Multi-agent systems in epidemiology: A first step for computational biology in the study of vector-borne disease transmission

Supplementary Materials

1 Biological analyses

1.1 Dynamical properties of the spatial model: Importance of space

To understand the net impact of the spatial features in a vector-borne disease model, we have analyzed how variations in the moving capacity of vector and reservoir individuals could impact on the intensity of disease persistence. We chose our ecological and epidemiological parameters in order to generate an epidemic as indicated in the legends to figure.

Simulations show that the reservoir moving capacity does not seem to influence the intensity of disease transmission (Figure S1), which is represented here by the maximal number of infectious vectors. Indeed, reservoir individuals are not the organisms that “make contact” between vector and reservoir individuals. If reservoir individuals are within the range of a vector’s moving capacity, the moving capacity of the reservoirs will not change the frequency of contact and hence the probability of infection.

In contrast, vectors’ moving capacity clearly plays a greater role. An increase of vector moving capacity boosts the number of available reservoir individuals that can be reached and bitten. However, a moving capacity’s threshold can be observed (Figure S1). The world used in this example had a 100 pixel size on each side. Although this threshold value is not enough for vectors to reach all reservoir individuals, it seems sufficient, with the support of well-mixed vector individuals, to reach a maximal value in the ratio between vector abundance and reservoir abundance and, thus, to create an epidemic. The existence of a moving capacity’s threshold suggests that vectors’ moving capacity should be reduced below a given threshold to effectively decrease the intensity of disease transmission.

These first results indicate that vectors’ spatio-temporal dynamics may impact more largely on disease transmission than those of reservoir species. However, the majority of the modeling studies on vector-borne disease, like for instance malaria, try to cope with complex modeling of the reservoir host displacements. Our study suggests that it could be better, in the case of vector-borne diseases, to focus on the spatio-temporal dynamics of vector species rather than on the moving characteristics of the reservoir host individuals.
1.2 Impact of spatial configuration

After having identified moving capacity’s core mechanisms, we could analyze how the spatial distribution of host species influences disease dynamics. Indeed, in natural systems, species are not well mixed as previously assumed, and only some (host) species have contacts with some other (host) species [1]. Here, we looked at which could be the consequences of different spatial configurations in terms of spatial distribution of species in the landscape. Hence, we used a fractal landscape to characterize real landscapes. In landscape ecology, fractal landscape is assumed to be a ‘neutral’ model [2]. This model does not have a pure formalization of mechanisms. Thus, we decided to use the classical Middle Displacement Point algorithm (MDP [3]) to generate our fractal landscape in order to make it slightly more complex. This fractal landscape yields continuous values and we have to apply an identification cluster every time we want to define a discrete landscape within a given number of classes which will be also analyzed. This operation is classically done in epidemiological studies involving GIS [4]. Finally, to dispatch vector and reservoir species within the fractal landscape, we assumed that vectors are set in odd-numbered habitats and reservoirs in even-numbered habitats. The surprising result is that neither fractal dimension nor the number of classes seem to have a significant impact on disease dynamics (Figure S2). In fact, despite different spatial configurations, the total number of vector and reservoir individuals is kept constant, and the surface area for each species is similar. In this study, we assume that each individual could reach any locations within its habitat and its tolerance to go out from its habitat. Hence, the increase of landscape structure complexity decreases area of contacts between individuals located within different habitats, which could, intuitively, decrease disease transmission.
transmission.

Figure 2: Impact of fractal dimensions and number of possible habitats on disease dynamics. Time series represent the number of infectious vectors across time, and the small inset within each plot illustrates the landscape generated. These two parameters do not impact strongly on disease dynamics: increasing the fractal dimension of landscape and the number of possible habitats will not change the habitat surface area for each vector and reservoir species or the abundance of each species. This phenomenon leads to a constant disease transmission. Parameters used here are identical to those used previously.

1.3 Importance of adding (reservoir) species

The last part of this theoretical study concerns the impact of the addition of a new host reservoir species in space. An interesting effect due to species addition, i.e., the “dilution effect”, was found in empirical [5], experimental [6] and theoretical context [7]. The “dilution effect” theory stipulates that disease prevalence in the vector population decreases with the increase of the abundance of low-competent reservoir species. Hence, it might be important to understand how the dilution effect is linked to the spatial dimension.

To this aim, we considered a situation which always leads to disease persistence. We then analyzed how the introduction of one low-competent reservoir species (competence=1%) can buffer disease transmission and lead to lower disease transmission. The size of the newly introduced reservoir species plays a determinant role in disease dilution (Figure S3). Conversely, as it was expected from previous results, an increase of its moving capacity does not. Once again, since vectors “make the contact” in disease transmission, the higher moving capacity of the new reservoir species does not increase the probability for vector individuals to find a poorly competent reservoir individual. On the other hand, a larger population of poorly competent reservoir individuals increases this probability, and hence the “dilution effect” phenomenon.
Figure 3: Impact of the introduction of a new reservoir species on the intensity of disease transmission. The population size of the new reservoir species strongly influences disease transmission. This is a direct application of the “dilution effect” as explained previously. On the contrary, the moving capacity of the new species does not influence disease transmission because this parameter does not increase the number of available vectors for each reservoir individual. Parameters used here are identical to those used previously. Competence of the introduced species is 1% and ecological parameters for this species are identical to those of the other reservoir species.

2 Algorithms of function

Host with landscape: Moving function

IF BitingRate==1 OR 1/BitingRate > RAND()
  INTEGER MovingArea=((MovingCapacity+1)*2)^2
  INTEGER IndexNewPosition=CEIL(RAND()*MovingArea)
  NewX=(x-MovingArea+ROUND(IndexNewPosition/MovingArea))
  NewY=(y-MovingArea+ROUND(IndexNewPlace%MovingArea))
  IF NewX AND NewY are not in species Habitat
    NewX and NewY receive closest values in habitat from NewX and NewY
  END IF
  nextX=NewX
  nextY=NewY
END IF
ELSE
  ReservoirList=ModelSwarm.LookingForReservoir(x,y,MovingCapacity,IndexHabitat,HabitatTolerance)
  IF ReservoirList is not empty
    ReservoirChosen=CEIL(RAND()*ReservoirList.size())
    NewX=ReservoirChosen.x
    NewY=ReservoirChosen.y
    Infection(ReservoirChosen.parasite)
    ReservoirChosen.Infection(Parasite)
  END IF
END ELSE
FutureX=NewX
FutureY=NewY
Host with parasite: Infection function
Parameters: PARASITE pParasite
IF pParasite!=NULL
   IF CurrentParasite==NULL and NextParasite==NULL and PastParasite==NULL
      IF RAND()<Competence
         NextParasite=pParasite
      END IF
   END IF
END IF

Main algorithm
FOR Time=1 TO Time=TimeMax
   FOR EACH Host Objects
      HostObjet.STEP()
   END FOR EACH
   FOR EACH Host Objects
      HostObjet.UPDATE()
   END FOR EACH
END FOR

Function STEP
Move()

Function UPDATE
IF currentParasite==NULL ET nextParasite!=NULL
   IF 1/latencyPeriod <RAND()
      currentParasite=nextParasite
      nextParasite=NULL
   END IF
END IF
IF currentParasite!=NULL ET previousParasite==NULL
   IF 1/infectiousPeriod <RAND()
      previousParasite=currentParasite
      currentParasite=NULL
   END IF
END IF
x=nextX
y=nextY
BIRTH()
DEATH()
3 Files input

To fill in all characteristics of host species, parasite species and landscape structure, different input files were needed. The first one was the "Community" file which contains the species characteristics: "Index Species", "Offspring Size", "Lifespan", "Biting Rate" and "Competence" (in order of appearance). Each line represents a new species. The second file was the "Habitat" file which describes the spatial distribution of habitats. Its structure is similar to that of a PPM image file (Portable PixMap file format). The header of the file contains the resolution, in x-axis and y-axis, in the first line, and the number of habitats of the file in the second line. After this header, each line represents a line (in x way) of the picture and pixels on each line are split by tabulation. Finally, the last file was the "Correspondence" file which makes the connection between the "Habitat" and "Community" files. Each line integrates the "Index Species", "Index Habitat", "Moving Capacity" and "Habitat Tolerance" values. Hence, we could take into account the fact that some species have different moving capacities and tolerance values regarding their habitat. This allowed us to characterize the spatial distributions of all species.
4 Communication protocol for distributed behavior

| Order                    | Description                                                                 |
|--------------------------|-----------------------------------------------------------------------------|
| GET PLACE HAB            | Get place of host’s index in the habitat                                   |
| GET LEVEL HAB            | Get habitat value of input pixel                                           |
| ADD PEOPLE               | Add a new people to the output file for abundance                          |
| ADD INFECTIOUS           | Add a new infectious to the output file                                     |
| ADD HOST                 | Add a new host into the system                                             |
| GET PIXEL HABITAT        | Get habitat value for a given pixel                                        |
| GET NB                   | get number of host for a given pixel                                       |
| GET BITE                 | Get ”BitingRate“ value for a given individual                              |
| GET HOST                 | Get ”Host“ object for a given individual                                    |
| HOST GENERATION          | Create a new ”Host“ object with input parameters                           |
| HOST REMOVE              | Remove ”Host“ object found in input                                        |
| GET PARASITE             | Get ”Parasite“ object for a given individual, NULL if any                  |
| END STEP                 | Indicating to the ”ModelSwarmServer“ Object that ”ModelSwarmClient“ object has finished ”Step“ function |
| END UPDATE               | Indicating to the ”ModelSwarmServer“ Object that ”ModelSwarmClient“ object has finished ”Update“ function |
References

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