account, in a stylized way, three main peculiarities of the SARS-CoV-2 transmission: (i) the silent transmission of the pathogen through the young portion of the population, (ii) the large potential for the spatial dissemination of the pathogen provided by the mobility patterns of adults, and (iii) the severe pathology caused by COVID-19 in the elderly that yields to a dramatic increase of medical and hospital demands. Thus, the model can be viewed as three coevolving spreading processes with different spatiotemporal scales.

The analysis of the model and the main results are presented as follows. First, we show the main features of the dynamical model in Sec. II, describing first how epidemiological, clinical, and social data are incorporated into the metapopulation framework and, finally, showing its validation comparing with data corresponding to the COVID-19 outbreak in Spain. Once the model is defined and validated, we tackle in Sec. III the analytical derivation of the effective reproduction number. In Sec. IV, this expression is analyzed in detail to unravel the nonlinear dependency between the mobility restrictions and the epidemic incidence. Finally, in Sec. V, we round off the article by summarizing the main findings of the work and discussing the implications in public health policies aimed at facing COVID-19.

II. METAPOPULATION MODEL

We present below the details of the metapopulation framework tailored for capturing the spread of SARS-CoV-2 and the evolution of COVID-19 patients. This presentation is divided into four parts: the explanation of the compartmental dynamics for each age stratum in a single patch, the description of contagion processes governed by social contacts and the mobility among patches, the incorporation of contention measures aimed at reducing human mobility and increasing social distance, and, finally, the validation of the model with data corresponding to the epidemic wave in Spain.

A. Compartmental dynamics

The epidemic model is built upon previous work on the method known as the microscopic Markov chain approach (MMCA) [29,30]. It is designed for the modelization of epidemic spreading dynamics but is also applied to other types of processes, e.g., information spreading and traffic congestion [31–36].

In the original MMCA formalism, which analyzes a susceptible-infected-susceptible dynamics in a single population (i.e., each node representing an individual), the description of the dynamics is made using as variables the

ALEX ARENAS et al. PHYS. REV. X 10, 041055 (2020)
probabilities \( p_i(t) \) of each node \( i \) being infected at time \( t \). The MMCA equations express how these probabilities evolve in time [29]: A node is infected at a time step if it was previously infected and did not recover or was susceptible and became infected by one of its neighbors: 
\[ p_i(t + 1) = p_i(t)(1 - \mu) + [1 - p_i(t)]\Pi_i(t). \]
These equations just rely on probabilities, i.e., sums of probabilities of disjoints events, and products of probabilities of (approximately) independent events. Moreover, they are deterministic; thus, they can be iterated from any desired initial condition to know the evolution of the \( p_i(t) \) of each node.

The capacity of describing the epidemics at the level of nodes explains the qualification of MMCA as a microscopic methodology. This characteristic distinguishes MMCA from the more common mean-field approaches, which are based on differential equations that express the rate of change of the fraction of infected individuals: The variables, the equations, and even time are analyzed differently.

The adaptation of MMCA to the study of epidemics in structured metapopulations, with heterogeneous agents, and subject to recent mobility patterns, is performed by some of the authors in previous works [20–24]. The main difference with respect to standard MMCA is that variable \( \rho^{m,g}(t) \) represents the probability that individuals, with residence at node \( i \), are in state \( m \). An alternative interpretation for them is as the fraction of individuals living at the node that are in the given state. However, the equations of the dynamics are better understood with the probabilistic semantics.

The evolution of these probabilities \( \{\rho^{m,i}(t)\} \) determines the dynamical state of a single patch \( i \) and is dictated by the underlying compartmental model. Here, this compartmental model is composed of the following epidemiological and clinical compartments: susceptible (S), exposed (E), asymptomatic infectious (A), symptomatic infectious (I), to be admitted in ICU (prehospitalized in ICU, \( P_H \)), fatal prognosis (predeceased, \( P_D \)), admitted in ICU that will recover \( (H_R) \) or decease \( (H_D) \), recovered \( (R) \), and deceased \( (D) \). Compartments \( P_H \) and \( P_D \) are introduced to account for the delays observed for individuals before they are admitted in ICU or before they die outside ICU, respectively. Similarly, the compartments \( H_R \) and \( H_D \) represent the individuals in ICU but separated according to their final state (recovered or deceased, respectively), which is reached at different rates.

As anticipated above, the population of a patch is further divided into \( N_G \) age strata. Although we present the model in a general form, its application to COVID-19 makes use of only the three age strata mentioned above \((N_G = 3)\): young people \( (Y) \), with age up to 25; adults \( (M) \), with age between 26 and 65; and elderly people \( (O) \), with age larger than 65. This differentiation is very relevant in light of the observation of a large fraction of infected asymptomatic individuals at ages \(<25\), in contrast with the abundance of severe and critical cases reported for people at older ages \(>65\).

Let us suppose we have a population of \( N \) individuals distributed in \( N_p \) regions, with \( n_i \) individuals residing in region \( (patch) \) \( i \). We also consider that individuals belong to one of \( N_G \) different age strata, in such a way that \( n^g_i \) individuals of age strata \( g \) live in region \( i \). Thus,
\[ N = \sum_{g=1}^{N_G} \sum_{i=1}^{N_p} n^g_i = \sum_{i=1}^{N_p} n_i = \sum_{g=1}^{N_G} n^g, \]
where \( n^g \) is the total population of age strata \( g \). Our system is completely characterized by variables \( \rho^{m,g}(t) \), which account for the probabilities that individuals of age stratum \( g \) assigned to patch \( i \) are in state \( m \) at time \( t \), where \( m \in \{S, E, A, I, P_H, P_D, H_R, H_D, R, D\} \) and \( g \in \{Y, M, O\} \).

The temporal evolution of these quantities is given by
\[
\begin{align*}
\rho^{S,g}_i(t + 1) &= \rho^{S,g}_i(t)[1 - \Pi^g_i(t)], \\
\rho^{E,g}_i(t + 1) &= \rho^{S,g}_i(t)\Pi^g_i(t) + (1 - \eta^g)\rho^{E,g}_i(t), \\
\rho^{A,g}_i(t + 1) &= \eta^g\rho^{E,g}_i(t) + (1 - \alpha^g)\rho^{A,g}_i(t), \\
\rho^{I,g}_i(t + 1) &= \alpha^g\rho^{A,g}_i(t) + (1 - \mu^g)\rho^{I,g}_i(t), \\
\rho^{P_H,g}_i(t + 1) &= \mu^g\theta^g\rho^{I,g}_i(t) + (1 - \zeta^g)\rho^{P_H,g}_i(t), \\
\rho^{P_D,g}_i(t + 1) &= \mu^g(1 - \theta^g)\rho^{I,g}_i(t) + (1 - \lambda^g)\rho^{P_D,g}_i(t), \\
\rho^{R,g}_i(t + 1) &= \mu^g(1 - \theta^g)(1 - \gamma^g)\rho^{L,g}_i(t) + \chi^g\rho^{H,g}_i(t) + \rho^{R,g}_i(t), \\
\rho^{H,g}_i(t + 1) &= \lambda^g\rho^{H,g}_i(t) + (1 - \psi^g)\rho^{H,g}_i(t), \\
\rho^{H,D,g}_i(t + 1) &= \lambda^g(1 - \omega^g)\rho^{P_H,g}_i(t) + (1 - \chi^g)\rho^{H,D,g}_i(t), \\
\rho^{D,g}_i(t + 1) &= \zeta^g\rho^{P_D,g}_i(t) + \psi^g\rho^{H,D,g}_i(t) + \rho^{D,g}_i(t).
\end{align*}
\]
These MMCA equations correspond to a discrete-time dynamics, in which each time step represents a day. Note that the sum over all \( \rho^{m,g}_i(t) \) for a given patch \( i \) and age group \( g \) equals 1 for each time step \( t \).

The rationale of the former compartmental dynamics is the following. Susceptible individuals get infected by contacts with asymptomatic and symptomatic infected agents, with a probability \( \Pi^g_i(t) \) (described in Sec. II B) becoming exposed. Exposed individuals turn into asymptomatic at a certain rate \( \eta^g \), which, in turn, become infectious at a rate \( \alpha^g \). Once infected, three paths emerge, which are reached at an infectious rate \( \mu^g \). The first one is death without being hospitalized in ICU, with probability \( \theta^g \), after a latency period governed by rate \( \chi^g \). Otherwise, with probability \( \gamma^g \) the individuals are hospitalized in ICU, which are reached at rate \( \lambda^g \), while with probability \( 1 - \gamma^g \)
they recover. Individuals in ICU have a fatality probability $\omega^g$, which is reached at a rate $\psi^g$, whereas the recovery is reached at a rate $\chi^g$. See Fig. 1 for a sketch of the compartmental model with all the transitions and Tables I–III for the list of the parameters and their values to simulate the spreading of COVID-19 in Spain, which are discussed in Appendix A.

B. Social contacts and mobility among patches

To understand the geographical diffusion of the disease, as a result of human-human interactions in small geographical patches, one has to combine the contagion process with the long-range disease propagation due to human mobility across different spatial scales. For the case of epidemic modeling, the usual metapopulation scenario assumes that the individuals within each patch are well mixed; i.e., pathogens can be transmitted from an infected host to any of the healthy agents placed in the same patch with the same probability. The second aspect of the metapopulation approach concerns the mobility of agents. Each host is allowed to change its current location and occupy another patch, thus fostering the spread of pathogens at the system level. Mobility of agents between different patches is usually represented in terms of a contact network where nodes are locations while a link between different patches is usually represented in terms of a contact matrix $C$. Each host is allowed to change its current location and choose destination $j$ while living in region $i$ with probability $\pi_{ij}^g$, whereas $P^g_i(t)$ denotes the probability that those agents get infected by the pathogen inside patch $i$. This way, the first term in the rhs of Eq. (12) denotes the probability of contracting the disease inside the residence patch, whereas the second term contains those contagions taking place in any of the neighboring areas. Furthermore, we assume that the number of contacts increases with the density of each area according to a monotonically increasing function $f$. Finally, we introduce an age-specific contact matrix $C^g$, whose elements $C^g_{ij}$ define the fraction of contacts that individuals of age stratum $g$ perform with individuals belonging to age stratum $h$. With the above definitions, $P^g_i(t)$ reads

$$P^g_i(t) = 1 - \prod_{h=1}^{N_h} \prod_{j=1}^{N_j} \prod_{m \in \{A,I\}} \left(1 - \beta_m^g\right) e^{\zeta^g} f(n_i^g / s_i) C^g_{ij} \left(n^g_{ji}(t) / R_i^g\right).$$

and age stratum $g$ are captured in the Markovian equations through the probability $\Pi^g_i(t)$. The value of $\Pi^g_i(t)$ encodes the probability that a susceptible agent belonging to age stratum $g$ and patch $i$ contracts the disease. Under the model assumptions, this probability is given by

$$\Pi^g_i(t) = (1 - p^g) P^g_i(t) + p^g \sum_{j=1}^{N_j} R^g_{ij} P^g_j(t), \quad (12)$$

where $p^g$ denotes the degree of mobility of individuals within age stratum $g$, $R^g_{ij}$ is the mobility matrix (fraction of individuals of group age $g$ that choose destination $j$ while living in region $i$), and $P^g_i(t)$ denotes the probability that those agents get infected by the pathogen inside patch $i$. Finally, we introduce an age-specific contact matrix $C^g$, whose elements $C^g_{ij}$ define the fraction of contacts that individuals of age stratum $g$ perform with individuals belonging to age stratum $h$. With the above definitions, $P^g_i(t)$ reads

$$P^g_i(t) = 1 - \prod_{h=1}^{N_h} \prod_{j=1}^{N_j} \prod_{m \in \{A,I\}} \left(1 - \beta_m^g\right) e^{\zeta^g} f(n_i^g / s_i) C^g_{ij} \left(n^g_{ji}(t) / R_i^g\right).$$

(13)
Parameters $\beta_A$ and $\beta_I$ correspond to the infection probabilities for contacts of a susceptible individual with asymptomatic and symptomatic infected individuals, respectively. The exponents represent the number of contacts made by an agent of age stratum $g$ in patch $i$ with infectious individuals—compartments $A$ and $I$, respectively—of age stratum $h$ residing at patch $j$. Accordingly, the double product expresses the probability for an individual belonging to age stratum $g$ not being infected while staying in patch $i$.

The term $z^g(k^g)f(\tilde{n}_i/s_i)$ represents the overall number of contacts (infectious or noninfectious) that an agent from age group $g$ makes inside patch $i$. These contacts, as mentioned above, increase monotonically with the density of patch $i$ following function $f(\tilde{n}_i/s_i)$, where $s_i$ is the area of patch $i$ measured in square kilometers and $\tilde{n}_i$ represents the effective population inside patch $i$. Likewise, the term $z^g$ is a normalization factor introduced to ensure that the average degree of population belonging to age group $g$ is $\langle k^g \rangle$. Therefore (see details in Appendix B),

$$z^g = \frac{n^g}{\sum_{i=1}^{N_g} f(\tilde{n}_i/s_i)\tilde{n}_i^g},$$

where the effective population at patch $i$, i.e., the number of people present at patch $i$ when commuting takes place, is given by

$$\tilde{n}_i = \sum_{g=1}^{N_g} \tilde{n}_i^g,$$

which is distributed in age strata of size

$$\tilde{n}_i^g = \sum_{j=1}^{N_h} M_{ji}^g n_j^g.$$  \hspace{1cm} (15)

For convenience, we define the effective mobility matrices $M_{ji}^g$:

$$M_{ji}^g = (1 - p^g)\delta_{ij} + p^g R_{ji}^g,$$ \hspace{1cm} (17)

that take into account both the degree of mobility of the population and the transition probabilities to the neighboring patches.

From now on, we use the tilde notation to indicate variables measured while the commuting is active, to distinguish them from variables when all the population is in its residence patch.

The function $f(x)$ governing the influence of population density is selected, following Ref. [41], as

$$f(x) = 1 + (1 - e^{-5x}).$$  \hspace{1cm} (18)

The last term of the exponents in Eq. (13) contains the probability that these contacts are contagious, which is proportional to $n_{j-i}^{m,h}$, the expected number of individuals of age stratum $h$ in the given infectious state $m$ (either $A$ or $I$) which transit from region $j$ to region $i$:

$$n_{j-i}^{m,h}(t) = n_j^h \rho_j^m(t) M_{ji}^h.$$  \hspace{1cm} (19)

The discrete-time nature of this model allows for an easy computation of the time evolution of all the relevant variables, providing information at the regional level.

C. Modeling confinement measures

Finally, this formalism allows for quantifying the performance of different confinement measures aimed at reducing the impact of COVID-19. In this section, we put our focus on the effect of extended lockdown policies promoted across the entire system, but other interventions such as modifying the morphology of the mobility network or isolating specific age groups can also be assessed with our framework.

To incorporate confinement policies in our formalism, we consider that a fraction of a given age stratum $\kappa_0^g(t)$ is under lockdown at time $t$. In this sense, let us remark that the time series $\kappa_0^g(t)$ allows us to tune the strength of those confinement measures proposed to contain COVID-19. In this sense, $\kappa_0^g = 1$ corresponds to a total lockdown scenario in which all activity is paralyzed, whereas $\kappa_0^g = 0$ reflects a less strict policy not affecting the population. Moreover, we introduce the effects of social distancing measures taken by the population as a reduction $\delta$ of the number of contacts made by the nonconfined population. We assume the social distancing $\delta$ to be constant in time, while the confinement level $\kappa_0^g(t)$ is time dependent.

Let us denote as $t_c$ the initial time from which lockdown interventions are deployed. The incorporation of lockdown policies in the formalism demands some modifications of the equations. First, population confinement implies a reduction of the mobility within the population. Therefore, the degree of mobility $p^g$ becomes time dependent and now reads

$$p^g(t) = [1 - \kappa_0^g(t)\Theta(t - t_c)]p^g,$$ \hspace{1cm} (20)

where $\Theta(x)$ is the Heaviside function, that is 1 if $x \geq 0$ and 0 otherwise. As a by-product, $\tilde{n}_i^g$ and $z^g$ also become dependent on time; see Eqs. (14)–(16). Second, the confined population reduces their interactions to the members of their households. Equally, independently of whether they are confined or not, the contacts of individuals belonging to age strata $O$ are reduced to their household contacts, since they do not form part of the working population. Therefore, in the lockdown scenario, the average degree of elderly population is $\langle k^O \rangle = k^O_h$, with $k^O_h$ being the contacts taking place inside their households [42]. In contrast, given the nature of the interventions deployed by the authorities, the
contacts outside the household of the nonconfined population belonging to age strata $Y$ and $M$ are restricted to the workplace and shaped by social distancing. To incorporate both aspects, in the lockdown scenario, we define the average degree of individuals belonging to age group $g \in \{Y, M\}$ at time $t$, $k^g(t)$, as

$$
<k^g(t)> = k^g_h(t) + [1 - k^g_h(t)](1 - \delta)k^g_{h+w},
$$

(21)

where $k^g_h$ and $k^g_{h+w}$, which encode the contacts made by confined (at home) and nonconfined (at home and at work) population, respectively, are obtained from Ref. [42]. This way, the average number of daily contacts $\langle k^g \rangle$ of agents belonging to each group $g \in \{Y, M, O\}$ should be updated to account for this dependence:

$$
\langle k^g \rangle(t) = [1 - \Theta(t - t_c)]\langle k^g \rangle + \Theta(t - t_c)\langle k^g \rangle(t_c).
$$

(22)

The time dependence of the number of interactions affects the probability that an individual from age group $g$ and patch $i$ becomes infected, $P^g_i(t)$, which is now given by

$$
P^g_i(t) = 1 - \prod_{h=1}^{N_G} \prod_{j=1}^{N_p} \prod_{m=\{A,I\}} (1 - \beta_m) e^{\sigma(t)\langle k^g \rangle(t)\Theta(t)\langle k^g \rangle(t)/\bar{n}^g_i(t)}.
$$

(23)

Finally, a perfect confinement would entail the complete shielding of fully susceptible households against the disease. Therefore, the confinement of the population modifies the pool of susceptible individuals reachable by their infectious counterparts [43]. Unfortunately, in practical terms, complete isolation of households is impossible, since their members are required to go out for essential activities such as buying groceries, drugs, etc. Therefore, we introduce a household permeability $\phi$ accounting for the social mixing among members from different households in these situations. We fix the household permeability $\phi$ as constant in time. While the interactions among members of different confined households clearly shape household isolation, we assume that they do not substantially modify the average number of contacts of the population.

In this scenario, a relevant indicator to quantify the efficiency of the policy is the probability of one individual living in a household, inside a given municipality $i$, without any potentially infectious individual. Assuming that confinement is implemented at time $t_c$, this quantity, denoted in the following as $CH_i(t_c)$, is given by

$$
CH_i(t) = \left(\frac{1}{n_i^g} \sum_{g=1}^{N_G} [1 - \rho_i^{S,g}(t) - \rho_i^{A,g}(t) - \rho_i^{I,g}(t)]n_i^g\right)^\sigma.
$$

(24)

This confinement strategy is introduced in the dynamical Eqs. (2)-(11) by modifying Eqs. (2) and (3) for the time after $t_c$:

$$
\rho_i^{S,g}(t+1) = [\rho_i^{S,g}(t) + \rho_i^{CH,g}(t)]\Pi^g_i(t)
\times \left[1 - \Theta(t - t_c)(1 - \phi)\kappa^g_0(t)CH_i(t)\right],
$$

(25)

$$
\rho_i^{A,g}(t+1) = \rho_i^{A,g}(t),
$$

$$
\rho_i^{I,g}(t+1) = [\rho_i^{I,g}(t) + \rho_i^{CH,g}(t)]\Pi^g_i(t)
\times \left[1 - \Theta(t - t_c)(1 - \phi)\kappa^g_0(t)CH_i(t)\right],
$$

(27)

$$
\rho_i^{CH,g}(t+1) = \rho_i^{CH,g}(t),
$$

(28)

where we add a new compartment $CH$ to hold the individuals under household isolation after applying confinement and returning them to the susceptible pool as confinement is relaxed. Confinement also affects the average number of contacts; thus, we must also update Eq. (13).

In summary, we introduce a confinement strategy driven by social distancing and the isolation of a certain fraction of the population, which, in turn, reduces the average number of contacts, the mobility, and, as a result, the probability of becoming infected, thus reducing the overall prevalence of the disease. Therefore, confinement policies are captured in the formalism by the time series describing confinement $\kappa^g_0(t)$, the social distancing factor $\delta$, and household permeability $\phi$. See Table IV for the list of the confinement parameters and their values to simulate the spreading of COVID-19 in Spain, which are discussed in Appendix A.

### D. Validation of the model

The metapopulation model is calibrated with real data of fatalities caused by COVID-19 in Spain until April 8, using approximate Bayesian computation [45]. Because of the absence of more specific data, we take the reduction in the work mobility as a proxy for the temporal evolution of the confined population in Spain. Furthermore, since displacements not related to work activities were banned as the lockdown was enforced, the evolution of the work...
mobility reflects well the variations in the population under confinement. The confinement measures are introduced into the model from March 14 onward (before this date, no intervention was made). We obtain the time series $\kappa^0_0(t)$ from official reports [44] by estimating the work mobility reduction with respect to a baseline day, November 19, 2019. We assume that these time series are equal for all age groups and refer in the following to $\kappa^0_{Y,A,O}(t)$ as $\kappa_0(t)$.

Further details on the data used and model calibration are shown in Appendixes A and C.

Once calibrated, we validate our model using two datasets: the evolution of daily fatalities in Spain [Fig. 2(a)] and the evolution of the daily number of newly detected symptomatic individuals [Fig. 2(b)]. The results in these figures show a qualitative and quantitative agreement in all the phases corresponding to the different containment interventions implemented by the government of Spain. Note that the time series of new symptomatic individuals is rescaled to account for case underreporting. For the sake of visualization, Figs. 2(a) and 2(b) incorporate the lockdown stages depicted as horizontal bars that correspond to the different interventions enforced in Spain. In LM, Spain was under lockdown but laboral mobility was allowed. In a stricter scenario (EM), only essential mobility (that corresponding to the commuting of essential workers) was allowed. In ECS, in addition to permitting laboral mobility,
III. EFFECTIVE REPRODUCTION NUMBER

The stage of an ongoing outbreak is typically quantified by studying the effective reproduction number $R(t)$, which captures the number of secondary cases that an individual infected at time $t$ will produce during the duration of its infectious period. To compute the evolution of $R(t)$ using our model and understand its dependencies, we present an incremental rationale from the most rough approximation to the most accurate formula.

Let us consider, first, a scenario (known as mean-field in physics) in which an infected subject $i$ makes $\langle k \rangle$ contacts each time step. Assuming an infection probability $\beta$ per contact, the expected number of individuals infected by $i$ at each time step is $\beta(\langle k \rangle)\rho_S$, where $\rho_S$ is the fraction of susceptible individuals on the population. We can estimate how many individuals are infected by subject $i$ over an infectious period of duration $\tau$ as

$$R = \tau\beta(\langle k \rangle)\rho_S.$$  \hspace{1cm} (28)

This very simple mean-field approach is already very informative, as it reveals the main dependencies of $R$ on the variables of the epidemic propagation, indicating the actionable variables we can resort on to reduce $R$. Acting on the infection probability $\beta$ can be attained with the use of face masks and physical distance or through pharmacological interventions via prophylactic treatments. Another actionable variable is $\langle k \rangle$, which can be modified by human behavioral changes such as confinement. Also, the infectious period $\tau$ can be modified by implementing a fast testing policy to confirm and isolate suspected cases. Finally, the value of $\rho_S$, that can be directly estimated from the model, is highly dependent on the extent of lockdown policies, on the level of immunity reached in the population, or on the degree of vaccinated individuals when this pharmacological intervention is available.

The previous approach, however, neglects many important social, demographic, and mobility details key to shaping the way in which secondary infections spread through a geographically distributed heterogeneous population. Likewise, it obviates the temporal evolution of the group of individuals susceptible to contracting the disease. This naive approach can be largely improved by leveraging the information of the proposed age-structured mobility-driven metapopulation epidemic model.

A. Determination of the effective reproduction number from the dynamics

Let us define $R^g_i$, the effective reproduction number of geographical patch $i$ and age stratum $g$, as the number of secondary cases produced by an infected individual belonging to patch $i$ and age stratum $g$ [47]. Mathematically, $R^g_i$ is expressed as

$$R^g_i = \tau^{\text{d}}(\langle \beta^g \rangle) \sum_{h=1}^{N_p} \sum_{j=1}^{N_{g_j}} k_{ij}^{gh} S^g_j. \hspace{1cm} (29)$$

In the above expression, the following quantities interplay: the duration of the infectious period of each age stratum $g$, $\tau^{\text{d}}$; the average infectivity during the infectious period, $\langle \beta^g \rangle$, per age stratum $g$; the average number of daily contacts an individual belonging to patch $i$ and age stratum $g$ makes with individuals in patch $j$ belonging to age stratum $h$, $k_{ij}^{gh}$; and the fraction of susceptible individuals present in patch $j$ belonging to age stratum $h$, $\rho_j^{gh}$. In summary, the structure of $R^g_i$ resembles the mean-field expression presented in Eq. (28), but it takes explicitly into account the mixing of individuals according to their age strata and mobility patterns ($k_{ij}^{gh}$ and $\rho_j^{gh}$). Nevertheless, this expression is still a static snapshot at time $t$.

We can extend this approach to include both the temporal dependence and the heterogeneity of contacts induced by the daily commuting patterns, making use of all the probabilities provided by our model equations. Moreover, here, we aim at studying the temporal evolution of the effective reproduction number beyond the early stage of the disease. In this sense, we define the reproduction number of patch $i$ and age stratum $g$ as the number of infections observed if we seed an infected individual in the aforementioned patch and age stratum [47]. This definition involves considering that this infected individual, with residence in patch $i$, may commute to patch $j$, where it will be able to contact and infect susceptible individuals coming from any other patch $k$. 
First, we calculate the expected number of susceptible individuals of age stratum \( h \) which move from region \( k \) to region \( j \) as
\[
\rho^{S,h}_{k \rightarrow j}(t) = n^h_k \rho^{S,h}_{k}(t) m^{h}_{kj}(t).
\]
(30)

The fraction of susceptible individuals of each age stratum \( g \) present at time \( t \) in patch \( j \) is expressed by
\[
\tilde{\rho}^{S,h}_{j}(t) = \frac{1}{N_j} \sum_{k=1}^{N_p} \rho^{S,h}_{k \rightarrow j}(t).
\]
(31)

Next, we compute the number of susceptible contacts made by an individual of age stratum \( g \) and patch \( i \), which can be expressed as
\[
Q^g_i(t) = \sum_{j=1}^{N_p} \sum_{h=1}^{N_h} \rho^{S,h}(t) \langle k^{h}(t) \rangle f \left( \frac{\tilde{\rho}^{S,h}(t)}{s_j} \right) C^{g,h} \tilde{\rho}^{S,h}_{j}(t) M^{g}_{ij}(t).
\]
(32)

Using this expression and assuming that we have an individual from patch \( i \) and age stratum \( g \) that has become infectious, we can compute the effective reproduction number for each patch and age stratum as
\[
\mathcal{R}_i^g(t) = \left( \frac{\beta_A}{\alpha^g} + \frac{\beta_I}{\beta^g} \right) Q^g_i(t).
\]
(33)

In Eq. (33), we assume that the fraction of susceptible individuals as well as the average number of contacts is constant during the infectious period. However, the fraction of susceptible individuals obviously changes during that time. More importantly, as containment measures come into play, the number of contacts varies as well. To account for the temporal variability of these quantities, we calculate the contributions to \( \mathcal{R}_i^g(t) \) of the infections produced since the individual became exposed at time \( t \). The contributions can be determined through the probabilities to be exposed, \( \zeta^{E,g}(t) \), asymptomatic, \( \zeta^{A,g}(t) \), or symptomatic, \( \zeta^{L,g}(t) \), time steps after an infection. The evolution of these probabilities is given by
\[
\zeta^{E,g}(t+1) = (1 - \eta^g) \zeta^{E,g}(t),
\]
(34)
\[
\zeta^{A,g}(t+1) = \eta^g \zeta^{E,g}(t) + (1 - \alpha^g) \zeta^{A,g}(t),
\]
(35)
\[
\zeta^{L,g}(t+1) = \alpha^g \zeta^{A,g}(t) + (1 - \mu^g) \zeta^{L,g}(t).
\]
(36)

Since the seed is initially placed as exposed, the initial conditions are given by \( \zeta^{E,g}(0) = 1 \) and \( \zeta^{A,g}(0) = \zeta^{L,g}(0) = 0 \). Accordingly, solving explicitly the recursive Eqs. (34)–(36), we find
\[
\zeta^{E,g}(t) = (1 - \eta^g)^t, \quad \zeta^{A,g}(t) = \frac{\eta^g}{\alpha^g - \eta^g} [(1 - \eta^g)^t - (1 - \alpha^g)^t],
\]
(37)
\[
\zeta^{L,g}(t) = \frac{\eta^g}{\alpha^g - \eta^g} \frac{1}{(\alpha^g - \eta^g)(\mu^g - \eta^g)} \times \left[ (\mu^g - \alpha^g)(1 - \eta^g)^t + (\eta^g - \mu^g)(1 - \alpha^g)^t + (\alpha^g - \eta^g)(1 - \mu^g)^t \right].
\]
(39)

Incorporating these probabilities into Eq. (33), we obtain the following expression for the effective reproduction number of individuals of age stratum \( g \) in patch \( i \):
\[
\mathcal{R}_i^g(t) = \sum_{s=1}^{\infty} \left[ \zeta^{A,g}(s-t) \beta_A + \zeta^{L,g}(s-t) \beta_I \right]
\]
\[
\times \sum_{j=1}^{N_p} \sum_{h=1}^{N_h} k^{g,h}_{ij}(s) \tilde{\rho}^{S,h}_{j}(s),
\]
(40)

where
\[
k^{g,h}_{ij}(s) = \zeta^{g}(t) \langle k^{h}(t) \rangle f \left( \frac{\tilde{\rho}^{S,h}(t)}{s_j} \right) C^{g,h} \tilde{\rho}^{S,h}_{j}(t).
\]
(41)

The variables \( \mathcal{R}_i^g(t) \) tell us the effective reproduction number of one newly infected individual from age stratum \( g \) with residence in \( i \).

To compute the temporal evolution of the global effective number \( \mathcal{R}(t) \) or the effective number \( \mathcal{R}^g(t) \) associated to a given age stratum \( g \) at time \( t \), we must make a weighted average taking into account the distribution of newly infected individuals across patches and age strata. Therefore, these indicators read
\[
\mathcal{R}(t) = \frac{\sum_{m \in \{S.C.H.\}} \sum_{i=1}^{N_p} \sum_{g=1}^{N_h} [\rho^{m,g}_{i}(t-1) - \rho^{m,g}_{i}(t)] n^g_i \mathcal{R}_i^g(t)}{\sum_{m \in \{S.C.H.\}} \sum_{i=1}^{N_p} \sum_{g=1}^{N_h} [\rho^{m,g}_{i}(t-1) - \rho^{m,g}_{i}(t)] n^g_i}
\]
(42)

and
\[
\mathcal{R}^g(t) = \frac{\sum_{m \in \{S.C.H.\}} \sum_{i=1}^{N_p} \rho^{m,g}_{i}(t-1) - \rho^{m,g}_{i}(t)] n^g_i \mathcal{R}_i^g(t)}{\sum_{m \in \{S.C.H.\}} \sum_{i=1}^{N_p} [\rho^{m,g}_{i}(t-1) - \rho^{m,g}_{i}(t)] n^g_i},
\]
(43)

respectively. To show the validity of the former analytical expression for the effective reproduction number \( \mathcal{R}(t) \), we perform in Appendix E a comparison of its time evolution with the results obtained by using EpiEstim [48,49] applied to both the incidence in our model and the daily incidence data observed in Spain.
B. Monitoring the effect of contention measures in Spain

To check the effect of the policies aimed at reducing the impact of COVID-19 in Spain, we compute the expression of $R(t)$ according to Eq. (42) using the reported variations of $\kappa_0$ in this country from March 14. In Fig. 3, we represent $R(t)$ along with the time series of the degree of confinement $\kappa_0(t)$ obtained from the Instituto Nacional de Estadística (INE) of Spain [44]. Note that the confinement during the progressive exit of the lockdown state lies between 0.7 and 0.6 approximately, stabilizing $R(t) < 1$. However, there is a clear tendency of $R(t)$ to grow as the confinement level was relaxed in May (see also Fig. 9 in Appendix E). This growing tendency may be one of the reasons leading to the second epidemic wave Spain is facing during the months of July and August 2020.

Furthermore, in the plot, we obtain from Eq. (43) the effective reproduction number by age strata, that is, the $R^g(t)$ of the young, adult, and elderly population (being $g \in \{Y, M, O\}$). There, we notice that relevant heterogeneities exist among the reproduction number of the different age strata. In particular, at the end of the curve, we observe that a supercritical scenario for the adult population [$R^M(t) > 1$] is compatible with a global subcritical regime. This valuable information opens up the possibility of designing suppression strategies that allow a certain age stratum to have $R(t) > 1$, provided the average value of the whole population is below the threshold $R(t) < 1$. The same conclusion applies in terms of geographical areas instead of ages. This result offers a perspective on epidemic control that could be very useful to design strategic interventions targeting individual age stratum or particular geographical areas in heterogeneously distributed populations.

The analytical expression for $R^g(t)$ in relation to confinement $\kappa_0$, social distancing $\delta$, and household permeability $\phi$ can be exploited to ascertain the implications of non-pharmacological interventions. Let us assume that the aim of the intervention is that of keeping the epidemic under control, in the absence of any other tool besides confinement and social distance. In this case, the strong goal must be that of keeping of $R^g(t) \leq 1$ for all $i, g$, or the weaker goal $R(t) \leq 1$. We can use the analytical dependence of $R(t)$ to estimate the minimal, constant intervention in $\kappa_0$ and $\delta$ such that the goal is achieved.

In Fig. 4, we present, for the case of Spain, the lower bounds of confinement needed, assuming fixed social distancing, to achieve $R(t) < 1$. This analysis makes use of $R(t)$ to obtain the minimum possible confinement that, while damaging the socioeconomic structure as little as possible, allows the epidemic to be controlled and gradually mitigated. Note that the previous goal can be relaxed, or changed, according to any other demands of the authorities, as, for example, maintaining the number of ICUs in the health system below a certain threshold, minimizing the number of fatalities, or achieving suppression in the minimum time. Figure 4 highlights the relevance of timely and strict lockdown policies, to avoid the breakdown of the health system, in the absence of other non-pharmacological interventions. We observe that small variations in the fraction of confined population $\kappa_0$ entail

![FIG. 3. Effect of lockdown policies on the effective reproduction number. Temporal evolution of the effective reproduction number $R(t)$ for each age stratum and its average computed according to Eq. (42). Inset: The black line represents the temporal evolution of the degree of confinement $\kappa_0(t)$ extracted from data daily updated by INE [44]. The blue line shows the average over a 7-day time window to smooth out fluctuations observed in the data. To improve the visibility of the inset, degrees of confinement from March 14 until March 28 are not represented.](image-url)
FIG. 4. Relation between the confinement level and time needed for epidemic extinction. Top: Time to epidemic extinction from May 15 as the reproduction number is fixated from this day onward. The reproduction number is kept constant by adjusting the confinement level through time. We define epidemic extinction as a daily incidence of fewer than ten cases. Bottom: Average confinement level necessary to keep the reproduction number constant. At each time step, we correct the confinement level by steps of 0.1% until the effective reproduction number \( R(t) \) differs less than 1% from the envisaged value. Significant changes in the time needed to mitigate an epidemic wave.

IV. QUANTIFYING THE EFFECT OF CONTAINMENT MEASURES OVER \( R(t) \)

The expression for the global effective number \( R(t) \) [Eq. (42)], while useful for computing the impact that containment measures have on the spreading dynamics, does not allow a direct interpretation of how these policies, in particular, confinement and social distancing, influence the epidemic detriment. Here, starting from the complete expression derived for \( R(t) \), we consider some simplifying assumptions that allow us to unravel the contribution of these measures and, based on these approximations, derive an expression for the critical confinement needed to bend the epidemic curve.

A. Phase diagram of \( R(t) \) as a function of confinement and social distancing

To reveal the explicit dependence of \( R(t) \) on confinement and social distancing interventions, we now simplify Eq. (40) by, first, neglecting the heterogeneities among different subpopulations and, second, considering a static view in which the pool of susceptible individuals from each age stratum \( \langle \rho S,\theta \rangle \) remains constant during the intervention (constant \( \kappa_0 \)). The latter assumption enables us to recover Eq. (28) which, after introducing the aforementioned confinement parameters, reads

\[
R^\beta(t_c) \approx \left( \frac{\beta_A}{\rho^g} + \frac{\beta_I}{\rho^g} \right) [1 - \kappa_0(1 - \phi)] \langle CH \rangle \\
\times [\kappa_0 k_{h}^g + (1 - \delta)(1 - \kappa_0) k_{h+w}^g] \\
\times \sum_{k=1}^{N_g} C_{g}^{\phi}(\rho^{S,h}(t_c)), \tag{44}
\]

where \( \langle CH \rangle \) denotes the average fraction of households without any infected member.

Equation (44) shows an explicit quadratic dependence of \( R^\beta(t_c) \) on \( \kappa_0 \) that helps to understand the effectiveness of lockdown interventions. In Fig. 5(a), we plot a phase diagram illustrating how the global effective reproduction number is shaped by the fraction of confined population \( \kappa_0 \) and the household permeability \( \phi \). We highlight the phase transition occurring at \( R(t_c) = 1 \) (white solid line) separating two different regimes, \( R(t_c) > 1 \) (flattening) and \( R(t_c) < 1 \) (bending). Note also that, as the permeability of confinement \( \phi \) increases, the transition point is reached for larger values of confinement \( \kappa_0 \), this transition being lost for very high values of \( \phi \geq 0.83 \).

To shed light onto the qualitative differences between the regimes separated by the transition described above, in Fig. 5(b), we plot the number of daily new cases for different levels of confinement, \( \kappa_0 \), while keeping the social distancing as calibrated, \( \delta = 0.207 \) (95% credible interval Crl: 0.053–0.359). The confinement is applied at time \( t_c \) corresponding to March 14. It is observed that, as confinement is applied to a small fraction of the population \( \kappa_0 > 0 \), the curve for the number of new cases per day starts to broaden and the maximum shifts forward in time. Such a mitigation strategy is known as flattening the curve, in which containment delays and lowers the incidence peak. The consequence of the flattening scenario is that the impact over health systems is ameliorated at the expense of a larger epidemic period. However, as confinement is increased (higher \( \kappa_0 \)), there is a sharp change in the behavior of the epidemic curve for \( \kappa_0 \) between 0.4 and 0.6, signaling that the transition point \( R(t_c) = 1 \) lies between these two values. In particular, for values of \( \kappa_0 \) large enough, the epidemic curve reaches its maximum soon after confinement is put in place, achieving a completely new scenario, referred to as bending the curve, in which incidence decreases steadily and the epidemic wave is shortened.

The bending scenarios shown in Fig. 5(b) correspond qualitatively to the trend observed in Fig. 2(b). In fact, from the sequence of values \( \kappa_0(t) \) in the inset in Fig. 3, we observe that the degree of confinement achieved clearly places the epidemic in the bending regime. Also, from the evolution \( \kappa_0(t) \), we notice that after April 14 (the day when the gradual lifting of lockdown starts) a clear relaxation of confinement is observed.
If we impose the condition for mitigation and suppression of the epidemics $R \leq 1$, the critical confinement value $\kappa_0^c$ must fulfill the equation

$$
(k_0^c)^2(1 - \phi)(CH)[(k_{h+w})(1 - \delta) - \langle k_h \rangle] - \kappa_0^c[(k_{h+w})(1 - \delta) + (1 + \langle CH \rangle)(1 - \phi) - \langle k_h \rangle] + (k_{h+w})(1 - \delta) - \frac{1}{\rho^2(\frac{\rho^2}{\alpha} + \frac{\rho^2}{\mu})} = 0.
$$

Defining the following coefficients:

$$
A = (1 - \phi)(CH)[(k_{h+w})(1 - \delta) - \langle k_h \rangle],
$$

$$
B = (k_{h+w})(1 - \delta)[1 + \langle CH \rangle(1 - \phi)] - \langle k_h \rangle,
$$

$$
C = (k_{h+w})(1 - \delta) - \frac{1}{\rho^2(\frac{\rho^2}{\alpha} + \frac{\rho^2}{\mu})},
$$

we can write Eq. (46) as

$$
A(k_0^c)^2 - Bk_0^c + C = 0,
$$

whose solution is given by

$$
k_0^c = \frac{B \pm \sqrt{B^2 - 4AC}}{2A}.
$$

Note that only the physically meaningful solution $k_0^c \in [0, 1]$ must be retained.

To further analyze the relevance of the mechanisms characterizing the confinement of individuals in our model, we illustrate in Fig. 6 the effects of social distance encoded in $\delta$ and the household permeability $\phi$ on the critical confinement.
confined population needed to keep an outbreak under control is notably reduced.

V. CONCLUSIONS

As the COVID-19 pandemic affects more and more countries and threatens to overload the critical capacity of health systems, the absence of a SARS-CoV-2 vaccine requires forceful nonpharmacological containment interventions by governments. Among these interventions, the confinement of populations and social distancing have been implemented by many countries in an attempt to reduce the impact on health systems and save the time necessary to try more efficient treatments against this emerging pathogen.

To mount the strategy against SARS-CoV-2 and anticipate its trajectory, it is necessary, in addition to correctly modeling its epidemiological characteristics, to take into account as closely as possible the influence of the variability of social contacts that the pathogen uses to spread. Here, we have presented a metapopulation model to track the spatiotemporal evolution of COVID-19. This framework incorporates the essential factors of the SARS-CoV-2 transmission and clinical outcome of COVID-19 patients by considering three group of ages: young individuals (being mainly asymptomatic carriers of the pathogen), adult people (characterized by a working activity that involve recurrent mobility patterns), and the elderly (displaying the most severe and critical COVID-19 cases). Although the age partition can be further split into more groups, the former partition appears to be the minimal one to capture the main characteristics of the current epidemics. Apart from the recurrent mobility between patches (in the current work municipalities), contacts at the local level are modeled using a mean-field approach that incorporates a density-dependent function for the contact patterns between age groups. This assumption allows the straightforward consideration of household confinement and also enables one to explore contention strategies aimed at reducing the sociability within and between age-specific groups. The model constitutes an stylized framework to explore early nonpharmacological interventions when massive testing and contact tracing are not available, as was the case of the first epidemic wave in Spain and many other countries around the globe.

Analyzing in detail this model, we have shown that it is possible to construct the expression of the effective reproduction number $R(t)$ capturing both the epidemiological characteristics of COVID-19 and those social patterns that facilitate its expansion. This expression enables an accurate evaluation of the spreading potential of SARS-CoV-2 on a given population and, more importantly, the assessment of the effects of implementing or lifting nonpharmacological interventions in advance. It can be used to assess the extent of nonpharmacological interventions on a given territory seeking for the less harming intervention in terms of confinement level and time needed to epidemic suppression.

Focusing on the outbreak in Spain, we have analyzed the effects of different degrees of confinement, $k_0$, using the expression for $R(t)$ evaluated at the time $t_c$ when containment is applied. This way, we can accurately determine the effects on the slowdown and suppression of the epidemic caused by different degrees of confinement. Calculating the value $R(t_c)$, we observe a transition as the set of confined inhabitants decreases and the consequent social mixing increases. This transition defines a critical confinement $k_0^*$ that separates a supercritical scenario, in which $R(t_c) > 1$ flattens the curve, to another subcritical one, $R(t_c) < 1$, that bends the epidemic curve due to the profound change in the social contacts structure, making it almost impossible for the virus to spread.

The exact critical confinement separating the supercritical and subcritical regimes is highly dependent on the underlying social structure and the intrinsic mobility patterns of each population. Furthermore, it depends on the time it is applied, since it depends on the available pool of susceptible agents that can be culled out from the system. The generality of the expression for $R(t)$ provided here makes it possible to apply it to any population and to use with any metapopulation epidemic model, paving the way for implementing timely and well-founded epidemiological and socially based nonpharmacological responses.

ACKNOWLEDGMENTS

We thank Gourab Ghoshal, Silvio Ferreira, Carlos Bouthelier, Sergio Faci, and Miguel Hernan for useful
discussions. We also thank Matteo Rini for his suggestions. A. A., B. S., and S. G. acknowledge financial support from Spanish MINECO (Grant No. PGC2018-094754-B-C21), Generalitat de Catalunya (Grant No. 2017SGR-896), and Universitat Rovira i Virgili (Grant No. 2019PFR-URV-B2-41). A. A. also acknowledges support from Generalitat de Catalunya ICREA Academia and the James S. McDonnell Foundation (Grant No. 220020325). J. G.-G. and D. S.-P. acknowledge financial support from MINECO and FEDER funds (Project No. FIS2017-87519-P) and from the Departamento de Industria e Innovación del Gobierno de Aragón y Fondo Social Europeo (FENOL group E-19). C. G. acknowledges financial support from Juan de la Cierva-Formación (Ministerio de Ciencia, Innovación y Universidades). B. S. acknowledges financial support from the European Union’s Horizon 2020 research and innovation program under the Marie Skłodowska-Curie Grant Agreement No. 713679 and from the Universitat Rovira i Virgili (URV). W. C. acknowledges financial support from the Coordenação de Aperfeiçoamento de Pessoal de Nível Superior, Brasil (CAPES), Finance Code 001.

APPENDIX A: PARAMETERS FOR THE MODELIZATION OF THE SPREADING OF COVID-19

We now describe the choice of our parameters to study the epidemic outbreak in Spain, whose values are enumerated in Tables I–IV.

Throughout the manuscript, the patches of the metapopulation correspond to municipalities which constitute the lowest administrative divisions in Spain. Regarding the population structure, we obtain the population distribution, population pyramid, and average household size at the municipality level from Instituto Nacional de Estadística [50], whereas the age-specific contact matrices are extracted from an international social study [42]. At the beginning of the pandemic, the entire population is assumed to follow their usual mobility patterns, so $p^g = 1$ for every

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>Estimates for $g \in {Y, M, O}$ in Spain</th>
</tr>
</thead>
<tbody>
<tr>
<td>$n_i^g$</td>
<td>Region population</td>
<td>Data provided by INE</td>
</tr>
<tr>
<td>$s_i$</td>
<td>Region surface</td>
<td>Data provided by INE</td>
</tr>
<tr>
<td>$R_{ij}^g$</td>
<td>Mobility matrix</td>
<td>Data provided by INE</td>
</tr>
<tr>
<td>$(k^g)$</td>
<td>Average total number of contacts</td>
<td>(11.8,13.3,6.8)</td>
</tr>
<tr>
<td>$(k_h^g)$</td>
<td>Average number of contacts at home</td>
<td>(3.1,3.2,3.3)</td>
</tr>
<tr>
<td>$(k_w^g)$</td>
<td>Average number of contacts at work</td>
<td>(1.8,5.2,0.0)</td>
</tr>
<tr>
<td>$C_{gh}$</td>
<td>Contacts-by-age matrix</td>
<td>(0.5980, 0.3849, 0.0171)</td>
</tr>
<tr>
<td>$\xi$</td>
<td>Density factor</td>
<td>0.01 km$^2$</td>
</tr>
<tr>
<td>$p^g$</td>
<td>Mobility factor</td>
<td>(1.0,1.0,1.0)</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>Average household size</td>
<td>2.5</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>States</th>
<th>COVID-19 Spain estimates</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\beta_I$</td>
<td>Infectivity of symptomatic</td>
<td>$S \rightarrow E$</td>
<td>0.075</td>
<td>Calibrated</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>95% CrI: 0.068–0.082</td>
<td></td>
</tr>
<tr>
<td>$\beta_A$</td>
<td>Infectivity of asymptomatic</td>
<td>$S \rightarrow E$</td>
<td>0.5$\beta_I$</td>
<td>Assumed</td>
</tr>
<tr>
<td>$\eta^g$</td>
<td>Exposed rate</td>
<td>$E$</td>
<td>2.444 days$^{-1}$</td>
<td>Calibrated</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>95% CrI: 1.823–3.064</td>
<td></td>
</tr>
<tr>
<td>$\alpha^g$</td>
<td>Asymptomatic rate</td>
<td>$A$</td>
<td>(5.671, 2.756, 2.756) days$^{-1}$</td>
<td>Calibrated</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>95% CrI$^2$: 5.046–6.356</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>95% CrI$^4$: 2.135–3.377</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>95% CrI$^6$: 2.135–3.377</td>
<td></td>
</tr>
<tr>
<td>$\mu^g$</td>
<td>Infectious rate</td>
<td>$I$</td>
<td>(1.3,915, 3.915) days$^{-1}$</td>
<td>$Y$: Assumed</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>95% CrI: 3.470–4.360</td>
<td>$M$, $O$: Calibrated</td>
</tr>
</tbody>
</table>
To construct the daily mobility network, we first assume that the elderly and young population do not leave their residential municipality, so their corresponding mobility matrices read \( R_{Y,M,O}^{i,j} = \delta_{ij} \). Finally, the mobility matrix governing adult population movements is estimated from extensive surveys conducted by Instituto Nacional de Estadística in 2011 [50], which capture the commuting patterns of the population living inside each municipality.

Regarding epidemiological parameters, the incubation period is reported to be \( \eta^{-1} + \alpha^{-1} = 5.2 \) days [2] in average which, in our formalism, must be distributed into the exposed and asymptomatic compartments. In principle, if one does not expect asymptomatic transmissions, most of this time should be spent inside the exposed compartment, the asymptomatic infectious compartment thus being totally irrelevant for disease spreading. However, along the line of some works [51–53], we find that the unfolding of COVID-19 cannot be explained without accounting for infections from individuals not developing any symptoms previously. In particular, after calibrating the model, we estimate \( \alpha^{-1} = 2.756 \) (95\% CrI: 2.135–3.377) days as the asymptomatic infectious period; see Appendix C for the details of the calibration methodology. In turn, the infectious period while being symptomatic is calibrated to \( \mu^{-1} = 3.915 \) days (95\% CrI: 3.470–4.360), except for the young stratum, for which we reduce it to 1 day, assigning the remaining days as asymptomatic; this reduction is due to the reported mild symptoms in young individuals, which may become unnoticed [28]. Furthermore, we assume asymptomatic individuals are half as infectious as symptomatic ones, \( \beta_A = 0.5\beta_S \), as in Refs. [54,55].

Regarding the clinical parameters, we fix the fatality rate \( \omega = 30\% \) for ICU patients according to official reports [37]. We incorporate from previously published studies the typical time from ICU admission to death as \( \gamma^{-1} = 7 \) days [39] and the stay in ICU for those overcoming the disease as \( \psi^{-1} = 21 \) days [40]. In turn, the parameters \( \gamma^0 \) and \( \theta^0 \), controlling the infected population requiring ICU beds and dying without occupying any of them, are estimated by correcting the real observed values [37] with the estimated underreporting from Ref. [38] (see next paragraph). Likewise, the parameters \( \zeta^0 \) and \( \lambda^0 \) are computed by assuming that times between onset of symptoms and ICU occupation or fatal outcome are 8 and 11 days [37] on average, respectively.

To account for underreporting, we compare the seroprevalence estimates of the extensive seroepidemiological extensive survey [38] made by the Spanish authorities with that derived from the official reported data. In particular, 0.53\% of the Spanish population was diagnosed as COVID-19 cases by May 4.
whereas the actual seroprevalence observed was 5.0% (95% confidence interval CI: 4.7–5.4). Therefore, we estimate that 89.4% (95% CI: 88.7–90.2) of COVID-19 cases have not been diagnosed in Spain and use this value to rescale the time series for the daily new symptomatic cases shown in Fig. 2(b).

Finally, regarding the parameters controlling the efficiency of the measures taken by the authorities, we calibrate those depending on social awareness and individual behavior, such as social distancing $\delta$ and household permeability $\phi$, due to the lack of data. In contrast, the fraction of the population under confinement $\kappa_0(t)$ is

![Calibration analysis](image_url)
straightforwardly incorporated from data on the mobility reduction daily updated by INE which are shown in Fig. 3.

Figure 7 contains the posterior distribution for the calibrated epidemiological parameters along with the covariance matrix showing the relations between those parameters involved in the calibration.

APPENDIX B: NORMALIZATION FACTOR FOR EACH AGE GROUP

Here we want to explain the rationale behind the normalization factor \( z^g \), first introduced in Eq. (14), which shapes the number of contacts made by individuals belonging to age group \( g \). In the main text, we state that \( z^g \langle k^g \rangle s_i n_i \) represents the total number of contacts that an agent from age group \( g \) makes inside patch \( i \). To construct this term, let us start by assuming that the number of contacts within a patch increases monotonically with the density according to a given function \( f \). Therefore, in general terms, we can write the number of interactions of one individual belonging to age-group \( g \) inside patch \( i \), \( k^g_i \), as

\[
    k^g_i = x^g f \left( \frac{n_i}{s_i} \right),
\]

being \( x^g \) an (in principle) arbitrary constant which may differ for the different age-groups \( g \). In fact, \( x^g \) allows for re-scaling the contacts such that the average number of interactions of individuals belonging to strata \( g \) across the entire population is \( \langle k^g \rangle \). This constraint imposes that

\[
    \sum_{i=1}^{N_P} \tilde{n}_i^g x^g f \left( \frac{n_i}{s_i} \right) = n^g \langle k^g \rangle.
\]

This way, \( x^g \) can be expressed as

\[
    x^g = z^g \langle k^g \rangle,
\]

where \( z^g \), which is defined as the normalization factor throughout the manuscript, is given by

\[
    z^g = \frac{n^g}{\sum_{i=1}^{N_P} \tilde{n}_i^g f \left( \frac{n_i}{s_i} \right)}.
\]

APPENDIX C: MODEL CALIBRATION

Several procedures are proposed in the literature to calibrate the mathematical models aimed at reproducing the evolution of COVID-19 across a given country. Calibration using indicators such as ICU occupation, daily incidence, or the number of fatalities rely on the quality of the data obtained. Unfortunately, in Spain, the data acquisition process is very messy, given the different protocols that every Comunidad Autónoma follows. For example, the number of officially reported cases does not constitute a reliable indicator in Spain because of delays in reporting to the National Health system as well as temporal variations in the number of tests. This unreliability makes reported cases a very noisy variable to fit. In front of this important problem, we finally decide to calibrate the model with the number of daily fatalities, data that are curated by the Spanish government.

The calibration procedure consists of the following steps:

1. Most of the parameters governing the clinical outcome of the disease are incorporated from either previous studies or official reports and do not take any part in the calibration of the model. Therefore, the calibration is mostly focused on the epidemiological parameters governing the unfolding of the disease, whose values remain still uncertain. In particular, the parameters subjected to calibration are the number of initial infectious seeds \( A_0 \), the infectivity \( \beta_i \), the asymptomatic infectious period \( \alpha^{-1} \), and the infectious symptomatic period \( \mu^{-1} \). Finally, regarding the confinement, household permeability \( \phi \) and social distancing \( \delta \) need also to be calibrated due to the lack of data, whereas the values for \( k^0_0(t) \) are directly incorporated in the model from mobility reduction estimations published by INE.

2. The initial date from which our model runs is February 9. To spatially distribute the initial seeds, the reported cases by March 3 are backtracked until February 9 and let the model run. Once the initial seeds are set, we calibrate the epidemiological part of the model by performing an approximate Bayesian computation (ABC) [45] including real data about the new daily fatalities at a national scale until April 8.

We use the logarithmic least squares error as an objective function for the ABC. The daily fatalities from April 8 onward [Fig. 2(a)] as well as the daily new cases for the whole time span [Fig. 2(b)] serve as validation. As stated in Sec. A, the time series for the daily new cases provided by the health authorities is rescaled according to the findings of the nationwide serological survey to correct for the underreporting.

3. Our model is able to predict the spatiotemporal propagation of COVID-19 triggered by community contagions and human mobility. To leverage the full potential of the formalism, we also check the accuracy of the formalism in predicting the fatalities at a regional level. We observe that the introduced seeds corresponding to the reported cases by March 3 are enough to predict the fatalities observed in some areas but fail in capturing the evolution of COVID-19 in other regions, possibly due to initial underreporting of cases there. To solve this discord-
ance, we introduce the minimal set of infectious individuals in those largest populated cities of each Comunidad Autónoma to ensure a fair correspondence between the data and model, which is optimized by using the Nelder-Mead method.

APPENDIX D: SENSITIVITY ANALYSIS

Here, we perform a sensitivity analysis to check the robustness of our results concerning the time evolution of the effective reproduction number. As the value for $R(t)$ is independent from the clinical parameters, we restrict the sensitivity analysis to those epidemiological parameters assumed to be constant when calibrating the model. In particular, we focus on studying how the values of $R(t)$ in both supercritical and subcritical regimes are affected by changes in the incubation period $\eta^{-1} + \alpha^{-1}$, the infectivity of individuals without symptoms $\beta_A$, and the infectious symptomatic period of young population $(\mu^Y)^{-1}$.

Figure 8 compares the original distribution for the effective reproduction number before and after confinement is enforced with those obtained by introducing a 10% relative modification in each of the aforementioned parameters. The mean and the shape of the distribution remain almost unaltered in both supercritical and subcritical regimes, thus proving the robustness of the results reported throughout the manuscript.

APPENDIX E: VALIDATION OF EFFECTIVE REPRODUCTION NUMBER

The validation of our analytical expression Eq. (42) for the effective reproduction number is performed by comparing it with the case reproduction number obtained using EpiEstim [48,49]. We apply EpiEstim to both the incidence in our model (EpiEstim model) as well as the real incidence data (EpiEstim data); see Fig. 9. The incidence in our model corresponds to the one of the compartment exposed ($E$). The real incidence data, collected until May 15, are shifted by 6 days due to the average lag between symptom onset and report [37]. The data are then subsequently shifted another 5 days accounting for the time in our model between becoming exposed and developing symptoms. Estimations of the reproduction number are then performed on both datasets by averaging over 50 randomly chosen parameter sets of our posteriors. For each parameter set, the generation time distribution is fixed correspondingly. Additionally, we perform a rolling average of 10 days on all curves. Please note that, for the EpiEstim data curve, the confidence interval is narrower than the line’s thickness. The curve is plotted until the last day EpiEstim is able to perform an estimation.

We observe a good agreement between the analytical expression and the estimation by EpiEstim on the incidence of our model. Qualitatively, also the estimation using real incidence data matches the analytical expression. In particular, the difference on which day the two curves drop below one is only around 2 days. The fact that the

![FIG. 8. Sensitivity analysis. Distribution of effective reproduction number $R$ values while exploring the parameter space by tuning parameters $\beta_A/\beta_I$, $\eta^{-1} + \alpha^{-1}$, and $(\mu^Y)^{-1}$ as described in Appendix D on (a) a supercritical regime (2020-03-01) and (b) a subcritical regime (2020-04-01). As a reference, we provide a baseline distribution obtained fixing the aforementioned parameters to the values reported in Table III.](image)

![FIG. 9. Validation of effective reproduction number. Reproduction number from the analytical expression compared to an estimation of the case reproduction number obtained using EpiEstim [48,49] on the incidence in our model (EpiEstim model) as well as the real incidence data (EpiEstim data).](image)
estimation on real incidence data results in a higher reproduction number at the beginning of the epidemics is probably due to the quickly increasing test capacity as the first cases were detected in Spain. Instead of observing the actual growth of the epidemic, tests performed in the beginning discovered to what extent the epidemic had already spread in the population. Unfortunately, there are no data available on how the number of performed tests evolved until April 23.

APPENDIX F: CODE AVAILABILITY

The model described above is implemented in the open source Julia programming language [56]. The source code is freely available online in Ref. [57], distributed under the open source licence GNU AGPL-3. The documentation can be read in Ref. [58].

For the ABC calibration described in Appendix C, we use the AproxBayes Julia package [59], that implements the approximate Bayesian computation method defined in Ref. [45].


[59] M. J. Williams, *ApproxBayes.jl (v0.3.2)*, https://github.com/marcjwilliams1/ApproxBayes.jl.
S1. EPIDEMIC SCENARIOS

Apart from capturing the past course of a real pandemic and forecasting its future evolution, the framework here presented constitutes an useful tool for evaluating different interventions of interest for policy making. In this regard, the complexity of the equations governing the evolution of an age-stratified metapopulation, allows for incorporating realistic interventions and assessing their performance in reducing the impact of the disease over the population. Taking advantage of this possibility, we now study four different scenarios: the constraint of the inter-territorial mobility, the closure of the epicenter of the pandemic in Spain, and the selective confinement to elderly and young population and the extension of the strict lockdown regime.

A. Constraining mobility among provinces

First, we aim at modifying the morphology of the mobility network to unveil the role that human flows over different territories have played in fostering the spread of COVID-19. In particular, we are interested in revealing the relevance of the mobility network for the dissemination of the first cases in Spain until lockdown was enforced.

For this purpose, we compare the evolution of the spatial diffusion of COVID-19 with the picture provided by an alternative scenario in which the mobility between provinces (the administrative division of Comunidades Autónomas in Spain) is forbidden since the beginning of the disease. In mathematical terms, this policy is reflected in the mobility network by turning every flow connecting different provinces into self-loops, so that the population following these mobility paths stay inside their municipality. It is important to stress that this mobility restriction does not imply any kind of household confinement.

B. Closing the epicenter of the pandemic

Madrid became the epicenter of the pandemic during the first epidemic wave in Spain. The international connectivity facilitating the arrival of imported cases along with a massive community transmission driven by the high population density existing in Madrid lead to the rapid unfolding of COVID-19 epidemics there. In the presence of clear epidemic centers, one typically discussed intervention consists in isolating these zones from the rest of the territory. To have quantitative information about the impact of such intervention, we remove the flows in the mobility network involving Madrid and analyze the course of the epidemic.

In Fig. S2 we represent the time evolution of the relative difference number of cases when Madrid is isolated with its corresponding to the original mobility network. Positive values indicate a beneficial effect of the policy whereas negative values encode a detrimental impact of Madrid closure. Overall, restricting Madrid mobility has a positive impact in the course of the epidemic. Nonetheless, note that the effect of the geographical closure of the most affected areas is not very pronounced, for it should be combined with strong local control policies implemented inside them to efficiently control the outbreak.

C. Selective confinement of non-active population

A crucial factor to understand the disparate COVID-19 numbers across similar countries is the timeliness of

Figure S1 shows the distribution across Spain's territory of the municipalities with more than 10 cases as of March 15 when no mobility restriction is at work (left) and when the inter-provincial mobility is blocked (right). The maps in this figure clearly reveal the relevance of inter-provincial mobility at the early stages of the disease. In particular, we check that those connections disseminated the initial infectious seeds across more municipalities, thus making more difficult to concentrate efforts to fight local outbreaks.
the policies promoted to keep its propagation under control. One typical example can be found in Europe during the first epidemic wave: while countries such as Austria or Portugal deployed early interventions, thus alleviating the impact of COVID-19, other countries such as Spain or Italy acted late when the disease was already widespread across their territories. In this sense, the fear of dismantling the socio-economic fabric of the countries when isolating the active population constitutes one of the most important conditioning factor preventing these countries from acting earlier.

In light of this problem, one alternative intervention, aimed at keeping most of the country functionality and improving the control of the epidemic, would have been to enforce an early selective confinement of the non-active population. To capture this intervention in our formalism, we restrict the contacts of the young and elderly population to those taking place inside their households at a given time $t_c$ before the state of emergency was declared in Spain (March 14, 2020). After this day, the lockdown of young and elderly patterns is again coupled to the one of the active population.

In Fig. S3 we illustrate the huge benefit of cutting possible transmission chains by performing timely selective confinements at the early stages of the disease, i.e., when the number of cases grows exponentially. In particular the results are obtained when this age-selective confinement is applied on February 28.

### D. Extension of strict lockdown period

Finally, we address the benefits of prolonging the second lockdown period in Spain when every activity with
Figure S3. Effect of enforcing a selective confinement to non-active population (see text for details) on the course of the pandemic in Spain. The course is characterized by the time evolution of the cumulative number of cases (a) and new daily cases (b) for both the baseline and the intervened (YO) scenarios. Colored ribbons cover a 95% prediction interval.

<table>
<thead>
<tr>
<th>Scenarios</th>
<th>Cases</th>
<th>Daily new cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Real</td>
<td>[2,060,526 - 3,347,466]</td>
<td>[2,720 - 19,273]</td>
</tr>
<tr>
<td>1 week</td>
<td>[2,037,360 - 3,175,669]</td>
<td>[1,700 - 11,731]</td>
</tr>
<tr>
<td>2 weeks</td>
<td>[1,994,002 - 3,047,854]</td>
<td>[968 - 6,845]</td>
</tr>
<tr>
<td>4 weeks</td>
<td>[1,977,928 - 2,970,631]</td>
<td>[449 - 3,437]</td>
</tr>
</tbody>
</table>

Table S1. Predictive interval on the number of cases and daily new cases for the different scenarios considered in Fig. S4.

the exception of essential services was banned (EM window in Fig. 2 of the main text). For this purpose, we depict the evolution of the number of cases and daily new cases under the observed mobility and when increasing the duration of the EM lockdown. To do so, we extend the time series \( \{\kappa_0(t)\} \) characterizing the evolution of the mobility reduction over the EM lockdown (see Fig. 3 of the main text) during three different periods: 1 week, 2 weeks or 4 weeks. In each of the three cases, after the extra lockdown window, the evolution of the mobility is kept as it was observed after the original EM period. The predictions show a reduction in both metrics as we increase the weeks of lockdown, showing a clear decreasing trend in the number of daily new cases on the most restrictive scenario. Remarkably, simulations also show considerable shrinking of the uncertainty range for daily new cases as we add weeks to the confinement policy, see Table S1 and Fig. S4. Namely, the upper bound goes from 19,273 daily new cases for the real scenario to just 3,437 for the four-week lockdown extension scenario. Although a complete stop of the spreading is almost impossible to achieve without a vaccine or clinical treatment, reducing the incidence to a manageable value through an extension on the lockdown duration could have had an essential role in mitigating, or at least delaying the current second wave.
Figure S4. Evolution of the number of cases (a), and daily new reported cases (b) as function of time. Lines represent different scenarios where the total lockdown was extended starting on April 10 one week, two weeks, four weeks. For comparison purposes we added the real scenario. Colored ribbons covers a 95% prediction interval.