

Modelling plant resistance deployment: the R package *landsepi*

The R package *landsepi* provides a general modelling framework to help compare plant resistance deployment strategies and understand the impact of epidemiological, evolutionary and genetic factors for a wide range of pathosystems.

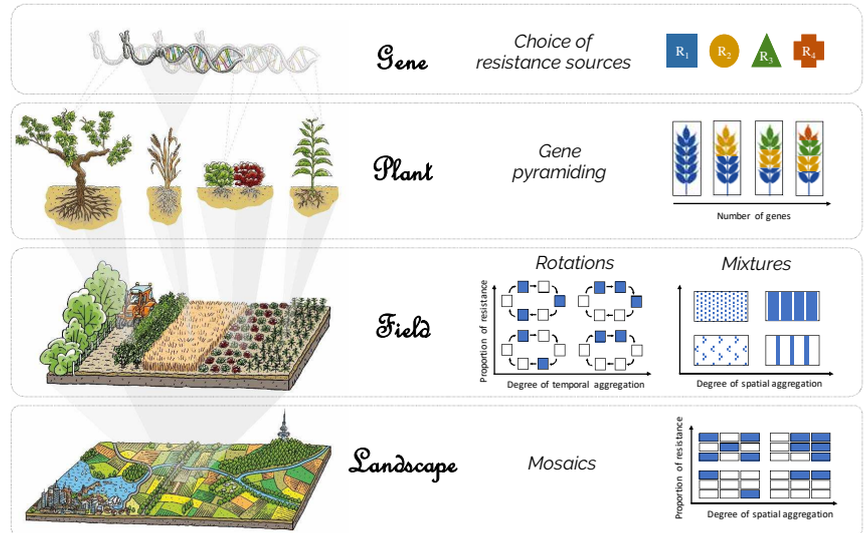


Strategies to improve plant resistance management rely on careful selection of resistance sources and their combination at various spatio-temporal scales.

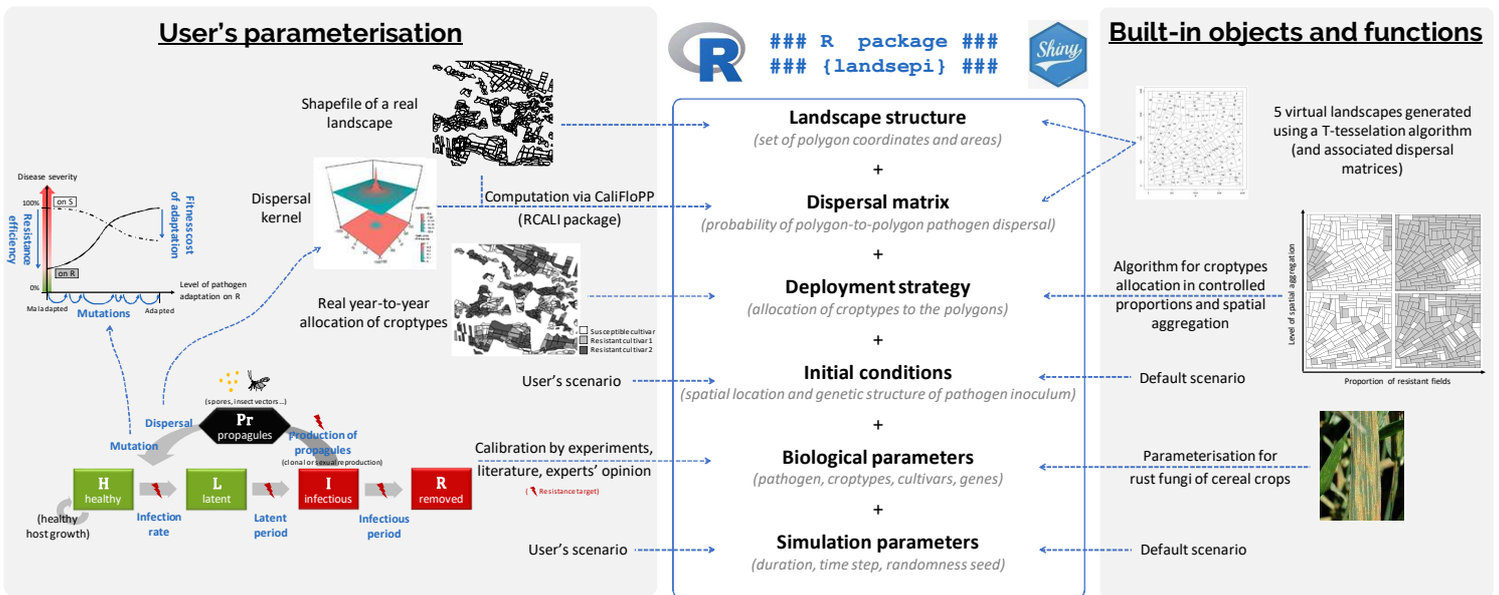
In *landsepi*, the landscape is a **dynamic mosaic of fields** cultivated with **croptypes**. Each croptype is composed of either a pure cultivar or a mixture; and each **cultivar** may carry one or several **resistance genes**. Each resistance gene targets one or several **pathogenicity traits**, with complete or partial **efficiency**, and may be expressed from the beginning of the season or later (e.g. APR gene). The pathogen may adapt to these genes (restoring its pathogenicity), possibly associated with a **fitness cost**.

The model is based on a **spatial geometry** for describing the heterogeneous landscape and allocating different cultivars, a **dispersal kernel** for the dissemination of the pathogen, and a **stochastic SEIR** ('Susceptible-Exposed-Infectious-Removed') structure with a discrete time step for the description of the host-pathogen interaction. Cropping seasons are split by host harvests which impose potential **bottlenecks** to the pathogen. The model accounts for pathogen evolution (via **mutation**, **sexual reproduction**, **selection** and **drift**) and provides **epidemiological**, **evolutionary** and **economic outputs** to assess the performance of the simulated strategies.

The package also includes a **shiny interface** for pedagogical purposes.



Adapted from Rimbaud L., Fabre F., Papaix J., Mouy B., Lannou C., Barrett L. and Thrall P. (2021). Models of plant resistance deployment. *Annual Review of Phytopathology*, 59(1):125-152.



Average computational time: 30 seconds (for a 30-year simulation with 3 cultivars and 2 major resistance genes)

Spatial unit: The spatial unit is a polygon. An agricultural field (i.e. a piece of land cultivated by the same croptype) may be composed of a single or several polygons.

Individual: A host 'individual' is an infection unit and may correspond to a given amount of plant tissue (where a local infection may develop, e.g. fungal lesion) or a whole plant (e.g. systemic viral infection). In the first case, plant growth increases the amount of available plant tissue (hence the number of individuals) during the cropping season.

Computation of specific outputs from raw results

Outputs

- Evolutionary: **Resistance durability**
ability to limit pathogen evolution and delay resistance breakdown
(Pathotype and genotype frequencies, Time to resistance breakdown)
- Epidemiological: **Disease control**
ability to reduce disease impact
(AUDPC, GLA)
- Economic: **Cost efficiency**
ability to overcompensate epidemic losses and management costs
(Yield, Product, Operational cost, Margin)

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PUBLICATIONS

- Rimbaud L., Fabre F., Papaix J., Mouy B., Lannou C., Barrett L. G. and Thrall P. H. (2021). Models of plant resistance deployment. *Annu. Rev. Phytopathol.* 59(1):125-152.
- Rimbaud L., Papaix J., Rey J.-F., Barrett L. G. and Thrall P. H. (2018). Assessing the durability and efficiency of landscape-based strategies to deploy plant resistance to pathogens. *PLoS Comput. Biol.* 14:e1006067.
- Rimbaud L., Papaix J., Barrett L. G., Burdon J. J. and Thrall P. H. (2018). Mosaics, mixtures, rotations or pyramiding: What is the optimal strategy to deploy major gene resistance? *Evol. Appl.* 11:1791-1810.

LINKS

- Homepage: <https://csiro-inra.pages.biosp.inrae.fr/landsepi/>
Web app: https://shiny.biosp.inrae.fr/app_direct/landsepi/
R package: <https://cran.r-project.org/web/packages/landsepi/index.html>

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