



UNIVERSITÀ DEGLI STUDI DI TORINO

## AperTO - Archivio Istituzionale Open Access dell'Università di Torino

### Development of SSR markers and assessment of polymorphism by High Resolution Melting Analysis in populations of chestnut nut rot agent Gnomoniopsis castanea

### This is the author's manuscript

Original Citation:

Availability:

This version is available http://hdl.handle.net/2318/1551809

since 2016-06-24T17:22:47Z

Terms of use:

**Open Access** 

Anyone can freely access the full text of works made available as "Open Access". Works made available under a Creative Commons license can be used according to the terms and conditions of said license. Use of all other works requires consent of the right holder (author or publisher) if not exempted from copyright protection by the applicable law.

(Article begins on next page)



# UNIVERSITÀ DEGLI STUDI DI TORINO

This is an author version of the contribution:

Questa è la versione dell'autore dell'opera: [Sillo F., Lione G., De Cesare S., Zampieri E., Giordano L., Gonthier P., 2015. Journal of Plant Pathology, 97, S38-S39]

> *The definitive version is available at:* La versione definitiva è disponibile alla URL: [http://sipav.org/main/jpp/index.php/jpp]

#### DEVELOPMENT OF SSR MARKERS AND ASSESSMENT OF POLYMORPHISM BY HIGH RESOLUTION MELTING ANALYSIS IN POPULATIONS OF CHESTNUT NUT ROT AGENT *GNOMONIOPSIS CASTANEA*. F. Sillo, G. Lione, S. De Cesare, E. Zampieri, L. Giordano, P. Gonthier. *Department*

of Agricultural, Forest and Food Sciences (DISAFA), University of Torino, Largo Paolo Braccini 2, I-10095 Grugliasco (TO), Italy. Email: paolo.gonthier@unito.it

In the complex phytosanitary situation of European chestnut, a relevant role is played by the recently described chestnut nut rot agent Gnomoniopsis castanea. In order to elucidate the epidemiology of this pathogen through a population genetic study, five SSR loci were isolated starting from four genomic libraries enriched in SSR sequences prepared and screened through the Microsatellite Amplified Library (MAL) method. To asses their polymorphism, a recently developed analysis called High Resolution Melting Analysis (HRMA) was used on 132 G. castanea isolates, collected in Italy, France and Switzerland. This approach allowed to distinguish different alleles based not only on repeat number but also on melting temperature differences among amplicons. Based on HRMA results, isolates were grouped in different clusters, each representing an allelic variant. Clusterization was confirmed by sequencing and alignment of the representative alleles. These results indicate that HRMA is an efficient, rapid and sensitive alternative to traditional electrophoresis-based method for SSR markers. It also allows to detect polymorphisms present in the SSR flanking regions, thus allowing to discriminate a larger number of haplotypes in fungal genetic population studies.