OVERVIEW, DESIGN CONCEPTS AND DETAILS (ODD) – Unstructured Habitat

The model description follows the Overview, Design concepts and Details protocol for describing individual and agent-based and other computation models (Grimm et al., 2006, 2010; Railsback and Grimm, 2019).

1. Purpose

The purpose of our model is to analyse if the cost experienced by transconjugant cells may be evolutionarily advantageous to donor cells and that plasmid transfer constitutes spiteful behaviour in an **unstructured habitat**. The model aims to identify the role of the frequency of plasmid donor bacteria relative to recipient bacteria, the conjugation rate, and the costs associated with the plasmid, in the ability of the donor bacteria to exert a harmful behaviour.

2. Entities, state variables and scales

The entities included in this model are bacteria.

Bacteria are characterized by: type ([1-6]), plasmid fitness cost ([0, 0.05, 0.2, 0.2, 0.4, 0.6]), and adaptation period [(70, 400)].

The parameter type defines whether the bacterium is recipient (1), donor (2), transconjugant (3), adapted transconjugant that lost the plasmid (4), non-adapted transconjugant that lost the plasmid (5), or donor that lost the plasmid (6).

The plasmid fitness cost is the cost associated with the presence of the plasmid. Transconjugant cells pay a fitness cost for some generations, after which the cost ameliorates to the same value as that of the original donor cells.

The adaptation period is the number of generations during which the conjugants pay the fitness cost.

In this model, a time step corresponds to a possibility for a bacterium to duplicate and/or conjugate (in case it has a plasmid).

3. Process overview and scheduling

In each simulation cycle of the unstructured habitat, **a bacterium** is randomly chosen to verify whether the "bacterial_growth" submodel is performed. If the bacterium has a plasmid, it is verified if the submodel "conjugation" is performed.

If the "bacterial_growth" submodel is activated, is added a new bacterium, with the corresponding characteristics of the originating bacterium. If the "conjugation" submodel is activated, the characteristics of the bacterium receiving the plasmid are updated accordingly. At the end of each time step, it is checked whether the number of bacteria present is at least 950,000 bacteria. If this condition is met, the number of each type of bacteria present is stored in a variable, and then bacteria are randomly eliminated so that only 50% of bacteria remains.

This process is repeated 1073 times. After that, the data with the number of each type of bacteria at each time is written to a .csv-file in the submodel "files_writing".

4. Design concepts

We took the following design concepts into account:

 <u>Duplication</u>. One bacterium gives rise to a new bacterium, while the other characteristics remain the same.

- <u>Segregation</u>. In the "bacterial_growth" submodel, the new bacterium originated by a plasmid-containing bacterium loses the plasmid.
- <u>Conjugation</u>. A bacterium with a plasmid transfers the plasmid to a bacterium that does not have one.
- <u>Cell death</u>. Bacteria are eliminated randomly.

5. Initialisation

The "bacteria_distribution" submodel **creates the initial bacterial lists** according to the number of donor cells [10, 99, 5000, 9901] and the number of recipient cells [9990, 9901, 5000, 99]. In this way, the initial cell number is always 10 000 cells.

In the submodel "bacteria_distribution", each of the bacteria is assigned inherent characteristics, namely: **type**, **plasmid fitness cost**, **and adaptation period**.

6. Input data

There is no external input of data.

7. Submodels

Table 1 – Model parameters.			
Entities	Parameter range	Description	
maximum_number_bacteria	950 000 [constant]	Maximum number of bacteria	
remaining_proportion_bacteria	0.5 [constant]	Proportion of bacteria remaining when bacteria are randomly removed	

number_plasmid_free_bacteria	9990, 9901, 5000,	Initial number of bacteria not
	99	carrying plasmid
donor_bacteria	10, 99, 5000,	Initial number of bacteria that
	9901	carry plasmid
maximum_growth_rate (ψ^{max})	1 [constant]	Maximum bacterial growth rate
maximum_conjugation_rate	1 [constant]	Maximum bacterial conjugation
(γ^{max})		rate
theta (θ)	0.6, 0.8, 1	Value of theta (bacterial growth)
theta_1 (θ_1)	0.2 [constant]	Value of theta 1 (conjugation)
theta_2 (θ_2)	0.3 [constant]	Value of theta 2 (conjugation)
initial_plasmid_cost	0.2, 0.4, 0.6	Cost that bacterium have when
		receiving the plasmid
permanent_plasmid_cost	0, 0.05, 0.1	Cost associated with the presence
		of the plasmid in donors and
		adapted transconjugants
adaptation_time	70, 400	Number of duplications that
		bacteria need until the initial
		plasmid cost changes to
		permanent plasmid cost
segregation_probability	0.001	Probability of a bacterium losing
		the plasmid at the moment of its
		duplication

Bacterial_growth

Bacterial growth depends on the growth rate (ψ). Therefore, for each selected bacterium, we obtain the growth probability according to the following function:

$$\psi(\mathsf{C}) = \begin{cases} \psi^{max}, & \text{if } \mathsf{C} \geq \theta \\ \psi^{max} \frac{\mathsf{C}}{\theta}, & \text{if } \mathsf{0} \leq \mathsf{C} < \theta \end{cases}$$

Note that $\psi^{max} = 1$ for plasmid-free cells and that $\psi^{max} = 1 - cost$ for plasmid donor cells, with "cost" being the plasmid cost (note that, in our model, the fitness cost differs between bacteria and can even evolve). The C value is the proportion of existing bacteria over the maximum capacity (1 000 000).

If a random number is equal to or less than the growth rate, we add a new bacterium.

If the original bacterium has plasmid, and if a random number is less than the segregation probability, the resulting bacterium will have no plasmid, retaining the remaining characteristics of the original bacterium. Otherwise, the new bacterium will have all the characteristics of the original bacterium. If it is an unadapted bacterium, the adaptation time of the original bacterium and the new bacterium will decrease by one. If the adaptation time is zero, the plasmid cost of both bacteria is updated to the permanent cost.

Conjugation

Conjugation depends on the encounter probability and the conjugation rate (γ). Therefore, for each plasmid carrying bacterium selected, we check the encounter probability given by: PBC/N*PFC/N, where PBC is plasmid-bearing cell, PFC is plasmid-free cell, and N is the total number of cells in the system. If a random number is less or equal to the encounter probability, we obtain the conjugation rate according to the following function:

$$\gamma(C) = \begin{cases} \gamma^{max}, & \text{if } C \ge \theta_2 \\ \gamma^{max} \frac{C - \theta_1}{\theta_2 - \theta_1} & , \text{if } \theta_1 \le C < \theta_2 \\ 0, & \text{if } C < \theta_1 \end{cases}$$

If a random number is equal to or less than the conjugation rate, the characteristics bacterial type, plasmid cost and adaptation time are updated in the plasmid-free bacterium. Note that, a segregant bacterium may have become segregant due to: (i) a donor bacterium that lost the plasmid; (ii) an adapted transconjugant bacterium that lost the plasmid; or (iii) an unadapted transconjugant bacterium that lost the plasmid.

Thus, if the plasmid-free bacterium is segregant, when it receives the plasmid, it will have the same characteristics as the originating bacterium. Once again, the C value is the proportion of existing bacteria over the maximum capacity (1 000 000).