

## Contribution to phenomics for quantitative disease resistance in a germplasm collection of Capsicum spp.: a multivariate analysis by self-organizing map

Melissa Cantet, Guillaume Marie, Anne-Marie Sage-Palloix, Anne Massire, Alain Palloix, Véronique Lefebvre

## ▶ To cite this version:

Melissa Cantet, Guillaume Marie, Anne-Marie Sage-Palloix, Anne Massire, Alain Palloix, et al.. Contribution to phenomics for quantitative disease resistance in a germplasm collection of Capsicum spp.: a multivariate analysis by self-organizing map. 9. Solanaceae Conference (SOL2012), Aug 2012, Neuchâtel, Switzerland. 250 p. hal-02745281

## HAL Id: hal-02745281 https://hal.inrae.fr/hal-02745281v1

Submitted on 3 Jun 2020

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

## Contribution to phenomics for quantitative disease resistance in a germplasm collection of capsicum spp.: a multivariate analysis by self-organizing map

**Mélissa Cantet,** Guillaume Marie, Anne-Marie Sage-Palloix, Anne Massire, Alain Palloix and Véronique Lefebvre

INRA, Avignon, France

Large-scale phenotyping of genetic resources for agronomic traits is useful to structure collections, identify valuable progenitors, and define core-collections for association studies. Plant diseases are one of the main threats for crop production, and resistant genitors are constantly needed for breeding. Many traits such as partial resistance to *Phytophthora capsici* in pepper (*Capsicum* spp.) exhibit a continuous range of responses evolving over time, and large collections are difficult to reliably assess for quantitative traits. To phenotype a pepper collection of 1186 lines for *P. capsici* resistance, we measured 6 variables along a 21 day-time-course post inoculation. By a Self-Organizing Map multivariate analysis, an undemanding clustering method that locates lines and weights variables on a precise map, we defined 10 clusters of distinct resistance levels. The most resistant one gathers 7.5% of the collection, including 6 well-known genitors, confirming the relative rarity of resistance. Other clusters highlight distinction between early and late resistances. Resistant lines originate mostly from *C. annuum*, and a majority of *C. pubescens* lines were resistant. Our results reveal that resistance phenotype is highly frequent in Asia, a secondary centre of diversification of pepper. We are now constructing a trait-specific core-collection and achieving an association study by sequencing strategic positions on a locus of interest to detect candidate SNPs.