

# Emerging dynamics from high-resolution spatial numerical epidemics

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**Simulating nationwide realistic individual movements with a detailed geographical structure is urgently needed to optimize public health policies. However, existing tools have limited resolution or can only account for a limited number of agents. We introduce Epidemap, a new framework that can capture the daily movement of more than 60 million people in a country at a building-level resolution in a realistic and computational-efficient way. By applying it to the case of an infectious disease spreading in France, we uncover hitherto neglected effects, such as the emergence of two distinct peaks in the daily number of cases or the importance of local density in the timing of arrival of the epidemic. Finally, we show that the importance of super-spreading events strongly varies over time.**

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21 A major limitation to the applicability of mathematical models is that they need to resort to  
22 strong assumptions. In the case of infectious disease epidemiology, for instance, it is easy to  
23 point out that the canonical Susceptible-Infected-Recovered model (1) neglects stochasticity,  
24 misses spatial structure or ignores contact patterns. Although these aspects, and many others,  
25 can be incorporated with the appropriate hypotheses and techniques (2–4), the resulting models  
26 usually remain too abstract to help improve local policies.

27 Agent-Based Simulation (ABS), where individuals are modelled explicitly, represent a seduc-  
28 ing option to achieve a high degree of realism, from a biological and environmental point of  
29 view (5), but their routine use in public health faces three major limitations. First, ABS are  
30 very computationally demanding (6), which restrains the total number of agents that can be  
31 simulated. The second limitation comes from the model dimensionality and the introduction  
32 of numerous parameters, many of which are poorly informed and set in an *ad hoc* manner,  
33 to capture individual heterogeneity. A third limit resides in the way geographic structure is  
34 implemented into the simulation (but see (7)).

35 Here, we introduce Epidemap, a novel numerically efficient agent-based method that addresses  
36 the current limitations of ABS platforms. Epidemap simulates infectious diseases epidemic  
37 scenarios at the scale of a whole country by combining detailed geographical structure, de-  
38 mographic information, and mobility statistics. Practically, this is achieved by using High-  
39 Performance Computing (HPC) techniques, high-resolution spatial data from the OpenStreetMap  
40 (OSM) project (8), and sophisticated mobility models (9). Overall, these simulations only re-  
41 quire 6 parameters (see Supplementary Methods).

42 To illustrate the power and flexibility of the platform, we study the transmission dynamics of  
43 an (uncontrolled) epidemic of respiratory infections in France. The biological features of the  
44 epidemiological model originate from a discrete-time SARS-CoV-2 transmission model param-  
45 eterised with national hospital data (10). We analyse the output of 100 stochastic simulations

46 from epidemic emergence to extinction, *i.e.* approximately one year. A typical simulation gen-  
47 erates daily mobility patterns for 65 million individuals at the scale of buildings and lasts less  
48 than 2 hours on a 12-core AMD Ryzen 9 CPU, with less than 64 GB of RAM.

49 In the simulation (Figure 1), each individual is assigned to a residency building following the  
50 national density estimates at the district level, the French ‘canton’ level (11). Every day, we  
51 assume that individuals visit 0, 1, or 2 buildings other than their own residence (12). These  
52 buildings are chosen at random based on a distance kernel. If more than one individual vis-  
53 its the building the same day, transmission can occur with a probability  $b = 6\%$  (13), which  
54 is weighted by the number of days since the onset of the infection in the donor host. As in  
55 previous studies, this weight is based on the SARS-CoV-2 serial interval (10, 14). The simula-  
56 tion tracks the age of the individuals, which is distributed geographically according to national  
57 demographics data (15). For infected individuals, disease progression follows that described  
58 in (10) and a fraction  $\theta$  of infected individuals develop severe symptoms requiring hospitalisa-  
59 tion, which is simulated by moving them to the nearest hospital building from their residence.  
60 For all recovered individuals, we assume perfect immunity for the duration of the simulation.  
61 All simulations are initialised with 15 infected individuals in Paris to avoid premature random  
62 epidemic extinction.

63 Figure 2 shows the output of the dynamics at the national level for 100 stochastic simulations.  
64 For optimal readability, the dates are aligned based on the day where ICU-occupancy reaches  
65 a value of 700. With our minimal parameterisation, we see that the basic reproduction number,  
66 which is denoted ( $R_0$ ) and corresponds to the average number of secondary infections caused  
67 by an infected individual (16), is in the order of 3, which is consistent with estimates for the  
68 French epidemic (10, 17). Note that this value is here computed directly at the individual level,  
69 by counting how many infections an individual causes. Furthermore, the daily estimates for  
70 the temporal reproduction number calculated in the same manner are very similar to those esti-

71 mated using the daily case incidence data (dashed blue curve) and the method from (18). These  
72 uncontrolled epidemics last 308 days (95% confidence interval (CI): [286;345] days) and the  
73 final total epidemic size is  $q = 61.1\%$  (95%CI [60.1%;61.9%]) of the initial susceptible popu-  
74 lation. As described by earlier studies (19), this proportion is lower than the prediction from a  
75 mean-field model that is given by the well-known equation  $qR_0 + \log(1 - q) = 0$  (1), which  
76 yields 93% for  $R_0 = 3$ . This shows that geographical structure greatly impacts the unfolding of  
77 the epidemic.

78 The national prevalence data uncovers a bimodal structure of the epidemic, which can be under-  
79 stood by moving to the regional scale (Figure 2b). The first peak corresponds to the spread in the  
80 region where the outbreak emerges (here the Ile-de-France), whereas the second corresponds  
81 to the sum of the epidemic peaks in the other regions. This bimodal structure is particularly  
82 pronounced given the population density in the region of origin but it is a direct consequence of  
83 the detailed geographical information in the simulation platform.

84 The resolution of the Epidemap simulations allows us to perform analyses at the district level  
85 (see Supplementary Videos 1 and 2). In Figure 3a, we show that the date of onset of the epi-  
86 demic in an area strongly depends on its distance from the origin (here assumed to be Paris).  
87 Furthermore, there is an additional effect of density such that denser areas are infected first.  
88 Interestingly, at the departmental level, these trends are not significant, further showing the im-  
89 portance of a fine-grain simulation level. Furthermore, the total proportion of inhabitants of a  
90 district who have been infected at the end of the epidemic strongly increases with density. The  
91 pandemic propagation velocity also increases with time. This can be explained by the fact that,  
92 when incidence is high, long-distance dispersion events are not rare anymore, which biases the  
93 mean distance of contamination towards higher values. For densely-populated districts (Figure  
94 3b), this proportion converges towards the mean-field prediction from well-mixed models men-  
95 tioned above (1). However, if the population density is low, this proportion is more variable

96 showing the limit of classical assumptions.

97 The individual-based nature of our simulations allows us to follow transmission chains (Supple-  
98 mentary Video 3), which has direct applications. For instance, we can count, at the end of each  
99 infection, how many secondary infections were caused. The distribution of these individual re-  
100 production numbers is particularly important in the context of emerging epidemics because the  
101 more disperse, the more the spread relies on super-spreading events (20). Early in the epidemic,  
102 the distribution is tightly centred around the  $R_0$  value (Figure 4a). As the epidemic unfolds, the  
103 distribution changes with a mode that decreases towards 0, and a wider dispersal. This pattern  
104 can be formalised by assuming that the distribution of individual reproduction numbers follows  
105 a negative binomial distribution (20). In Figure 4b, we show that the mean of this distribution  
106 (in blue) follows the pattern estimated in Figure 2a. The dispersal parameter ( $k$ ) indicates that  
107 super-spreading events reach a peak at the end of the first national epidemic wave. During the  
108 end of the epidemic, stochasticity is strong but there is a general trend towards a decrease in  
109 super-spreading events. Note that in these simulations we do not introduce host heterogeneity,  
110 which means we only capture the dimension of superspreading that originates from spatial het-  
111 erogeneity. This is already sufficient to show that studies attempting to quantify the importance  
112 of superspreading events should account for the stage of the epidemic they analyse.

113 Simulating the daily activity of millions of individuals at a build-level resolution with a realistic  
114 mobility model significantly improves our understanding of how epidemics unfold. Early stages  
115 appear to be consistent with stochastic and deterministic mean-field models. However, once  
116 local saturation effects cannot be neglected anymore, Epidemap reveals striking patterns with  
117 an unprecedented resolution. First, we show that at a national level, a two-waves epidemic  
118 pattern is to be expected, largely because of a wide regional structure. Second, we find that  
119 districts are affected by the epidemic depending on how far they are from the epicentre, but also  
120 depending on their density. The latter effect is absent at the departmental level, which shows the

121 added value of a detailed geographical structure. Furthermore, we show that, as expected, the  
122 simplistic estimate of the final epidemic size as a function of  $R_0$  does not apply at the national  
123 level. However, we also show that it does yield relevant results at the district level if the density  
124 is sufficiently high. Finally, individual follow-ups allow us to show how super-spreading events  
125 becoming increasingly important as the epidemic unfolds, but that their role decrease as the role  
126 of stochastic processes increases.

127 Being able to perform these detailed simulations for so many agents at a national scale has to be  
128 traded-off against some simplifying assumptions. The major one is that individual movements  
129 are based on a distance kernel centred in their residency home. As shown by earlier work on  
130 influenza dynamics, although this assumption is relevant for France, it might be too simplistic  
131 for countries like the United States, where air traffic represents a greater proportion of the  
132 travels (21). A second limitation is that, because of the lack of relevant data, we assumed all  
133 individuals to have the same average behaviour, *e.g.* in terms of the number of buildings visited  
134 per day. However, it can readily be made a function of the age of infection status (or of the  
135 individual) in the simulations.

136 These results have immediate applications for public health. For instance, they allow authorities  
137 to derive a risk-factor per district to help control an epidemic and prevent outbreaks. This can  
138 also be instrumental to optimise vaccine coverage, or other control strategies (mask-wearing,  
139 lockdown, vaccination). Many insights can also be gained from the ability to follow individual  
140 trajectories and transmission chains (Supplementary Video 3), and superspreading events. Fur-  
141 ther developments in Epidemap will incorporate building meta-data, for instance, to simulate  
142 children attending the closest school from the residency home or study the local saturation of  
143 hospital emergency care services. Finally, we focused here on SARS-CoV-2 but an asset of  
144 this platform is its versatility. It can readily be deployed to study other infectious agents to  
145 investigate a variety of control scenarios with high resolution at a national level.

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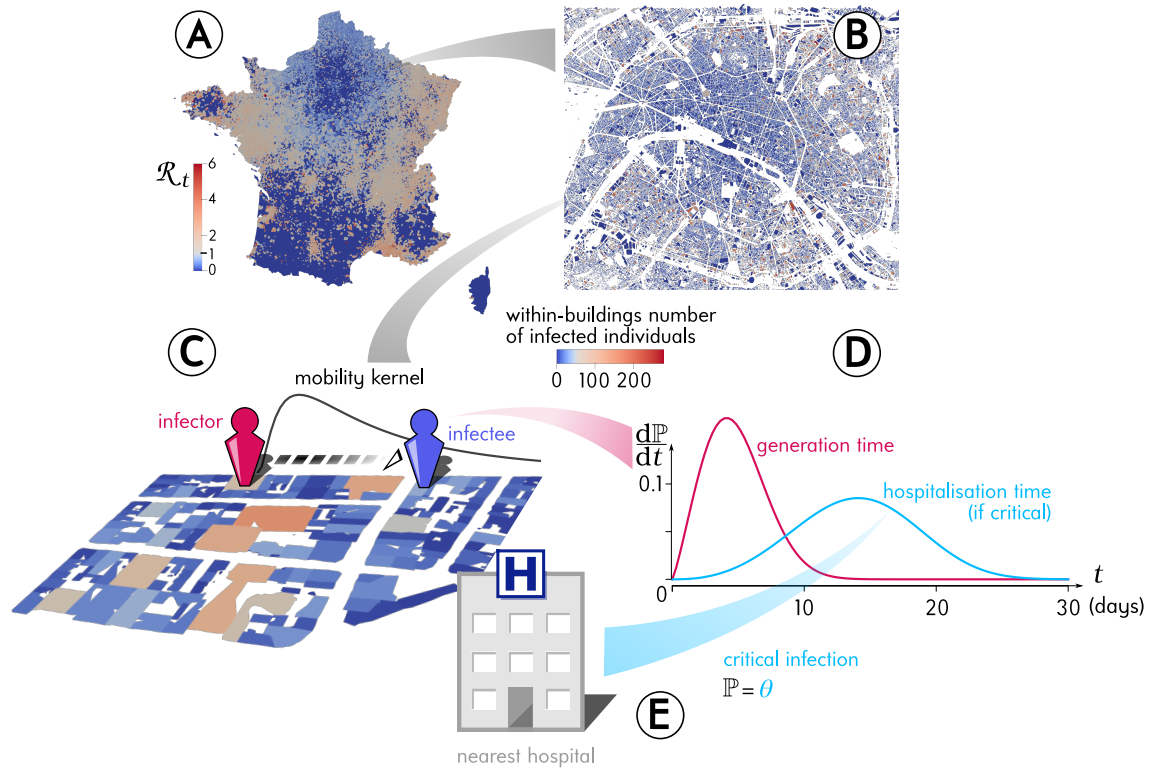


Figure 1: **Outline of the Epidemap simulation framework.** **A)** The 65 millions inhabitants of metropolitan France are explicitly mapped to housing buildings following cartographic and demographic data. **B)** At each time point of the simulation, the number of infected individuals in each building of the country is recorded, as well as the time past since each got infected (the panel shows the Paris area). **C)** Every day, individuals can randomly move from their home to other buildings according to a mobility kernel and meet other people. If an infected individual encounters a susceptible host, a transmission event can occur. **D)** The contagiousness of a infected individuals varies depending on the time since infection. **E)** A small fraction  $\theta$  of infected individuals develop a critical form of the disease that requires their hospitalisation to the nearest facility.

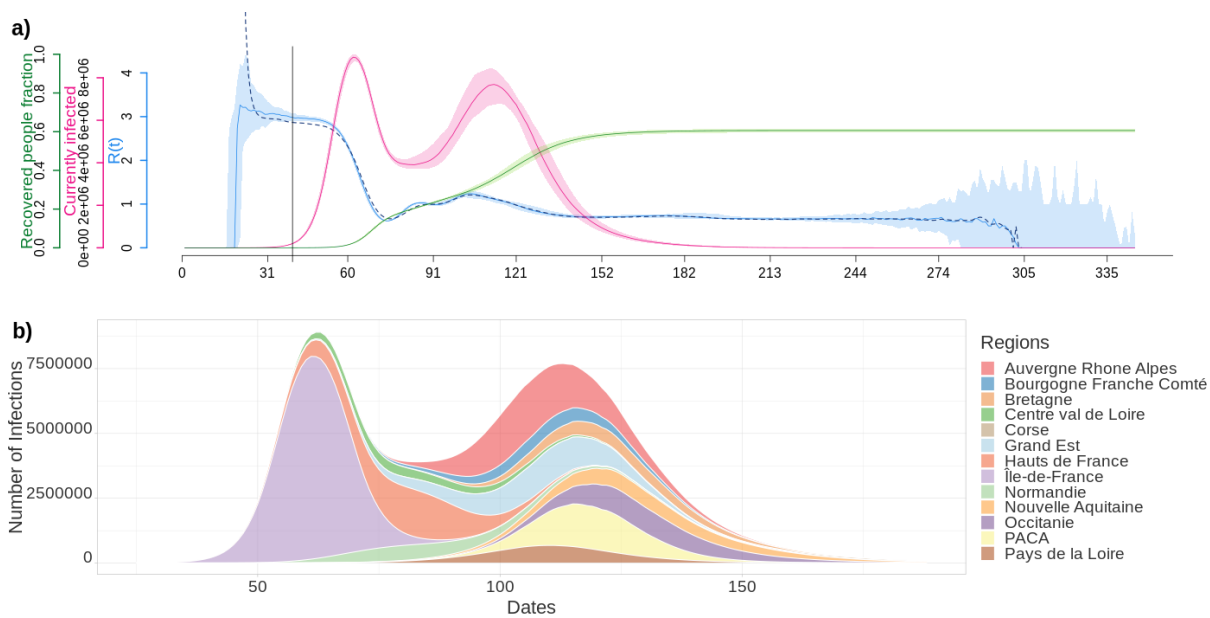


Figure 2: **Epidemiological dynamics at the national (a) and regional (b) level.** a) The daily prevalence (number of infected individuals) is in red, the temporal reproduction number ( $R_t$ ) in blue, and the cumulative number of recovered individuals in green. Shaded areas show the 95% sample quantiles of 100 stochastic simulations. The dashed line shows the median  $R_t$  calculated on the case incidence data. b) Each color shows the prevalence in a French region.

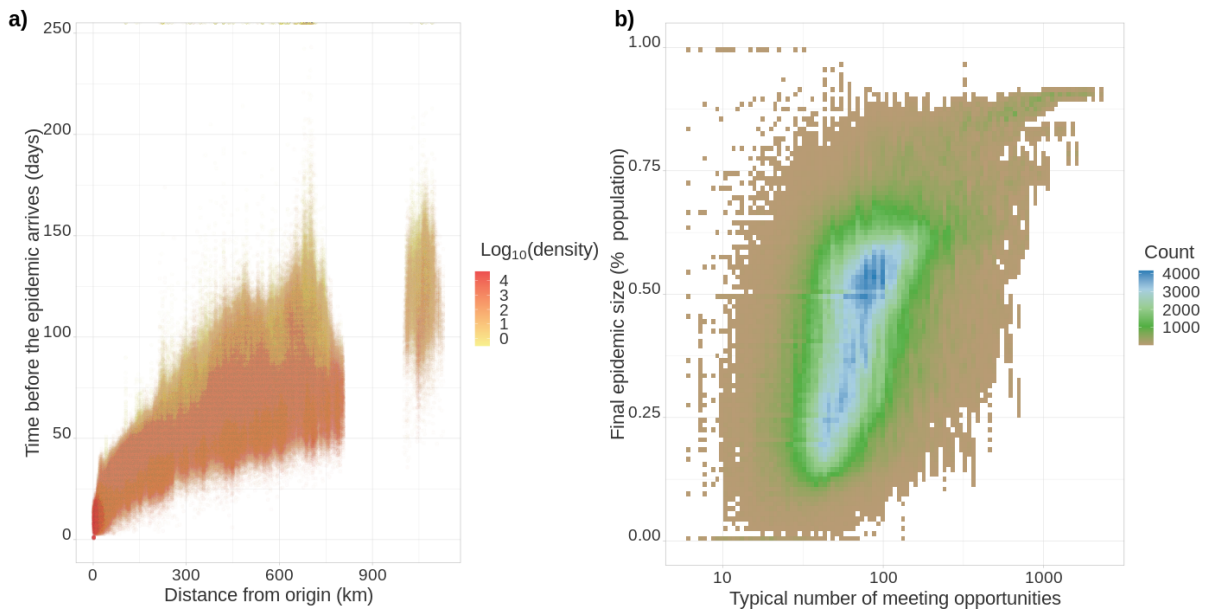


Figure 3: **Epidemic arrival date and size at the district level.** a) Effect of the distance from the epicentre on the number of days until the epidemic begins in a district. The color indicates the population density in the district (number of inhabitants divided by the district's surface). b) District final epidemic size as a function of the characteristic distance between two individuals normalised by the average dispersal distance. The later is computed as  $2E[X]\sqrt{\text{density}}$ , where  $X$  is the log-normal distribution of daily individual covered distance. Both panels show the value for 35,234 French districts and 100 stochastic simulations.

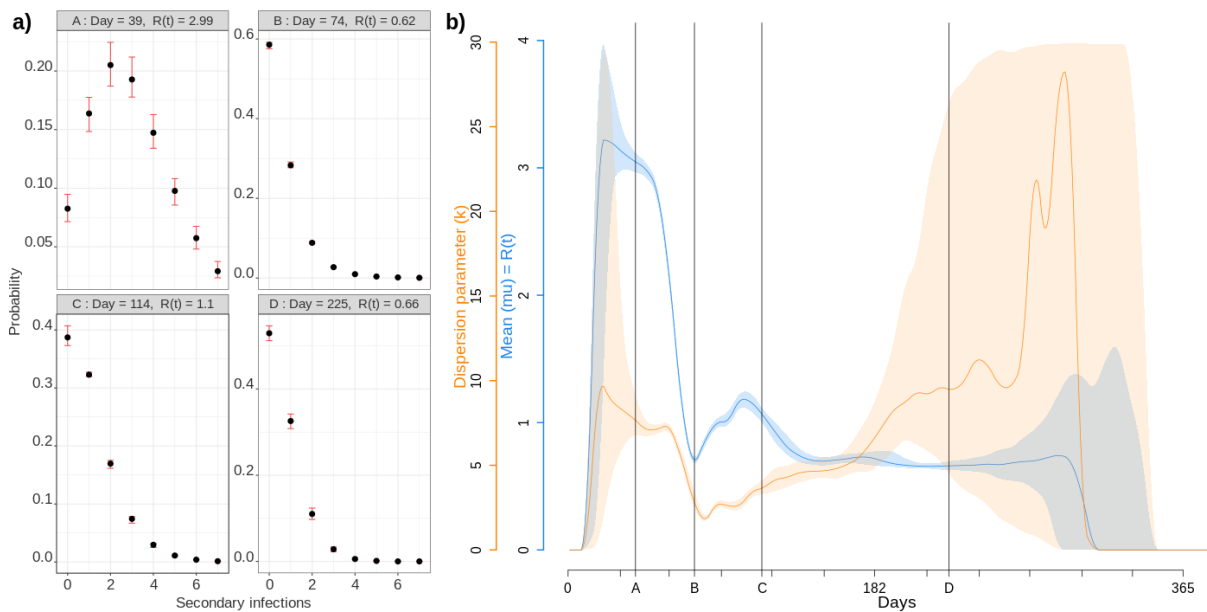


Figure 4: **Individual reproduction number dynamics.** a) Distribution obtained over the whole population on 4 different days post outbreak. b) Daily variation of the mean (in blue) and dispersion (in orange) of the distribution of individual reproduction numbers, which is assumed to follow a Negative Binomial distribution (20). Shaded areas show the 95% CI.