Proceedings of the LXIV SIGA Annual Congress Online, 14/16 September, 2021 ISBN: **978-88-944843-2-8** 

Poster Communication Abstract - 7.20

## FUNCTIONAL STUDY OF LIPOXYGENASE-MEDIATED RESISTANCE TO FUNGAL PATHOGENS IN MAIZE AND GRAPEVINE

GUCHE M. D.\*\*\*\*\*\*, PILATI S.\*\*\*, DALLA COSTA L.\*\*\*, MOSER C.\*\*\*, GUELLA G.\*\*\*\*, TRENTI F.\*\*\*\*, LANUBILE A.\*, MAROCCO A.\*

\*) Department of Sustainable Crop Production, Università Cattolica del Sacro Cuore, Via Emilia Parmense 84, 29122, Piacenza, Italy. \*\*) C3A - Centro Agricoltura Alimenti Ambiente, Via Edmund Mach, 1 - 38010 San Michele all'Adige, Italy.
\*\*\*) Research and Innovation Centre, Fondazione Edmund Mach, via E. Mach 1, San Michele all'Adige, 38098, Italy.
\*\*\*\*) Department of Physics University of Trento, Via Sommarive,14 - 38123 Povo, Italy.

lipoxygenase, pathogen resistance, CRISPR/Cas9, Zea mays, Vitis vinifera

Fungal pathogens represent a big concern in maize and grapevine production, economically relevant crops for Italian agriculture. two Maize is challenged by the mycotoxigenic fungi Aspergillus flavus and Fusarium verticillioides, while grapevine production is affected by Erysiphe necator the causal agent of powdery mildew disease. Plant lipoxygenases genes ( LOXs) synthesize oxylipins that play a crucial role in the regulation of defense mechanisms against pathogens and influence the outcome of pathogenesis. Their genetic manipulation results in the alteration of plant resistance or susceptibility to certain pathogens. The role of LOX in host resistance against these fungi was investigated using in silico and in planta approaches. The phylogenetic analysis of grapevine and maize LOXs including well-characterized Arabidopsis and apple homologs showed the separation of 9-LOX and 13-LOX and several orthologous with a stronger clustering tendency in dicot species. Moreover, several duplication events leading to paralogous groups were inferred. In addition, in silico analysis of grapevine transcriptomic data on VESPUCCI platform was carried out to examine grapevine LOXs expression pattern against multiple fungal infections. As regards maize, a mutant carrying transposon insertion in the ZmLOX4 gene (UFMulox4) together with inbred lines W22, Mo17 and Tzi18 were

tested for the resistance to Aspergillus seedling rot (ASR) caused by A. flavus as well as for the aflatoxin production at 3- and 7-days post inoculation (dpi). Mo17 and W22 showed increased fungal susceptibility and higher levels of aflatoxin contamination compared to Tzi18 and UFMulox4. Moreover, the expression of genes involved in the LOX and jasmonic acid pathways (as *ZmLOX4*, hydroperoxide lyase 1 and acyl-coenzyme A oxidase) were induced earlier in Tzi18, whereas the genes 12-oxo-phytodienoic acid (12-OPDA) reductase (*ZmOPR8*) and *ZmLOX10* were upregulated at 7 dpi in infected samples. Interestingly, from lipid analysis an increased accumulation of the compound 10-oxo-11-phytoenoic acid was observed in the Tzi18 ears after *F. verticillioides* infection in field. In grapevine, controlled infection experiment of *E. necator* was performed using a susceptible and a resistant genotype (Teroldego and NY95 x Eger99 39, respectively) and gene expression pattern was analysed in treated and control leaves. Grapevine VviLOX2 and VviLOX12 were upregulated at 12 hours post inoculation (hpi), while upregulation of VviLOX7 and VviLOX9 occurred at 48 hpi in infected samples of the resistant genotype. Conversely, VviLOX13 was upregulated in infected samples of the susceptible genotype. Currently, field infection experiments are being carried out in maize against A. flavus and F. vorticellids to assess ZmLOXs expression patterns and associated lipid changes, while gene knock-out (CRISPR/Cas9 system) and over-expression experiments are underway for in-depth functional characterization of promising grapevine LOXs.