## EUROPEAN AND MEDITERRANEAN PLANT PROTECTION ORGANIZATION ORGANISATION EUROPEENNE ET MEDITERRANEENNE POUR LA PROTECTION DES PLANTES Summary sheet of validation data for a diagnostic test

The EPPO Standard PM 7/98 Specific requirements for laboratories preparing accreditation for a plant pest diagnostic activity describes how validation should be conducted. It also includes definitions of performance criteria.

Laboratory contact details	Institute for Sustainable Plant Protection via Amendola, 122/D, 70126 Bari, Italy	
Short description of the test	detection of Xylella fastidiosa by Molecular real time PCR in insect vectors using the automatized DNA extraction protocol by Promega (Maxwell® RSC PureFood GMO and Authentication Kit AS1600)	
Date, reference of the validation report	2021-01-30 - INTERLABORATORY COMPARISON EUXF-IC-2020-03. Evaluation of molecular methods for the detection of Xylella fastidiosa	
Validation process according to EPPO Standard PM7/98?	yes	
Is the lab accredited for this test?	no	
Was the validated data generated in the framework of a project?	Other_project	
If yes, please specify	XF-ACTORS	
Description of the test		
Organism(s)	Xylella fastidiosa (XYLEFA)	
Detection / identification	detection	
Method(s)	Extraction Molecular Extraction DNA RNA Molecular real time PCR	
Method: Extraction		
Reference of the test description		
As or adapted from an EPPO diagnostic protocol	no	
New test being considered for inclusion in the next version of the EPPO diagnostic protocol?	yes	
As or adapted from an IPPC diagnostic protocol	no	
Reference of the test	INTERLABORATORY COMPARISON EU-XF-IC-2020-03 Evaluation of molecular methods for the detection of Xylella fastidiosa, January 20201	
Is the test modified compared to the reference test	no	
Other information		

Method: Molecular Extraction DNA RNA		
Reference of the test description		
As or adapted from an EPPO diagnostic protocol	no	
New test being considered for inclusion in the next version of the EPPO diagnostic protocol?	yes	
As or adapted from an IPPC diagnostic protocol	no	
Reference of the test	INTERLABORATORY COMPARISON EU-XF-IC-2020-03 Evaluation of molecular methods for the detection of Xylella fastidiosa, January 20201	
Is the test modified compared to the reference test	no	
Kit		
Is a kit used	yes	
Manufacturer name	PROMEGA	
Specify the kit used	Maxwell® RSC PureFood GMO and Authentication Kit	
Kit used following the manufacturer's instructions?	no A single insect head, should be homogenized in a 2 mL tube with one or two 5-mm tungsten carbide beads (for a maximum of 15-20 s at a frequency of 24 cycles s-1, in Mill300 mixer/Tissue Lyser II (Qiagen) or similar equipment). Five hundred microlitres of CTAB buffer (Promega), 10μl of RNase A Solution (to eliminate RNA) and 20μl of Proteinase K (PK) Solution are added to each microcentrifuge tubes. The tubes are tapped, inverted and vigorously vortexed until the sample is resuspended. Tubes are placed in a heat block at 65°C for 30 minutes. During the incubation, cartridges are prepared according to manufacturer instructions. After incubation, tubes with the lysate are inverted or vortexed to mix thoroughly. Tubes are centrifuged at ≥16,000 g for 10 minutes to separate any oils and solids. 300 μl of clear lysate are transferred into well #1 of the reagent cartridge by avoiding pipetting any solid material from the bottom of the tube or on the surface of the liquid and avoiding putting oil on the surface (these materials may inhibit downstream tests). If necessary, transfer the cleared lysate to a new tube and centrifuge again to avoid oils and solids. Purify on the Maxwell® Instrument according to the manufacturer's instructions. For elution, 50 μl of Elution Buffer are used.	
Other information		
Method: Molecular real time PCR  Reference of the test description		
As or adapted from an EPPO diagnostic protocol	yes	

New test being considered for inclusion in the next version of the EPPO diagnostic protocol?	no	
EPPO Diagnostic Protocol name	PM 7/024 Xylella fastidiosa (version 4)	
Name of the test	Real-time PCR - simplex (Harper et al., 2010; erratum 2013)	
As or adapted from an IPPC diagnostic protocol	yes	
IPPC diagnostic Protocol name	(version )	
Is the test modified compared to the reference test	no	
Kit		
Is a kit used	no	
Other information		
Reaction type	Simplex	
Performance Criteria :		
Organism 1.:	Xylella fastidiosa(XYLEFA)	
Analytical sensitivity		
What is smallest amount of target that can be detected reliably?	10^2 cfu/ml	
Diagnostic sensitivity		
Proportion of infected/infested samples tested positive compared to results from the standard test, see appendix 2 of PM 7/98	98.75% (on 3 replicates for each 10-fold serial dilutions from 10^6 to 10 CFU/ml of bacterial suspension spiked in Xylella-free P. spumarius homogenate)	
Standard test(s)	CTAB-based extraction protocol (diagnostic sensitivity 95.11%)	
Diagnostic Specificity		
Proportion of uninfected/uninfested samples (true negatives) testing negative compared to results from a standard test	97.92% (on 3 replicates of Xylella-free P. spumarius)	
Specify the test(s)	CTAB-based extraction protocol (diagnostic sensitivity 88.89%, repeatability 96.30%, reproducibility 94.07 %)	
Reproducibility		
Provide the calculated % of agreement for a given level of the pest (see PM 7/98)	98.61% (on 3 replicates for each 10-fold serial dilutions from 10^6 to 10 CFU/ml of bacterial suspension spiked in Xylella-free P. spumarius homogenate; on 3 replicates of Xylella-free P. spumarius; tested in 16 different laboratories)	
Repeatability		
Provide the calculated % of agreement for a given level of the pest (see PM 7/98)	97.22% (on 3 replicates for each 10-fold serial dilutions from 10^6 to 10 CFU/ml of bacterial suspension spiked in Xylella-free P. spumarius homogenate; on 3 replicates of Xylella-free P. spumarius)	

Test performance study		
Test performance study?	no	
Brief details of the test performance study and its output.It available, link to published article/report	TPS organized within the interlaboratory comparison EU-XF-IC-2020-03), in the framework of the activities related to the experimental plan foreseen in WP4/WP9 of the Horizon 2020 project "XF-ACTORS – 727987", and follows the previous European proficiency testing EU-XF-PT-2017-02 carried out in 2017. https://www.xfactorsproject.eu/wp-content/uploads/2021/01/EU-XFIC-2020-03-Report-V2-1.pdf	
The following complementary files are available online:	• Report IC	

Creation date: 2021-12-14 12:47:31 - Last update: 2022-01-25 14:46:02