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A metapopulation network model for the spreading of SARS-CoV-2: Case study for Ireland

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ABSTRACT

We present preliminary results on an all-Ireland network modelling approach to simulate the spreading the Severe Acute Respiratory Syndrome CoronaVirus 2 (SARS-CoV-2), commonly known as the coronavirus. In the model, nodes correspond to locations or communities that are connected by links indicating travel and commuting between different locations. While this proposed modelling framework can be applied on all levels of spatial granularity and different countries, we consider Ireland as a case study. The network comprises 3440 electoral divisions (EDs) of the Republic of Ireland and 890 superoutput areas (SOAs) for Northern Ireland, which corresponds to local administrative units below the NUTS 3 regions. The local dynamics within each node follows a phenomenological SIRX compartmental model including classes of Susceptibles, Infected, Recovered and Quarantined (X) inspired from Science 368, 742 (2020). For better comparison to empirical data, we extended that model by a class of Deaths. We consider various scenarios including the 5-phase roadmap for Ireland. In addition, as proof of concept, we investigate the effect of dynamic interventions that aim to keep the number of infected below a given threshold. This is achieved by dynamically adjusting containment measures on a national scale, which could also be implemented at a regional (county) or local (ED/SOA) level. We find that in principle dynamic interventions are capable to limit the impact of future waves of outbreaks, but on the downside, in the absence of a vaccine, such a strategy can last several years until herd immunity is reached.

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1. Introduction

On 30th January, 2020, the World Health Organisation (WHO) declared the newly emergent novel coronavirus pathogen, Severe Acute Respiratory Syndrome CoronaVirus 2 (SARS-CoV-2), to be a public health emergency of international concern (World Health Organization, 2020a, 2020b). This characterisation was later updated to pandemic status on 11th March 2020 (World Health Organization, 2020c). Following its initial detection in a cluster of patients experiencing acute respiratory symptoms in the city of Wuhan, in the Hubei province of China (Chen et al., 2020), the spread of SARS-CoV-2, which causes the coronavirus disease 2019 (COVID-19), to 218 countries and territories in the months since has led to unprecedented social and economic disruption (Douglas et al., 2020; Worldometer, 2020b). Despite widespread efforts to contain the virus through a series of non-pharmaceutical interventions, global figures show reported cases standing at over 63 million, with deaths from the virus totalling 1.5 million up to late November 2020 (Dong et al., 2020; Johns Hopkins University, 2020). It is clear that there is an urgent and ongoing need to understand how the virus spreads in order to effectively protect populations.

As of yet there are no significantly effective pharmaceutical measures available that prevent the spread of the disease. Vaccination programmes commenced on 14th December, 2020 and in the first month, approximately 35 million doses have been administered in 49 countries worldwide (Bloomberg, 2021). Immunisation of billions of people around the world will be a logistical tour de force, and the time frame for a significant level of coverage is still on the order of many months or more. Until such measures are widely implemented, the best available mitigation strategies include the use of face masks, sanitisation, and social distancing. Authorities have implemented lockdowns to promote these measures. Mathematical models are invaluable in providing us with an insight into the processes by which the disease spreads (Al-Khani et al., 2020; Blyuss & Kyrychko, 2020; Kyrychko et al., 2020; Liu et al., 2020; Maier & Brockmann, 2020; Pinotti et al., 2020) and these models can help decide on the best course of action to take in the form of non-pharmaceutical interventions (Ames et al., 2020; Prem et al., 2020; Pullano et al., 2020; Schlosser et al., 2020).

The model proposed in this discussion paper and the numerical results arising from it are to be understood as a contribution to the scientific and public discussion of possible spreading scenarios and the impact of intervention measures including realistic levels of compliance. We will clearly state the model ingredients and assumptions and elaborate on conclusions that can be drawn based on these limitations; some of which are unlikely for any practical purpose, but worth exploring, e.g., for worst-case scenarios.

We start with an introduction of the model equations for the local and networked dynamics in sections 2 and 3, respectively. This includes a discussion of the local dynamics from a geometric point of view, where we sketch the trajectories and equilibria in phase space. Then, we continue with a case study of Ireland in section 4 including realistic mobility data and assumptions and elaboration on conclusions that can be drawn based on these limitations; some of which are unlikely for any practical purpose, but worth exploring, e.g., for worst-case scenarios.

We start with an introduction of the model equations for the local and networked dynamics in sections 2 and 3, respectively. This includes a discussion of the local dynamics from a geometric point of view, where we sketch the trajectories and equilibria in phase space. Then, we continue with a case study of Ireland in section 4 including realistic mobility data and compliance levels with respect to movement restrictions. Besides an unlikely, worst-case scenario, we consider different levels of increased awareness during the post-lockdown period in section 4.2. In addition, in section 4.3, we explore effects of dynamic interventions to keep the prevalence below a given level. These dynamic lockdowns are considered at different levels of spatial granularity.

2. Local dynamics

We start with the model for the local, well mixed dynamics at a single node as the basis for the network model:

\[
\begin{align*}
\dot{S} &= -\beta \frac{SI}{N} \quad (1a) \\
\dot{I} &= \beta \frac{SI}{N} - \mu I - \alpha I - \kappa I \quad (1b) \\
\dot{X} &= \kappa I - \mu X - \alpha X \quad (1c) \\
\dot{R} &= \mu I + \mu X \quad (1d) \\
\dot{D} &= \alpha I + \alpha X. \quad (1e)
\end{align*}
\]

The above model is a simple extension of the original Kermack and McKendrick SIR model (Kermack & McKendrick, 1927), inspired by Ref. (Maier & Brockmann, 2020). There are five compartments, \( N \) individuals, and each individual may belong to just one compartment at any time:

- **Susceptible** \( S(t) \) Individuals who have not been infected by the disease.
- **Infected** \( I(t) \) Individuals who are infected with the disease and are capable of transmission.
- **Quarantined** \( X(t) \) Individuals who are infected with the disease, but are quarantined or self-isolating and thus not transmitting.
- **Recovered** \( R(t) \) Individuals who have recovered from the disease and are considered immune.
- **Dead** \( D(t) \) Individuals who have died from the disease.
The above system of ordinary differential equations (ODEs) assume homogeneous mixing of the population, which can be a good approximation for small communities, and small number of deaths. The parameters of the ODE model (1) refer to the rates of moving from one compartment to another and population size:

\[\begin{align*}
\beta & \text{ probability of infection on contact.} \\
c & \text{rate of contact/mixing.} \\
\mu & \text{rate of recovery.} \\
\kappa & \text{rate of moving from infected to quarantined.} \\
a & \text{rate of death.} \\
N & \text{total population size } N = S(t) + I(t) + X(t) + R(t) + D(t) = \text{const.}
\end{align*}\]

Much insight into the network dynamics can be gained from geometric analysis of the local, single-node dynamics. In the \((S, I, X, R, D)\) phase space, system (1) has a whole family of stationary solutions (equilibria). They are given by

\[I_e = X_e = 0 \quad \text{and} \quad S_e + R_e + D_e = N,\]

correspond to disease-free conditions, and form a triangular surface shown in Fig. 1. This family includes the important pre-disease equilibrium

\[S_e = N, I_e = X_e = R_e = D_e = 0,\]

to discuss stability of equilibria, it is convenient to introduce the basic reproduction number:

\[R_0 = \frac{\beta c}{a + \kappa + \mu}\]

when \(0 < R_0 < 1\), all equilibria are stable, meaning that no disease outbreaks can occur. When \(R_0 > 1\), the pre-disease equilibrium turns unstable, and disease outbreaks become possible. To be more specific, the family of equilibria is divided into the (pink) unstable part with \(S_0 < S_e \leq N\), which contains the pre-disease equilibrium, and the (blue) stable part with \(0 \leq S_e < S_0\), shown in Fig. 1. The instability threshold separating the two parts is given by

\[S_0 = \frac{N}{R_0}.\]

guided by these observations, we define herd immunity, whether achieved naturally or by means of vaccination, in terms of stable equilibria as follows:

**Definition 1.1.** In the local SIXRD model (1), we define herd immunity as a disease-free state of the system such that a small increase in infected \(I\) decays monotonically to zero (no outbreak):

---

**Fig. 1.** Sketch of the outbreak dynamics in the local SIXRD model (1) for \(R_0 > 1\), shown as a (red) trajectory from the disease-free equilibrium \(S_e = N\) to one of the stable equilibria with \(0 < S_e < S_0\) in the projection of the \((S, I, X, R, D)\) phase space onto the \((S, I, R)\) subspace. \(S_0\) is the herd immunity threshold separating the triangular family of equilibria into the (pink) unstable and (blue) stable parts.
0 \leq S < S_0 \quad \text{and} \quad I = X = 0.

In other words, herd immunity is represented by a stable equilibrium. We call $S_0 = N/R_0$ the herd immunity threshold.

In Fig. 1, a disease outbreak is depicted as the (red) trajectory from the pre-disease equilibrium, giving a peak in infected $I(t)$ around the herd immunity threshold $S_0$, to one of the (blue) herd immunity equilibria that lie below $S_0$. The lower the $S_0$, the higher the peak in $I(t)$. Thus, bringing the herd immunity threshold $S_0$ closer to $N$ for a period of time and then back to its original position, e.g., via imposing temporary public-health restrictions, can modify the outbreak trajectory so that ultimate herd immunity is achieved in a controlled way, that is with a peak in $I(t)$ that remains below some desired level.

### 3. Network model

While the ODE model (1) can be justified for small communities, it fails at capturing the dynamics that occurs at large spatial scales and cannot account for heterogeneities in a population and interaction between metapopulations via mixing/commuting (Belik et al., 2011). For this reason, we develop a network framework by splitting the population up into a number of smaller communities based on their geographic location. Ideally, these communities are small enough to justify a well-mixed ODE model at the local level. Every community corresponds to a node in the network. Two nodes are connected by a link, which represent the number of people travelling between the involved communities. The number of people travelling can vary with time or remain static. The resulting network model is given by

\[
\begin{align*}
\dot{S}_i &= -\beta c \left( S_i \sum_{j=1, j \neq i}^{n} S_{ij} \right) \frac{I_i + \sum_{k=1, k \neq i}^{n} (I_{ki} - I_{ik})}{N_i + \sum_{k=1, k \neq i}^{n} (N_{ki} - N_{ik})} - \beta c \left( \sum_{j=1}^{n} S_{ij} \right) \frac{I_j + \sum_{k=1}^{n} (I_{kj} - I_{jk})}{N_j + \sum_{k=1}^{n} (N_{kj} - N_{jk})} - \beta c \left( \sum_{j=1}^{n} S_{ij} \right) \frac{I_j + \sum_{k=1}^{n} (I_{kj} - I_{jk})}{N_j + \sum_{k=1}^{n} (N_{kj} - N_{jk})} - \mu I - aI - \kappa I \\
\dot{I}_i &= \beta c \left( S_i \sum_{j=1, j \neq i}^{n} S_{ij} \right) \frac{I_i + \sum_{k=1, k \neq i}^{n} (I_{ki} - I_{ik})}{N_i + \sum_{k=1, k \neq i}^{n} (N_{ki} - N_{ik})} + \beta c \left( \sum_{j=1}^{n} S_{ij} \right) \frac{I_j + \sum_{k=1}^{n} (I_{kj} - I_{jk})}{N_j + \sum_{k=1}^{n} (N_{kj} - N_{jk})} - \mu I - aI - \kappa I \\
\dot{X}_i &= \kappa I_i - \mu X_i - \alpha X_i \\
\dot{R}_i &= \mu I_i + \mu X_i \\
\dot{D}_i &= aI_i + \alpha X_i.
\end{align*}
\]

For notational convenience, the compartments are now labelled by a subscript, i.e., $S_i, I_i, X_i, R_i,$ and $D_i,$ to denote the number of individuals belonging to the respective compartments of $i$-th node in the network ($S_i + I_i + X_i + R_i + D_i = N_i, i = 1, \ldots, n$). Similarly, the number of individuals travelling from node $i$ to node $j$ belonging to a given compartment is denoted by $S_{ij}, I_{ij}, X_{ij},$ and $R_{ij}$, and $D_{ij}$ resulting in $N_{ij} = S_{ij} + I_{ij} + X_{ij} + R_{ij} + D_{ij}$. The proposed modelling framework accounts for travel as commuting pattern, that is, individuals return to their original location (cf. Appendix D). The local dynamics is included via the single-subscript quantities. Hence, $S_i = I_i = X_i = R_i = D_i = 0$. Note that the network model includes interactions between nodes only in the susceptible and infected compartments. Accordingly, we account for the number of (i) individuals that remain in their home node and (ii) those who travel to other nodes in the first and second term of Eqs. (4a) and (4b), respectively. For example, the term $\sum_{k=1}^{n} (I_{ki} - I_{ik})$ in Eq. (4a) refers to the net flux of infected individuals, that is, the difference between individuals leaving node $i$ to $k$ and arriving from node $k$ to $i$.

Given a number of travellers $N_{ij}$ between nodes, the network model (4) assumes that the number of travellers belonging to the respective compartments is determined by a relation such as $S_{ij} = N_{ij} \frac{S_i}{N_i}$. Thus, the number of travelling susceptibles is proportional to the number of susceptibles in the home node. The number of travelling individuals in the other compartments is computed in the same fashion. $N_{ij}$ may be randomly generated at each time step. The distance travelled between locations often follows a power-law or heavy-tail distribution and can be described by a gravity or radiation population model (Brockmann et al., 2006; Brockmann & Hufnagel, 2007; Grauwin et al., 2017; Simini et al., 2012; Song et al., 2010a, 2010b). Alternatively, empirical movement data can inform the model. From these data one can either build a static network using, e.g., the jump-size distributions, or randomly generate a network at each time-step. Note that the proposed network model does not account for any importation from outside the considered network.

For the fully networked case [cf. Eq. (4)], the basic reproduction number $R_0$ can be computed from the system’s next generation matrix $\mathbb{G}$ (van den Driessche & Watmough, 2002). This takes into account the couplings between the nodes when computing the stability of the disease-free equilibria. The $(i, j)$ entry of this matrix corresponds to the expected number of new
infections in node $i$ caused by an infected individual introduced in node $j$. The next generation matrix, as derived in Appendix A, is given as

$$
G_{ij} = \frac{\beta c}{\mu + \alpha + \kappa} \left[ \left( S_i - \sum_{k=1 \atop k \neq i}^{n} S_{ik} \right) \delta_{ij} \left( 1 - \sum_{k=1 \atop k \neq i}^{n} \frac{N_{ik}}{N_i} \right) + \frac{N_j}{N_i} \right] + \sum_{k=1 \atop k \neq j}^{n} \frac{S_{jk}}{N_k} \delta_{jk} \left( 1 - \sum_{l=1 \atop l \neq j}^{n} \frac{N_{jk}}{N_j} \right) + \frac{N_{ik}}{N_i} \right],
$$

(5)

where $\delta_{ij}$ denotes the Kronecker delta: $\delta_{ij} = 1$ for $i = j$ and zero otherwise. Then, the effective reproduction number $R_0$ for the networked system is given by the spectral radius $\rho$ of the next generation matrix,

$$
R_0 = \rho(G). \tag{6}
$$

This provides a condition necessary for the stability all disease-free equilibria ($I_i = X_i = 0, S_i + R_i + D_i = N_i$ for all $i = 1, \ldots, n$). It also gives a natural definition for herd immunity in the network, as the networked reproduction number $R_0$ is now a function of all susceptible populations (cf. the definition of herd immunity for the local dynamics in Sec. 2). All scenarios discussed in the next sections are simulated using our library for modelling epidemics on networks, EPIGRAPH (Humphries, 2020), which is implemented in C++ and freely available on GITHUB.

4. Case study: Ireland

For the remainder of this discussion paper, we elaborate on the spreading of SARS-CoV-2 in Ireland as a case study. The network model given by Eq. (4) is informed by publicly available data detailed in the next section.

4.1. Network data

For the Republic of Ireland, each of the 3440 nodes in the network represents an electoral division as defined by the central statistics office (CSO), for which small area population statistics are published from the 2016 census (Census 2016 Open Data Site, 2016). The links in the network are the number of commuters between each electoral division according to CSO data (Central Statistics Office, 2016). Similarly, for Northern Ireland, nodes refer to 890 superoutput areas according to Northern Ireland Statistics and Research Agency (NISRA) (Northern Ireland Statistics and Research Agency, 2016). This spatial resolution corresponds to the local administrative units below the category-3 regions of the Nomenclature of Territorial Units for Statistics (NUTS) (Eurostat, 2018).

4.2. 5-phase roadmap without additional countermeasures

In order to demonstrate the general feasibility of the proposed modelling framework, we consider the 5-phase roadmap for reopening society & business by reducing lockdown restrictions as outlined by the Irish Government in May 2020 (Department of the Taoiseach and Department of Health, 2020).

Our model attempts to follow the historic evolution of the disease up until phase 2. Then, we allow the model to run under the predicted effects of the remaining lockdown phases. See Tab. 1 for implications on travel and gathering restrictions. Appendix B provides a comparison to reported case numbers. It is important to note that at this stage, we do not consider any reintroduction or roll back to earlier phases during the course of the simulated outbreak. This corresponds to a worst-case scenario and is highly unlikely from a practical point of view, because interventions will be reintroduced, if case numbers rise again.

The parameters are chosen such that the model results match the actual number of deaths in Ireland. See Tab. 2 for details. It is important to note, however, that this model is in the early stages of testing and has not undergone any sensitivity training on the parameters. The effective infection probability $\beta_{\text{eff}}$ is computed as $\beta_{\text{eff}} = \beta c$. As defined in Eq. (2), the basic reproduction number of the local dynamics [cf. Eqs. (1)] exceeds the critical value of $R_0^{\text{crit}} = 1$ during pre lockdown and after phase 3. Conversely, only initial lockdown, phases 1 and 2 have a $R_0$ value well below unity and — neglecting network-related effects — exhibit a burn out of an outbreak and self-guided decrease of the number of infected.

We introduce a small number of diseased individuals to the Dublin area on 22nd January, 2020, that is, 50 days before the initial lockdown, and run the model such that the disease spreads naturally and without control. On the 12th of March, we introduce the initial lockdown measures, which take some time to reach full effect. Individuals’ movement is limited to 2 km and this is adhered to with 80% compliance. This level of compliance is supported by preliminary findings of a nation-wide series of phone interviews accounting for essential/justified and non-essential travel (HRB project, 2020). The level of compliance to movement restrictions is kept constant at 80% in the simulations (cf. Fig. 11 for the impact of compliance levels during regional interventions). We account for social distancing and self-isolation (X compartment) by reducing the rate of mixing $c$ within communities (cf. Table 2 for the respective parameter values). During phases 1 and 2/3, the travel radius
increased to 5 km and 20 km, respectively, and we assume the same level of compliance. Travel restrictions are removed in phases 4 and 5 (cf. Table 1). Each of the numbered phases have a duration of 21 days. See Appendix D for further details on the impact of movement restrictions on the mobility. In the post-lockdown period, the social mixing parameter is not reset to $c = 1$, but to a lower level of $c = 0.75$ to account for a residual effect of an increased level of awareness compared to pre-lockdown. This is justified, for instance, by ongoing closure of schools and universities and higher levels of home office than at the start of the year.

Fig. 2 depicts the time series of all five compartments — aggregated over all nodes — in semi-logarithmic scale. One can see two distinct peaks: (i) a small peak that occurs before the lockdown and (ii) a second peak that occurs once all lockdown measures are lifted. It is clear that unless some of the measure are maintained, then the lockdown will just have the effect of delaying the epidemic and result in a second wave with a prevalence that measures are lifted. It is clear that unless some of the measure are maintained, then the lockdown will just have the effect of delaying the epidemic and result in a second wave with a prevalence that — in this unlikely worst-case scenario — is 100 times higher than the first one and exceeds any realistic capacity of the health-case system. The number of infected eventually reaches zero due to a depletion of the pool of susceptibles. Following the geometric analysis of the local dynamics (cf. Sec. 2), the trajectory lands at one of disease-free equilibria $I_0 = 0$. A change in parameters, e.g., rate of mixing $c$, or a rescaling of the population to the number of remaining susceptibles can render this state unstable again. Then, a perturbation to the networked system, for instance, introduction or importation from outside, has the potential to trigger an additional outbreak.

For the simulation run given in Fig. 2 we compute the next generation matrix (5) at each time-step and find its spectral radius giving rise to the network’s effective basic reproduction number $R_0$, which is not only a function of the model parameters but also the number of susceptibles. For details, see Appendix A. $R_0$ is plotted as black dashed curve in Fig. 3. For comparison, the local/simplistic value of $R_0$ given by Eq. (2) is shown as (dash-dotted) red curve. The value of $R_0$ shows the potential for the disease to spread at each phase in the simulation. From Fig. 3 we can see clearly that when $R_0$ falls below unity, the number of infected also begins to fall. We also note the point of herd immunity is reached is in when the number of susceptibles crossed the point where the $R_0$ goes below 1.

As stated above, the parameters are chosen to match the actual number of deaths in Ireland. This, in turn, results in a larger first peak compared to the reported numbers of confirmed cases on the 12th of March ranging from 27 according to (Health Protection Surveillance Centre (HPSC), 2020a) to 70 according to (Worldometer, 2020a) (cf. Fig. 13 in Appendix B for a detailed comparison to reported data). The simulated results, however, are not meant to reflect this range of confirmed cases, but following the logic of the model, refer to all infected individuals including asymptomatic, non-diagnosed and untested.

Fig. 4a and b shows spatially resolved snapshots of the prevalence of the disease per km$^2$ at times $t = 50$d (March 12) and $t = 320$d (December 7), respectively (cf. (Health Protection Surveillance Centre (HPSC), 2020b) for a map based on data of reported, confirmed cases). Fig. 4a corresponds to the disease prevalence at the first peak before the lockdown. The disease remains in the East of the country for the most part with the exception of big towns and cities. In contrast, at $t = 320$d (Fig. 4b), the outbreak covers the entire country and affects every electoral division. See Fig. 2 for a full time series of the aggregated variables.

Fig. 5 shows the effect of different levels of awareness during the post-lockdown phase, which is modelled by different values for the mixing parameter $c$. For convenience, the plots only depict the time series of the aggregated infected compartment $I$ in a linear scale. A reduction to 70% of the original probability of infection significantly leads to a significantly smaller and delayed second peak (red curve), while the basic reproduction number of the local dynamics remains above unity (cf. Table 2).

In order to show how the heavy-tail distribution of travel across the network causes a lack of sensitivity in initial conditions, we run the model for a number of random initial seeds where for each run only a single individual is placed in one electoral division or superoutput area at the start of the simulation. The resulting time series are shown in Fig. 6.

One can see that both amplitude and timing of the large, second peak are stable and it is impossible to visually differentiate between any of the curves for different initial conditions. In the presented 4 cases, the initial location is selected uniformly at random from all network nodes. We note that deliberately selecting urban or rural locations will have no qualitative impact. To further corroborate the results, initial infections of a significant, finite fraction of the population will be explored. This might account for a considerable amount of unreported cases at the start of the recording of SARS-CoV-2 cases.

| Phase       | Date     | Mobility                        | Social distancing         |
|-------------|----------|---------------------------------|---------------------------|
| pre lockdown| 22-01-20 | no restrictions                 | no restrictions           |
| school closure| 12-03-20 | no restrictions                 | recommended reduction     |
| initial lockdown| 27-03-20| travel with 2 km radius         | essential meetings only   |
| 1           | 18-05-20 | travel with 5 km radius         | essential meetings only   |
| 2           | 08-06-20 | travel with 20 km radius        | maximum of 4 people       |
| 3           | 29-06-20 | travel with 20 km radius        | maximum of 4 people       |
| 4           | 20-07-20 | no restrictions                 | small gatherings           |
| 5           | 10-08-20 | no restrictions                 | large gatherings           |
| post lockdown| 31-08-20| no restrictions                 | no restrictions           |

Table 1
Travel and gathering restrictions during the 5 phases of the Irish roadmap.
Table 2
Parameters for the 5 phases of the Irish roadmap. $\beta_m$ and $R_0$ are computed from the system parameters.

| Phase             | $\beta$ | $c$   | $\mu$ | $\kappa$ | $\alpha$ | $\delta_m$ | $R_0$ |
|-------------------|---------|-------|-------|----------|----------|------------|-------|
| pre lockdown      | 0.37    | 1     | 0.1   | 0.1      | 0.0028   | 0.37       | 1.82  |
| school closure    | 0.37    | 0.7   | 0.1   | 0.1      | 0.0028   | 0.26       | 1.28  |
| initial lockdown  | 0.37    | 0.4   | 0.1   | 0.1      | 0.0028   | 0.15       | 0.73  |
| 1                 | 0.37    | 0.4   | 0.1   | 0.1      | 0.0028   | 0.15       | 0.73  |
| 2                 | 0.37    | 0.5   | 0.1   | 0.1      | 0.0028   | 0.19       | 0.91  |
| 3                 | 0.37    | 0.6   | 0.1   | 0.1      | 0.0028   | 0.22       | 1.09  |
| 4                 | 0.37    | 0.65  | 0.1   | 0.1      | 0.0028   | 0.24       | 1.19  |
| 5                 | 0.37    | 0.7   | 0.1   | 0.1      | 0.0028   | 0.26       | 1.28  |
| post lockdown     | 0.37    | 0.75  | 0.1   | 0.1      | 0.0028   | 0.28       | 1.37  |

4.3. Dynamic interventions

Finally, we discuss the likely scenario of a flexible lockdown policy that monitors the prevalence of the disease within the island and aims to keep it below the capacity of the health-care system. As a simple measure, we define some threshold $I_{th}$ as the maximum number of infected that can be present before we go back into lockdown and go through earlier phases again. Such a procedure corresponds to dynamic (Chowdhury et al., 2020; Killeen & Kiware, 2020; Kissler et al., 2020) or active (Ames et al., 2020) interventions. Specifically, we simulate until phase 3 as before to maintain the agreement with reported cases. Afterwards, if the condition $I_{th} > \sum I_i$ is met, the system goes back to phase 1 and then progresses from there until the condition is met again.

**Island Interventions.** In detail, we consider two scenarios with different threshold parameters, which are inspired by the reported, maximum number of hospitalized cases during the first wave in April 2020 (Health Protection Surveillance Centre (HPSC), 2020b): (i) $I_{th} = 10^4$, as shown in Fig. 7 and (ii) $I_{th} = 10^3$ as depicted in Fig. 8. In both figures, the worst-case scenario of Fig. 2 is added as dashed curves. Note that the curves of recovered $R$ (orange) and deaths $D$ (red) are only increasing at a low rate. For the case of $I_{th} = 10^4$, for instance, these values arrive at $10^5$ after two years and thus, way below any effect of herd immunity.

Under the assumptions the considered network model, it would take many years for a significant depletion of the pool of susceptibles. Extrapolating from an extended run of the model similar to Fig. 7, we find that it would take roughly 27 years for the number of recovered to reach half the population, leading to a total number of approximately 92,000 deaths over the course of many waves. Even with a large threshold $I_{th} = 10^4$ for the maximum number of infected cases, the feasibility of such a strategy is questionable due to the time it would take to reach herd immunity of on the island.

The re-introduction of lockdown measures to account for rising numbers of infected can change the stability of the disease-free equilibria $I_c = 0$. However, as soon as the restrictions are relaxed, e.g., during phases 3 to 5, an unstable family of these equilibria re-emerges and the outbreak pattern repeats. Suppressing the occurrence of unstable disease-free equilibria requires other forms of interventions. One could consider, for example, a fishing-type extension of the model (Amann, 2020; Beverton & Holt, 1993; Christensen & Walters, 2004). This will basically turn the rate $\kappa$ in the local dynamics (1) into a function of the number of infected, e.g., $\kappa(I) = h I_{th} I_c$ with a consumption/fishing rate $h$ (here: rate of isolation) and a reference capacity $I_c$ for $\kappa(I) = h/2$. This $I$ dependence will account for contact-tracing effects, which are especially effective for low numbers of infected. As a consequence, a high rate $h$ of isolation will be able to push the basic reproduction number [cf. Eq. (2) or Eq. (6)] below the critical value of $R_0^{crit} = 1$ and change the stability of the equilibria $I_c = 0$.

**Regional Interventions.** Next, we consider another scenario that accounts for regional lockdowns at the level of counties. In our dynamic intervention model this corresponds to each county having its own $I_{th}$. Thus, once the county surpasses this threshold, another lockdown is triggered sending the county back to phase 1. When lockdown phases differ between counties, we assume that the travel restrictions in and out are followed with the compliance of the county being travelled to. When we run the simulations using the regional interventions, the model operates at the level of electoral divisions and superoutput areas but their travel and interaction dynamics are consistent across counties. The parameters used for each phase are the same as those given in Tables 2 and 1. For this run we took the maximum number of infected per county $I_{th}^{count} = 1400$ and as usual the model is allowed to run without intervention until phase 3 to match the historic number of deaths.

From Fig. 9 we see some interesting dynamics when we view the evolution of the number of infected in each county. After Dublin re-enters phase 1 due to dynamic lockdown after only a few days the numbers start rising again. This is not due to its own dynamics, but due to its interaction with neighbouring counties, which are not yet in lockdown. It is only once all the other counties enter phase 1 lockdown do we see the numbers in Dublin and Cork begin to drop. We conclude that treating such counties in isolation is not a good strategy due the strong coupling between each county that is not eliminated when some counties remain free to travel even only within their own borders. Fig. 10 shows the geographic distribution of the disease during the largest peak of Dublin seen in Fig. 9 at the level of electoral divisions and superoutput areas in panel (a) and at the level of counties in panel (b).

Next, we investigate the impact of compliance to movement restrictions. Fig. 11 depicts the time series of the $I$ compartment for 100%, 99%, 90%, and 70% compliance (cf. Fig. 9 for the case of 80% compliance). One can see that already a
Fig. 2. Number of individuals (aggregated over the entire population/network) belonging to each compartment as stated in the legend over time with a log scale on the y-axis. Parameters as in Table 2 and an 80% compliance to movement restrictions. Each successive lockdown phase is indicated by the differently colored shaded regions on the plot. After the initial lockdown (violet), phases 1–5 last 3 weeks each.

Fig. 3. The basic and effective reproduction numbers $R_0$ (black dashed) and $R_0^*$ (red dash-dotted) over time as calculated from the basic reproduction number and spectral radius of the next generation matrix given by Eq. (5), respectively, together with the number of infected (blue solid) in the network in the doomsday scenario as shown in Fig. 2.

Fig. 4. The spatial distribution of infected individuals per km$^2$ around the two distinct peaks shown in Fig. 2 at (a) Day 50 (b) Day 320. Parameters as in Table 2.
small amount of non-compliance results in outbreaks with numbers above the threshold. For example, 99% compliance shown in panel (b) triggers numbers of infected as high as during the first wave in March/April 2020.

Fig. 12 shows the time evolution of infected aggregated over both Dublin and Cork county for different levels of compliance (cf. Figs. 9 and 11). These examples show that lower compliance yields higher peaks after the introduction of regional interventions.

5. Conclusion

We have proposed a dynamical network model to study the spreading of SARS-CoV-2 that accounts for travel or commuting between nodes of the network. As a case study, we have presented results from numerical simulations for Ireland, where the model is informed from publicly available data on geographical regions, population, and mobility in both the Republic of Ireland and Northern Ireland. The network model is flexible in terms of spatial granularity and parameter selection. We have focused on the governmental 5-phase roadmap for reopening society & business and considered different initial conditions, parameters, and mobility data including realistic levels of adherence to movement restrictions. We have observed that the roadmap will lead to a decline of case numbers. However, once the restrictions are lifted, the numbers will rise again, even if an increased level of awareness is considered. This is in excellent agreement with reported case numbers (cf. Appendix B).

Finally, we have elaborated on a procedure of dynamic interventions that re-introduce lockdown measures, if a certain threshold of infection is reached. Following such a protocol, the overall number of infected cases can be controlled and safely kept below a prescribed threshold, which can be selected according to the capacity of the health-care system. We have
**Fig. 7.** The number of individuals belonging to each compartment over time (using the parameters in Table 2) under a dynamic lockdown strategy taking $k_{th} = 10^4$. The dashed lines are the equivalent model without the dynamic lockdown strategy shown in Fig. 2.

**Fig. 8.** The number of individuals belonging to each compartment over time (using the parameters in Table 2) under a dynamic lockdown strategy taking $k_{th} = 10^3$. The dashed lines are the equivalent model without the dynamic lockdown strategy shown in Fig. 2.

**Fig. 9.** Number of individuals (aggregated over counties) belonging to the $I$ compartment. Parameters as in Table 2. Each successive lockdown phase is indicated by the differently colored shaded regions on the plot up until the lockdown phases for each county become desynchronised. After the initial lockdown (violet), phases 1–5 last 3 weeks each. The four largest counties by population are explicitly denoted in the legend.
considered that containment measures would be re-introduced on a country-wide level, but a more regional approach (county level) appears feasible as well. It is likely that in the long term, the disease will be contained best by vaccination and not by the development of natural herd immunity (Alwan et al., 2020). While we have not considered the effects of vaccination in this paper, in our model naturally acquired herd immunity, or population resistance, will develop slowly over a long period of time. This is true even with our assumption that infection confers long-lasting resistance, and it is far from clear that this actually the case. Therefore the model supports the view that effective herd immunity will best be obtained with widespread immunisation (Alwan et al., 2020) although factors such the duration of immunity, efficacy in different age demographics or ethnic groups, likely vaccination uptake, effects of different vaccines, as well as efficacy against novel variants are poorly understood. Our model could be used to study these variables but this is outside the scope of this paper.

Fig. 10. The geographic distribution of infected individuals in the simulation shown by the time-series in Fig. 9 on the 7th December, 2020. Panel (a) shows the number of infected per km² in each electoral division and superoutput area. Panel (b) shows the number of infected in each county.

Fig. 11. Number of individuals (aggregated over counties) belonging to the I compartment for compliance of 100%, 99%, 90%, and 70% in panels (a), (b), (c), and (d) respectively. Other parameters as in Table 2. Each successive lockdown phase is indicated by the differently colored shaded regions on the plot up until the lockdown phases for each county become desynchronised. After the initial lockdown (violet), phases 1–5 last 3 weeks each. The four largest counties by population are explicitly denoted in the legend.
We hope that the results presented in this discussion paper will stimulate scientific and public discussion beyond the island of Ireland about the purpose and effect of lockdown measures in the months ahead (cf. Appendix C). These are plausible scenarios given the time frame of vaccination campaigns that have just begun.

Authors’ contributions

RH, KM, SW, MOR, and PH designed the study. SW and RH developed the theory. RH and MS performed the numerical simulations and analyzed the data. All authors discussed the results and contributed to the writing of the article. All authors read and approved the final version of the manuscript.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Next Generation Matrix

In this appendix, we provide a detailed derivation of the next generation matrix given Eq. (5). We start by splitting the networked SIXRD equation (4) into the parts that cause new infections and all other changes. We shall denote the rate of appearance of new infections at node $i$ in compartment $U$ by $F_{U,i}(x)$. Similarly, let $V_{U,i}(x)$ be the rate of change of individuals in compartment $U$ and node $i$ by all other means such that $\dot{U}_i = F_{U,i} - V_{U,i}$. This yields the following equations:

$$F_{S,i} = 0 \quad (7a)$$

$$F_{I_i} = \beta e \left( S_i - \frac{S_i}{N_i} \sum_{\substack{j = 1 \atop j \neq i}}^n N_{ij} \right) \frac{I_i \sum_{j=1}^n \frac{I_j}{N_j} N_{ij} - \frac{I_i}{N_i} \sum_{j=1, j \neq i}^n N_{ij}}{N_i \sum_{j=1, j \neq i}^n (N_{ji} - N_j)}$$

$$+ \beta e \frac{S_i}{N_i} \sum_{\substack{j = 1 \atop j \neq i}}^n N_{ij} \frac{I_j \sum_{k=1, k \neq j}^n \frac{I_k}{N_k} N_{kj} - \frac{I_j}{N_j} \sum_{k=1, k \neq j}^n N_{kj}}{N_j \sum_{k=1, k \neq j}^n (N_{kj} - N_k)} \quad (7b)$$

$$F_{X,i} = 0 \quad (7c)$$

Fig. 12. The time-series for the evolution of the $I$ compartment in (a) Dublin and (b) Cork extracted from Figs. 11 and 9 for different values of compliance: 70% (blue), 80% (orange), 90% (green), 99% (red), and 100% (blue).
it is an important note that the movement of individuals form I to X does not count as new infections. Details on the impact of the travel statistics during the different phases of the roadmap can be found in Appendix D.

\[
\begin{align*}
\mathcal{F}_{R,i} &= 0 \\
\mathcal{F}_{D,i} &= 0.
\end{align*}
\] (7d) (7e)

\[\nu_{S,i} = \beta c \left( S_i - \frac{S_i}{N_i} \sum_{j=1, j \neq i}^N N_{ij} \right) \frac{I_i + \sum_{j=1, j \neq i}^N I_j N_{ij}}{N_i + \sum_{j=1, j \neq i}^N (N_{ij} - N_{ji})} \] (8a)

\[\nu_{I,i} = \mu I_i + aI_i + kI_i \] (8b)

\[\nu_{X,i} = -kI_i + \mu X_i + aX_i \] (8c)

\[\nu_{R,i} = -\mu I_i - \mu X_i \] (8d)

\[\nu_{D,i} = -aI_i - aX_i. \] (8e)

for the above equations, we replace the coupling variables with their actual values, i.e. we take \( S_{ij} = \frac{S_i}{N_i} N_{ij}, I_{ij} = \frac{I_i}{N_i} N_{ij}, \ldots \) etc. Writing these equations in vector form we obtain

\[
\mathcal{F} = \begin{bmatrix}
\mathcal{F}_{I,1} \\
\mathcal{F}_{I,n} \\
\mathcal{F}_{X,1} \\
\mathcal{F}_{X,n} \\
\mathcal{F}_{S,1} \\
\mathcal{F}_{S,n} \\
\mathcal{F}_{R,1} \\
\mathcal{F}_{R,n} \\
\mathcal{F}_{D,1} \\
\mathcal{F}_{D,n}
\end{bmatrix}, \quad \nu = \begin{bmatrix}
\nu_{I,1} \\
\nu_{I,n} \\
\nu_{X,1} \\
\nu_{X,n} \\
\nu_{S,1} \\
\nu_{S,n} \\
\nu_{R,1} \\
\nu_{R,n} \\
\nu_{D,1} \\
\nu_{D,n}
\end{bmatrix}
\] (9)

the disease-free equilibria are all solutions that satisfy

\[
x_0 = (I_1, \ldots, I_n, X_1, \ldots, X_n, S_1, \ldots, S_n, R_1, \ldots, R_n, D_1, \ldots, D_n) \] (10)

\[
= (0, \ldots, 0, 0, \ldots, 0, S_1, \ldots, S_n, R_1, \ldots, R_n, D_1, \ldots, D_n) \] (11)

taking the derivatives of \( \mathcal{F} \) and \( \nu \) at the disease-free equilibria, we find that

\[
D\mathcal{F}(x_0) = \begin{bmatrix}
F & 0 \\
0 & 0
\end{bmatrix} \quad \text{and} \quad D\nu(x_0) = \begin{bmatrix}
\nu & 0 \\
J_1 & J_2
\end{bmatrix}
\] (12)

as shown in (van den Driessche & Watmough, 2002). We obtain
\[ F_{ij} = \beta c \left[ \left( S_i - \frac{S_i}{N_i} \sum_{k=1, k \neq i}^{n} N_{ik} \right) \frac{\delta_{ij} \left( 1 - \sum_{k=1, k \neq i}^{n} \frac{N_{ik}}{N_j} \right) + \frac{N_{ii}}{N_j}}{N_i + \sum_{k=1, k \neq i}^{n} \left( N_{ii} - N_{ik} \right)} + \frac{S_i}{N_i} \sum_{k=1, k \neq i}^{n} N_{ik} \frac{\delta_{jk} \left( 1 - \sum_{l=1, l \neq j}^{n} \frac{N_{jl}}{N_j} \right) + \frac{N_{jk}}{N_j}}{N_k + \sum_{l=1, l \neq k}^{n} \left( N_{ik} - N_{kl} \right)} \right] \] (13)

\[ V_{ij} = (\mu + \alpha + \kappa) \delta_{ij} \]. \] (14)

where \( \delta_{ij} \) denotes the Kronecker delta: \( \delta_{ij} = 1 \) for \( i = j \) and zero otherwise. Finally, following [40], the next generation matrix is given by

\[ F V_{ij}^1 = \frac{\beta c}{\mu + \alpha + \kappa} \left[ \left( S_i - \frac{S_i}{N_i} \sum_{k=1, k \neq i}^{n} N_{ik} \right) \frac{\delta_{ij} \left( 1 - \sum_{k=1, k \neq i}^{n} \frac{N_{ik}}{N_j} \right) + \frac{N_{ii}}{N_j}}{N_i + \sum_{k=1, k \neq i}^{n} \left( N_{ii} - N_{ik} \right)} + \frac{S_i}{N_i} \sum_{k=1, k \neq i}^{n} N_{ik} \frac{\delta_{jk} \left( 1 - \sum_{l=1, l \neq j}^{n} \frac{N_{jl}}{N_j} \right) + \frac{N_{jk}}{N_j}}{N_k + \sum_{l=1, l \neq k}^{n} \left( N_{ik} - N_{kl} \right)} \right] \] (15)

### Appendix B. Comparison to Real-World Data

Fig. 13 shows a comparison of model results for a countrywide dynamic lockdown scenario with daily and cumulative figures for reported deaths and infections from the Health Surveillance Centre (HPSC). Following the initial lockdown and phased reopening as outlined in Table 1, the threshold of infection was chosen to be 10000 in these simulations, before a national lockdown was reintroduced. This figure is adjustable based on current public health guidelines. Panels 13(a) and 13(b) compare the model to daily and cumulative reported figures for infections, respectively. Moreover, panels 13(c) and 13(d) compare the model results to daily and cumulative figures for deaths. The blue curves correspond to the model results, and the red curves refer to the data reported by HPSC. A 14-day moving average has been applied to data from the HPSC. Note that the infected compartment in the model does not distinguish between confirmed/undetected symptomatic/asymp- tomatic cases. This explains the larger number of infections in the model in which we see a clear qualitative agreement in the number of infections over time. The number of deaths, however, is more reliable and the model reproduces the actual numbers with large agreement. Overall, we find that the model results compare favourably to these real-world data.
Appendix C. Projected Post-Christmas Lockdown

As a final scenario, we finish with a projection on the possible post-Christmas projection during which we expect to see the third wave of lockdown. Fig. 14 considers the current period of the second wave and the corresponding lockdown with level-5 movement restrictions in the Republic of Ireland. After a partial lifting of these measures scheduled for 1st December, 2020, we find that the case numbers will rise again. Note that the numbers remain significantly higher than during summer 2020. This is illustrated by the maps in Fig. 15, which compares the situation by the end of June and 1st December, 2020. We project that the numbers will reach a high level similar to the introduction of the current lockdown by late-January 2021 and thus will require another lockdown of similar duration.

Fig. 14. Simulating the evolution of the disease such that the historic deaths are matched up until level 5 and projected from then on. There are two subsequent periods of lockdown: the current lockdown that started mid-November and is expected to lift 1st December, 2020. The next lockdown, which occurs in January 2021, is projected based on the number of the numbers of the November lockdown. Other parameters are as given in Table 2.
Fig. 15. The geographic distribution of infected individuals per km² in each electoral division and superoutput area in the simulation shown by the time-series in Fig. 14. Panel (a) shows the geographic distribution during the lowest incidence of infected during the summer and panel (b) shows the geographic distribution at the end of the lockdown projected to end the 1st December, 2020. Other parameters are as given in Table 2.

Appendix D. Movement data

Fig. 16. The complementary cumulative proportion of individuals travel distance with the 2 km (orange dashed), 5 km (green dash-dotted), and 20 km (red dotted) restrictions in place for a compliance level of 80%. The original, pre-lockdown data, which are used during periods with no travel restrictions, is shown as blue solid curve. The inset shows the full range in a semi-log scale.

In this appendix, we demonstrate how travel restrictions affect the movement of individuals in the simulations. We extracted the number of travelling individuals \(N_{ij}\) between locations \(i\) and \(j\) from CSO data and generated a probability distribution of travelling from one electoral division/super-output area to another as a function of population density and distance. Once the travel restrictions are put in-place, the probabilities of travelling to areas outside this distance are scaled by the compliance factor making individuals less likely to travel there. Each \(N_{ij}\) was then generated from these probability distributions inferred from real commuting data within the country.

The difference in these movement patterns can be seen in Fig. 16 where the cumulative distribution of travel distances is plotted for each level of travel restrictions (2 km, 5 km, and 20 km) and the original data prior to the first lockdown period, which are used during periods with no travel restrictions (cf. Table 1). The inset shows the original distribution for the full range in a semi-logarithmic scale. The kinks in the distribution of the restricted movement occur at the respective radii. Larger distances still occur due to non-perfect compliance (here: 80%).
Fig. 17. A visual representation of the journeys out of a particular electoral division in the Dublin area (red area marked by the arrow) to surrounding electoral divisions with different restrictions in place with a compliance of 80%. Panels (a), (b), (c), and (d) correspond to travel restrictions of 2 km, 5 km, 20 km, and none, respectively.

In addition, we show in Fig. 17 how the travels out of a particular area (here an electoral division in Dublin; red area marked by the arrow) are affected by the different travel restrictions. As the travel radius is reduced, the movement is confined to a more local vicinity. Note that travel beyond the travel radius is still present due to non-compliance.

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