Strawberry cultivar and breeding lines susceptibility to Phytophthora crown and root rot in Huelva (Spain)

C. Borrero¹, A. Refoyo², C. Sanz³ and M. Avilés¹

¹E.T.S.I.A. Universidad de Sevilla, Sevilla, Spain, ²Fresas Nuevos Materiales S.A, Huelva, Spain, ³Instituto de la Grasa, CSIC, Sevilla, Spain.

Abstract
Phytophthora crown and root rot of strawberries is caused by Phytophthora cactorum. Breeding programs for resistance to this disease are in progress in Spain. In a previous work, the different susceptibility of the most important strawberry cultivars used in Huelva (Spain) was observed. In the present work, 2 cultivars and 3 breeding lines were included in assays of susceptibility with artificial inoculation. Eighteen plants of each cultivar were inoculated with a root bath in a suspension with P. cactorum propagules. After inoculation, plants were cultivated in a growth chamber in a randomized design with three repetitions of six plants each. The assays were conducted two times from September to December in 2014 and 2015. Results showed 4 susceptibility groups: 'Fortuna' the most susceptible cultivar; 'Primoris' with moderate susceptibility; 'A11-407P-3' and 'A10-48-3' with lower susceptibility and 'A10-207P'-8 the lowest susceptible breeding line. 'A10-207P-8' had 56.4% less disease severity than the most susceptible cultivar. Therefore, the use of these three breeding lines will be an interesting resource to control this disease in an integrated disease management program.

Keywords: Breeding, Fragaria × ananassa, soilborne, disease, Phytophthora cactorum.

INTRODUCTION
Crown and root rot caused in strawberry (Fragaria × ananassa Duch.) caused by Phytophthora cactorum was first reported in northern Germany in 1952 (Deutschmann, 1954). Since then, this disease has become a common problem in strawberry areas over the world (Semüller, 1984; Martínez et al., 2010). Spain is one of the five top strawberry producers in the world. In 2013 strawberry yield reached at 312,500 t (FAO, 2016). This culture is mainly localized in Huelva coast. Phytophthora crown and root rot of strawberries in Huelva, as other soil fungal diseases, is becoming worse every year during the phase-out of methyl bromide (Miranda et al., 2012) because fumigants authorized for soil to control disease are less effective. In this context, non-chemical approaches must be considered and developed. The use of strawberry plants resistant or tolerant against this and other soilborne pathogens, as a part of integrated control strategies, is a realistic alternative that must be seriously taken into account. Breeding programs towards resistance against important soilborne diseases are in progress in different countries (Faedi et al., 2002) including Spain.
This work summarized the study to characterize the response of some strawberry cultivars used in Huelva and new advanced selections into the Breeding program developed by Fresas Nuevos Materiales S.A. in collaboration with the University of Seville.

MATERIALS AND METHODS

Inoculation Methods
Five isolates obtained from affected plants in Huelva (Spain) were selected for inoculation. Isolate selection was different for each year in order to use fresh isolates. Inoculum was obtained blending 7 days-V8 growing colonies from that isolates in 100 ml of sterilized water per plate. Root plants were submerged in this suspension during one hour with orbital agitation 125 rpm. After that, 200 ml of this suspension was poured per pot with six plants. Plants were incubated at 100% humidity during two weeks.

Susceptibility Assay
This assay included 2 cultivars (‘Fortuna’ and ‘Primoris’) and 3 breeding lines (A10-84-3A10-207P-8 and A11-407P-3). Strawberry runner plants growing in 1.5 litres pots with peat growth medium were used for assay. Plants were cultivated in a growth chamber at 25 ºC during the day and 24 ºC at night (photoperiod: 10 h day and 14 h night) with a randomised complete block design with three repetitions (pots). The assays were conducted two times from September to December in 2014 and 2015.

Severity Assessment
Disease severity was monitored once per week after inoculation and was scored based on a symptom severity scale where: 0 = asymptomatic plants; 1 = infected plants (necrotic leaves); 2 = dead plants. Disease severity was expressed as proportion of the maximum possible disease severity. The area-under-the-disease-progress-curve-standardized (AUDPCs) per plant was calculated by disease severity integrated between symptoms onset and bioassay final time and dividing by the total epidemic time (days).

Statistical Analysis
Data collected from the trials were analyzed with Statgraphics Plus, Version 6 (SGS, 1999). Cultivars and year effect on severity (AUDPCs) were analyzed with a factorial ANOVA. Significant means were compared by Fisher LSD test (P<0.05).

RESULTS AND DISCUSSION
Cultivars showed different susceptibility to P. cactorum (P<0.001). Year factor also had significant differences. Severity was higher in 2014 than in 2015 (P<0.001, data not shown). On the contrary, the interaction between cultivar and year did not showed significant difference (P>0.05, data not shown). The different severity in both years can be due to the variable virulence for each group of isolates per year. Phytophthora isolates storage can induce the loss of virulence (Shaw, 1988). On the other hand, physiological condition of the host also greatly affects susceptibility, which can fluctuate considerably throughout the year (Duncan, 2002). Therefore, the different severity between years can be explained for virulence attenuation after isolates storage or for variations in plant development at the beginning of assays.

Results showed 4 susceptibility groups: ‘Fortuna’ the most susceptible cultivar; ‘Primoris’ with moderate susceptibility; ‘A11-407P-3’ and ‘A10-48-3’ with similar susceptibility than ‘Primoris’ and ‘A10-207P-8’ and finally ‘A10-207P-8’ had lower susceptibility than ‘Primoris’. ‘A10-207P-8’ had 56.4% less disease severity than the most susceptible cultivar (Fig. 1). The high susceptibility in ‘Fortuna’ cultivar to crown root rot
recorded in these bioassays is confirmed with the high crown root rot incidence in ‘Fortuna’ plants in Huelva fields as observed by farmers and technicians. This cultivar is the most cultivated in the last two years (Gómez-Mora et al., 2015). Therefore, the use of these three breeding lines will be an interesting resource to control this disease in an integrated disease management program.

ACKNOWLEDGEMENTS
This research was supported by grants from Corporación Tecnológica de Andalucía (CTA: 12/633). We thank S. Castillo, A. Gata and S. Pérez for excellent technical assistance.

Figure 1: Disease severity recorded during the bioassays in five cultivars.

*AUDPCs: Area-under-the-disease-progress-curve-standardize per plant was calculated by disease severity integrated between symptoms onset and bioassay final time and dividing by the total epidemic time (days). Data were recorded from two bioassays performed in two different years. Analysis of variance was performed with transformed data with $x^{-1/2}$. Bars with the same letters were not significantly different according to LSD test at $P<0.05$. Standard error of mean is indicated by a vertical line (n=6).

**Literature cited**


