

The StructuraLEP project: structural and functional characterization of Leptosphaeria maculans effectors and of their interactants

Isabelle Fudal-Grolier Fudal, Yohann Petit, Francoise F. Blaise, Bénédicte B. Ollivier, Clémence Plissonneau, Michel M. Meyer, Julie Gervais, Thierry T. Rouxel, Marie-Helene Balesdent, Karine Blondeau, et al.

▶ To cite this version:

Isabelle Fudal-Grolier Fudal, Yohann Petit, Francoise F. Blaise, Bénédicte B. Ollivier, Clémence Plissonneau, et al.. The StructuraLEP project: structural and functional characterization of Leptosphaeria maculans effectors and of their interactants. 8th Effectome meeting, Sep 2015, Lauret, France. , 2015. hal-02800676

HAL Id: hal-02800676 https://hal.inrae.fr/hal-02800676

Submitted on 5 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.



The StructuraLEP projet: structural and functional characterization of *Leptosphaeria maculans* effectors and their interactants

INR isbV

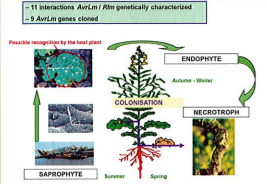
S UNIVERSITÉ PARIS SUD

Isabelle Fudal ¹, Yohann Petit ¹, Françoise Blaise ¹, Bénédicte Ollivier ¹, Clémence Plissonneau ¹, Michel Meyer ¹, Julie Gervais ¹, Thierry Rouxel ¹, Marie-Hélène Balesdent ¹, Karine Blondeau ², Noureddine Lazar ², Ines Gallay ², Herman van Tilbeurgh ²

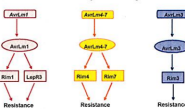
Context

Fungal effector genes are very diverse and typically encode small proteins, predicted to be secreted, with no or low homology in databases, and absence of known motif. As such their function or role in pathogenesis is mostly unknown and structure information provides an elegant way to resolve functional traits. Recently, three-dimensional (3-D) structures of several fungal and comycete avirulence effectors have been determined and have provided key advances in our understanding of plant-pathogen interactions including: the identification of structural similarities in effectors that were not visible in the sequence data, the identification of protein functions that were not apparent from sequences alone, and the visualization of molecular interfaces of relevance to pathogen virulence and plant immunity [1-2]. On these bases, the project StructuraLEP aims at elucidating the involvement of *L. maculans* effectors in pathogenicity through the structural and functional characterization of a few major effector proteins and the determination

Leptosphaeria maculans displays complex interactions with oilseed rape resistance genes



The Dothideomycete Leptosphaeria maculans causes stem canker, one of the most devastating diseases of oilseed rape (Brassica napus) worldwide. Control of fungal crop diseases necessitates a global understanding of fungal pathogenicity determinants and of their evolution. Genetic studies demonstrated gene-for-gene relationships between L. maculans and B. napus and allowed us to genetically identify eleven avirulence (AvrLm) genes in the pathogen and eleven corresponding resistance (Rlm) genes in the host plant [3]. We are investigating five L. maculans effectors chosen for their biological significance (involvement in fungal fitness, cognate R gene identified) or because they may represent novel modes of interaction with their plant target (two AVR genes necessary to be recognized by a specific R gene) [4-5; Plissonneau, Petit, Degrave, Rouxel and Balesdent, pers. comm.].





I. Structural and functional characterization of L. maculans effectors

Structural and functional characterization of *L. maculans* effectors will include production of effectors in heterologous systems (*Escherichia coli and / or Pichia pastoris*) and determination of their 3-D structure, determination of *in planta* localisation and identification of the cellular processes targeted by effectors.

Heterologous production in E. coli or P. pastoris and determination of 3D structure



Biological function of effectors

Localisation in planta (transient expression in tobacco leaves, immuno-cytolocalisation in oilseed rape during infection by L. maculans



Cellular processes targeted by effectors



II. Screening of plant proteins and molecules interacting with L. maculans effectors

Pull-down



Plant proteins targeted by effectors





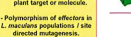
Validation (BiFC,

The search for molecules interacting with effectors will be performed using a Fluorescent Thermal Shift assay with a ligand library (including metals and co-factors). The search for host proteins interacting with effectors will be performed by yeast two hybrid screens using a cDNA library of oilseed rape infected by L. maculans. In parallele, pull-down assays using effector proteins produced in heterologous system against a set of plant proteins will be initiated (or, in case heterologous proteins are not available, effectors will be transiently expressed with a tag in Nicotiana bentamiana). Interesting candidates will be validated through independent methods such as colocalisation in N. benthamiana coupled to FRET / FLIM, Bimolecular Fuorescence Complementation (BiFC) or Co-immunoprecipitation (CoIP)

III. Functional and physical understanding of the identified interactions

The most interesting interactions identified will be characterized through co-crystallisation of interactants coupled to directed mutagenesis and through manipulation of plant targets in A. thaliana.

- Co-crystalisation effector / plant target or molecule.





Protein regions or amino-acids involved in interaction

Conclusions

Integration of knowledge on effector 3-D structure, localization, targeting of host proteins and host processes by *L. maculans* effectors studied in the StructuraLEP project will give insight into the fundamental processes governing the close association of a pathogenic fungus with its host plant. Progresses in these fields are necessary for sustainable crop production through the development of novel strategies to control fungal diseases.





¹ UMR BIOGER, INRA / AgroParisTech, Thiverval-Grignon, France ² UMR I2BC, Université Paris-Sud, / CEA / CNRS, Orsay, France.

Reference

- 1. Boutemy L.S., King S.R., Win J., Hugues R.K., Clarke T.A., Blumenschein T.M.A., Kamoun S. and Banfield M.J. (2011). J Biol Chem 286(41): 35834-35842.
- 2. Blondeau K, Blaise F., Graille M., Kale S.D., Linglin J., Ollivier B., Labarde A., Lazar N., Daverdin G., Balesdent M.H., Choi D.H.Y., Tyler B., Rouxel T., van Tilbeurgh H. and Fudal I. (2015). *Plant J* 83(4): 610-624.
- 3. Delourme, R, Pilet-Nayel, M L, Archipiano, M, Horvais, R, Tanguy, X, Rouxel, T, Brun, H, Renard, M, and Balesdent, M H (2004). Phytopathology 94: 578-583.
- 4. Gout L, Fudal I, Kuhn ML, Blaise F, Eckert M, Cattolico L, Balesdent MH and Rouxel T (2006). Mol Microbiol 60: 67-80.

 5. Parlange F, Daverdin G, Fudal I, Kuhn ML, Balesdent MH, Blaise F, Grezes-Besset B, Rouxel T (2009). Mol Microbiol 71(4):851-63.