Since January 2020 Elsevier has created a COVID-19 resource centre with free information in English and Mandarin on the novel coronavirus COVID-19. The COVID-19 resource centre is hosted on Elsevier Connect, the company's public news and information website.

Elsevier hereby grants permission to make all its COVID-19-related research that is available on the COVID-19 resource centre - including this research content - immediately available in PubMed Central and other publicly funded repositories, such as the WHO COVID database with rights for unrestricted research re-use and analyses in any form or by any means with acknowledgement of the original source. These permissions are granted for free by Elsevier for as long as the COVID-19 resource centre remains active.
Towards predicting COVID-19 infection waves: A random-walk Monte Carlo simulation approach

D.P. Mahapatra, S. Triambak

Abstract

Phenomenological and deterministic models are often used for the estimation of transmission parameters in an epidemic and for the prediction of its growth trajectory. Such analyses are usually based on single peak outbreak dynamics. In light of the present COVID-19 pandemic, there is a pressing need to better understand observed epidemic growth with multiple peak structures, preferably using first-principles methods. Along the lines of our previous work [Physica A 574, 126014 (2021)], here we apply 2D random-walk Monte Carlo calculations to better understand COVID-19 spread through contact interactions. Lockdown scenarios and all other control interventions are imposed through mobility restrictions and a regulation of the infection rate within the stochastically interacting population. The susceptible, infected and recovered populations are tracked over time, with daily infection rates obtained without recourse to the solution of differential equations.

The simulations were carried out for population densities corresponding to four countries, India, Serbia, South Africa and USA. In all cases our results capture the observed infection growth rates. More importantly, the simulation model is shown to predict secondary and tertiary waves of infections with reasonable accuracy. This predictive nature of multiple wave structures provides a simple and effective tool that may be useful in planning mitigation strategies during the present pandemic.

1. Introduction

In the midst of the current COVID-19 pandemic, there is a continuing need to accurately model region-specific infection and mortality data, so that intervention methods and containment strategies can be planned accordingly. The simplest picture of epidemic growth is at most times provided by phenomenological models that are based on logistic growth [1–7]. However, real-time interventions that may affect the trajectory of the growth curve are not incorporated in such models. More commonly, epidemiological modeling uses compartmentalized populations based on the SIR model [8] and its variants (see, for example [9–11]) that follow the time evolution of susceptible (S), infected (I) and recovered (R) populations, among others [12]. Usually such deterministic models do not involve a stochastic formulation, which takes into account the random aspects of human mobility. Based on the early work by Bartlett [13], such randomness can be incorporated through the equations [14–16]

\[
\frac{\partial S}{\partial t} = D_S \nabla^2 S - \beta SI \tag{1}
\]

\[
\frac{\partial I}{\partial t} = D_I \nabla^2 I + \beta SI - \mu I. \tag{2}
\]

Here, \(\nabla^2 = \left(\frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2}\right)\), \(S(x,y,t)\) and \(I(x,y,t)\) are the spatial densities of the susceptible and infectious components of the population. The \(D_S \nabla^2(S,I)\) diffusion terms represent the spatial movement of both susceptibles as well as infectives. The parameters \(\beta\) and \(\mu\) represent the infection and recovery rates respectively. Such formulations have also been extended to biased [17] and Lévy [18] random walks. More recently, stochastic models have been developed using Markovian chains [19,20], contact and community networks [21,22], Bayesian modeling [23] etc. The effects of mobility restrictions have also been studied using Monte Carlo techniques [24–26] that inherently include the diffusion terms mentioned above. In most studies, the analyses that are solely based on single epidemic growth curves do not adequately explain the multiple wave structures seen in global COVID-19 data. For example, it was recently proposed that a superposition of epidemic
waves [27] could be used to describe COVID-19 growth curves. However this does not describe the observed multiple waves of infection for individual countries, that are well separated in time.

The present work uses an uncorrelated random walk approach to study COVID-19 infection spread via contact interactions. Our simulations show that the number of successive waves of COVID-19 infection in specific countries depend on their underlying effective population density, the intermixing rate, and most importantly the timing and the duration of the control interventions imposed on/by the population. The effects of these interventions are visible as modulations in the infection rate. As test cases, we compare our simulation results with reported data from India, South Africa, Serbia, South Africa and USA. We show that multiple peak structures of the pandemic waves are reasonably well reproduced by the simulations.

2. The random walk Monte Carlo method

Our random walk simulation model is described in our previous paper [25] and similar in approach to the work reported in Ref. [28]. Briefly, people belonging to a region of population $N_i$ are described as points that execute random walks on a 2-dimensional plane. The speed of infection growth depends on the jump distance $l$ for each point, which we assume is a multiple of the mean separation $\langle r \rangle$ between points. In Cartesian coordinates, the jump components are simply $\cos \theta$ and $\sin \theta$, with $\theta$ generated randomly between 0 and $2\pi$ from a uniform distribution. Unlike other approaches [29,30], we do not consider a lattice or apply periodic boundary conditions. Instead, if a jump takes a point out of the area considered, it is reflected back into the system in a random direction. In all cases we consider individuals in a 1 km$^2$ unit area element. Therefore we can use the words population and population density interchangeably. For $N$ random walkers per unit area, $\langle r \rangle = \sqrt{2N}$ in units of km. A separation distance of $\leq 2$ m between individuals is taken as the ‘contact’ distance [25].

The simulation starts with the introduction of an infected individual in state $I_l$. The disease then spreads through contact interactions, resulting in a drop in the number of susceptible points, $S$, which at time $t = 0$ equals $N - 1$. Each time step corresponds to one random jump executed by all the points and we take that as one day. The number of infected points increase with such steps, based on the number of contacts between susceptible and infectious individuals. The populations in $S$, $I$ and $R$ states are $N_S$, $N_I$ and $N_R$ respectively, so that $N = N_S + N_I + N_R$ is preserved at all times. This results in an SIR compartmentalization.

In the next step of the simulations, a recovery rate was incorporated in a slightly different manner than used previously in Ref. [25]. We assume that 80% of the infected population (picked at random) ultimately recover, becoming immune, and keep track of the number of time-steps (days) taken by an infected point before it goes into a recovered state. This time period is taken to be 35 days, and determined from a comparison of our simulated results with data. This choice of $\mu = 1/35$ day$^{-1}$ is not unfounded. The incubation period of the coronavirus disease (after which symptoms develop) for infected individuals is found to be in the range of 8.2–15.6 days at the 97.5th percentile level [31]. Furthermore, recent studies have shown that COVID-19 recovery times have an average value of about 25–28 days [32,33]. Given that these recovery times are evaluated after symptoms develop, a total of 35 days after infection is a reasonably accurate estimate.

From here on, one can follow two equivalent approaches to incorporate the various intervention and mitigation strategies, usually employed after the start of an epidemic such as COVID-19. A decisive criterion is the strict imposition of a lockdown that places significant mobility constraints on a majority of the population. This effectively moderates the growth in $N_i$, so that it is compensated by the recovery rate $\mu$. Other interventions such as vaccinations, mask mandates and behavioral changes by the population (social distancing, bubbles, etc.) additionally contribute to stalling the epidemic growth.

In our earlier work [25], we studied contact-interactions between random walkers on a plane, with $\beta$ equal to unity. This approach can be modified to incorporate the effects of all control interventions, exclusively via mobility restrictions on the stochastic agents. Such restrictions would impede the growth of the epidemic curve, during which time the number of recoveries continue to rise. However this is never a permanent solution. In more realistic scenarios the restrictions are relaxed from time to time (such as with lockdowns). In such a situation the epidemic growth continues, usually with a steep rise in the number of infections that is shifted in time.

An alternative approach would be to assume $\beta$ values that are $< 1$ for specific cases under consideration, during the times that the control interventions and mitigation strategies are followed. For example, vaccinations, mask mandates, etc. lead to a drop in the number of susceptible individuals that can be infected. The effect of this drop can also be incorporated within the parameter $\beta$. This fractional $\beta$ is implemented by invoking a random number $r$ uniformly between 0 and 1, such that a change in state from $S \rightarrow I$ only occurs when $r \leq \beta$. We show below that both approaches yield similar results with regard to predicting multiple infection waves. However we prefer the second approach, as the former may mislead the reader in terms of the difference between actual lockdowns imposed on a population and effective mobility restrictions on the random walkers in the simulations.

The simulations were performed for comparison with reported data for India, South Africa, Serbia and USA. The adjustable parameters were the population density (two values were used for this work, $N = 5k$ and $10k$), $\beta$, the percentage of mobile walkers, the duration during which mobility was restricted and the jump length $l$. The recipe for the simulations was as follows. We consider the start date of the growth curve to be $D_0$. The intervention and mitigation strategies during this initial phase would effectively produce a first wave peak in $N_i$ on date $D_1$, which is obtained from reported data. The parameters in the simulation are then adjusted, so that the Monte Carlo results reproduce the observed first wave peak in the fractional daily infection rate. The simulated results are obtained from $\frac{N_i(t + \Delta t) - N_i(t)}{N_i(t)}$, for $\Delta t = 1$ day. Here, it should be noted that since $\frac{\partial N_i}{\partial t} \propto N_i$, the peaks in $N_i$ roughly coincide with the daily infection rate peaks along the time axis. These simulated results are compared to scaled-down data from the World Health Organization (WHO) [34]. This comparison is then used to infer if secondary or tertiary pandemic waves subsequently appear at later dates, due to the control interventions imposed in previous time windows. This is based on the premise that the control interventions and mitigation strategies would reduce the slope of the growth curve in the time duration that they are imposed (or followed). At the end of each such duration (when restrictions, mandates etc. are relaxed) there would be growth again. This effectively results in growth curves with multiple peaks in daily infection rates, that are shifted in time and interpreted as epidemic waves. We discuss a few country specific results below.

3. Analysis

3.1. India

We first consider the case of India, since the reported data show two distinct pandemic peaks. Furthermore, there has been a lot of speculation regarding the appearance of a large third wave.

\[1\text{ We only consider events where the disease trajectory extends beyond day 500.}\]
following the devastating second wave in 2021. The data show a first wave peak around Sep 15, 2020 and a second wave peak around May 06, 2021. These two structures contain in them all effects of interventions and mitigation strategies, including imposed lockdowns. We take D0 to be April 22, 2020, with the first peak (D1) around day 150 (September 15, 2020). Since this was the first country that we studied, we performed several simulations to better understand the general shapes of the curves, obtained for different parameter values. The first two sets of simulations used jump steps of \( l = 1(r) \) and \( 2(r) \), with \( \beta = 1 \). In the third set, we used \( l = 1(r) \) from D0 to D1 and \( l = 2(r) \) beyond D1. The fractional \( N_i \) results for different parameter values and \( N = 10k \) are shown in Fig. 1. As evident in the figure, a larger amplitude infection peak is obtained when \( l = 2(r) \) step sizes are used, with a peak value of approximately 0.45 around day 140. For this case the number of infections drop to negligible levels after about 200 days. In comparison, \( l = 1(r) \) produces a broader and smaller peak that is shifted to a later date (near day 230). On the other hand, the \( l = 2(r) \) and \( l = 1(r) \) combination yields two distinct peak structures, as shown in the figure. In the next set of simulations we used this combination, together with \( \beta \) values < 1 that account for intervention/mitigation strategies, which would lead to a drop in the \( N_i \) fraction and consequently the daily infection rate. These results are also shown in Fig. 1, for different values of \( \beta \) during the time period D1 to D2 (the falling part of the first curve), with the later date representing the beginning of the second wave. Beyond D2 we assume \( \beta = 1 \) for the rise of the second wave. Guided by reported data we take D2 to be day 300.

We observe that the shapes of the fractional \( N_i \) curves, obtained with \( \beta = 0.3 \) from dates D1 to D2 (magenta curve in Fig. 1) are very similar to scaled down daily infection rates obtained from the WHO, including the relative amplitudes for the two peaks. This is further validated in Fig. 2, which shows a comparison between the reported daily rates and the simulated results (assuming \( \beta = 0.3 \)).4 The excellent agreement between the simulations and the reported data for the two waves is noteworthy. Similar agreement is not obtained from other parameter value combinations (such as the use of only \( l = 1(r) \) or a population density \( N = 5k \) for the \( 1(r) + 2(r) \) jump-step combination. The results from the latter simulations are shown in Fig. 2 for comparison). The infection rate \( \beta = 0.3 \), used for time periods when effective mitigation/intervention measures are assumed in the simulations, is consistent with those obtained with a deterministic SIOQR model that also took into account the quarantined and confirmed recovered portions of the population (c.f. Table 2 in Ref. [33]).

It is also important to point out here that in addition to the aforementioned agreement with data, the second wave peak in Fig. 2 emerges naturally from our simulations, as a result of interventions imposed previously, mainly through the parameter \( \beta \). Furthermore, the simulations also show no indication of a third peak that may appear on applying additional interventions (through \( \beta \)) in the second wave. This is supported by Fig. 3, which shows the time development of the individual S, I and R components for these data. One can see that after day 500 \( N_i \) drops to nearly zero, while \( N_i \) saturates at \( \sim 90\% \). It is apparent that at this point there are not enough infectious agents remaining to drive the curve and infect the small remainder of the susceptible population. This does not indicate a significant third wave. However, it may be noted that we assume all recovered individuals to be immune in our simulations. During the course of this work, several countries reported a surge in COVID-19 infections due to the emergence of the SARS-CoV-2 Omicron variant [35]. This variant is characterized by an unusually large number of mutations in the spike protein and an ability to escape vaccine induced immunity [36]. In light of this recent development, we performed additional simulations that allow reinfections. Our preliminary results show that a small fraction of reinfections (\( \lesssim 0.5\% \)) would increase the susceptible population

---

4 All data discussed in this work were smoothed over the transition periods when a change in \( \beta \) was incorporated.
significantly, enough to cause a third wave for India. Further investigations that delve into this aspect and take into consideration the use of a different infection probability $\beta$ (due to the waning of vaccine-induced immunity) will be useful in this regard.

### 3.2. Serbia

Serbia is an interesting case study. The data show that the pandemic trajectory comprises two small peaks followed by two large peaks [34]. The country was grappling with another wave of infections during the time of this work. It was therefore interesting to see whether the last wave is a consequence of the interventions imposed earlier. Here we take $D_0$ as March 04, 2020, following which there are four pandemic peaks in the daily rates (excluding the latest wave). In our simulations these correspond to time domains where $\beta$ values $< 1$ are used. We identify these to be days 50–100, 130–180, 250–320 and 380–450. Since Serbia has a much lower population density compared to India, we used $N = 5k/\text{unit area and } l = 2(r)$. Guided by the results for India, we again assumed $\beta = 0.3$ for the periods that correspond to systematic drops in the observed rates, following each observed peak in the epidemic curve.

The simulation results are shown in Fig. 4, together with corresponding scaled down data reported by the WHO. One can clearly see reasonable agreement between the two, with the simulations adequately predicting the peak of the fifth wave that was ongoing at the time of this work.

### 3.3. South Africa

Next we looked at the data for South Africa, whose growth trajectory is interesting to follow, as the Omicron variant was first identified in the region [35].

We took $D_0$ to be Mar 04, 2020. There are three pandemic waves, with peaks around days 150, 300 and 500. To simulate the initial wave we used $N = 5k$ and effective interventions ($\beta = 0.3$) from day 150 to 250, with $l = 2(r)$. On imposing a second period of lower infections from day 320 to day 420, we obtain consistent results for the first two waves. These are plotted in Fig. 5. In comparison, the peak around day 500 was found to be too broad compared to the reported data from the WHO. We note that a further intervention period from day 500 to day 600 (again with $\beta = 0.3$)

![Fig. 4. Simulated daily rates of Serbia, compared with scaled down WHO data. The simulation results were averaged over 10 sets. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)](image1)

![Fig. 5. Simulated results for South Africa, compared with scaled down WHO data. The simulations assumed $N = 5k$ and were averaged over 10 sets. The blue curve is generated assuming no control interventions beyond day 500. The green curve assumes such interventions ($\beta = 0.3$) from day 500–600. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)](image2)

![Fig. 6. Simulated rates for USA compared with scaled-down WHO data. These simulations assumed $N = 10k, l = 0.7(r)$ and were averaged over 5 runs. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)](image3)

yields daily rates for the third wave that are in reasonably good agreement with the WHO data. These results also show a clear fourth wave, whose position depends on the imposed intervention in the previous wave. This Omicron driven fourth COVID-19 wave has already been reported for South Africa [37]. It is interesting to note that our predicted fourth wave is rather broad, similar to the blue curve obtained on assuming $\beta = 1$ beyond day 500. Although we have not carried out simulations with imposed control interventions beyond day 700, this similarity suggests the possibility of a fifth wave for South Africa.

### 3.4. USA

Our final set of simulations were for USA, which presents a different challenge as each of its 50 states follow independent mitigation and containment strategies. There are five observed waves of infections, with the third peak being significantly larger than the others. A scaled down version of the reported data is shown in Fig. 6. In our simulations we use $N = 10k$ and take $D_0$ to be January 1, 2020. Based on the WHO data, we introduce three intervention periods, between days 130–180, 230–280, and 400–550. For these time periods we used $l = 0.7(r)$ and $\beta = 0.3$. As Fig. 6 shows, apart from the small peak around day 470, all other features in the data are reasonably well reproduced by the simulations. The simulations also correctly predict a wave after day 500, which was ongoing during the time of this work. On the basis of our results

---

3 For example, our preliminary simulations show that 0.2% of reinfections would cause a third wave peak for India around day 600.

4 South Africa also has a relatively lower population density, similar to Serbia.
one can also anticipate a fresh wave of infections following the peak after day 600. Similar to the other cases studied, we observe that contact interventions following an increase in the daily rate of infections invariably lead to a new epidemic wave later in time, provided there are enough agents available to infect the remaining susceptible fraction.

4. The equivalence between mobility restrictions on the random walkers and imposition of a lower infection rate

One can get similar results as above by using the other approach that assumes $\beta = 1$ at all times. This is obtained by applying mobility restrictions at different time ranges on most of the random walkers, so that the primary infection peaks are reproduced. However, it is important to keep in mind that these time ranges are not the same as periods of imposed lockdowns by governing bodies. We discuss below the results obtained using this analogous approach, for the case of India.

Here again two different sample populations with $N = 5k$ and 10k were used to simulate the data for different jump lengths $l$. All calculations were carried out for $\mu = 1/35$ day$^{-1}$ with a 20% mobile population during the time period spanning from day 120 to day 300. The remaining 80% remain frozen at their positions. Fig. 7 shows the fractional daily rates obtained for the two populations for the intervention period mentioned above, together with a jump distance of $l = 2\langle r \rangle$. The 10k results show excellent agreement with reported data for the second wave. Similarly consistent results are obtained for the other countries studied in this work.

5. Discussion

The results presented in this work show that it is possible to predict the trajectory of an epidemic using a random walk Monte Carlo approach that includes time periods of control interventions. Two equivalent simulation approaches (either using $\beta < 1$ or keeping a percentage of the population immobile when effective interventions were in place) can be followed, as they yield similar results. We observe that for populous countries such as India and USA reasonably good results are obtained on using a higher population density (10k) in the simulations. In comparison, for countries such as Serbia and South Africa (whose population densities are much smaller) a $N = 5k$ works best for the simulations. Conservatively, both these numbers take into account higher floating densities at localized places. In all cases, we show that the use of step-sizes $\leq 2\langle r \rangle$ yield optimal results. The bound on this parameter value can be justified as below.

In our earlier work [25] we showed that there are essentially two limits of growth. If the jump step is too small then the growth is purely quadratic in time [10] and if the jump step is too large the growth is exponential due to homogeneous mixing [38]. The country-specific growth modes described here occur between these two limits when there is sub-exponential behavior [9] due to the containment strategies employed. The power-law nature of the growth is lost for jump steps $\geq 3\langle r \rangle$. On the other hand when the density becomes too high, since our simulations use jump lengths that are proportional to the average inter-person separation, the epidemic growth approaches the quadratic (lower) limit. In such a scenario an alternative approach, such as in Ref. [24] may be better suited for studying the sub-features in a single wave. This may offer an explanation as to why our simulated results for large jump lengths or population densities fail to reproduce the observed data.

It must also be added that in the present formalism the exact total number of cases are not meaningful. Instead, the focus is on the rapid time-shifted increase in the number of infections following intervention periods. These correspond to new waves of infection. On this basis the simulations are used to predict pandemic peaks at later dates, based on earlier data. If one chooses to use a calibration factor based on previously observed peaks, it is also possible to roughly estimate the total number of people infected.

6. Conclusions

To conclude, we use a random walk Monte Carlo simulation technique to better understand contact-based epidemic spread. The model of independent random walkers as infection carriers in two-dimensional space is intuitive and shows promise in better understanding infectious disease outbreaks [25]. It is more accessible than other models and has the ability to capture random interactions that are missed through more conventional approaches. The simulations are shown to predict reasonably accurate disease trajectories in terms of secondary and tertiary waves, for COVID-19 data from four countries with vastly different features.

Declaration of Competing Interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

Smarajit Triambak reports financial support was provided by National Research Foundation.

CRediT authorship contribution statement

D.P. Mahapatra: Conceptualization, Formal analysis, Data curation, Writing – original draft. S. Triambak: Formal analysis, Data curation, Writing – original draft.

Acknowledgment

We are grateful to Prof. Niranjan Barik for several helpful discussions and to Dr. Ara Chutjian for useful remarks. ST acknowledges funding support from the National Research Foundation (South Africa) under grant number 85100.

5 Although not shown explicitly, we find that a higher population of 20k per unit area leads to poor results.

References

[1] Richards FJ. A flexible growth function for empirical use. J Exp Bot 1959;10(2):290–301. doi: 10.1093/jxb/10.2.290.
[2] Roosa K, Lee Y, Luo R, Kirpich A, Rothenberg R, Hyman JM, Yan P, Chowell G. Real-time forecasts of the COVID-19 epidemic in China from February 5th to February 24th, 2020. Infect. Dis. Model. 2020;5:256–63. doi:10.1016/j.idm.2020.02.002.

[3] Wu K, Darcey D, Wang Q, Sornette D. Generalized logistic growth modeling of the COVID-19 outbreak: comparing the dynamics in the 29 provinces in China and in the rest of the world. Nonlinear Dyn 2020;101(3):1561–81. doi:10.1007/s10042-020-06962-6.

[4] Singer HM. The COVID-19 pandemic: growth patterns, power law scaling, and saturation. Phys Biol 2020;17(5):055001. doi:10.1088/1478-3975/ab9b55.

[5] Shen CY. Logistic growth modelling of COVID-19 proliferation in China and its international implications. Int. J. Infect. Dis. 2020;96:582–9. doi:10.1016/j.ijid.2020.04.085.

[6] Chowell G, Tariq A, Hyman JM. A novel sub-epidemic modeling framework for short-term forecasting epidemic waves. BMC Med 2019;17(1):1667–80. doi:10.1186/s12916-019-1466-6.

[7] Triambak S, Mahapatra DP, Mallick N, Sahoo R. A new logistic growth model applied to COVID-19 fatality data. Epidemics 2021;37:100051. doi:10.1016/j.epidem.2021.100015.

[8] Kernack WO, McDendrick AG. A contribution to the mathematical theory of epidemics. Proceedings of the Royal Society of London A: Mathematical, Physical and Engineering Sciences 1927;115(772):700–21.

[9] Maier BF, Brockmann D. Effective containment explains subexponential growth in recent confirmed COVID-19 cases in China. Science 2020;368(6492):742–6. doi:10.1126/science.abb4557.

[10] Brandenburg A. Piecewise quadratic growth during the 2019 novel coronavirus epidemic. Infect. Dis. Model. 2020;5:681–90. doi:10.1016/j.idem.2020.08.014.

[11] He S, Peng Y, Sun K. SEIR modeling of the COVID-19 and its dynamics. Nonlinear Dyn. 2020;101(3):1567–80. doi:10.1007/s11071-020-05743-y.

[12] Agrawal M, Kanitkar M, Vidyasagar M. SUTRA: An Approach to Modelling Pandemics with Asymptomatic Patients, and Applications to COVID-19. arXiv preprint arXiv:2011.09258.

[13] Bartlett MS. Deterministic and stochastic models for recurrent epidemics. In: Neyman J. editor. Proceedings of the Third Berkeley Symposium on Mathematical Statistics and Probability, Volume 4: Contributions to Biology and Problems of Health. 1956.

[14] Fofana AM, Hurford A. Mechanistic movement models to understand epidemic spread. Philosophical Transactions of the Royal Society B: Biological Sciences 2017;372(1719):20160086. doi:10.1098/rstb.2016.0086.

[15] Bailey NTJ. The mathematical theory of infectious diseases and its applications. Charles Griffin and Company Limited; 1975.

[16] Noble J V. Geographic and temporal development of plagues. Nature 1974;250:726–9. doi:10.1038/250726a0.

[17] Codling E, Plank M, Benhamou S. Random walks in biology. J R Soc Interface 2008;5:813–34. doi:10.1098/rsif.2008.0014.

[18] Raichlen DA, Wood BM, Gordon AD, Mabulla AZP, Marlowe FW, Pontzer H. Evidence of Lévy walk foraging patterns in human hunter–gatherers. Proc. Natl. Acad. Sci. 2013. doi:10.1073/pnas.1318616111.

[19] Wang Z, Brockard M, Mignan A, Sornette D. The dynamics of entropy in the COVID-19 outbreaks. Nonlinear Dyn. 2020;101(3). doi:10.1007/s10042-020-06971-5.

[20] Gourieroux C, Jasiak J. Time varying Markov process with partially observed aggregate data: An application to coronavirus. J Econometrics. 2020. doi:10.1016/j.jeconom.2020.09.007.

[21] Karaiskou A. A social network model of COVID-19. PLoS ONE 2020;15(10):1–33. doi:10.1371/journal.pone.0248078.

[22] Hoen AG, Hladish TJ, Eggo RM, Lenczner M, Brownstein JS, Meyers LA. Epidemiologic wave dynamics attributable to urban community structure: A theoretical characterization of disease transmission in a large network. J. Med. Internet Res. 2015;17(7):e169. doi:10.2196/jmir.4720.

[23] Dehning J, Zierenberg J, Spitzzier FF, Wibral M, Neto JP, Wilczek M, et al. Inferring change points in the spread of COVID-19 reveals the effectiveness of interventions. Science 2020;369(6500):eaab9789. doi:10.1126/science.abab9789.

[24] de Sousa LE, de Oliveira NPH, Filho DADS. Kinetic Monte Carlo model for the COVID-19 epidemic: Impact of mobility restriction on a COVID-19 outbreak. Phys. Rev. E. 2020;102:032133. doi:10.1103/PhysRevE.102.032133.

[25] Triambak S, Mahapatra DP. A random walk Monte Carlo simulation study of COVID-19-like infection spread. Physica A 2021;574:126014. doi:10.1016/j.physa.2021.126014.

[26] Kaguzzu H, Berker A. The effect of weekend curfews on epidemics: A Monte Carlo simulation. Turk. J. Biol. 2021;45:436.

[27] Kolotsova EM, Kurkina ES, Vasketsy AM. Superposition of waves for modeling COVID-19 epidemic in the world and in the countries with the maximum number of infected people in the first half of 2020. arXiv preprint arXiv:2007.02283.

[28] Frasca M, Buscarino A, Rizzo A, Fortuna L, Boccaletti S. Dynamical network model of infective mobile agents. Phys. Rev. E 2006;74:036110. doi:10.1103/PhysRevE.74.036110.

[29] Chu A, Huber G, McGeever A, Veytsman B, Yllanes D. A random walk-based epidemiological model. Sci. Rep. 2021;11(1):19308. doi:10.1038/s41598-021-96211-5.

[30] Kelker D. A random walk epidemic simulation. J. Am. Stat. Assoc. 1973;68(344):821–3. doi:10.2307/2284506.

[31] Lauer SA, Grantz KH, Bi Q, Jones FK, Zheng Q, Meredith HR, Azman AS, Reich NG, Lesser J. The incubation period of coronavirus disease 2019 (COVID-19) from publicly reported confirmed cases: Estimation and application. Ann. Intern. Med. 2020;172(9):577–82. doi:10.7326/M20-0504. PMID: 32150748.

[32] Barman MP, Rahman T, Bora K. Borgohain C. COVID-19 pandemic and its recovery time of patients in India: A pilot study. Diabetes and Metabolic Syndrome: Clinical Research and Reviews 2020;14(5):1205–11. doi:10.1016/j.dsx.2020.07.004.

[33] Chae SY, Lee K, Lee HM, Jung N, Lee QA, Mafwiele BJ, et al. Estimation of infection rate and predictions of disease spreading based on initial individuals infected with COVID-19. Front Phys 2020;8:311. doi:10.3389/fphy.2020.00311.

[34] World Health Organisation https://covid19.who.int/WHO-COVID-19-global-data.csv.

[35] Callaway E, Ledford H. How bad is Omicron? What scientists know so far. Nature 2021;601:197. doi:10.1038/d41586-021-03614-z.

[36] Pulliam JRC, van Schalkwyk C, Govender N, von Gottberg A, Cohen C, Groome MJ, et al. Increased risk of SARS-CoV-2 reinfection associated with emergence of the Omicron variant in South Africa. medRxiv 2021. doi:10.1101/2021.11.21.21266068.

[37] Roelf W. South Africa hit by fourth COVID wave driven by Omicron. https://www.reuters.com/business/healthcare-pharmaceuticals/south-africa-health-ministers-says-country-entering-4th-wave-covid-infections-2021-12-03.

[38] Buscarino A, Fortuna L, Frasca M, Latora V. Disease spreading in populations of moving agents. EPL (Europhysics Letters) 2008;82(3):38002. doi:10.1209/0295-5075/82/38002.