

Key Interactions and Exit Mechanisms for Covid-19

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Abstract

We develop a minimalist compartmental model to analyze the impact of mobility restrictions in Italy during the Covid-19 outbreak. Our findings show that early lockdowns shifts the epidemic in time and that beyond a critical value of the lockdown strength, the epidemic restarts after lifting the restrictions. We investigate the effects of different lockdown scenarios and exit mechanisms by accounting for two fundamental sources of heterogeneity within the model: geography and demography. First, we consider Italian Regions as separate administrative entities, in which social interactions between age cohorts occur. Due to the sparsity of the mobility matrix, epidemics tend to develop independently in different regions. Second, we show how disregarding the specific structure of social contacts between age classes could lead to severe underestimation of a second wave of infection at the end of the lockdown period, while measures which take into account the age of the population can sustain the mitigation of rebound effects. Our model is general, and it highlights the effects of key parameters on non-pharmaceutical mitigation mechanisms for epidemics.

1 Introduction

Different epidemic models and approaches contribute to identify specific mechanisms relevant for policy design [1]. At present, although the World Health Organization (WHO) organizes regular calls for Covid-19 modelers to compare strategies and outcomes, policymakers barely handle the discrepancies between the proposed models.¹

To contain the Covid-19 epidemic, governments worldwide have adopted severe social distancing policies, ranging from partial to total population lockdown [2]. Restrictions have led to a sudden stop of economic activities in many sectors, while the majority of Covid-19 infections affect active population (i.e. the age cohort between 15–64 years) [3]. Overall, the impact of contagion and lockdown measures on health and on economic activities is substantial and pervasive.

Against this background, we introduce a model-based scenario analysis for Covid-19, highlighting the role of non-pharmaceutical variables in the epidemic spreading and how those dimensions are likely to impact on lockdown policies and exit strategies [4].

The general behavior of our model holds for the vast class of epidemic models where transmission rate is proportional to the number of susceptible people times the density of infected. We focus on the determinants of short-term interventions in response to an emerging epidemic when geographic and demographic compartments are included in the model. Our goal is general in nature, since we focus on two relevant decomposability conditions, under which partial dynamics influence the overall configuration of the system (see, e.g., [5, 6, 7, 8]). We study how mobility restriction measures and the timing of the lockdown release affect the total fraction of infected, the peak prevalence, and, possibly, the delay of the epidemic. Our analysis identifies two fundamental sources of heterogeneity in the diffusion process: regional boundaries and age cohorts [4]. We show how such dimensions can shape policy interventions aiming at containing the epidemic, irrespective of any detailed quantitative predictions. Then, we point out how disregarding such dimensions would affect exit scenarios.

With this paper, we contribute to the extant literature on trade-offs between mitigation, aimed to slow down the epidemic contagion, and suppression, consisting of temporarily lowering the risk of contagion [4, 9, 10]. Notwithstanding detailed quantitative information and relevant medical issues are not addressed here, our work still reveals how a simple and parsimonious compartmental model based on geographical and age classes uncovers relevant aspects which may help guiding decision makers in balancing the restrictions of the lockdown phase and the timing of its release. First, we show that early lockdowns shift the epidemic in time and that the delay is proportional to the anticipation time with an intensity which grows with the strength of the lockdown. Beyond a critical threshold, the epidemic would tend to fully recover its strength as soon as the lockdown is lifted. Second, we show how the sparsity of the matrix representing mobility flows across administrative regions accounts for the delays of

¹See, e.g.: <https://www.sciencemag.org/news/2020/03/mathematics-life-and-death-how-disease-models-shape-national-shutdowns-and-other>

the contagion and implies that once the epidemic has started, it then develops independently. Finally, we consider social contact heterogeneity between age cohorts and we find that the structure of social contacts is of primary importance to estimate post-lockdown effects so that age based strategies are a key ingredient to mitigate rebound effects.

2 Model

To analyze mobility-restriction policies, we introduce a minimalist compartmental model [10, 11]. Although many models, both mechanistic, statistic and stochastic [12], have been proposed for the Covid-19 infection, data collected from the national healthcare systems suffer from the lack of homogeneous procedures in medical testing, sampling and data collection [13]. Not to mention the difficulties in assessing the impact of variability in social habits during the epidemics [10, 14]. Moreover, especially in the early phases of the epidemic – i.e. the ones characterised by an exponential growth – different models sharing a given reproduction number R_0 can fit the data with equivalent accuracy (see Sec. 8.1). For these reasons, our aim is to focus on some fundamental qualitative scenarios and not on detailed predictions. We adapt the *SIR* model, the most basic epidemic model for flu-like epidemics, to the observed data available in the Italian case.

The model relies on four compartments, namely: S, I, O, R . Hence, S (usceptible) individuals can become I (nfective) when meeting another infective individual, I (nfectives) either become O (bserved) – i.e. present symptoms acute enough to be detected from the national health-care system – or are R (emoved) from the infection cycle by having recovered; also O (bserved) individuals are eventually R (emoved) from the infection cycle (see Fig. 9 for a visual representation of the model workflow). Notice that, it is not still clear if there is an asymptomatic phase [15, 16]; we are implicitly assuming that asymptomatics are infective and their recovery time is the same of the I class. The model is described by the following differential equations:

$$\begin{aligned}\partial_t S &= -\beta S \frac{I}{N} \\ \partial_t I &= \beta S \frac{I}{N} - \gamma I \\ \partial_t O &= \rho \gamma I - h O \\ \partial_t R &= (1 - \rho) \gamma I + h O\end{aligned}\tag{1}$$

$N = S + I + O + R$ is the total number of individuals in a population, the transmission coefficient β is the rate at which a susceptible becomes infected upon meeting an infected individual, γ is the rate at which an infected either becomes observable or is removed from the infection cycle. Like the *SIR* model, the basic reproduction number is $R_0 = \beta/\gamma$; the extra parameters of the *SIOR* model are ρ , the fraction of infected that become observed from the national

health-care system, and h , the rate at which observed individuals are removed from the infection cycle. Notice that we consider that O (bserved) individuals not infecting others, being in a strict quarantine.

3 The Italian Lockdown

The Italian lockdown measures of the 8th and 9th of March [17, 18] were intended to change mobility patterns and to reduce de-visu social contacts, through quarantine measures and to an increased awareness of the importance of social distancing. We analyze an extensive data set on Facebook mobility data² [19]; our analysis confirms that the lockdown has reduced both the travelled distance and the flow of travelling people.

We consider the effects of such lockdown measures on the parameters of our model. Lockdowns are non-pharmaceutical measures; hence the rate γ is the most unaffected, since it is related to the “medical” evolution of the disease. Analogous arguments apply to the rate h of exiting a condition serious enough to be observed and to the probability ρ of being observed by the national healthcare system (although ρ could be influenced variations in testing schemes and alert thresholds). On the other hand, the transmission coefficient β can be thought as the product $C\lambda$ of a contact rate C times a disease-dependent transmission probability λ . Hence, if we assume that the speed of Covid-19 mutation is irrelevant on our timescales, lockdown strategies mostly influence β by reducing the contact rate C among individuals.

To adapt the *SIOR*’s parameters to the Italian data [20], we compare the reported cumulative number of Covid-19 cases Y^{Obs} with the analogous quantity $Y^{\text{model}} = \int \rho \gamma I dt$ in our model. We want to stress that our model fitting is not aimed to produce an accurate model for detailed predictions, but to work in a realistic region of the parameter space.

We first estimate model’s parameters by least square fitting on the pre-lockdown period. Since in such range the data Y^{Obs} show an exponential growth trend, we are possibly observing a very early phase of the epidemic, where $\beta - \gamma$ equals the growth rate of Y^{Obs} (see Sec. 8.1). For fixed $\beta - \gamma$, the time of the epidemic start (that we conventionally assume as the time t_0 where the number of infected is 1) and the fraction ρ of serious cases observed by the national healthcare service, allows to vary the values of β and γ as long as their difference is fixed. Hence, estimating *medical* parameters as the rate γ of escaping the infected state is paramount for calibrating mathematical models.

In response to the outbreak of Covid-19, several estimates of model parameters have been proposed in the literature, revealing a certain amount of uncertainty about some fundamental variables of the epidemic contagion. The European Centre for Disease controls reports an infection time duration τ_I between 5 and 14 days [21]; in our model, we will use $\tau_I = 10$ (i.e. $\gamma = \tau_I^{-1} = 1/10 \text{ days}^{-1}$).

²Those data are part of the Facebook project “Data for Good”, and illustrate mobility patterns of fb users, who allowed the social network to track their location. See <https://dataforgood.fb.com/docs/Covid-19/>

$\beta = 0.35 \text{ day}^{-1}$	$\gamma = 10^{-1} \text{ day}^{-1}$	$h = 1/9 \text{ day}^{-1}$
$t_0 = -30 \text{ days}$	$\rho = 40\%$	$\alpha = 0.49$

Tab. 1. Standard parameters used for the *SIOR* model in the paper.

According to a report of ISS, the Italian National Health Institute, the time from the start of serious symptoms (i.e. when one gets “observed” from ISS) to the resolution of the symptoms can be estimated as $\tau_H \sim 9 \text{ days}$ [22], corresponding in our model to a value $h = 1/9 \text{ days}^{-1}$. Notice the analysis of 12 different models [12] reports varying estimates for the basic reproduction number R_0 , ranging from 1.5 to 6.47, with mean 3.28 and a median of 2.79.

From fitting the 15 days of Y^{obs} (pre-lockdown phase) and by performing a bootstrap sensitivity analysis of the parameters, we obtain $\beta - \gamma \sim 0.25 \pm 0.01$ and $t_0 = -30 \pm 5 \text{ days}$ by assuming that $\rho = 40\%$. Varying ρ in $[10\%, 100\%]$ varies $\beta - \gamma$ in $[0.22, 0.27]$. On the other hand, for fixed $\beta - \gamma$, R_0 would vary linearly with τ_I ; as an example, R_0 varies in $[2.5, 4.5]$ for the literature parameters $\tau_I \in [5, 14]$; accordingly, to adjust the difference in growth rate, t_0 varies in $[26, 32]$. However, despite the variability of the parameter range, the qualitative behavior of the model – and hence our analysis of the key factors of the epidemic evolution – is unchanged.

We then assume that, after the lockdown day $t_{\text{Lock}} = 15$ (corresponding to the 9th of march), contact rate drops down by a factor α and hence $\beta \rightarrow \alpha\beta$. By fitting the observed data Y^{obs} for a symmetric period of 15 days after t_{Lock} , and by performing a bootstrap sensitivity analysis, we find $\alpha = 0.49 \pm 0.01$, i.e. a $\sim 50\%$ reduction in infectivity and hence in R_0 . Our figure is in line with the observed reduction in R_0 in response to the combined non-pharmaceutical interventions, that across several countries has an average reduction of 64% compared to the pre-intervention values [23]. Notice that Facebook mobility data show a post-lockdown reduction in mobility of 15% at regional level and of 73% at inter-regional level; however, as we will point out later, mobility is much more related to the beginning of epidemics in new regions/countries than to the evolution of the epidemics in a region/country.

In the following of the paper, we will use the exemplified parameters of Tab. 1 corresponding to a basic reproduction number $R_0 = 3.5$. Moreover, since patients in intensive care represent the highest burden for the national health systems, in the graphs of the paper we will indicate the number of patients in intensive care, estimated as 3.5% of the total patients by using the figures reported by ISS [20].

4 National scenarios and exit mechanisms

Since we are interested on the factors driving the exit dynamics from lockdown, and not on the detailed analysis or production of realistic scenarios, we will consider several lockdown scenarios where the lockdown is abruptly released and

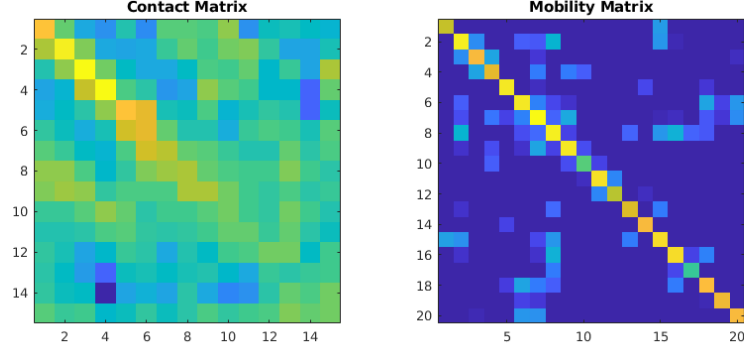


Fig. 1. Left Panel: social contact matrix, from [24]. Right panel: inter-regional mobility matrix, from the Facebook project “Data for Good”. The intensity of a color maps the strength of a matrix element (light colors: high values; dark colors: low values). The inter-age social mixing matrix is dense; hence age classes dynamics are strongly coupled. The inter-regional mobility flows is very sparse (i.e. off diagonal elements are order of magnitudes lower than diagonal elements): this mean that most of the people travel within the same region of origin; hence, the regional dynamics can be considered “almost” decoupled.

the system let return to the pre-lockdown parameters. Such an approach amplifies the possible perils and represent a worst-case estimate of the exit strategies. Hence, we consider several simplified scenarios where we use the *SIOR* model of Eq. 1 with the parameters of Tab. 1: First, in the simple case of a *SIOR* model fitted on Italian data, we analyze how the post-lockdown dynamics changes according to different starting dates and to different levels of the restrictions implemented by the national authorities. Then, we study the effect of explicitly considering Italy as a collection of separate administrative entities (Regions) independently evolving; finally, we consider the effects of social interactions across age cohorts.

Interestingly, mobility flows [19] and inter-age social mixing [24] lie at the two opposite range of modelling. In fact, the regional social contact matrix is dense (Fig. 1, left panel), indicating that age classes dynamics are strongly coupled. On the other hand, the inter-regional mobility matrix is very sparse (Fig. 1, right panel), indicating that regions have their own independent dynamics.

We first consider a simple exit strategy consisting in releasing the lockdown at a time t_{Unlock} after the peak of O has occurred. For instance, we hypothesize that infection proceeds uncontrolled up to time t_{Lock} ; in the following lockdown period $[t_{\text{Lock}}, t_{\text{Unlock}}]$, the transmission coefficient β is reduced by a factor α ; finally, β returns to its initial value and herd immunity is responsible for the dampening of the epidemics.

Our results show that the lockdown lowers the peak of O - i.e. the individuals with noticeable symptoms - to $\sim 70\%$ of the free epidemic one, but also doubles

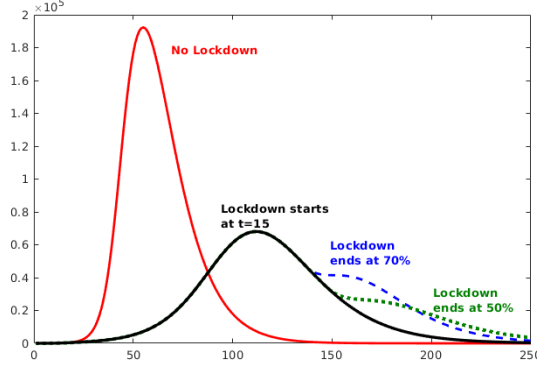


Fig. 2. Comparison of the scenarios where the lockdown is relaxed after the percentage of people with visible symptoms (O) is reached the 70% and the 50% of the reported cases peak. Releasing the lockdown earlier has the epidemics disappear faster, but has higher impact on the number of hospitalized and intensive care patients; moreover, lifting the lockdown too early can result in a rebound of the number of cases.

the time of its occurrence from ~ 1.9 months to ~ 3.8 months: an extremely obnoxious effect for the sustainability conditions of the economy of a country. However, since the number of hospitalized patients and - most importantly - the number of patients in intensive care is only a fraction of O , lowering the peak puts less stress on the healthcare system. The ideal situation would be to have accurate data, an accurate model and accurate estimates of the parameters; as an example, in our model lifting the lockdown when the number of infected people per unit time $\beta S(t)I(t)/N$ is lower than the average number of recovering people $\gamma I(t)$ would ensure that the number of infections would continue to decrease. In real life, situations are more fuzzy: having not enough information, we could decide to resort on some heuristics, like lifting the lockdown after the observed people O have dropped to a suitable percentage of the maximum peak. As an example, after ~ 4.7 months the peak has reduced to 70% of its initial value, while after ~ 5.2 months to 50%, i.e. ~ 0.5 months later. Notice that, the earlier the lockdown is lifted, the faster O decays to zero even if it starts from higher figures and could even experience a rebound. All such effects are shown in Fig. 2.

Our framework sustains the identification of several mechanisms. The first is related to the timeliness of the lockdown, i.e. to the choice of anticipating t_{Lock} . As expected, early lockdown (i.e. well before the “free” infection peak) reduces the height of the peak without much moving it forward in time. Conversely, lifting the lockdown too soon can make epidemic start again and reach values higher than the ones before the release. A peculiar and counter-intuitive effect can be generated if the lockdown is anticipated: in fact, a too early lockdown

delays the start of the epidemic without attenuating its severity (see Sec. 8.2). In other words, an early lockdown buys time, but it postpones the problem without mitigating its severity.

Another effect is the impact of extreme quarantine measures on the post-lockdown dynamics. Increasing the strength α of the lockdown (i.e., reducing the social contacts), not only delays the time at which the lockdown can be lifted, but it also induces a stronger re-start of the epidemic in the post-lockdown (see Sec. 8.2). Moreover, such an approach could result to be not sustainable for the economy, motivating a gradual lifting of the lockdown measures to lessen the extent of further peaks.

An additional counter-intuitive mechanism must be considered. Since to an attenuation α corresponds an effective reproduction number $R_0^{\text{eff}} = \alpha R_0$, at the critical value $\alpha_{\text{crit}} = 1/R_0$ the epidemic neither grows nor decreases³. Thus, after t_{Lock} the system stays stationary until the quarantine is released at t_{Unlock} ; at this point the epidemic starts growing again as it was before the lockdown. In general, if $\alpha < \alpha_{\text{crit}}$, the system looks to ameliorate (infected, hospitalized, all the infective compartments go down) but as soon as the lockdown is lifted, the epidemic starts again to reach its full extent (see Sec. 8.2). Nevertheless, our estimate $\alpha \sim 0.5 > \alpha_{\text{crit}} \sim 0.3$ for the Italian lockdown gives us hope that, perhaps, it will not be necessary to follow a repeated seek-and-release strategy in the post-lockdown phase. On the other hand, if it can be attained a lockdown strength $\alpha \sim \alpha_{\text{crit}}$ without disrupting the economy, the epidemic could be contained until the creation, production and distribution of a vaccine.

5 Regional Scenarios

Starting with the first confirmed cases in Lombardy on 21 February, by the beginning of March the Covid-19 outbreak had spread to all regions of Italy. While the delay in the beginning of the infection is due to the different interaction among regions, once an epidemic has started it grows exponentially and the intake of external infected people becomes quickly irrelevant (see Sec. 8.6). As a consequence, the growth curves of the epidemic variables should tend to the same shape (see Sec. 8.5). Coherently, looking at the regional info graphics released by the Italian National Healthcare Institute (ISS) [20], one may notice that they have a similar shape but different starting date (see Fig. 3). Such observation can be justified as follows: Italian regions are independent administrative units, where most of the population tend to work inside the resident region [25]. Hence, epidemics propagate from region to region via the fewer inter-regional exchanges (notice that Lombardy is among the Italian regions most involved in international trade connections [26], therefore it appears as one of the most probable candidate for the outbreak of the Italian epidemic). More practically, we estimate these delays by minimizing the distance among the observed curves (see also Sec. 8.4); results are reported in Tab. 2. Notice

³To be precise, the decrease becomes sub-exponential, thus taking a practically infinite time in populations as big as a country or even a megacity

Tab. 2. Regional delays (in days)

Lombardia	0.0	Molise	10.6
Emilia Romagna	3.1	Umbria	11.8
Marche	4.3	Abruzzo	13.1
Veneto	5.7	Lazio	14.5
Valle d'Aosta	6.4	Campania	15.0
P.A. Trento	6.6	Puglia	15.7
P.A. Bolzano	8.0	Sardegna	16.2
Liguria	8.1	Sicilia	16.6
Friuli Venezia Giulia	8.9	Calabria	17.2
Piemonte	9.0	Basilicata	19.2
Toscana	10.4		

that, assuming Lombardy has been the first region (i.e. delay=0), the resulting regional delays are mostly correlated to geographical distances.

We assume that the Covid-19 outbreak spreads independently in each region of Italy; as argued before, such an approximation is reasonable after the epidemic has started and is even more accurate under lockdown conditions. Hence, we apply the parameters for the whole Italy to regional cases⁴, where now the maximum number of individuals N_i is the population of the i^{th} region [CITAZIONE_PER_LA_DEMOGRAFIA]. Then, by summing up all the S_i, \dots, R_i , respectively, we obtain a proxy for the global evolution of Covid-19 epidemic throughout Italy. To evaluate the effect of heterogeneity in time delays, we compare the number of daily cases $O^{\text{Delay}} = \sum O_i^{\text{Delay}}$ (obtained by taking into account the regional delays t_i as reported in Tab. 2) with the number of daily cases $O^0 = \sum O_i^0$ we would observe by considering the epidemics starts at the same time t_0 in all regions. As expected, heterogeneity flattens the curve and shifts its maximum later in time. This is a first source of errors when fitting an heterogeneous dynamics with a global model.

Assuming that the right approach is the one with regional delays, we consider two possible exit strategies: in the first, that we call the *Asynchronous* scenario, each region i lifts the lockdown at the time t_i^{Unlock} when the peak of O_i^{Delay} decreases by 30%; in the second, that we call the *Synchronous* scenario, each region i lifts the lockdown at the same time t^{Unlock} , i.e., when the global peak of O^{Delay} decreases by 30%. Notice that, once an outbreak begins, the epidemic dynamics in a region i is essentially uncorrelated with the epidemic spreading of any other region $j \neq i$. Therefore, it could be safe and appropriate to decide the lockdown release time on a regional basis, instead of lifting restrictions throughout Italy at the same time. Indeed, it could be unreasonable to keep locked the regions where the epidemic started earlier; on the contrary, regions

⁴Again, we are exploring qualitative scenarios and not trying to predict the real evolution of the epidemics: in fact, Italian regions are different for social contact habits, mobility, healthcare quality and even for factors that could possibly affect the medical parameters like comorbidities, climate or pollution levels.

	Y	M	E
Y	2.35	0.44	0.67
M	0.47	0.59	0.50
E	0.50	0.55	0.80

Tab. 3. POLYMOD matrix aggregated for three age classes: *Young* (00 – 19), *Middle* (20 – 69) and *Elderly* (70+).

where the epidemic began with some delay could experience a strong rebound when subjected to a premature lockdown lifting. In Fig. 3 we show the effects of lifting the lockdown at both regional (*Async*) and national (*Sync*) level in Lazio and Lombardy. Since not only epidemics, but also the ruin of an economy is a non-linear process, the *Sync* scenario can turn out to be even more disruptive than the epidemic itself (see also Fig. 2). Notice that analogous arguments hold - mutatis mutandis - also for the world/countries scenario.

6 The role of Age

As we have already observed in the previous Section, heterogeneity strongly impacts on the results of a model [27]. Since the transmission coefficient is proportional to the contact rate of individuals, the rates of social mixing among different age classes could represent another important source of heterogeneity. This information has been estimated either through large-scales surveys [24] or through virtual populations modeling [28]. While the POLYMOD [24] matrices have been extensively used to estimate the cost-effectiveness of vaccination for different age-classes during the 2009 H1N1 pandemic [29, 30], here they are used to support the design of lockdown measures and exit strategies. Hence, to account for age classes, we extend our model by rewriting the transmission coefficient as βC (see Sec. 8.7), where β is the transmission probability of the infection, and C is the sociological matrix describing the contact patterns typical of a given country. For lack of further information, we assume β constant among age classes and C as in [24]. To simplify the analysis, we gather POLYMOD age groups into three classes: *Young* (00 – 19), *Middle* (20 – 69) and *Elderly* (70+) (see Tab 3). Such aggregation puts together the most “contactful” classes (00 – 19), the classes with the highest mortality risk (70+) [20], and a good approximation of the active population (20 – 69).

Fig. 4 shows how the percentage of people with visible symptoms (O) varies once the age class heterogeneity has been added into the model. Notice that, at difference of fig. 2, fully releasing the lockdown results in a conspicuous rebound of the epidemics, that reaches values even more severe than the pre-exit peak. Thus, models not considering heterogeneities could severely mis-forecast the post-exit dynamics. on the other hand, introducing the age structure in the model allows to set up different exit strategies based on age-targeted policies for dampening a possible upturn of contagion. Specifically, quarantine measures

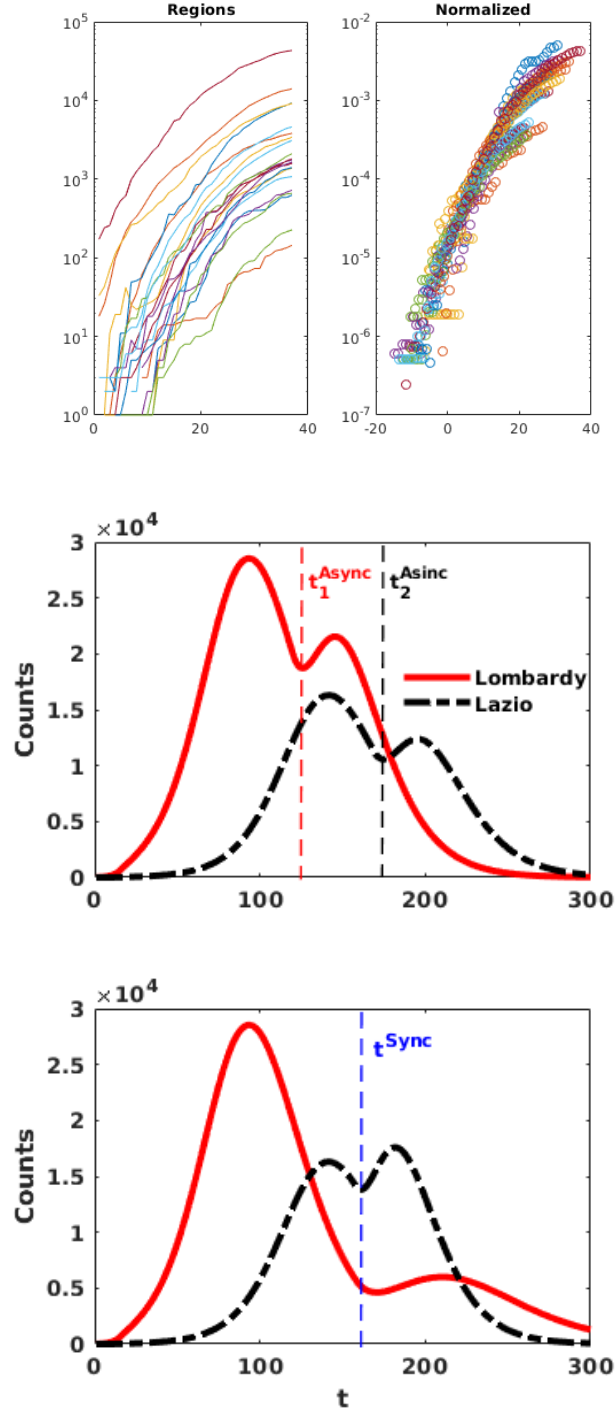


Fig. 3. Upper panels: analysis of time delays among the start of epidemics in different regions (see Tab. 2). Lower panels: sketch of an *Async*(hronous) exit strategy (i.e. each region releases the lockdown following its own policy) respect to a *Sync*(hronous) exit strategy (i.e. the lockdown release follows the same policy, but applied to a nation wide scale). In particular, t^{Sync} corresponds to releasing the lockdown in all the region after the peak has fallen by 30%, while t_i^{Async} corresponds to releasing the lockdown in the i^{th} region after the peak of such region has fallen by 30%.

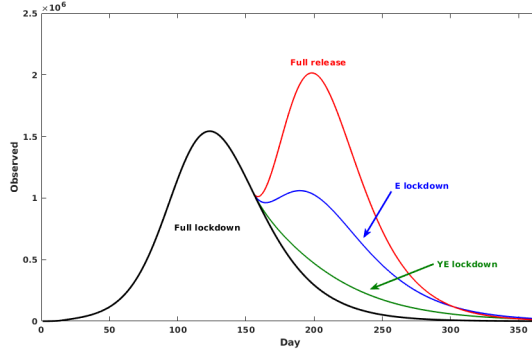


Fig. 4. Comparison of the scenarios where the lockdown is relaxed only for particular age class with respect to a full release policy. Strategies: YE = quarantine young and elderly, E = quarantine elderly. Notice that we have purposefully left the M class fully unrestrained, in order to show how maintaining a partial, age-based lockdown could deeply change the effectiveness of the exit strategy.

applied to only elderly people may limit the impact of a renewed upward phase, while maintaining lockdown restrictions to all classes except the middle age class (20-69) could be enough to smooth and lessen the propagation of contagion in the post-lockdown phase. Remember that we are using mock-up exit strategies that lead to worst-case scenarios: in real life, community measures and physical distancing, infection prevention and control in the community, personal hygiene habits and face mask usage could strongly contribute to the dampening of the epidemics [21].

7 Conclusions

In this work we propose and test a general framework to study an epidemic contagion through a compartmental model based on geographical groups and age classes. We reveal how the promptness of lockdown measures has a main effect on the timing of the contagion. Strict social distancing policies reduce the extent of contagion during the lockdown period, but full recover of the contagion occurs once such measures are relaxed. Moreover, we show how local dynamics at regional level can be inappropriately masked when observing the aggregate national system. Regional heterogeneity lowers and widens the curve of the contagion thus determining a shift forward in time for its peak at the aggregate national level. Moreover, by analyzing mobility data, we find that, due to the sparsity of interconnections across regions, contagion develops independently within each region once the epidemic has started. This, in turn, contributes to account for the delays observed in the alignment of the contagion curves across different geographical areas. The independence of regional dynamics allows both

to design regional exit strategies almost independently and to maintain – with the appropriate security measures – the movements of goods among the regions. The same reasoning valid for the Italy/regions system can be applied to the world/country system. Finally, we investigate the structure of social contacts to detect the role of different age classes in the spreading of contagion. Both young people (0-19) and elderly people (70+) are the most interconnected classes, and therefore their behavior significantly affects the post-lockdown phase. We show that while confirming quarantine policies only for elderly people can limit the possibility of a renewed upward phase in the contagion dynamics, relaxing the lockdown measures for the middle age class (20-69) can be enough to smooth and lessen the propagation of contagion in the post-lockdown phase.

Although our study is tuned on the Italian Covid-19 contagion, our modeling approach is general enough to help us understand the role of relevant dimensions, beside medical and pharmaceutical ones, in leveraging strategies to contain the epidemics and mitigate its effects. Our framework sustains an assessment of the trade-off between health outcomes and effects on wider economy. In particular, we show how the timeline of post-lockdown measures can benefit from the inclusion of compartmental aspects, such as geographical and age classes. This feature is general, and it can sustain simulations on specific restrictions, such as those targeting specific age cohorts, especially the most fragile (70+), enforcing social distancing while containing the overall burden on the economy.

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8 Supplementary Information

8.1 Initial parameters estimation

In the early phases of the epidemic, observed quantities follow an approximately exponential growth $Y^{\text{Obs}} \sim Y_0 e^{gt}$ as expected in most epidemic models. To understand what happens in our model, we notice that for $I/S \ll 1$ we can linearize Eq. 1 resulting in $I \sim I_0 e^{(\beta-\gamma)t}$ and in $O \sim \rho\gamma I$. Hence, minimizing the difference between O and Y^{Obs} in such time range would yield estimates for β, γ such that $\beta - \gamma \sim g$ and the basic reproduction number $R_0 \sim 1 + g/\gamma$ would increase linearly with the characteristic time $\tau_I = \gamma^{-1}$ for exiting the infective phase. Notice that most of the compartmental models based on ordinary differential equation will show an initial exponential growth phase with the same exponent (see Fig. 5); hence, in the early stage of the epidemic it is possible to successfully fit the “wrong” variables.

8.2 Effects of lockdown time and lockdown strength

By increasing the strength α of the lockdown (where α is the ratio between the transmission β after and before the lockdown) the height of the peak lowers but shifts to farther times. On the other hand, slowing down the epidemic implies that lifting the lockdown would bring back the infection. In the left panel of Fig. 6, we show what happens by releasing the lockdown when the peak is fallen by 30%: stronger lockdowns induce a stronger reprise of the epidemic. An analogous effect can be observed by varying the lockdown time: anticipating the lockdown ameliorates the peak by decreasing its height, but shifts it to later time and retards the end of the epidemic.

Contrary to what could be naively expected, an early imposition of the lockdown does not ameliorate the epidemics: in fact, anticipating too much the lockdown just shifts the timing of the epidemics, leaving its evolution unchanged (see Fig. 7). This is to be expected every time extreme measures of social distancing are applied in the very early, exponentially growing, stages. In fact, let us consider two countries A and B , having the same number of inhabitants, the same contact matrix, and the same number of infected people. If A and B decide to put a lockdown of strength α at time t_A and t_B , respectively, at time t any quantity y would have grown as $y_A(t) \sim y^0 e^{R_0 t_A} e^{\alpha R_0 (t-t_A)}$ and as $y_B(t) \sim y^0 e^{R_0 t_B} e^{\alpha R_0 (t-t_B)}$. If there exists a t' such that $y_A(t) = y_B(t')$, the epidemics in A and in B will proceed in parallel (even in the non-linear phase) with a delay $t' - t$. Therefore, if the epidemic dynamics of A and B are still well approximated by exponential distributions at times $< \max\{t, t'\}$, then $t' - t \propto -(t_A - t_B)$, i.e, the country that has started the lockdown before will experience the same epidemic of the other country, just delayed in time. In particular, for identical initial conditions, we have that:

$$t - t' = -\frac{1 + \alpha}{\alpha} (t_A - t_B) \quad (2)$$

as long as all the times are before the initial exponential regime ends.

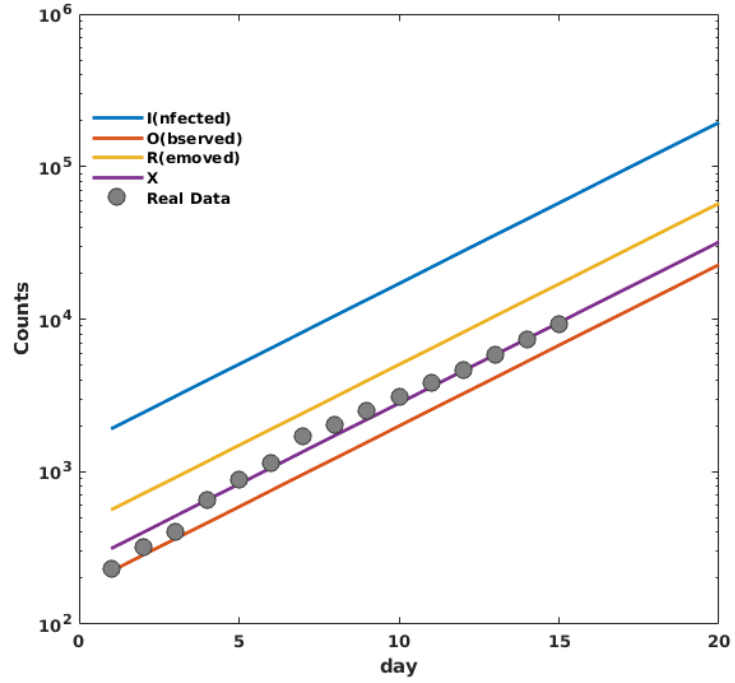


Fig. 5. In the initial stage, most of the quantities experience an exponential growth with the same exponent; hence, it would be possibly to “successfully” fit the wrong variables. In the panel, we show the pre-lockdown growth of the number of $I(nfected)$, $O(bserve)$, $R(removed)$ individuals in our model (1). Full circles are the experimental counts of confirmed Covid-19 cases in Italy; X is the cumulative variable we use to fit the experimental data.

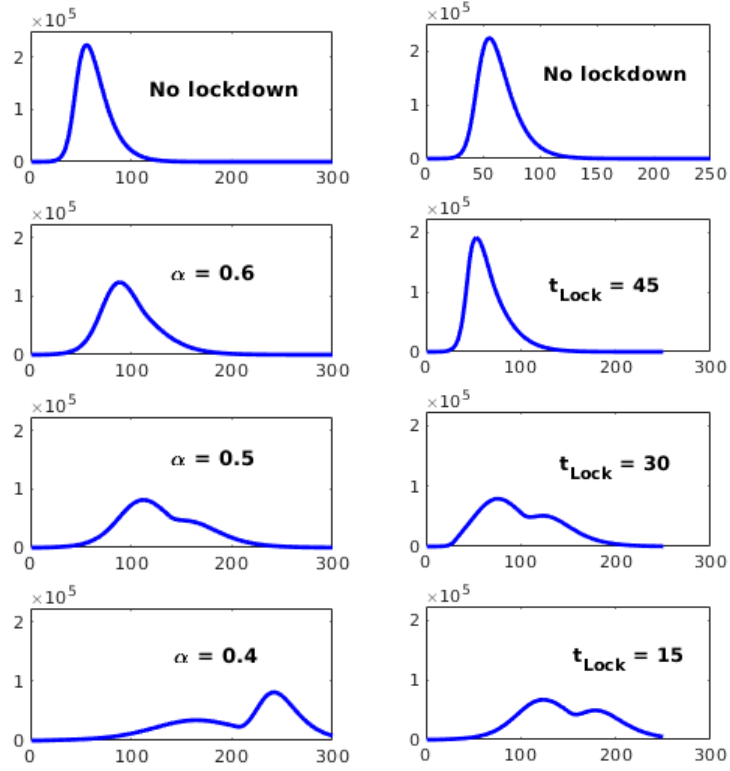


Fig. 6. Left panel: variation of the behavior of the model by varying the lockdown strength α . Lockdown starts at $t_{\text{Lock}} = 15$ and is fully lifted when the peak has fallen by 30%. Right panel: variation of the behavior of the model by delaying the lockdown time t_{Lock} . Lockdown strength is fixed at $\alpha = 0.5$ and is fully lifted when the peak has fallen by 30%.

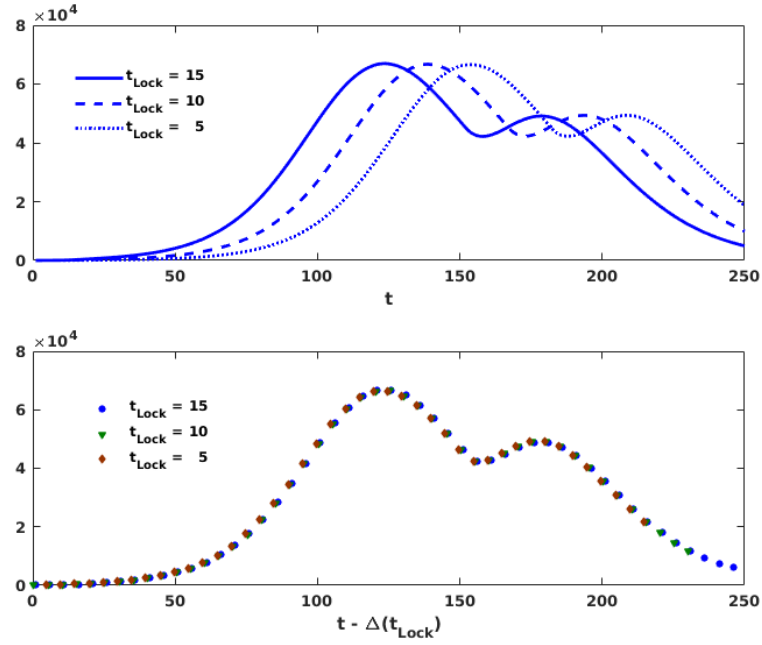


Fig. 7. Upper panel: variation of the behavior of the model by anticipating the lockdown time. Notice that anticipating the lockdown leaves unchanged the behaviour of the epidemics, just shifting all the times of an amount proportional to how much the lockdown is anticipated. Lockdown strength is fixed at $\alpha = 0.5$ and is fully lifted when the peak has fallen by 30%. Lower panel: by applying the Eq. 2, we show how the curves in the upper panel collapse on each other.

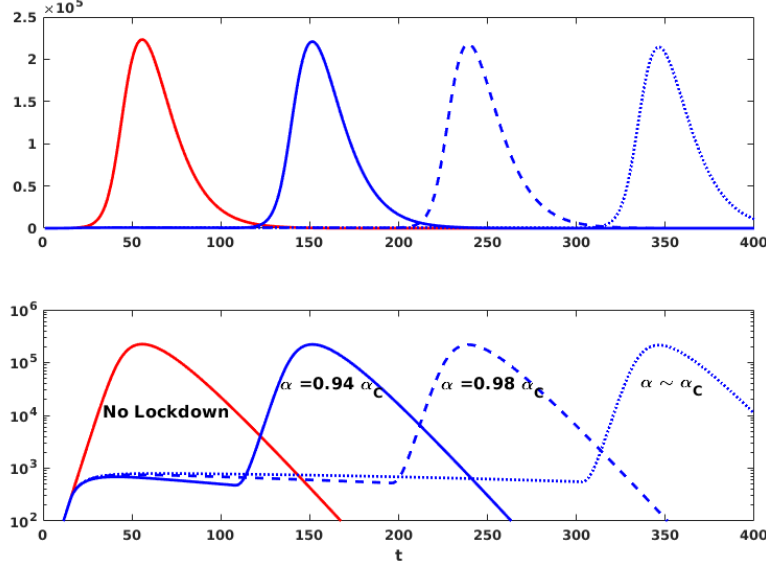


Fig. 8. Upper panel: variation of the behavior of the model for lockdown strengths $\alpha < \alpha_{\text{crit}} = 1/R_0$. Notice that the height of the peaks after the lockdown is released is almost unchanged if compared with the no lockdown scenario. Lockdown time is fixed at $t_{\text{Lock}} = 15$ and is fully lifted when the peak has fallen by 30%. Lower panel: for better clarity, the plot is also reported in log-linear scale.

Finally, we notice that to each lockdown strength α corresponds an effective reproductive number $R_0^{\text{eff}} = \alpha R_0$; hence, for $\alpha \sim \alpha_{\text{crit}} = 1/R_0$, the epidemics is expected to stay in a quiescent state where it does not either grow or decay sensibly. On the other hand, for $\alpha < \alpha_{\text{crit}}$ the epidemics decreases; nevertheless, since this happens before a sufficient number of recovered individuals has built up herd-immunization, the height of the peaks after the lockdown lifting are almost unchanged if compared with the no lockdown scenario. Again, a “too good” intervention risks to postpone the problem without attenuating it. Notice that, if one applies lockdowns with $\alpha < \alpha_{\text{crit}}$, it could be necessary to switch back and forth to lockdown to avoid the peak go beyond the capacity of a national healthcare system (see Fig. 8).

8.3 Stationary state of the *SIOR* model

Let $X = O + R$. Then, $\partial_X S = -R_0 S$ and $S(t \rightarrow \infty) = N e^{-R_0 X(t \rightarrow \infty)}$. Since $O(t \rightarrow \infty) = I(t \rightarrow \infty) = 0$ and hence $R(t \rightarrow \infty) = N - S(t \rightarrow \infty)$, we recover the same solution of the *SIR* model: $S(t \rightarrow \infty) = N e^{-R_0 [N - S(t \rightarrow \infty)]}$. For $R_0 \in [2.5, 4.5]$, we have that the final fraction of uninfected population varies

between 10% and 1%.

8.4 Estimation of the experimental time delays

We first normalize the observed data by dividing the number of non-zero observations in a region for the population of the region. Let y_i be the normalized observations for the i^{th} region. For each pair of regions i, j , we define the variation interval $\Delta_{ij} = [\min_{ij}, \max_{ij}]$ that contains the maximum number of points of both y_i and y_j , i.e. $\min_{ij} = \max\{\min(y_i), \min(y_j)\}$ and $\max_{ij} = \min\{\max(y_i), \max(y_j)\}$. The delay t_{ij} between the epidemics start in i and j , respectively, is calculated by minimizing the square norm of $\|(\Delta_{ij} \cap y_i(t)) \setminus (\Delta_{ij} \cap y_j(t - t_{ij}))\|$, where $\Delta_{ij} \cap y$ denotes the values of y falling in the interval Δ_{ij} . Denoting with T_i the times corresponding to the observation in $\Delta_{ij} \cap y_i$, it is easy to verify that $t_{ij} = \langle T_i \rangle - \langle T_j \rangle$, where $\langle T \rangle$ is the average value of the times contained in T .

8.5 Equivalence of normalized curves

Eq. 1 referred to region k becomes:

$$\begin{aligned}\partial_t S_k &= -\beta S_k I_k / N_k \\ \partial_t I_k &= \beta S_k I_k / N_k - \gamma I_k \\ \partial_t O_k &= \rho \gamma I_k - h O_k \\ \partial_t R_k &= (1 - \rho) \gamma I_k + h O_k\end{aligned}\tag{3}$$

where N_k is the population of the region. By rewriting Eq. 3 in terms of normalized quantities $s_k = S_k / N_k, \dots, r_k = R_k / N_k$, we obtain the same equation for all the regions:

$$\begin{aligned}\partial_t s &= -\beta s i \\ \partial_t i &= \beta s i - \gamma i \\ \partial_t o &= \rho \gamma i - h o \\ \partial_t r &= (1 - \rho) \gamma i + h o\end{aligned}\tag{4}$$

Hence, for similar initial conditions, by normalizing the experimental observations by the population, one should obtain similar time behaviors.

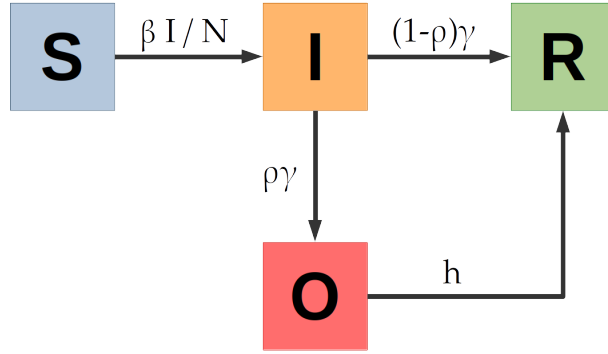


Fig. 9. The *SIOR* compartmental model: workflow of the epidemic process. A *S*(usceptible) individual becomes *I*(nfective) when meeting an infected person. An *I*(nfective) either become *O*(bserved), with symptoms acute enough to be detected from the national health-care system, or is *R*(emoved) from the infection cycle by having recovered. An *O*(bserved) individual can also be *R*(emoved) from the infection cycle having become immune. The parameter β defines the rate at which a susceptible becomes infected, γ represents the rate at which infected either become observable or recover, ρ is the fraction of infected that become observed from the national health-care system and h is the rate at which observed individuals are removed from the infection cycle.

8.6 Regional metapopulation model

Let us assume that we know the fraction T_{kl} of people commuting from region k to region l , Eq. 4 becomes:

$$\begin{aligned}\partial_t s_k &= -\beta s_k \sum_l T_{kl} i_l \\ \partial_t i_k &= \beta s_k \sum_l T_{kl} i_l - \gamma i_k \\ \partial_t o_k &= \rho \gamma i_k - h o_k \\ \partial_t r_k &= (1 - \rho) \gamma i_k + h o_k\end{aligned}\tag{5}$$

From mobility data, we know that $\epsilon_k = \sum_{l \neq k} T_{kl}/T_{kk} \ll 1$ and $T_{kk} \sim 1$; in particular, from Facebook mobility data we can estimate $\langle \epsilon_k \rangle \sim 10^{-3}$. If all the neighbors of a given region k are fully infected (i.e. $i_l = 1 \forall l \neq k$) and $i_k(t_0) = 0$, then the variation of i_k can be approximated as $\partial_t i_k \sim \epsilon_k + (\beta - \gamma) i_k$. Namely, as soon as $i_k > \epsilon_k$, i_k will grow exponentially according to $\partial_t i_k \sim (\beta - \gamma) i_k$ and ϵ_k will become irrelevant; that is to say, the dynamics of the regions will decouple. On the other hand, if epidemic is decaying everywhere, then $i_l \ll 1 \forall l \neq k$; thus $\sum_{l \neq k} T_{kl} i_l \ll \epsilon_k$ and equation again decouple, having each region followed Eq. 4 separately. In Tab. 4 we confront regions ordered by simulating an hypothetical epidemics starting from Lombardy and propagating with Eq. 5, with regions ordered by the estimated delays of Tab. 2. It is reasonable to assume that inter-regional mobility has had a role in the regional delay structure; however, many other factors come to play in the long range propagation of epidemics: as an example, both airline transportation network [31, 32] and individual work commutes [33, 34] have played important roles in understanding the spread of infectious diseases.

8.7 Social mixing

To take account for social mixing, we rewrite the transmission coefficient as the product of a transmission probability β times a contact matrix C whose element C_{ab} measure the average number of (physical) daily contacts among an individual in class age a and an individual in class age b . Notice that the probability that a susceptible in class a has a contact with an infected in class b is the product of the contact rate C_{ab} times the probability I_b/N_B that individual in class b is infected. Hence, denoting with S^a, \dots, R^a the number of

Mobility Matrix	Experimental Delays
Lombardia	Lombardia
Emilia Romagna	Emilia Romagna
Piemonte	Marche
Veneto	Veneto
Valle d'Aosta	Valle d'Aosta
Trentino Alto Adige	Liguria
Lazio	Friuli Venezia Giulia
Liguria	Piemonte
Toscana	Trentino Alto Adige
Campania	Toscana
Marche	Molise
Friuli Venezia Giulia	Umbria
Abruzzo	Abruzzo
Umbria	Lazio
Sardegna	Campania
Sicilia	Puglia
Molise	Sardegna
Basilicata	Sicilia
Puglia	Calabria
Calabria	Basilicata

Tab. 4. Region ordered by simulations using the mobility matrix (left column) and by the delays obtained by rescaling experimental data (right column).

$S(\text{usceptibles}), \dots, R(\text{removed})$ individuals in class age a , we can rewrite Eq. 1 as:

$$\begin{aligned}
\partial_t S^a &= -\beta S^a \sum_b C_{ab} \frac{I^b}{N_b} \\
\partial_t I^a &= \beta S^a \sum_b C_{ab} \frac{I^b}{N_b} - \gamma I^a \\
\partial_t O^a &= \rho \gamma I^a - h O^a \\
\partial_t R^a &= (1 - \rho) \gamma I^a + h O^a
\end{aligned} \tag{6}$$

Although the form of Eq. 6 is similar to Eq. 3, here it is not possible to consider separate evolutions for the different age classes since, differently than the inter-regional mobility matrix T , the off diagonal elements of the social matrix $C_{a,b}, a \neq b$, measure the interaction among different age classes and are of the same magnitude of the diagonal elements C_{aa} measuring the interaction among individuals of the same age class.

Notice that Eq. 6 can be summed up, and the resulting equation can be obtained by substituting $\beta \rightarrow \beta C^{\text{eff}}$ in Eq. 1, where $C^{\text{eff}} = \frac{\sum_{ab} C_{ab} S^a I^b / N_b}{SI/N}$ is the average contact value among infected and susceptible individuals of all age classes.