EUROPEAN AND MEDITERRANEAN PLANT PROTECTION ORGANIZATION ORGANISATION EUROPEENNE ET MEDITERRANEENNE POUR LA PROTECTION DES PLANTES (11-17239)

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.

Target Organism	Erwinia amylovora		
Short description	Detection of Erwinia amylovora from plant material by Commercial lateral flow device Pocket Diagnostics		
Laboratory contact details	Bacteriology. Instituto Valenciano de Investigaciones Agrarias CV-315, km. 10.7, 46113 Moncada, Spain		
Date and reference of the validation report	2012-03 - Not specified		
Validation process according to EPPO Standard PM 7/98:	Yes		
Reference of the test description	PM 7/020(1) To be included in the revision Pocket Diagnostics (Forsite Diagnostics, York, UK).		
Is the test the same as described in the EPPO DP?	No Pocket Diagnostics (Forsite Diagnostics, York, UK).		
Is the lab accredited for this test?	No		
Plant species tested (if relevant)	Several plant species from the Rosaceae family		
Matrices tested (if relevant)	Shoots, leaves		
List of methods used			
Method for extraction / isolation / baiting of target organism from matrix			
Molecular methods, e.g. hybridization, PCR and real time PCR			
Serological methods: IF, ELISA, Direct Tissue Blot Immuno Assay	Х	Commercial lateral flow device	
Plating methods: selective isolation			
Bioassay methods: selective enrichment in host plants, baiting, plant test and grafting.			
Pathogenicity test			
Fingerprint methods: protein profiling, fatty acid profiling & DNA profiling			

Morphological and morphometrical methods intended for identification				
Biochemical methods: e.g. enzyme electrophoresis, protein profiling				
Other				
Analytical sensitivity (= limit of detection)				
What is smallest amount of target that can be detected reliably?	10^5-10^6 CFU/mL plant extract			
Diagnostic sensitivity				
Proportion of infected/infested samples tested positive compared to results from the standard test , see appendix 2 of PM 7/98	Proportion of true positives/total number of samples: 0.13 (in samples from 1 to 10^6 CFU/mL and healthy samples in ring test 2010)			
Specify the standard test				
Analytical specificity				
Specificity value	-			
Number of strains/populations of target organisms tested				
Number of non-target organisms tested				
Cross reacts with (specify the species)				
Diagnostic Specificity				
Proportion of uninfected/uninfested samples (true negatives) testing negative compared to results from a standard test	Proportion of true negatives/total number of samples: 0.93 (in samples from 1 to 10^{6} CFU/mL and healthy samples in ring test 2010)			
Specify the standard test				
Reproducibility				
Provide the calculated % of agreement for a given level of the pest (see PM 7/98)	94% wher	n tested with different operators, in IVIA assays		
<u>Repeatability</u>				
Provide the calculated % of agreement for a given level of the pest (see PM 7/98)	96% In IV	'IA assays		
Test performance study				
Test performance study?	Yes			
Include brief details of the test performance study and its output.It available, provide a link to published article/report	Yes (14 laboratories from Europe, Morocco, USA and New Zealand) analysed 12 samples each (from 1 to 10^6 CFU/mL plant extract and healthy samples). Details about ring test protocol available.			
Other information				
Any other information considered	Recomme	nded only for symptomatic samples for its low		

useful	sensitivity but high specificity.
e.g. robustness, ease of performing the test, etc.	