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Impact of quantitative plant resistance on within-host viral demo-genetic dynamics

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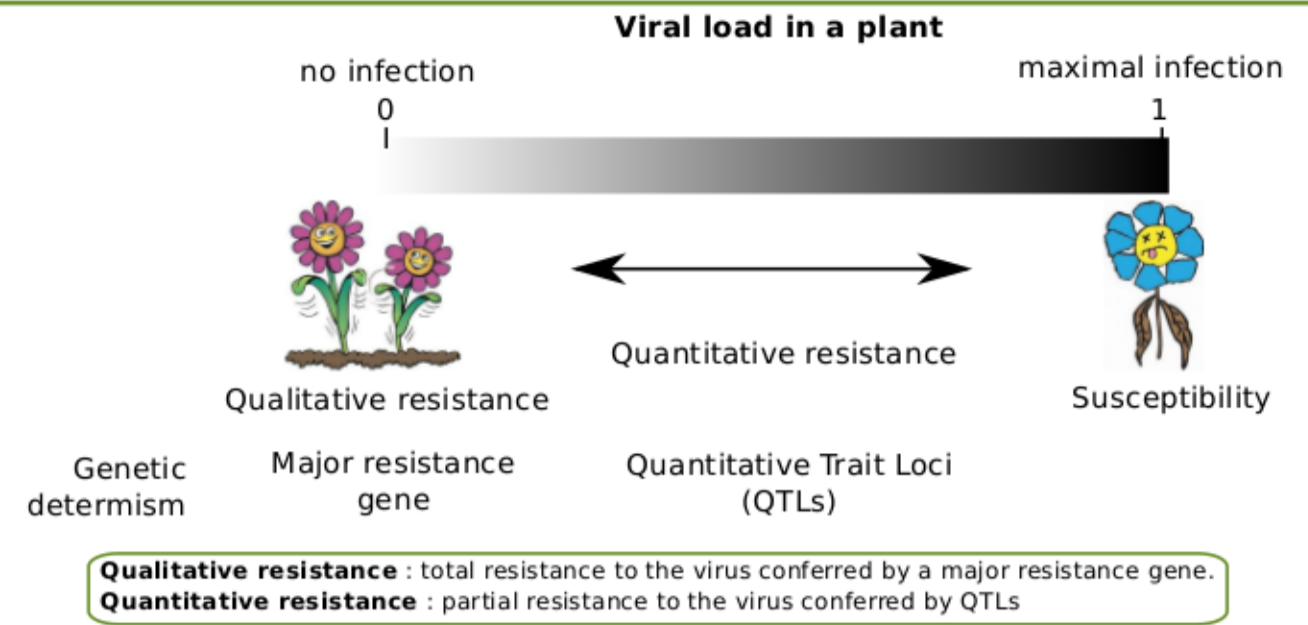
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Introduction

The deployment of virus-resistant plants often leads to the emergence of **resistance-breaking** (RB) mutants that suppress the yield benefit provided by the resistance.

Although breakdowns are well known for **qualitative resistances**, they are still poorly understood for **quantitative resistances**.

Furthermore, it has been proved for several pathosystems that **combining qualitative and quantitative resistances can increase the sustainability of the qualitative resistance** (Palloix *et al.* 2009, New Phytol, Brun *et al.* 2010, New Phytol).

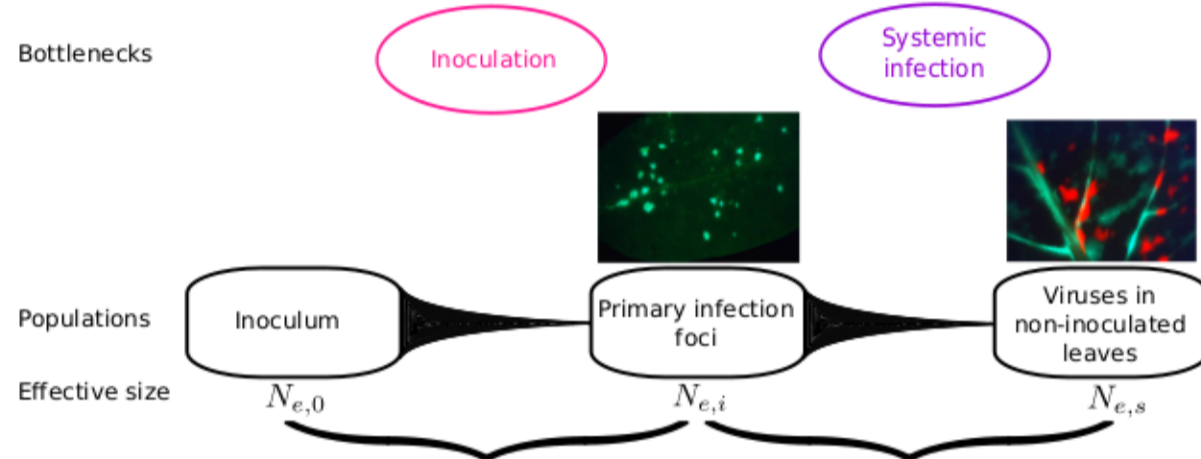


Objective: To analyze the **effect of quantitative resistances**, in terms of genetic drift and selection, on the **sustainability of qualitative resistances** by coupling experimental and modelling approaches.

Quantitative resistances are associated to a **decrease of the fixation rate of RB mutants**. Two possible explanations:

- **decrease of the selection differential**
- **increase of the genetic drift** (not studied in the context plant resistances to pathogens)

Experimentation: strength of genetic drift on viruses during plant infection



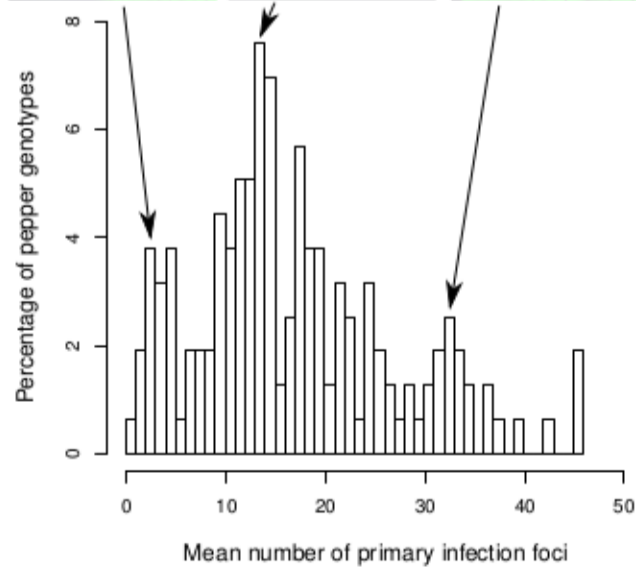
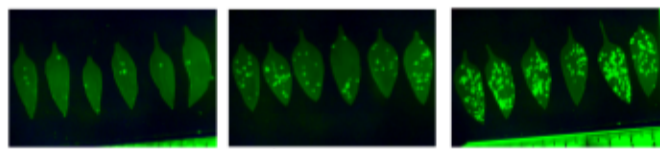
Exp 1: genetic drift due to inoculation

One virus particle is responsible for each primary infection focus. Hence, effective population size N_e following inoculation is well estimated by the number of primary infection foci. (Zwart *et al.* 2011, PLoS Pathog)

Strategy:

Inoculation of 158 pepper genotypes carrying a **major resistance gene** ($pvr2^3$) and several combinations of quantitative trait loci (QTLs), with Potato virus Y (PVY) strains carrying the Green Fluorescent Protein (GFP) fluorescent marker.

Counts of the primary infection foci.



We observed a **large variability** in the **number of primary infection foci** depending on the pepper genotype.

Objective:

Detection of **QTLs** associated to **effective population size** variation after **inoculation**.

Exp 2: genetic drift due to systemic infection

Strategy:

Inoculation of pepper genotypes exhibiting **contrasted effective population size** following **inoculation** (cf. Exp 1) with a **1:1 mix of PVY-GFP and PVY-mCherry**, carrying a green and a red fluorophore, respectively.

Observations:

- t_1 : relative **frequency of primary infection foci** initiated by **PVY-GFP and PVY-mCherry** within **inoculated leaves**.

- t_2 : relative **frequency of PVY-GFP and PVY-mCherry** within **systemically infected leaves** (area of PVY-GFP and PVY-mCherry).

We will assess the **strength of genetic drift** due to **systemic infection** by comparing t_1 and t_2 frequencies.

Objective:

Detection of **QTLs** associated to **genetic drift** variation due to **systemic infection**.

What is the most appropriate estimator of the genetic drift due to systemic infection?

Is the estimation more accurate if we take more time points?

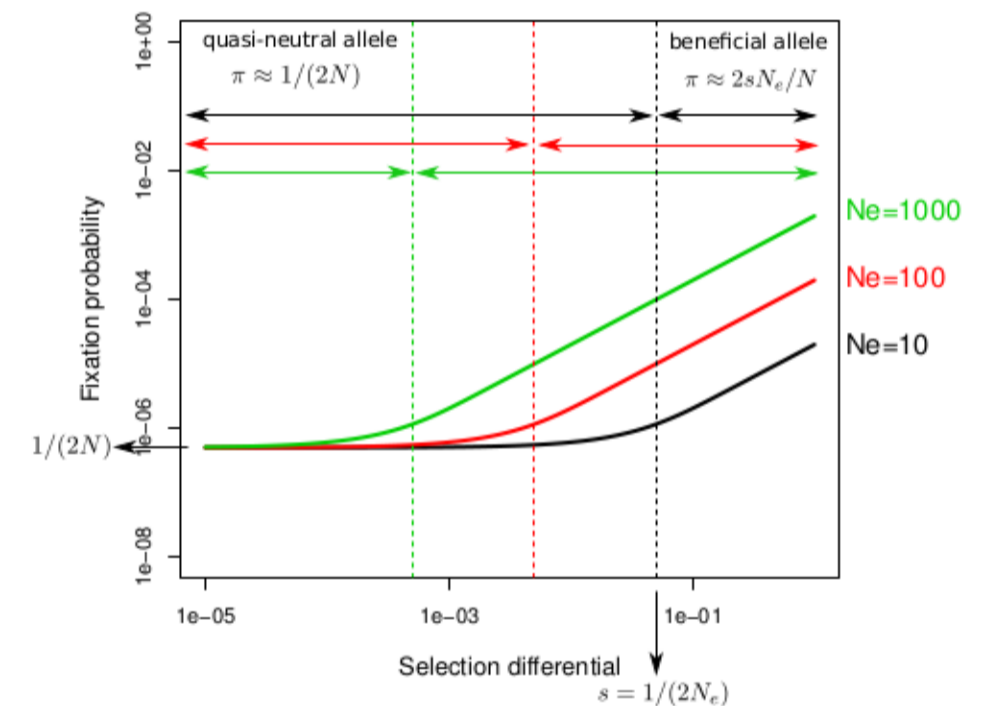
How can we estimate the genetic drift due to systemic infection if selection cannot be ignored?

Modelling: impact of genetic drift and selection on the durability of qualitative resistances

Ohta & Kimura (1970, Genetics) proposed a formula to approximate the **fixation probability** for a **beneficial mutation**, under the hypotheses:

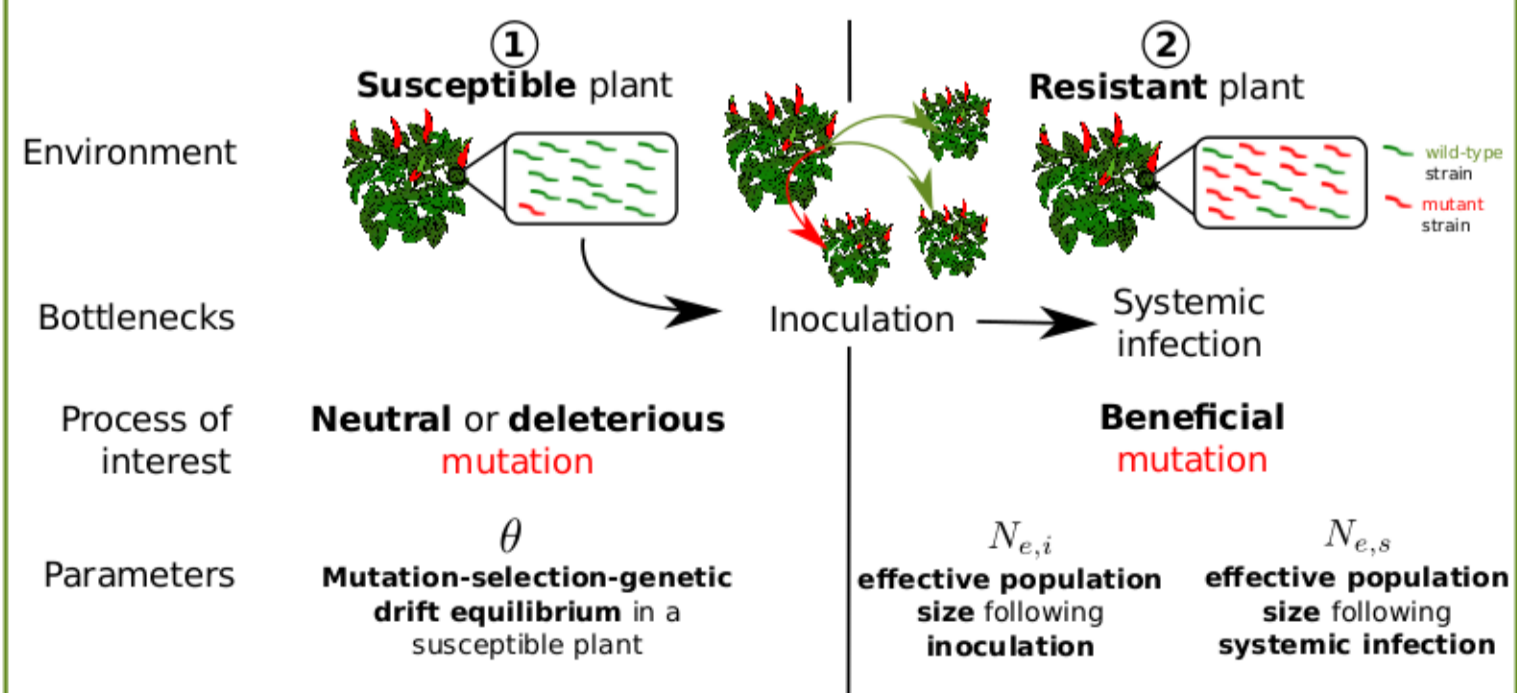
- of diploid individuals, with sexual reproduction, panmixia, and a large and constant population size,
- of discrete generations, and one mutant at a time (moderate mutation rate).

$$\pi = \frac{1 - \exp(-4spN_e)}{1 - \exp(-4sN_e)}$$



Strategy:

To study the emergence of RB mutants when resistant plants are deployed.



Objectives:

To determine the **fixation probability** π of **beneficial mutations** for **viruses** (haploid, varying size, high mutation rate (clonal interference), etc.).

$$\pi = f(\theta, N_{e,i}, N_{e,s}, \Delta s)$$

with Δs : selection differential between **wild-type** and **mutant** strains

Prospects

1 Experimentation and modelling: to follow the demo-genetic dynamics of PVY variants in different pepper genotypes by high-throughput sequencing, and to disentangle **genetic drift** and **selection** effects by fitting models to these data (Fabre *et al.* 2012, PLoS Pathog).

2 Experimental evolution: to record the **appearance** of virus **beneficial mutations** (against qualitative resistances) and their increase in frequency with time.

3 Modelling: to design and study an **epidemiological model** at the **landscape scale**, accounting for the **demo-genetic dynamics** of viruses at the **within-host scale**.

