# Detection of Pepino mosaic virus (PepMV) in Tomato Seed by Seed Extract RT-qPCR (SE-qPCR) 

Validation report, June 2023

## ISHI VALIDATION REPORTS

This ISHI validation study has been conducted to determine the fitness of the described method for its intended purpose according to the ISHI Guidelines for the Validation of Seed Health Methods ${ }^{1}$ and followed by an independent review of its outcome.

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ISF cannot guarantee that laboratories following the protocol described herewith will obtain similar results. Many factors, such as staff skills, laboratory equipment and conditions, reagents and sampling methods can influence the results. Consequently, in case of any litigation, ISF will not accept any liability on the use of these tests.

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# Detection of Pepino mosaic virus (PepMV) in Tomato Seed by Seed Extract RT-qPCR (SE-qPCR) 

## SUMMARY

Pepino mosaic virus (PepMV) is a Potexvirus known to cause a wide range of symptoms in cultivated tomato (Solanum lycopersicum) causing damage to fruit and crop. Seed is a proven pathway for the dispersal and the transmission of this virus. The ISHI initiated a project to develop and validate a seed extract (SE-) reverse transcriptase (RT) quantitative polymerase chain reaction (qPCR) assay for the detection of PepMV. The SE-qPCR assay is proposed as an (optional) prescreening step, as an alternative to the ELISA pre-screening step used in the current PepMV method and is therefore performed using phosphate buffered saline (PBS) buffer as RNA seed extraction buffer. The PepMV SE-qPCR assay involves collecting virus particles by macerating tomato seeds and this seed extract is used in a total RNA extraction. Samples are analysed for the presence of PepMV RNA in a multiplex RT-qPCR reaction.

In this validation study, the performance criteria analytical specificity, analytical sensitivity, selectivity, repeatability, reproducibility and diagnostic performance were validated to determine if the PepMV SE-qPCR assay is fit for the intended purpose of serving as a pre-screen prior to a bioassay. In validating the analytical specificity of the PepMV detecting qPCR assays, using NAKT05 and Bejop primers, the alignment of 130 PepMV whole genome sequences show that 15 accessions (11.5\%) have one or more mismatches with the NAKT05 and/or Bejop primers. Testing 25 PepMV isolates, representing the known five genotypes (CH2, EU, LP, PES and US1), and a synthetic ssDNA oligo containing the known mismatches show that the majority of the PepMV isolates will be detected by the two assays. The remaining isolates will at least be detected by one of the two PepMV assays. This showed a good inclusivity of the PCR primers. The tested 23 non-targets, consisting of 18 different virus/viroid species able to infect solanaceous crops, four virus-free plant material samples and nuclease free water, gave Cq values above the cut-off or no amplification with both PepMV primers, which showed that the exclusivity requirement is met for the assay. The analytical sensitivity of SE-qPCR was compared with ELISA and determined to be 1,000 -fold more sensitive. The LOD with a $100 \%$ confidence level was equivalent for both qPCR assays, showing an ability to detect around 19 to 65 PepMV copies $/ 5 \mu \mathrm{~L}$ eluate. Testing individual seeds from a PepMV infected seed lot in a background of healthy seeds has shown that the SE-qPCR assay is able to detect one PepMV infected seed in a subsample of 1,000 seeds. PepMV was detected in five matrices by SE-qPCR with comparable quantitative results, showing good selectivity of the assay. SE-qPCR results were repeatable, as shown with comparable detection of PepMV between replicate samples with $100 \%$ accordance of test results. A comparative test (CT) among eight ISHI laboratories showed a reproducibility of the SE-qPCR assay within the pre-set criterium. Six laboratories provided correctly scored data for the positive extraction control (PEC), negative process control (NPC) and positive process control (PPC) and were included in the concordance calculation. The two remaining laboratories obtained poor performance with the PEC resulting in no signal or reduced signal for the NPC. This failed control resulted in excluding these two laboratories from the concordance calculation. The concordance percentage for the six laboratories was with $98.6 \%$, with a false negative percentage of $0.7 \%$, above the required $90 \%$ indicating that the reproducibility criterium was met. For robustness, the seed sample set used in the CT was also tested with an alternative extraction method and compared with the extraction method described in the protocol. The samples processed with the
alternative extraction method, dry grinding, guanidine-based extraction buffer and RNA extraction kit suitable for KingFisher platform, showed an increased sensitivity in the detection of PepMV, but obtained the same qualitative positive/negative result. Although not fully validated, this alternative extraction method could be suitable for laboratories when increased throughput is needed. The SE-qPCR diagnostic performance in reference to ELISA is almost identical. All eight natural infected lots tested positive by ELISA were also tested positive by SEqPCR. From the 13 ELISA negatives lots, one tested positive by SE-qPCR. This shows that diagnostic sensitivity is $100 \%$ and diagnostic specificity $92.86 \%$, which is above the requirement of $90 \%$ for this criterium.

This report confirms that the PepMV SE-qPCR assay is a suitable pre-screen assay to the ISHI Bioassay, being able to detect a single PepMV contaminated seed in a background of 1,000 healthy seeds. The ISHI PepMV SE-qPCR assay is deemed fit for its intended purpose.

## 1. INTRODUCTION

Pepino mosaic virus (PepMV) is a Potexvirus, which is known to cause a wide range of symptoms in cultivated tomato (Solanum lycopersicum) causing damage to fruit and crop. Tomato seed is a proven pathway for PepMV (Hanssen et al., 2010), and PepMV is easily spread mechanically during crop handling, causing damage to fruit and crop such as fruit marbling, discoloration and the occurrence of 'open fruit' (Hanssen and Thomma, 2010). As stated in the ISF regulated pest list database (https://pestlist.worldseed.org/public/pestlist.jsp), the recommended management strategy is to evaluate seed productions by field inspection or seed testing of a representative sample of each seed lot. The current ISHI method for detecting PepMV in tomato seeds consists of i) inoculating Nicotiana benthamiana plants followed by a confirmatory ELISA assay or ii) prescreening seed extracts using an ELISA assay followed by a bioassay for the ELISA positive seed lots to determine if infectious PepMV virions are present (ISF, 2021).

ISHI has been working on the development and optimisation of a pre-screen assay based on seed extract (SE) reverse transcriptase (RT) quantitative polymerase chain reaction (qPCR) assay (SEqPCR) using phosphate buffered saline (PBS) buffer as RNA seed extraction buffer. This method may be useful in case of e.g., reduced availability of high-quality antibodies for the ELISA assays. The SE-qPCR assay described in the protocol in Annex A is proposed as an alternative to the ELISA assay used in the PepMV current method. The PepMV SE-qPCR can be used as a pre-screen assay to identify seed lots free from PepMV RNA. Remaining suspected seed lots can be tested by an indirect pre-screen ELISA assay and/or direct assay (bioassay) followed by an ELISA assay to detect and confirm the presence of infectious PepMV. Replacing the confirmation by ELISA with confirmation by RT-qPCR could be a next change in the method for the detection of Pepino mosaic virus on tomato seed (version 5, March 2021), but it was out of scope for this validation project. This is to reduce the complexity of the validation, e.g., including the matrix Nicotiana benthamiana leaf material.

The method process flow is shown in Figure 1. The pre-screen SE-qPCR assay is illustrated by the grey box and is the part of the method being validated in this project. If the SE-qPCR turns out to be negative for PepMV RNA, the seed lot can be considered free from PepMV. If the SE-qPCR is positive for PepMV RNA, the seed lot is considered suspect, and follow-up assays to detect and confirm the presence of infectious PepMV are required (marked in blue boxes).


Figure 1. Method process flow.

## 2. OBJECTIVES

In this validation report, the experiments to investigate whether the SE-qPCR pre-screen assay developed by ISHI is fit for purpose, are described, together with their results. The bioassay and ELISA detection and confirmation steps to test if a suspected sample contains infectious virions have already been described (ISF, 2021) and are, therefore, out of the scope of this project. The described experiments assess the analytical specificity, analytical sensitivity, selectivity, repeatability, reproducibility and diagnostic performance of the SE-qPCR pre-screen assay and were designed according to the ISHI Guidelines for the Validation of Seed Health Test (ISHI, 2020).

## 3. METHOD VALIDATION

### 3.1. Analytical specificity

Definition ISHI guidelines: The ability of an assay to detect the target(s) pathogens (inclusivity) while excluding non-targets (exclusivity).

The analytical specificity requirements will be met when no false negative results are found for the targets and no false positive results for the non-targets. A false negative means that the two PepMV specific targets in the qPCR assay (Bejop and NAKT05) score Cq>35 to no amplification and that the sample has been proven to be PepMV positive by either Sanger sequencing or a positive bioassay result.

Bacopa chlorosis virus (BaCV) is used as a positive extraction control (PEC) and should give no reaction with any of the target primer sets.

## Experimental approach

The following sources were used to compile an overview of specificity data:
1- In silico data analysis: The evaluation of the primer/probe sets was included in the experimental approach. Due to the possibility of having unreliable PepMV sequences in Genbank, which would make the interpretation of the results complicated when deviant isolate(s) cannot be tested in wet laboratory experiments, only full genome accessions were included in these analyses. Accessions displaying a $100 \%$ match with either Bejop or NAKT05 assay were accepted.

In the case of accessions with mismatches in both target assays, these were either tested in a wet laboratory experiment or ordered as a representative synthetic ssDNA oligo containing the deviant mismatch to determine the effect of this mismatch on detection. Deviant isolates with mismatches that give Cq values comparable or were more sensitive than the positive/negative outcome of ELISA were accepted.

2- Data from leaf and/or seed samples: Data from leaf and/or seed samples from various origins infected with the target virus PepMV or non-target virus/viroid of which tomato is a host were compiled. At least 20 different isolates of PepMV and 20 non-targets, including virus free plant material, were investigated.

## Statistical analysis

The data were analysed qualitatively (positive vs. negative).

## Results

Primer and probe positions (in base pair (bp)) are shown in Table 1. Both primer/probe sets were aligned to the 130 PepMV genomes available at NCBI in June 2021 and the complete NCBI nucleotide collection for viruses.

Table 1. Sequence of PepMV-specific primers.

|  | Fw primer | Probe | Rv primer | Amplicon length (bp) |
| :---: | :---: | :---: | :---: | :---: |
| Bejop PepMV | Location : RNA dependent RNA polymerase (87... 4406 bp ) |  |  |  |
| Bejop 134 | 195-214 | 232-259 | 261-278 | 84 |
| Bejop 135 |  |  |  |  |
| Bejop 136 |  |  |  |  |
| Bejop 137 |  |  |  |  |
| Bejop 138 |  |  |  |  |
| Bejop 139 |  |  |  |  |
| Bejop 140 |  |  |  |  |
| NAKT05 PepMV | Location: Triple gene block 2 protein (5117... 5488 bp ) |  |  |  |
| KL05_48 | 5129-5148 | 5184-5203 | 5216-5235 | 107 |
| KL05_49 |  |  |  |  |
| NAKT05_50 |  |  |  |  |
| KL05_51 |  |  |  |  |
| KL05_52 |  |  |  |  |

This in silico analysis of both Bejop and NAKT05 primer/probe sets showed that eight PepMV accessions have one or more mismatches with the NAKT05 assay, 14 PepMV accessions have only one mismatch with the Bejop assay and 15 PepMV accessions have one or more mismatches with both assays. A representation of these accessions is shown in Annex B. The 22 accessions with mismatches in the Bejop or NAKT05 assay have a $100 \%$ match with the complementary assay and are therefore expected to be detected by one of the primer/probe sets of this PCR method. The 15 accessions with mismatches in both assay targets were either validated by testing the isolate in wet lab experiments (e.g., Yur1.5 HG313807) or by testing and comparing representative synthetic ssDNA oligo with (PC_NAKT05m) and without (PC_NAKTO5) the deviant mismatches in the NAKT05 assay. The design of these two ssDNA oligonucleotides is shown in Annex C. These ssDNAs, with a concentration of $100 \mathrm{pmol} / \mathrm{\mu L}$, were separately diluted in IDTE (TE solution pH 8.0 solution (cat. n. 11-05-01-09, Integrated DNA Technologies, USA)) and tested
by RT-qPCR. The concentration is known; therefore, the number of target site copies per $5 \mu \mathrm{~L}$ of template used in a qPCR reaction was calculated. This experiment showed that the mismatches created in PC_NAKT5m ssDNA reduce the sensitivity of the assay by 10 -fold, causing a detection limit of 300 copies $/ 5 \mu \mathrm{~L}$ instead of the 30 copies $/ 5 \mu \mathrm{~L}$ in the ssDNA without mismatches, as is shown in Table 2.

Table 2. Comparison of the PepMV NAKT05 assay on two synthetic ssDNA oligos. Rep.: replicate, No amp.: no amplification.

| Concentration <br> (copies $/ 5 \boldsymbol{\mu L}$ ) | PC_NAKTO5 |  | PC_NAKTO5m |  |
| :---: | :---: | :---: | :---: | :---: |
|  | Rep. 1 (Cq) | Rep. 2 (Cq) | Rep. 1 (Cq) | Rep. 2 (Cq) |
| $3.01 \times 10^{8}$ | 11.70 | 11.57 | 14.56 | 14.39 |
| $3.01 \times 10^{7}$ | 15.17 | 14.88 | 18.20 | 18.00 |
| $3.01 \times 10^{6}$ | 18.43 | 18.26 | 21.53 | 21.28 |
| $3.01 \times 10^{5}$ | 21.50 | 21.58 | 24.94 | 24.76 |
| $3.01 \times 10^{4}$ | 25.29 | 25.24 | 28.40 | 28.29 |
| $3.01 \times 10^{3}$ | 28.64 | 28.47 | 31.52 | 31.71 |
| $3.01 \times 10^{2}$ | 32.35 | 31.58 | 34.47 | 34.74 |
| $3.01 \times 10^{1}$ | 36.43 | 35.68 | 39.51 | No amp. |
| $3.01 \times 10^{0}$ | No amp. | 37.95 | No amp. | No amp. |
| $3.01 \times 10^{-1}$ | No amp. | No amp. | No amp. | No amp. |
| $3.01 \times 10^{-2}$ | No amp. | No amp. | No amp. | No amp. |
| $3.01 \times 10^{-3}$ | No amp. | No amp. | No amp. | No amp. |

Company data from leaf and/or seed samples from various origins infected with the target virus PepMV or non-target virus/viroid was compiled (see Annex D, Table D. 1 and Table D.2). All 25 PepMV isolates, originating from 14 different countries, were detected with the NAKT05 assay but two isolates were not detected with Bejop. This includes PepMV isolate Yur1.5 (accession HG313807) of which was shown in the in silico analyses that isolates Yur1.5, Tor9 and Chi2.9 had mismatches for both assays. RNA from YUR1.5 was provided by Dr. Fernando García-Arenal Rodríguez (Plant-Virus Interaction and Co-evolution Research Group at Centro de Biotecnología y Genómica de Plantas UPM-INIA). Here, it was shown that these mismatches do not affect the detection with NAKT05 assay but result in no amplification with Bejop assay. The genotype of the 25 isolates was determined by Sanger sequencing of the amplicon generated by primer set Pep5-2 TACYTAGCHAAACCAATGGC and Pep6-2 GCWCCATCTTGKGATTGRTC (personal communication B. Woudt Syngenta, 2003, adapted), located on nucleotide position 3,441 to 3,761 of PepMV reference sequence NC004067. Two samples showed a mixed infection of isolates from the Ch2 and LP strains. The Yur1.5 isolate did not give any amplicon with this primer set. The remaining 22 isolates were aligned and organised in a phylogenetic tree (Figure 2), using the UPGMA construction method and Jukes-Canto nucleotide distance method with a bootstrap analyses of 500 replicates in CLC Main Workbench version 20.0.3., to determine if the isolates were genetically unique. These analyses showed that the 22 PepMV isolates used in the analytical specificity are genetically unique and represent the five known genotypes of PepMV; US1, CH2, EU, LP and PES (Moreno-Pérez et al., 2014).

All 22 non-target samples, consisting of 18 different virus/viroid species and four virus free plant material samples, resulted in Cq >35 or no amplification with both PepMV assays. The BaCV assay
that was used as inhibition control (IC) and positive extraction control (PEC) only reacted with the Bacopa chlorotic virus isolate and not with the PepMV isolates or other non-targets.


Figure 1. Phylogenetic tree of the 22 PepMV isolates used in the analytical specificity.

## Conclusion

The in silico analyses showed that there are multiple PepMV accessions with mismatches on the target locations of the assays. The wet lab experiments with the ssDNA containing the observed mismatches have shown that this can reduce the detection limit of the assays. Together with three RNA samples of PepMV isolates Tor-9, Chi2.9 and Yur1.5 with relevant mismatches have proven that this does not result in an overall false negative outcome. The compiled data of the assays ran on 25 targets and 22 non-targets has shown that the specificity requirements for the SE-qPCR were met. Although in the validation plan it was decided that the non-targets should have a Cq above 35, it is noteworthy that all non-targets were above the later determined cutoff of Cq 37 . For the validation criterium analytical specificity, the detection method is fit for purpose.

### 3.2. Analytical sensitivity

Definition ISHI guidelines: Smallest amount of the target pathogen that can be detected i.e., the limit of detection (LOD).

The analytical sensitivity requirements for the single seed detection will be met when a LOD equal to or lower than one infected seed in 1,000 seeds is obtained. The infection rate of the seed lot(s) was determined by ELISA.

## Experimental approach

Sensitivity data consists of two data sets:
1- Limit of Detection (LOD) data: Three independent dilution series of PepMV infected leaf material were prepared in healthy seed extracts of tomato (S. lycopersicum) and diluted in serial steps of 10 -fold. Each dilution was tested in triplicate by the SE-qPCR assay. The initial virus concentration was calculated using a dilution series of synthetic ssDNA of the PepMV assay
targets with a known number of copies as standard curve during the RT-qPCR. The lowest virus dilution in which PepMV can still be detected in all nine replicates is referred to as LOD and expressed as the number of copies per volume (copies $/ \mu \mathrm{L}$ ).

2- Single-seed detection data: Test samples of 1,000 seeds were prepared from three independent samples of a healthy seed lot spiked with 100, 10 and one single infected seed. The rate of infection and homogeneity of the PepMV positive seed lot used for spiking was determined by performing ELISA and SE-qPCR assays, in duplicate, on 30 individual seeds from the positive lot tested as 30 subsamples containing 249 healthy and one individual seed from the positive lot. The prepared seed samples of 1,000 seeds were tested in triplicate and the lowest seed dilution in which PepMV could still be detected in all nine replicates is referred to as LOD and expressed as the number of infected seeds per 1,000 seeds. Sensitivity of the test was performed with the highest expected seed subsample size of 1,000 . Assuming that if one wants to test smaller subsample sizes in the same buffer ratio, it does not negatively affect the detection of the target.

## Statistical analysis

The data were analysed qualitatively (positive vs. negative).

## Results

The compiled results of the LOD experiment are shown in Table 3 (raw data available in Annex E). They demonstrated that the LOD for the SE-qPCR lies at approximately dilution $10^{-7}$, which represents an average of 19 PepMV copies $/ 5 \mu \mathrm{~L}$ (Cq<37). The SE-qPCR was at least 1,000 -fold more sensitive than ELISA using the seed extraction buffer and the Qiagen RNeasy plant mini kit as described in the protocol. This is in-line with expectations since PCR includes a multiplication step, in contrast to ELISA, which leads to increased sensitivity.

Table 3. The LOD of PepMV in tomato seed extract with two different methods: ELISA and SE-qPCR. The average signal-to-noise ( $\mathrm{S} / \mathrm{N}$ ) extinctions ratios and Cq values, including the standard deviation, are displayed, respectively. Red cell: the LOD of the method used. No amp.: no amplification.

| Dilution $^{\mathbf{a}}$ | ELISA <br> Extinction <br> (OD) |  | S/N ratio | NAKT05 | Bejop | SE-qPCR (Cq) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Bejop |  |  |  |  |  |
| Undiluted | Not tested | Not tested | Not tested | Not tested | Not tested | Not tested |
| $10^{-1}$ | 0.778 | 16.2 | Not tested | Not tested | Not tested | Not tested |
| $10^{-2}$ | 0.309 | 6.6 | $18.92 \pm 0.37$ | $18.80 \pm 0.10$ | 1914444 | 2075556 |
| $10^{-3}$ | $0.091^{\text {c }}$ | $2.0^{\text {c }}$ | $22.22 \pm 0.07$ | $22.13 \pm 0.10$ | 202667 | 260111 |
| $10^{-4}$ | 0.049 | 1.1 | $25.60 \pm 0.39$ | $25.60 \pm 0.16$ | 21644 | 31800 |
| $10^{-5}$ | 0.046 | 1.0 | $28.75 \pm 0.18$ | $28.61 \pm 0.13$ | 2560 | 4862 |
| $10^{-6}$ | 0.043 | 0.9 | $33.00 \pm 0.23$ | $32.69 \pm 0.10$ | 150 | 406 |
| $10^{-7}$ | 0.042 | 0.9 | $36.18 \pm 0.64^{\mathrm{d}}$ | $35.72 \pm 0.24^{\mathrm{d}}$ | 19 | 65 |
| $10^{-8}$ | 0.041 | 0.9 | $38.45^{\mathrm{b}} \pm 0.20$ | $37.59^{\mathrm{b}} \pm 0.00$ | 4 | 22 |
| $10^{-9}$ | 0.042 | 0.9 | $38.46^{\mathrm{b}} \pm 0.00$ | No amp. | 4 | 0 |
| $10^{-10}$ | 0.043 | 0.9 | No amp. | No amp. | 0 | 0 |

[^1]The single-seed detection experiment on 30 individual seeds from seed lot SHR218, which was determined ELISA positive for three subsamples in 2018 (data not shown) and one subsample in 2022 (Table 10, diagnostic performance) out of 12 subsamples of 250 seeds each, demonstrated that virus titter of individual seeds is below the detection limit of the ELISA (Table 4 and Annex E). The SE-qPCR was able to detect 29 out of the 30 samples in two repeats with both assays. The only exception was for sample 17, which gave a Cq of 35.68 with the Bejop primer/probe set and a Cq of 37.49 / no amplification with the NAKT05 assay.

Table 4. The comparison of the results of 30 individual seeds from mixing one single seed from a PepMV positive seed lot together with 249 healthy seeds, with two different methods: ELISA and SEqPCR. The average signal-to-noise ( $\mathrm{S} / \mathrm{N}$ ) extinctions ratios and Cq values, including the standard deviation, are displayed, respectively. No amp.: no amplification, NPC: negative process control.

| Sample number | ELISA |  | SE-qPCR (Cq) |  | PepMV copies $/ 5 \mu \mathrm{~L}^{\text {a }}$ |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Extinction (OD) | S/N ratio | NAKT05 | Bejop | NAKT05 | Bejop |
| 1 | 0.043 | 0.94 | $31.23 \pm 0.10$ | $29.90 \pm 0.12$ | 545 | 1385 |
| 2 | 0.047 | 1.03 | $32.28 \pm 0.06$ | $31.13 \pm 0.32$ | 283 | 658 |
| 3 | 0.032 | 0.70 | $33.00 \pm 0.05$ | $31.31 \pm 0.00$ | 182 | 578 |
| 4 | 0.033 | 0.72 | $35.20 \pm 0.10$ | $33.80 \pm 0.59$ | 46 | 131 |
| 5 | 0.033 | 0.72 | $33.31 \pm 0.01$ | $31.16 \pm 0.04$ | 149 | 634 |
| 6 | 0.033 | 0.73 | $34.30 \pm 0.68$ | $31.31 \pm 0.11$ | 88 | 578 |
| 7 | 0.023 | 0.51 | $35.16 \pm 0.18$ | $33.16 \pm 0.05$ | 48 | 182 |
| 8 | 0.035 | 0.77 | $33.59 \pm 0.06$ | $31.34 \pm 0.18$ | 126 | 568 |
| 9 | 0.032 | 0.70 | $35.66 \pm 0.05$ | $32.63 \pm 0.57$ | 36 | 270 |
| 10 | 0.034 | 0.76 | $35.74 \pm 1.07$ | $34.26 \pm 0.22$ | 41 | 93 |
| 11 | 0.041 | 0.91 | $33.17 \pm 0.11$ | $31.46 \pm 0.06$ | 164 | 524 |
| 12 | 0.060 | 1.32 | $34.03 \pm 0.27$ | $32.03 \pm 0.73$ | 97 | 408 |
| 13 | 0.033 | 0.73 | $33.64 \pm 053$ | $31.06 \pm 0.51$ | 128 | 705 |
| 14 | 0.032 | 0.71 | $36.38 \pm 1.08$ | $33.96 \pm 0.15$ | 27 | 112 |
| 15 | 0.037 | 0.82 | $33.63 \pm 0.44$ | $31.64 \pm 0.08$ | 127 | 471 |
| 16 | 0.032 | 0.71 | $36.05 \pm 0.05$ | $34.83 \pm 0.70$ | 27 | 707 |
| 17 | 0.030 | 0.67 | 37.49/No amp. | $35.68 \pm 0.52$ | 6 | 41 |
| 18 | 0.034 | 0.76 | $33.49 \pm 0.70$ | $31.83 \pm 0.54$ | 147 | 442 |
| 19 | 0.032 | 0.71 | $34.94 \pm 0.12$ | $35.63 \pm 0.48$ | 54 | 41 |
| 20 | 0.034 | 0.76 | $35.90 \pm 0.56$ | $33.64 \pm 0.41$ | 32 | 140 |
| 21 | 0.056 | 1.24 | $33.31 \pm 0.58$ | $31.16 \pm 0.37$ | 63 | 650 |
| 22 | 0.033 | 0.73 | $37.15 \pm 0.39$ | $33.99 \pm 0.20$ | 14 | 110 |
| 23 | 0.032 | 0.70 | $34.65 \pm 0.09$ | $33.57 \pm 0.11$ | 65 | 142 |
| 24 | 0.034 | 0.74 | $33.15 \pm 0.38$ | $31.17 \pm 0.03$ | 170 | 631 |
| 25 | 0.032 | 0.71 | $33.58 \pm 0.45$ | $32.16 \pm 0.10$ | 132 | 340 |
| 26 | 0.031 | 0.68 | $29.67 \pm 0.06$ | $28.54 \pm 0.18$ | 1435 | 3245 |
| 27 | 0.030 | 0.67 | $34.39 \pm 0.35$ | $32.52 \pm 0.24$ | 78 | 276 |
| 28 | 0.033 | 0.72 | $35.48 \pm 0.23$ | $33.15 \pm 0.43$ | 39 | 191 |
| 29 | 0.033 | 0.72 | $32.24 \pm 0.27$ | $31.45 \pm 0.25$ | 294 | 534 |
| 30 | 0.033 | 0.72 | $33.02 \pm 0.05$ | $32.25 \pm 0.18$ | 179 | 373 |
| NPC | 0.045 | 0.99 | No amp. | No amp. | 0 | 0 |

${ }^{\text {a }}$ The PepMV concentration (copies $/ 5 \mu \mathrm{l}$ ) was calculated using a dilution series of synthetic ssDNA of the PepMV assay targets with a known number of copies as standard curve during the RT-qPCR.

Seed lot SHR218 was also used to create nine samples of 1,000 seeds of which three samples contained either one single seed, three samples with 10 positive seeds and three samples with 100 positive seeds from the positive lot, respectively, representing infection levels of 0.1\%, 1.0\% and 10\%. The results, compiled in Table 5 (full data available in Annex E), demonstrate that the SE-qPCR is able to detect approximately 28 copies per reaction originating from a single infected seed.

Table 5. The average Cq values of three subsamples of 1,000 seeds with three different PepMV infection levels and a sample of 1,000 healthy seeds tested with SE-qPCR. The Cq values, including the standard deviation, are displayed. No amp.: no amplification.

| Infection rate | SE-qPCR (Cq) |  |  | PepMV copies/5 $\boldsymbol{\mu L ^ { \mathbf { a } }}$ |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | NAKT05 | Bejop | PEC | NAKTO5 | Bejop |
| 0.1 | $35.68 \pm 0.66$ | $34.89 \pm 0.92$ | $32.30 \pm 0.67$ | 28 | 57 |
| 1.0 | $30.89 \pm 0.57$ | $29.68 \pm 0.62$ | $31.47 \pm 1.39$ | 626 | 1924 |
| 10.0 | $27.96 \pm 0.87$ | $26.95 \pm 1.19$ | $31.59 \pm 1.71$ | 4623 | 15847 |
| 0.0 | No amp. | No amp. | $31.71 \pm 0.39$ | 0 | 0 |

${ }^{\text {a }}$ The PepMV concentration (copies $/ 5 \mu \mathrm{l}$ ) was calculated using a dilution series of synthetic ssDNA of the PepMV assay targets with a known number of copies as standard curve during the RT-qPCR.

## Conclusion

The analytical sensitivity experiments have demonstrated that the SE-qPCR is 1,000 -fold more sensitive than ELISA and is able to detect around 19 to 65 copies per reaction when testing independent PepMV dilutions series in a tomato seed background. When testing single infected seeds in a subsample of 1,000 seeds, 28 to 57 copies per reaction were detected. These results prove that the SE-qPCR is able to detect a single low infected PepMV seed in a sample of 1,000 seeds and is more sensitive than ELISA. In conclusion, regarding the analytical sensitivity validation criterium, the SE-qPCR detection method is fit for purpose.

### 3.3. Selectivity

Definition ISHI guidelines: The effect of different seed matrices on the ability of the method to detect target pathogen(s).

The selectivity requirements will be met when:

- The Cq -values for both the spiked samples and the positive extraction control are within a range of 3 Cq between the different matrices,
- No positive results are obtained with the non-spiked samples, and
- No negative results are obtained with the spiked samples.


## Experimental approach

The seed extract of a single healthy seed samples of 1,000 seeds from five varieties, each belonging to a different tomato segment (including rootstock), was spiked with PepMV leaf extract and diluted in serial steps of 10 -fold in heathy seed extract until an equivalent of one infected seed, as determined by the analytical sensitivity criteria, was reached. All seed samples were processed according to the protocol described in Annex A. The different matrices consisted of three different tomato crops (beef tomato variety A produced in Chile in 2021, beef tomato variety T produced in Mexico in 2020 and truss tomato variety G produced in the Netherlands in 2021) and two rootstock varieties (rootstock variety $X$ and $Y$, both produced in the Netherlands in 2020).

## Statistical analysis

The analysis was done at a qualitative level per sample, viz. positive or negative result for PepMV.

## Results

A dilution was determined positive when all three replicates for both of the PepMV qPCR assays obtained a Cq<37. The selectivity experiment average data (Table 6; full data in Annex E) demonstrated that the LOD for the SE-qPCR lies at approximately dilution $10^{-7}$ for all five tomato and rootstock varieties. The Cq -values of each dilution of the spiked samples, the positive extraction control (PEC) and the negative process control (NPC) for each variety are within a range of 3 Cq between the different matrices.

Table 6. The matrices effect on PepMV in the seed extract of five different tomato varieties using SEqPCR method. The average Cq values, including the standard deviation, are displayed with the qualitative conclusion. Red cell: the LOD of the SE-qPCR, No amp.: no amplification, NPC: negative process control, NTC: negative template control, PEC: positive extraction control.

| Crop | Variety code | Dilution ${ }^{\text {a }}$ | SE-qPCR (Cq) |  |  | Qualitative conclusion |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | NAKT05 | Bejop | PEC |  |
| Tomato | A | $10^{-5}$ | $28.31 \pm 0.05$ | $28.54 \pm 0.13$ | $29.39 \pm 0.15$ | Positive |
|  |  | $10^{-6}$ | $31.64 \pm 0.13$ | $32.15 \pm 0.47$ | $28.94 \pm 0.17$ | Positive |
|  |  | $10^{-7}$ | $34.63 \pm 0.58$ | $35.84 \pm 0.73$ | $27.57 \pm 0.36$ | Positive |
|  |  | $10^{-8}$ | No amp. | 37.83 ${ }^{\text {b }}$ | $29.22 \pm 0.27$ | Negative |
| Tomato | G | $10^{-5}$ | $27.96 \pm 0.19$ | $28.02 \pm 0.25$ | $28.51 \pm 0.32$ | Positive |
|  |  | $10^{-6}$ | $31.13 \pm 0.11$ | $31.50 \pm 0.03$ | $28.33 \pm 0.04$ | Positive |
|  |  | $10^{-7}$ | $34.79 \pm 1.14$ | $35.37 \pm 0.79$ | $28.49 \pm 0.32$ | Positive |
|  |  | $10^{-8}$ | $36.77 \pm 0.01^{\text {b }}$ | $36.63^{\text {b }}$ | $28.25 \pm 0.12$ | Negative |
| Tomato | T | $10^{-5}$ | $28.26 \pm 0.12$ | $28.20 \pm 0.11$ | $28.64 \pm 0.12$ | Positive |
|  |  | $10^{-6}$ | $31.31 \pm 0.18$ | $31.84 \pm 0.31$ | $27.06 \pm 0.14$ | Positive |
|  |  | 10 | $34.76 \pm 1.68$ | $35.00 \pm 0.86$ | $28.53 \pm 0.07$ | Positive |
|  |  | $10^{-8}$ | $36.46 \pm 0.55{ }^{\text {b }}$ | $36.02^{\text {b }}$ | $29.42 \pm 0.15$ | Negative |
| Rootstock | X | $10^{-5}$ | $27.95 \pm 0.33$ | $28.49 \pm 0.18$ | $27.89 \pm 0.12$ | Positive |
|  |  | $10^{-6}$ | $31.14 \pm 0.08$ | $32.02 \pm 0.36$ | $28.50 \pm 0.22$ | Positive |
|  |  | $10^{-7}$ | $34.58 \pm 0.64$ | $34.91 \pm 0.56$ | $29.05 \pm 0.15$ | Positive |
|  |  | $10^{-8}$ | $36.96 \pm 0.09^{\text {b }}$ | No amp. | $29.27 \pm 0.04$ | Negative |
| Rootstock | Y | $10^{-5}$ | $28.12 \pm 0.11$ | $28.56 \pm 0.21$ | $29.54 \pm 0.25$ | Positive |
|  |  | $10^{-6}$ | $31.17 \pm 0.54$ | $31.38 \pm 0.24$ | $29.79 \pm 0.28$ | Positive |
|  |  | $10^{-7}$ | $34.87 \pm 0.62$ | $36.30 \pm 0.17$ | $30.02 \pm 0.12$ | Positive |
|  |  | $10^{-8}$ | No amp. | $35.93{ }^{\text {b }}$ | $29.65 \pm 0.07$ | Negative |
| Tomato | A | NPC | No amp. | No amp. | $29.94 \pm 0.65$ | Negative |
| Tomato | G | NPC | No amp. | No amp. | $28.44 \pm 0.10$ | Negative |
| Tomato | T | NPC | No amp. | No amp. | $30.16 \pm 0.26$ | Negative |
| Rootstock | X | NPC | $37.26^{\text {b }}$ | No amp. | $29.64 \pm 0.06$ | Negative |
| Rootstock | Y | NPC | No amp. | No amp. | $29.48 \pm 0.13$ | Negative |
| Water |  | NTC | No amp. | No amp. | No amp. | Negative |

${ }^{\text {a }}$ Five mg PepMV infected leaf in buffer, used to spike seed.
${ }^{\mathrm{b}}$ Includes six or more reactions without amplification.

## Conclusion

No matrices effect was observed when using the seed extraction buffer and the Qiagen RNeasy plant mini kit as described in the protocol. For the selectivity validation criterium, the SE-qPCR detection method is fit for purpose.

### 3.4. Repeatability

Definition ISHI guidelines: Degree of similarity in results of replicates of the same seed lots when the method is performed with minimal variations in a single lab.

The repeatability requirements will be met when accordance is $>90 \%$ and all subsamples yield consistent qualitative results between the three time points.

## Experimental approach

One technician using the same batch of material and equipment performed the SE-qPCR as described in Annex A at three different time points with three replicates of four different seed lots with a sample size of 1,000 seeds. The four lots consisted of a healthy seed lot (tomato seed lot A, Chile 2021), low-infected (0.5\%), medium-infected (1\%) and highly-infected (10\%) PepMV seed lot. The low and medium seed lot was made by adding five or ten seeds from naturally infected seed lot SHR218, to 995 or 990 seeds of the healthy seed lot, respectively. The highly infected seed lot was prepared by adding 100 seeds of a second naturally infected seed lot SHR203 to 900 seeds of the healthy seed lot. The samples were tested on the following time points, day 1 , day 8 and day 9 .

## Statistical analysis

The analysis was conducted on a qualitative level per sample, viz. positive or negative results for PepMV. The method of Langton et al. (2002) was used to evaluate the accordance (repeatability of data) of the SE-qPCR.

## Results

The results of the four different batches tested with three replicates on three different days are compiled in Table 7 and available in Annex E. The healthy batch resulted in no amplification for all nine replicates for the PepMV assays but did give amplification with the PEC. Concluding that this healthy batch is indeed repeatably negative. The three batches with a low, medium or high infection level of PepMV did give for all replicates an amplification for both PepMV assays and both $q P C R$ reaction repetitions. Although there is some natural variation in Cq values between replicates of the same batch, this did not result in a different outcome. All replicates of the three infected batches were repeatable and determined PepMV positive. All batches and their repeats gave the same consistent qualitative (positive or negative) result. The accordance for these batches was calculated as $100 \%$.

Table 7. The repeatability results of four different seed lots with different PepMV infection levels tested on three different days in triplicate. No amp.: no amplification, Rep: replicate.

| Infection rate | Time point | SE-qPCR (Cq) |  |  |  |  |  | Conclusion |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | NAKT05 |  | Bejop |  | PEC |  |  |
|  |  | Rep 1 | Rep 2 | Rep 1 | Rep 2 | Rep 1 | Rep 2 |  |
| Healthy | Day 1 | No amp. | No amp. | No amp. | No amp. | 30.39 | 30.27 | Negative |
|  |  | No amp. | No amp. | No amp. | No amp. | 31.11 | 30.96 | Negative |
|  |  | No amp. | No amp. | No amp. | No amp. | 30.88 | 31.04 | Negative |
|  | Day 8 | No amp. | No amp. | No amp. | No amp. | 30.59 | 30.53 | Negative |
|  |  | No amp. | No amp. | No amp. | No amp. | 27.15 | 27.26 | Negative |
|  |  | No amp. | No amp. | No amp. | No amp. | 30.68 | 30.67 | Negative |
|  | Day 9 | No amp. | No amp. | No amp. | No amp. | 30.71 | 30.50 | Negative |
|  |  | No amp. | No amp. | No amp. | No amp. | 30.80 | 30.63 | Negative |
|  |  | No amp. | No amp. | No amp. | No amp. | 30.30 | 30.14 | Negative |
| 0.5 | Day 1 | 29.01 | 28.73 | 27.42 | 27.31 | 27.42 | 27.41 | Positive |
|  |  | 29.09 | 29.00 | 27.56 | 27.36 | 28.87 | 28.77 | Positive |
|  |  | 32.65 | 32.34 | 30.77 | 30.57 | 29.46 | 29.54 | Positive |
|  | Day 8 | 33.18 | 33.41 | 32.81 | 33.12 | 30.02 | 30.25 | Positive |
|  |  | 26.40 | 26.33 | 26.48 | 26.32 | 30.14 | 30.17 | Positive |
|  |  | 31.61 | 31.91 | 30.76 | 30.55 | 27.48 | 27.77 | Positive |
|  | Day 9 | 33.20 | 32.79 | 32.49 | 32.47 | 31.25 | 31.13 | Positive |
|  |  | 26.36 | 26.32 | 26.31 | 26.22 | 30.18 | 29.97 | Positive |
|  |  | 32.42 | 32.37 | 31.90 | 32.18 | 30.49 | 31.06 | Positive |
| 1.0 | Day 1 | 30.16 | 29.81 | 28.38 | 28.66 | 29.77 | 29.32 | Positive |
|  |  | 27.87 | 27.79 | 26.59 | 26.38 | 30.70 | 30.96 | Positive |
|  |  | 28.90 | 29.17 | 27.13 | 27.17 | 30.14 | 30.14 | Positive |
|  | Day 8 | 30.50 | 30.50 | 30.13 | 30.17 | 31.31 | 31.03 | Positive |
|  |  | 30.78 | 30.71 | 29.93 | 29.62 | 31.42 | 31.49 | Positive |
|  |  | 31.09 | 31.02 | 30.34 | 30.43 | 31.59 | 31.31 | Positive |
|  | Day 9 | 27.31 | 27.36 | 26.69 | 26.62 | 29.07 | 29.40 | Positive |
|  |  | 30.50 | 30.38 | 29.63 | 29.57 | 30.59 | 30.79 | Positive |
|  |  | 27.49 | 27.64 | 26.10 | 26.32 | 31.48 | 31.89 | Positive |
| 10 | Day 1 | 22.84 | 22.63 | 21.05 | 20.96 | 31.65 | 31.62 | Positive |
|  |  | 26.13 | 25.76 | 24.04 | 23.86 | 27.20 | 26.94 | Positive |
|  |  | 27.15 | 27.09 | 25.15 | 25.17 | 31.61 | 31.30 | Positive |
|  | Day 8 | 22.43 | 22.28 | 20.74 | 20.58 | 29.79 | 29.40 | Positive |
|  |  | 26.07 | 25.64 | 24.20 | 23.93 | 31.12 | 30.05 | Positive |
|  |  | 26.03 | 25.96 | 24.12 | 24.13 | 30.13 | 30.10 | Positive |
|  | Day 9 | 26.92 | 26.86 | 24.99 | 24.94 | 31.28 | 31.33 | Positive |
|  |  | 26.09 | 26.14 | 24.16 | 24.13 | 30.64 | 31.02 | Positive |
|  |  | 26.67 | 26.52 | 24.75 | 24.66 | 30.61 | 30.27 | Positive |

## Conclusion

Testing these four batches with different PepMV infection levels by the same technician at three different time points resulted in consistent qualitative results and an accordance of $100 \%$. This is above the $90 \%$ that was set as the minimum requirement for this validation criterium. For the repeatability validation criterium, the SE-qPCR detection method is fit for purpose.

### 3.5. Reproducibility

Definition ISHI guidelines: Degree of similarity in results when the method is performed across labs with replicates of the same subsamples.

The reproducibility requirements will be met when there are at least six participating laboratories, and the concordance is $>90 \%$. In addition, the false negative rate should be $5 \%$ or less.

## Experimental approach

A comparative test (CT) was organized in ISHI with eight participating laboratories. Each participant received a total of 24 samples of 1,000 seeds each. The 24 samples consisted of four healthy samples from previously described tomato seed lot variety A (chapter 3.3.), six lowinfected ( $0.5 \%$ ) samples, six medium-infected (1\%) samples and eight highly-infected (10\%) samples. These samples were artificially created, as described in chapter 3.4. One healthy sample was coded as negative process control (NPC), and one highly infected sample was coded as positive process control (PPC). The remaining 22 samples were randomized and coded "A to V". All participants processed the seed samples according to the protocol described in Annex A.

In addition to the work described for reproducibility, the organizing laboratory of the CT had tested prior to the CT nine samples of 1,000 seeds of each of the seed lots with different infection levels used in the CT for homogeneity testing (the repeatability data in Table 7 of section 3.4 was used) and one CT set of 24 samples after receiving the results from all participants for stability testing. These data provided stability and homogeneity data of the samples.

## Statistical analysis

The analysis was done on a qualitative level per sample, viz., positive or negative results for PepMV. The method of Langton et al. (2002) was used to evaluate the concordance (reproducibility of data) of the SE-qPCR.

## Homogeneity results

To determine homogeneity of the samples used for the CT the organizing laboratory tested nine samples of 1,000 seeds of each of the four different PepMV infection rates, healthy ( $0 \%$ ), $0.5 \%$, $1 \%$ and $10 \%$ infected seeds, following the protocol as described in Annex A. The results of these 36 samples are shown in Figure 3. The raw data is shown in Table 7, Section 3.4.


Figure 2. Combined boxplot and violin plot (kernel density distributions) of Cq values obtained by the PepMV triplex SE-qPCR using BaCV as positive extraction control. FAM: NAKT05 assay, VIC: Bejop assay, TxR: BaCV (PEC) assay. *: Average Cq value.

## Comparative test results

The data of the eight participating laboratories were combined and qualitatively shown in Table 8 , and the quantitative data of each laboratory is presented in Annex E.

The CT results of participants 1, 2, 5 and 6 are $100 \%$ in line with the expected results.
Participant 4 did not detect one sample with an infection rate of $0.5 \%$ (sample S). The question is if this participant missed the PepMV in this sample or that they received a sample from this lot without PepMV positive seeds, because the infection rate of the used lot was not $100 \%$ but $96.67 \%$. According to random distribution, the probability of selecting five uninfected seeds from the used seed lot could lead to some of the $0.5 \%$ samples not containing any infected seeds. For the concordance calculations, the outcome of this sample will be seen as a missed PepMV detection (false negative).

Table 8. The qualitative results of all participating laboratories in random anonymous order. NPC: negative process control, PPC: positive process control.

| Infection rate | Sample name |  | Lab 1 | Lab 2 | Lab 3a/3b | Lab 4 | Lab 5 | Lab 6 | Lab ${ }^{\text {a }}$ | Lab 8 ${ }^{\text {a }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 10\% | 1 | A | Pos | Pos | Pos/Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ | Pos |
|  | 4 | D | Pos | Pos | Pos/Pos | Pos ${ }^{\text {b }}$ | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos |
|  | 7 | G | Pos | Pos | Pos/Pos | Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos |
|  | 12 | L | Pos | Pos | Pos/Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ |
|  | 14 | N | Pos ${ }^{\text {b }}$ | Pos | Pos/Pos | Pos ${ }^{\text {b }}$ | Pos | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ |
|  | 20 | T | Pos | Pos | Pos/Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ |
|  | 21 | U | Pos | Pos | Pos/Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ | Pos |
|  | 24 | PPC | Pos | Pos | Pos/Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ |
| 1.0\% | 2 | B | Pos | Pos | Pos/Pos | Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ |
|  | 6 | F | Pos | Pos | Pos/Pos | Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos |
|  | 10 | J | Pos | Pos | Pos/Pos | Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ |
|  | 13 | M | Pos | Pos | Pos/Pos | Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ |
|  | 18 | R | Pos | Pos | Pos/Pos | Pos | Pos | Pos | Failed | Pos ${ }^{\text {b }}$ |
|  | 22 | V | Pos | Pos | Pos/Pos | Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos ${ }^{\text {b }}$ |
| 0.5\% | 5 | E | Pos | Pos | Pos/Pos | Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Pos |
|  | 8 | H | Pos | Pos | Pos/Pos | Pos | Pos | Pos | Pos ${ }^{\text {b }}$ | Neg |
|  | 11 | K | Pos | Pos | Pos/Pos | Pos | Pos | Pos | Failed | Pos ${ }^{\text {b }}$ |
|  | 15 | 0 | Pos | Pos | Pos/Pos | Pos | Pos | Pos | Failed | Failed |
|  | 16 | P | Pos | Pos | Neg/Pos | Pos | Pos | Pos | Failed | Failed |
|  | 19 | S | Pos | Pos | Pos/Pos | Neg | Pos | Pos | Failed | Failed |
| 0\% | 3 | C | Neg | Neg | Neg/Neg | Neg | Neg | Neg | Failed | Neg |
|  | 9 | 1 | Neg | Neg | Neg/Neg | Neg | Neg | Neg | Failed | Neg |
|  | 17 | Q | Neg | Neg | Neg/Neg | Neg | Neg | Neg | Failed | Failed |
|  | 23 | NPC | Neg | Neg | Neg/Neg | Neg | Neg | Neg | Failed | Failed |

a Excluded from further analysis
${ }^{\mathrm{b}}$ Would be a failed test result (no signal or above Cq $\geqslant 33$ for PEC) if it had not been concluded PepMV positive.

Participant 3(a) also missed one sample with an infection rate of $0.5 \%$ (sample $P$ ), but it was also noticed that the Cq values for all samples for both PepMV assays (and not the PEC) were, on average, 6 Cq higher than the results of the other participants. The organizer of the CT contacted participant 3 to determine the cause of this Cq difference in PepMV. After several tests, participant 3 concluded that the results were caused by a specific batch of PepMV primers. After replacing the primers with a new batch and processing the RNA samples again, all their new results were $100 \%$ in line with the organizers expected results (Annex E, 3.5 Reproducibility Data, lab 3). All Cq-values were lowered with ~6 Cq, which caused seed sample $P$ to be detected as positive while it was detected negative in the first run. The change of primer batch did not result in false positive results of the negative seed lot samples C, I, Q and the NPC. Since the cause of the difference in Cq values between participant 3 and the other participants was identified and the problems solved, the results of the repeated experiment from participant 3 (3b) were used for the concordance calculation.

Due to the fact that all samples of participant 7 gave a failed test result, including no signal for PEC and the NPC, they contacted the organizer of the CT. After consultation between the organizer and participant 7, an additional test on their RNA samples was conducted. A selection of the RNA samples was additionally tested with the NAD5 assay (Menzel et al., 2002), which can detect endogenous tomato RNA. The organizing laboratory was able to detect NAD5 with Cq values between 25 and 28 on their own extracted RNA, while participant 7 did not obtain any amplification signal with their RNA. This is an indication that the RNA-extraction of the samples of participant 7 failed and not specifically the qPCR. Although the exact reason for the failure could not be explained, it was decided to exclude participant 7 for further analysis.

Participant 8 also had some issues with the PEC. Although all samples gave a PEC amplification signal, the average PEC value was 33.17 , with some extreme outgroup values of 26.70 and 37.82 . The standard deviation value for participant 8 was 2.44 , while for the other participants ( 1 to 6 ), this value was below 1. Meaning the other laboratories obtained reproducible PEC results with their samples. As participant 8 detected an important number of failed samples, including the NPC, the participant questioned their results and contacted the organizer of the CT. The PepMV Cq values were also 3 to 6 Cq higher as expected. To investigate whether the problem was the RNA-extraction or the qPCR, participant 8 and 2 exchanged RNA of their CT material and repeated the qPCR experiment on this RNA. Participant 2 obtained comparable results on the RNA received from participant 8 as the original results of participant 8 . The PEC values were higher when tested by participant 2. The average value of the PEC was 37.25 instead of the 33.17 , but the standard deviation of all the PEC values was 0.95 , which is a better reproducibility than the original value of 2.44 of participant 8 . When participant 8 tested again the RNA provided by participant 2 , their qPCR results for PepMV were as expected. The PEC-Cq values were 2 to 3 Cq higher than the original data from participant 2, inducing some FAILED results (PEC > Cq 33) for some of the negative samples and the NPC. The fact that participant 2 also had some increased PEC results in this additional experiment suggests that shipment on dry ice negatively affected PEC RNA, while this was not the case for the PepMV RNA. The experiment showed that the qPCR test of participant 8 was not the cause of their failed results for their samples but rather suggested an issue with the extraction of the samples. Therefore, it was decided to exclude participant 8 from the concordance calculations.

Based on the results of participants 1 to 6 (six participants was also the minimum stated in the validation requirements), the concordance of this CT was calculated to be $98.6 \%$. The percentage of false negatives obtained in the $10 \%$ and $1.0 \%$ infection level samples was $0 \%$. The percentage of false negatives in the $0.5 \%$ infection level samples was $2.8 \%$. The percentage of false negatives detected in all expected positive samples was $0.7 \%$. No false positives were found.

## Conclusion

It was stated that the reproducibility requirements will be met when performance criteria (concordance) are $>90 \%$, while the false negative rate remains below $5 \%$. Our results show that the concordance percentage is $98.6 \%$ and the false negative rate is $0.7 \%$, meaning that both requirements for passing the reproducibility criterium were met.

### 3.6. Diagnostic performance

Definition ISHI guidelines: An evaluation of the ability of the method to discriminate between positive and negative seed lots.

As the SE-qPCR assay precedes the bioassay with ELISA confirmation in the method workflow, the diagnostic sensitivity (no false negatives) is more important than the diagnostic specificity (no false positives), as false positives will be detected by the bioassay.

The diagnostic performance requirements will be met when positive samples with the ELISA prescreen assay or the bioassay with ELISA confirmation method (ISF 2021) also yield a positive result with the SE-qPCR. In case of a negative result by bioassay with ELISA confirmation method, a positive result with SE-qPCR is still possible due to non-infectious PepMV presence. The diagnostic performance requirements will therefore be met when diagnostic sensitivity reaches $100 \%$ and the diagnostic specificity is $\geqslant 90 \%$ when comparing the two indirect assays, SE-qPCR and ELISA.

## Experimental approach

Comparative data was generated by one laboratory in which 21 seed lots from different origins, infection levels and production years (Table 10) were tested by SE-qPCR as well as by ELISA assay (Annex F). All suspected lots identified as positive by either SE-qPCR or by ELISA assay were also tested by bioassay (Annex $G$ ). The SE-qPCR results of the diagnostic performance will be based on the SE-qPCR data obtained from an alternative described protocol, meaning pooling $25 \mu \mathrm{~L}$ of four different 250 -seed subsamples resulting in a $100 \mu \mathrm{~L}$ sample for extraction, as described in Annex A paragraph 1.2. By doing so, a comparison of the same seed subsamples between the three methods was possible. The previously described non-infected beef tomato variety A produced in Chile 2021 was used as NPC. The PPC was prepared by grinding naturally infected tomato leaf material in 1 mL of the NPC seed extract.

## Statistical analysis

The analysis was carried out on a qualitative level per sample, viz. positive or negative results for PepMV. The diagnostic sensitivity and diagnostic specificity were calculated according to the formulas presented in Table 9.

Table 9. The formulas used for diagnostic sensitivity and diagnostic specificity calculation.

|  | Validated method result / independent assessment |  |
| :---: | :---: | :---: | :---: |
|  | Positive | Negative |

## Results

The 21 seed lots were tested by ELISA and SE-qPCR and the suspect samples as detected by either ELISA or SE-qPCR were further processed by bioassay. The results of these 21 seed lots are shown in Table 10 and the raw data is provided in Annex E. From the 21 seed lots tested, eight were ELISA positive, nine qPCR positive, and none of the seed lots were bioassay positive. The PPC was ELISA, SE-qPCR and bioassay positive, while the NPC was negative for all three tests. The reason for negative bioassay results for all ELISA and/or qPCR positive seed lots may be the absence of infectious virus particles. Based on these results, including the PPC and NPC, the diagnostic sensitivity and diagnostic specificity were calculated for the SE-qPCR in comparison to the ELISA results. The diagnostic sensitivity was $100 \%$ and the diagnostic specificity was 92.86\%.

Table 10. Diagnostic performance data of the ELISA, qPCR and Bioassay. NPC: negative process control, nt: not tested, PPC: positive process control.

| Crop | Sample code | Production |  | Positive subsamples/total subsamples |  |  | Qualitative conclusion |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Country | Year | ELISA | $\begin{aligned} & \text { SE- } \\ & \text { qPCR } \end{aligned}$ | Bioassay | ELISA | qPCR | Bioassay |
| Rootstock | SHR039 | Chile | 2015 | 0/12 | 3/3 | 0/12 | Neg | Pos | Neg |
| Tomato | SHR126 | Chile | 2003 | 5/12 | 3/3 | 0/12 | Pos | Pos | Neg |
| Tomato | SHR202 | Tanzania | 2005 | 12/12 | 3/3 | 0/12 | Pos | Pos | Neg |
| Rootstock | SHR203 | Chile | 2006 | 12/12 | 3/3 | 0/12 | Pos | Pos | Neg |
| Tomato | SHR218 | Spain | 2018 | 1/12 | 3/3 | 0/12 | Pos | Pos | Neg |
| Tomato | SHR219 | Spain | 2018 | 10/12 | 3/3 | 0/12 | Pos | Pos | Neg |
| Tomato | SHR302a | Netherlands | 2022 | 7/7 | 2/2 | 0/7 | Pos | Pos | Neg |
| Tomato | SHR303 ${ }^{\text {a }}$ | Netherlands | 2022 | 10/10 | 3/3 | 0/10 | Pos | Pos | Neg |
| Tomato | SHR304 ${ }^{\text {a }}$ | Netherlands | 2022 | 7/7 | 2/2 | 0/7 | Pos | Pos | Neg |
| Rootstock | SHR601 | Chile | 2022 | 0/12 | 0/3 | 0/12 | Neg | Neg | nt |
| Rootstock | SHR602 | Chile | 2022 | 0/12 | $0 / 3^{\text {b }}$ | 0/12 | Neg | Neg | nt |
| Tomato | SHR603 | Netherlands | 2020 | 0/12 | 0/3 | 0/12 | Neg | Neg | nt |
| Tomato | SHR604 | Peru | 2019 | 0/12 | 0/3 | 0/12 | Neg | Neg | nt |
| Tomato | SHR605 | Mexico | 2020 | 0/12 | 0/3 | 0/12 | Neg | Neg | nt |
| Tomato | SHR606 | Peru | 2020 | 0/12 | 0/3 | 0/12 | Neg | Neg | nt |
| Tomato | SHR607 | Peru | 2019 | 0/12 | 0/3 | 0/12 | Neg | Neg | nt |
| Tomato | SHR608 | Netherlands | 2021 | 0/12 | 0/3 | 0/12 | Neg | Neg | nt |
| Tomato | SHR609 | Chile | 2021 | 0/12 | 0/3 | 0/12 | Neg | Neg | nt |
| Tomato | SHR610 | Netherlands | 2022 | 0/12 | 0/3 | 0/12 | Neg | Neg | nt |
| Tomato | SHR611 | Mexico | 2022 | 0/12 | 0/3 | 0/12 | Neg | Neg | nt |
| Tomato | SHR612 | Peru | 2021 | 0/12 | 0/3 | 0/12 | Neg | Neg | nt |
| Tomato | PPC |  |  | 4/4 | 4/4 | 4/4 | Pos | Pos | Pos |
| Tomato | NPC |  |  | 0/4 | 0/4 | 0/4 | Neg | Neg | Neg |

${ }^{\text {a }}$ The total amount of available seeds was below 3,000.
${ }^{\text {b }}$ PEC was above Cq 33, namely $>37$ for all three subsamples; therefore, no conclusion can be made for this seed sample.

## Conclusion

It was stated that the diagnostic performance requirements will be met when diagnostic sensitivity is $100 \%$ and the diagnostic specificity is $\geqslant 90 \%$. Our results show that the diagnostic sensitivity reached $100 \%$ and the diagnostic specificity was $92.86 \%$ for qPCR vs. ELISA. This means that both requirements for passing the diagnostic performance criteria are met.

### 3.7. Robustness

## Definition ISHI guidelines:

Robustness is not defined in the ISHI guidelines but was defined in this validation report as the capacity of a method to remain unaffected by small, deliberate variations in method parameters. This allows for the comparison of data in the case of an alternative extraction method as the one described in the protocol validated in this report.

## Experimental approach

Next to the standard protocol using PBS for RNA extraction that allows inoculation of the seed extract on an indicator plant, an additional set of 24 samples were tested using guanidine-based lysis buffer. The same set of samples as used for the CT as described in the reproducibility Section 3.5., was extracted using an alternative extraction method and compared to the quantitative and qualitative CT results of the laboratory performing this experiment. Samples were dry ground in a 50 mL conical tube with a 14 mm bullet inside using a GenoGrinder 2000 at 1300 rpm for 3 min. A volume of 20 mL of a guanidine-based lysis buffer spiked with BaCV was added to each tube of ground seeds and homogenized by vortexing. The seed extracts were incubated for 30 min at room temperature $\left(\sim 20^{\circ} \mathrm{C}\right)$ and afterwards centrifuged at $1,000 \times g$ for 1 min . From each of the tubes, 1 mL seed extract was mixed with $50 \mu \mathrm{~L}$ of 500 mM TCEP solution (catalogue number C4706, Sigma-Aldrich), which was used as an alternative for $\beta$-mercaptoethanol, and incubated for 15 min at $65^{\circ} \mathrm{C}$ and shaken at 850 rpm in a thermomixer. After incubation, $200 \mu \mathrm{~L}$ of this extract was used for RNA extraction using a KingFisher platform following the supplier's protocol for the Sbeadex maxi plant kit. The RNA was eluted in a final volume of $100 \mu \mathrm{~L}$ elution buffer. The RNA samples were tested in two repetitions with the same qPCR mixture and using the protocol as described in Annex A.

## Statistical analysis

The analysis were done on a quantitative and qualitative level per sample, viz. positive or negative result for PepMV.

Results
The results of the two extraction methods executed on two different comparative test sample sets were compared and the average Cq values of the replicates are shown in Table 11. Both methods were able to detect all samples with infected PepMV seeds. Both methods did not result in false positive results.

The samples with the same infection level contain natural variation; therefore, it is expected to observe variation between methods with the same sample name. For the comparison, the average Cq value of all the samples with the same infection rate was calculated and compared. The guanidine KingFisher method reached an average Cq value of $19.22,23.06$ and 24.32 with NAKT05 and $17.85,23.09$ and 24.56 with Bejop assay for the $10 \%, 1 \%$ and $0.5 \%$ infection level, respectively. The method validated in this report using the described seed extraction buffer and Qiagen kit as presented in Annex A obtained an average Cq value of 25.01, 29.26 and 30.63 with NAKT05 and 22.87, 28.61 and 29.52 with Bejop assay for the $10 \%, 1 \%$ and $0.5 \%$ infection level, respectively. This means that the Cq values of the guanidine KingFisher method are 5 to 6 Cq lower with this test set, which correlates to increased sensitivity by 32 to 64 times. The four samples with no infected PepMV seeds obtained for both methods Cq values $>37$ (cut-off value for positive/negative outcome) or a no amplification and are determined PepMV negative.

Table 91. Average PepMV quantitative results of two extraction methods on two comparative test sample sets tested by the same participant. NPC: negative process control, No amp: no amplification, PPC: positive process control.

| Infection rate | Sample name | Average Cq values |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Guanidine KingFisher |  |  | Seed Extraction Buffer Qiagen |  |  |
|  |  | $\begin{gathered} \text { NAKTO } \\ 5 \end{gathered}$ | Bejop | PEC/IAC | NAKT05 | Bejop | PEC/IAC |
| 10\% | A | 22.08 | 20.75 | 28.62 | 25.25 | 23.29 | 31.20 |
|  | D | 21.68 | 20.47 | 28.95 | 26.61 | 24.64 | 30.40 |
|  | G | 16.49 | 15.35 | 29.37 | 26.67 | 24.32 | 31.04 |
|  | L | 15.93 | 14.40 | 29.49 | 22.20 | 20.24 | 32.16 |
|  | N | 17.96 | 16.56 | 29.16 | 25.29 | 23.12 | 31.19 |
|  | T | 15.81 | 14.41 | 29.29 | 25.86 | 23.55 | 30.94 |
|  | U | 21.67 | 20.31 | 29.00 | 23.96 | 21.62 | 31.37 |
|  | PPC | 22.14 | 20.59 | 28.54 | 24.25 | 22.18 | 31.62 |
| 1\% | B | 24.78 | 25.07 | 28.73 | 27.60 | 26.74 | 29.54 |
|  | F | 24.79 | 25.09 | 28.95 | 30.06 | 28.61 | 32.07 |
|  | J | 23.38 | 23.37 | 28.71 | 28.75 | 27.79 | 31.27 |
|  | M | 22.46 | 22.17 | 29.02 | 29.93 | 29.50 | 30.89 |
|  | R | 21.90 | 21.91 | 29.04 | 26.50 | 26.18 | 30.43 |
|  | V | 21.06 | 20.96 | 28.88 | 32.71 | 32.85 | 31.34 |
| 0.50\% | E | 23.74 | 23.55 | 27.59 | 27.90 | 26.89 | 30.76 |
|  | H | 20.40 | 20.58 | 29.02 | 34.95 | 32.72 | 31.65 |
|  | K | 26.94 | 27.41 | 29.09 | 30.28 | 28.99 | 30.88 |
|  | 0 | 27.45 | 27.44 | 28.89 | 29.27 | 28.26 | 30.40 |
|  | P | 25.53 | 25.94 | 28.18 | 29.51 | 28.69 | 31.20 |
|  | S | 21.87 | 22.43 | 28.18 | 31.89 | 31.59 | 30.55 |
| 0\% | C | 38.56 | 37.50 | 29.06 | No amp. | No amp. | 30.23 |
|  | 1 | No amp. | No amp. | 27.89 | No amp. | No amp. | 30.87 |
|  | Q | No amp. | No amp. | 29.57 | No amp. | No amp. | 30.53 |
|  | NPC | No amp. | No amp. | 29.09 | No amp. | No amp. | 30.25 |

## Conclusion

The two compared extraction methods show differences in the quantitative results for the PepMV infected samples in which the guanidine extraction method was $\sim 64$ times more sensitive. The healthy samples raised a negative outcome for both extraction methods. This shows that the increased sensitivity does not lead to an increase in false positive results. Both methods obtained the same qualitative results. All samples expected to be positive were positive and the negative samples negative. The diagnostic performance criterium showed that the SE-qPCR with seed extraction buffer was already more sensitive than ELISA. Therefore, despite the improved sensitivity, there is no substantiated necessity to use the guanidine-based extraction method to prevent false negative results.

## 4. CONCLUSION

The data presented in this validation report shows that the SE-qPCR detection assay for PepMV in samples of tomato seed meets the requirements set for the validation criteria analytical specificity, analytical sensitivity, selectivity, repeatability, reproducibility, and diagnostic performance. It is concluded that the ISHI PepMV SE-qPCR assay is fit for its intended purpose to serve as an optional pre-screening assay for the ISHI PepMV bioassay.

## 5. REFERENCES

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## 6. ANNEXES

## Annex A. Protocol for detecting Pepino mosaic virus (PepMV) in tomato seed by SE-qPCR

## PRE-SCREEN BY SEED EXTRACT RT-qPCR

For PCR methods, in-house method optimization is often necessary, see Best Practices for PCR Assays in Seed Health Tests for information.

## Materials

- Seed extraction buffer (Table A.1)
- Grinder
- RT-qPCR mix, primers (Table A.2) and equipment
- Controls (Table A.3)
- Spike solution (see description below)
- RNA purification kit and equipment
- 1.5 mL RNase Free tube
- RNAse free water
- Centrifuge


## Sample Size

The recommended minimum sample size is 3,000 seeds with a maximum subsample size of 1,000 seeds.

## Spike solution

The spike solution is prepared by taking a leaf from a plant infected by Bacopa chlorosis virus ( BaCV ) and making an extract of it in seed extraction buffer (Table A1). The extract is diluted to obtain a suitable concentration usable as positive extraction control (PEC) and aliquots are stored at $-80^{\circ} \mathrm{C}$.

Note: Other organisms such as Dahlia latent viroid (DLVd) and Squash mosaic virus (SqMV) may also be used and should be shown to be compatible with the PepMV primers in a multiplex PCR.

Table A.1. Seed extraction buffer.

| Compound | Amount/L |
| :--- | :---: |
| NaCl | 8.0 g |
| $\mathrm{Na}_{2} \mathrm{HPO}_{4} .7 \mathrm{H}_{2} \mathrm{O}$ | 2.17 g |
| $\mathrm{KH}_{2} \mathrm{PO}_{4}$ | 0.2 g |
| KCl | 0.2 g |
| $\mathrm{Na}_{2} \mathrm{SO}_{3}{ }^{\mathrm{a}}$ | 1.0 g |

${ }^{a}$ Add after autoclaving. Use within 24 hours of preparation.
Note: If a different seed extraction buffer is used, it must be verified in a comparison using uniform positive control material. The use of a different seed extraction buffer can lead to a different limit of detection and could result in increased number of false positive or false negative outcomes.

Table A.2. Primer and probe sequences and references.

| Name | Sequence ( $5^{\prime}$ - $3^{\prime}$ ) | Source |
| :---: | :---: | :---: |
| Bejop 133 | ATC AAT TGT CCT TAT GCG CT | Bejo Zaden B.V., Netherlands |
| Bejop 134 | ATC AAT TGT CCT TAC GCG CT |  |
| Bejop 135 | ATC AAT TGT CCC TAT GCG CT |  |
| Bejop 136 | ATC AAC TGT CCT TAT GCG CT |  |
| Bejop 137 | VIC - ACA CCC TTG AGA ATC TTG GTG TCA CAA T - BHQ1 |  |
| Bejop 138 | VIC - ACA CCC TTG AGA ATT TAG GTG TCA CAA T - BHQ1 |  |
| Bejop 139 | GTT TGA ATT GCA TGA GGG TT |  |
| Bejop 140 | TTG GAT TGC ATG GGG RTT |  |
| KL05_48 | ACT CCT AGA GCT GAC CTC AC | Ling et al., 2007 |
| KL05_49 | ACT CCT AGA GCT GAT CTT AC |  |
| KL05_51 | TCT CCA GCA ACA GGT TGG TA |  |
| KL05_52 | TCA CCT GCA ACT GGT TGA TA |  |
| NAKT05_50 | 6FAM - TGT CAG CTT GCA TTT ACT TC - MGB NFQ | Naktuinbouw ${ }^{\text {a }}$, Netherlands |
| NEC BaCV F | CGA TGG GAA TTC ACT TTC GT | Naktuinbouw, Netherlands |
| NEC BaCV R | AAT CCA CAT CGC ACA CAA GA |  |
| NEC BaCV-P | TxR - CAA TCC TCA CAT GAT GAG ATG CCG - BHQ2 |  |

${ }^{a}$ Adapted from Ling et al., 2007.
Table A.3. Types of controls used.

| Control type | Description |
| :--- | :--- |
| Negative process control (NPC) | Tomato seed-free of PepMV |
| Positive process control (PPC) | Tomato seed with infectious PepMV |
| Positive amplification control (PAC) | PepMV RNA aiming for a Cq (Cycle quantification) value <br> between 28 and 32 or |
|  | PepMV oligo DNA (oligonucleotide (single-stranded DNA) for <br> all PepMV target sequences) aiming for a Cq value between <br> 28 and 32 or |
|  | PepMV cDNA aiming for a Cq value between 28 and 32 |
| Positive extraction control (PEC) | Spike solution added to the simple aiming for a Cq value <br> between 28 and 32 <br> The PEC serves as an internal amplification control (IAC) |
| Inhibition control (IC) | Dilution of the PEC in a non-infected seed extract aiming for <br> a Cq value between 28 and 32 <br> Note: a non-infected seed extract is preferred over a seed <br> extraction buffers, as a strong diluted infected leaf extract <br> may lead to a relatively high loss of RNA in the purification <br> process (i.e., no carrier RNA present) |
| Negative template control (NTC) | Contains all PCR reagents but no target or spike DNA, RNA <br> or PEC nucleic acids |

## 1. Seed extraction

1.1. Add the positive extraction control (PEC) to the seed extraction buffer (Table A.1).
1.2. Soak the 1,000 seeds of each subsample, the positive process control (PPC), and the negative process control (NPC) in 40 mL seed extraction buffer containing the PEC for 30 to 60 min .

Macerate the samples for 90 sec using an Interscience BagMixer or equivalent equipment. See Figure A. 1 for before and after macerating.

Notes: In case of a positive result when using this option, a fresh seed extract from a new sample of 3,000 seeds in subsamples of 250 seeds is required for the bioassay.

Alternatively, grind 12 subsamples of 250 seeds directly in 10 mL seed extraction buffer containing the PEC.
1.3. Process extracts immediately after grinding, place on ice for up to 1 hour or store at $-20^{\circ} \mathrm{C}$ for one day. Do not freeze if the extracts are to be used for the bioassay after RT-qPCR, but store at 4 to $7^{\circ} \mathrm{C}$. The bioassay must be completed within 20 hours after extraction. Store the PPC and NPC (Table A.3) under similar conditions as the samples to validate the results.


Figure A.1. A. 1,000 tomato seeds in seed extraction buffer after 30 min soaking. B. 1,000 tomato seeds after 90 sec of macerating with BagMixer.

## 2. RNA extraction

2.1. In case of 1,000 seed subsamples, use $100 \mu \mathrm{~L}$ from each subsample for further analysis.

In the case of 250 seed subsamples, combine $25 \mu \mathrm{~L}$ of four seed extract subsamples into a $100 \mu \mathrm{~L}$ combined sample. Use all three combined samples for further analysis.
2.2. Start RNA isolation within 1 hour after grinding.
2.3. Use the commercial RNA isolation kit RNeasy plant mini kit (Qiagen) for RNA isolation. Process the subsamples according to the supplier's instructions.
2.4. Eluate the RNA in $100 \mu \mathrm{~L}$ elution buffer.

## 3. Preparation of the RT-qPCR

3.1. Prepare the RT-qPCR mixture as indicated in Table A.4.

For each run, include a negative template control (NTC) and at least one positive amplification control (PAC) that give a Cq value between 28 and 32 .
3.2. Perform the PCR reaction in a real-time PCR instrument according to the PCR conditions mentioned in Table A.5. All samples and controls should be tested in duplicate as described in Best Practices for PCR Assays in Seed Health Tests.

Note: Good results have been obtained by ISHI member laboratories with the UltraPlex ${ }^{\text {TM }}$ 1Step ToughMix (QuantaBio). If different qPCR mixtures and amplification programs are used, it is necessary to verify their performance.

Table A.4. RT-qPCR PepMV mix.

| Component | Per reaction (in $\mu \mathrm{L})$ | Final concentration |
| :--- | :---: | :---: |
| Bejop 133 $(10 \mu \mathrm{M})$ | 0.25 | $0.1 \mu \mathrm{M}$ |
| Bejop 134 $(10 \mu \mathrm{M})$ | 0.25 | $0.1 \mu \mathrm{M}$ |
| Bejop 135 $(10 \mu \mathrm{M})$ | 0.25 | $0.1 \mu \mathrm{M}$ |
| Bejop 136 $(10 \mu \mathrm{M})$ | 0.25 | $0.1 \mu \mathrm{M}$ |
| Bejop 137 $(10 \mu \mathrm{M})$ | 0.50 | $0.2 \mu \mathrm{M}$ |
| Bejop 138 $(10 \mu \mathrm{M})$ | 0.50 | $0.2 \mu \mathrm{M}$ |
| Bejop 139 $(10 \mu \mathrm{M})$ | 1.50 | $0.6 \mu \mathrm{M}$ |
| Bejop 140 $(10 \mu \mathrm{M})$ | 1.50 | $0.6 \mu \mathrm{M}$ |
| KL05_48 $(10 \mu \mathrm{M})$ | 0.50 | $0.2 \mu \mathrm{M}$ |
| KL05_49 $(10 \mu \mathrm{M})$ | 0.50 | $0.2 \mu \mathrm{M}$ |
| NAKT05_50 $(10 \mu \mathrm{M})$ | 0.50 | $0.2 \mu \mathrm{M}$ |
| KL05_51 $(10 \mu \mathrm{M})$ | 0.50 | $0.2 \mu \mathrm{M}$ |
| KL05_52 $(10 \mu \mathrm{M})$ | 0.50 | $0.2 \mu \mathrm{M}$ |
| NEC BaCV F $(10 \mu \mathrm{M})$ | 0.3125 | $0.125 \mu \mathrm{M}$ |
| NEC BaCV R $(10 \mu \mathrm{M})$ | 0.3125 | $0.125 \mu \mathrm{M}$ |
| NEC BaCV-Probe $(10 \mu \mathrm{M})$ | 0.156 | $0.0625 \mu \mathrm{M}$ |
| Ultraplex 1-step Toughmix $(4 \times)$ | 6.25 | $1 \times$ |
| RNAse free water | 5.469 |  |
| Template RNA | 25.00 |  |
| Total |  |  |

Table A.5. RT-qPCR conditions.

| Step | Temperature | Duration |
| :--- | :--- | :--- |
| RT reaction | $50^{\circ} \mathrm{C}$ | 10 min |
| Denaturation | $95^{\circ} \mathrm{C}$ | 3 min |
| 39 cycles | $95^{\circ} \mathrm{C}$ | 10 sec |
|  | $60^{\circ} \mathrm{C}$ | 60 sec |

## 4. Interpretation and decisions

Determine the cut-off values. Cut-off values must be established by each laboratory for their positive and internal amplification controls prior to the assay being used on routine samples. For recommendations on setting cut-off values, see Real-time PCR pre-screening in seed health methods.

In the validation study a SE-qPCR result was regarded positive with a $C q$ value of $\leqslant 37$, and negative with a Cq value > 37 .

For interpretation and decision making, the results from all primer sets need to be taken into account, see Table A.6. Test results are only valid when all included controls presented in Table A. 3 give the expected results.

Table A.6. Interpretation and decision table for the SE-qPCR.

| Bejop | NAKT_05 | BaCV (PEC) | qPCR Result | Follow-up |
| :--- | :--- | :--- | :--- | :--- |
| Positive | Positive | Positive or <br> Negative | Target RNA for PepMV <br> detected | Bioassay for confirmation |
| Negative | Negative | Positive | No target RNA for PepMV <br> detected | Negative, no follow up |
| Negative | Negative | Negative | PEC/IAC failure | Repeat Extraction and/or <br> RT-qPCR. In case of <br> repeatable results, no <br> conclusion can be given for <br> this sample by PCR. <br> Continue with ELISA and/or <br> Bioassay |
| Positive | Negative | Positive or <br> Negative | Target RNA for PepMV <br> detected | Bioassay for confirmation |
| Negative | Positive | Positive or <br> Negative | Target RNA for PepMV <br> detected | Bioassay for confirmation |

## Annex B. Analytical specificity data from in-silico data analysis

Table B.1. Rate of identity of a representative selection of PepMV strains from NCBI with the ISHI PepMV SE-qPCR primers and probes.

| References ID | Isolate Name | Origin | Collection date | Host | Bejop |  |  | NAKT05 |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  | Fw | Pr | Rv | Fw | Pr | Rv |
| NC004067 | Sp-13 | Spain | <2002 | Solanum lycopersicum | 100 | 100 | 100 | 100 | 100 | 100 |
| AY509926 | US1 | USA | <2004 | Solanum lycopersicum | 100 | 100 | 100 | 100 | 100 | 100 |
| AY509927 | US2 | USA | <2004 | Solanum lycopersicum | 100 | 100 | 100 | 100 | 100 | 100 |
| DQ000985 | Ch2 | Chili | 2003 | Solanum lycopersicum | 100 | 100 | 100 | 100 | 100 | 100 |
| HQ663890 | SAR09 | Italy | 2009 | Solanum lycopersicum | 95 | 100 | 100 | 100 | 100 | 100 |
| MK133092 | PepMV-P12-3G | Germany | 2018 | Solanum lycopersicum | 100 | 96.4 | 100 | 100 | 100 | 100 |
| HQ650560 | P22 | Poland | 2005 | Solanum lycopersicum | 100 | 96.4 | 100 | 100 | 100 | 100 |
| MN549397 | Ca1A | Canada | 2019 | Solanum lycopersicum | 100 | 100 | 95 | 100 | 100 | 100 |
| HG976946 | CAM1-IT | Italy | 2010 | Solanum lycopersicum | 100 | 100 | 100 | 95 | 100 | 100 |
| MN038407 | MZ2 | Spain | 2014 | Solanum lycopersicum | 100 | 100 | 100 | 100 | 90 | 100 |
| JQ314461 | EU_EF09_60 | USA | 2009 | Solanum lycopersicum | 100 | 100 | 100 | 100 | 100 | 95 |
| HG313805 | Chi2.9 | Peru | 2008 | Solanum pimpinellifolium | 100 | 100 | 100 | 95 | 100 | 90 |
| HG313806 | Tor9 | Peru | 2008 | Solanum peruvianum | 100 | 100 | 100 | 95 | 100 | 90 |
| HG313807 | Yur1.5 | Peru | 2008 | Solanum peruvianum | 100 | 96.4 | 100 | 95 | 100 | 90 |
| MN038405 | MZ1 | Spain | 2014 | Solanum lycopersicum | 95 | 100 | 100 | 100 | 90 | 100 |
| HQ663891 | SIC1-09 | Italy | 2009 | Solanum lycopersicum | 95 | 100 | 100 | 95 | 100 | 100 |
| AJ606361 | LP-2001 | Peru | 2000 | Solanum peruvianum | 100 | 100 | 94.4 | 100 | 95 | 100 |
| AM109896 | SM74 | Peru | 1974 | Solanum muricatum | 100 | 100 | 94.4 | 100 | 95 | 100 |
| MF422612 | LP_BP0161 | Switzerland | 2014 | Solanum lycopersicum | 100 | 100 | 94.4 | 100 | 95 | 100 |

## Annex C. Sequences of the synthetic ssDNA PepMV oligonucleotides

PC_Bejop
CTTAAAGAATCTCTAATCAATTGTCCTTACGCGCTTAATGATTATGAAGCAGACACCCTTGAGAATCTTGGTGTCACAATTAACCCCCATGCAATCCAAACACACACACATG CCGC

PC_NAKT05
TGCCAGGTCTAACTCCTAGAGCTGACCTCACTGACACATACAAAATCATTGCCATTGCTTTCTTGTTGTCAGCTTGCATTTACTTCCAAAATAGCCACTACCAACCTGTTGC TGGAGACAACTTGCACCGT

PC_NAKT05m
TGCCAGGGCTGACCCCTAGAGCTGACCTCACTGACACATACAAGATAATTGCAATTGCCTTCCTGTTGTCAGCCTGCATTTACTTCCAGAACAGCCACTACCAACCAGTTGC TGGGGACAACTTGCATAGA

Annex D. Analytical specificity data from leaf and/or seed samples
Table D.1. Analytical specificity data from leaf and/or seed samples from various origins infected with PepMV. No amp.: no amplification, Rep.: replicate.

| Isolate name | Genotype ${ }^{\text {b }}$ | Origin | Year | Tissue type | Host | NAKTO5 |  | Bejop |  | BaCV |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  | Rep. 1 | Rep. 2 | Rep. 1 | Rep. 2 | Rep. 1 | Rep. 2 |
| 0148-10 | EU | Netherlands | 2010 | Leaf | Solanum lycopersicum | 20.09 | 20.17 | 19.47 | 19.56 | No amp. | No amp. |
| 0149-10 | LP | Netherlands | 2010 | Leaf | Solanum lycopersicum | 27.04 | 27.03 | N/A | N/A | No amp. | No amp. |
| 0004-11 | LP | Netherlands | 2011 | Leaf | Solanum lycopersicum | 15.39 | 15.33 | 14.79 | 14.74 | No amp. | No amp. |
| 0709-11 | CH2 | Netherlands | 2011 | Leaf | Solanum lycopersicum | 13.17 | 13.16 | 12.45 | 12.26 | No amp. | No amp. |
| 0776-11 | CH2 | Spain | 2011 | Leaf | Solanum lycopersicum | 15.07 | 14.90 | 13.98 | 13.96 | No amp. | No amp. |
| 0773-14 | CH2 | Hungary | 2014 | Leaf | Solanum lycopersicum | 24.81 | 24.97 | 23.37 | 23.46 | No amp. | No amp. |
| 0127-15 | CH2 | Italy | 2015 | Leaf | Solanum lycopersicum | 21.17 | 21.50 | 19.56 | 20.01 | No amp. | No amp. |
| 0615-16 | CH2 | Canada | 2016 | Leaf | Solanum lycopersicum | 17.89 | 17.67 | 17.10 | 17.24 | No amp. | No amp. |
| 0647-16 | CH2 | Chili | 2016 | Leaf | Solanum lycopersicum | 17.56 | 17.29 | 17.27 | 17.20 | No amp. | No amp. |
| 0245-17 | CH2 | South Africa | 2017 | Leaf | Solanum lycopersicum | 15.01 | 14.91 | 14.98 | 14.88 | No amp. | No amp. |
| 0938-17 | CH 2 and LP | UK | 2017 | Fruit | Solanum lycopersicum | 13.22 | 13.13 | 12.82 | 12.74 | No amp. | No amp. |
| 1340-18 | CH2 | Slovakia | 2018 | Leaf | Solanum lycopersicum | 12.87 | 12.89 | 12.08 | 11.99 | No amp. | No amp. |
| 7739-20 | LP | Brazil | 2020 | Leaf | Solanum lycopersicum | 27.75 | 27.81 | 27.30 | 27.33 | No amp. | No amp. |
| 8319-21 | CH2 | France | 2021 | Leaf | Solanum lycopersicum | 10.35 | 10.63 | 9.16 | 9.03 | No amp. | No amp. |
| 8564-21 | CH2 | New Zealand | 2021 | Leaf | Solanum lycopersicum | 21.15 | 21.26 | 19.29 | 19.34 | No amp. | No amp. |
| 8934-21 | US1 | Canada | 2021 | Leaf | Solanum lycopersicum | 15.15 | 15.36 | 13.73 | 13.70 | No amp. | No amp. |
| 9214-21 | CH 2 and LP | Poland | 2021 | Leaf | Solanum lycopersicum | 11.74 | 11.85 | 11.04 | 11.06 | No amp. | No amp. |
| DB1-10 | EU | Netherlands | 2010 | Leaf | Solanum lycopersicum | 19.51 | 19.57 | 18.05 | 18.12 | No amp. | No amp. |
| SHR030 | US1 | Chili | 2015 | Seed | Solanum lycopersicum | 26.47 | 26.43 | 24.84 | 24.82 | No amp. | No amp. |
| SHR125 | US1 | Chili | 2006 | Seed | Solanum lycopersicum | 28.15 | 28.34 | 26.44 | 26.63 | No amp. | No amp. |
| SHR126 | US1 | Chili | 2003 | Seed | Solanum lycopersicum | 23.84 | 23.46 | 19.33 | 19.12 | No amp. | No amp. |
| U287-21 | CH2 | Unknown | 2021 | Seed | Solanum lycopersicum | 27.67 | 27.98 | 27.03 | 27.08 | No amp. | No amp. |
| Tor-9a | PES | Peru | 2008 | Leaf | Solanum lycopersicum | 22.79 | 22.88 | 20.12 | 20.20 | No amp. | No amp. |
| CHI 2.9a | PES | Peru | 2008 | Leaf | Solanum lycopersicum | 21.72 | 21.65 | 18.76 | 18.69 | No amp. | No amp. |
| YUR 1.5 ${ }^{\text {a }}$ | PES ${ }^{\text {c }}$ | Peru | 2008 | Leaf | Solanum lycopersicum | 24.46 | 24.32 | No amp. | No amp. | No amp. | No amp. |

${ }^{a}$ RNA provided by Dr. Fernando García-Arenal Rodríguez (Plant-Virus Interaction and Co-evolution Research Group at Centro de Biotecnología y Genómica de Plantas UPM-INIA, Spain).
${ }^{\mathrm{b}}$ Determined based on Sanger sequence data comparison of nucleotide 3,441 to 3,761 of PepMV reference seq NC004067 (Pep5-2: 5'-TACYTAGCHAAACCAATGGC-3' and Pep6-2: 5'-GCWCCATCTTGKGATTGRTC-3').
${ }^{\text {c }}$ Genotype determined based on literature (Accession HG313807); no Sanger sequence data available, because sample was gel-PCR negative.

Table D.2. Specificity data from healthy leaf and/or seed samples from various origins or infected with a non-target virus/viroid. N/A: not applicable, No amp.: no amplification.

| Isolate name | Supplier | Target | Origin | Year | Tissue type | Host | NAKT05 | Bejop | BaCV |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  | Rep. 1\&2 | Rep. <br> 1\&2 | Rep. 1\&2 |
| 0710-09 | Field isolate | Potato virus $X$ | Australia | 2009 | Leaf | Capsicum annuum | No amp. | No amp. | No amp. |
| ToRSV | Prime diagnostics | Tomato ring spot virus | Unknown | Unknown | Leaf | Solanum lycopersicum | No amp. | No amp. | No amp. |
| PV-0215 | DSMZ | Arabic mosaic virus | Unknown | 2021 | Leaf | Chenopodium quinoa | No amp. | No amp. | No amp. |
| PV-0070 | DSMZ | Tomato black ring virus | Unknown | 1987 | Leaf | Solanum tuberosum | No amp. | No amp. | No amp. |
| TRSV | Prime diagnostics | Tobacco ringspot virus | Unknown | 2021 | Leaf | Solanum lycopersicum | No amp. | No amp. | No amp. |
| TMV | Field isolate | Tobacco mosaic virus | Unknown | 2021 | Leaf | Nicotiana tabacum | No amp. | No amp. | No amp. |
| 0406-05 | Field isolate | Tomato brown rugose fruit virus | Jordan | 2005 | Leaf | Solanum lycopersicum | No amp. | No amp. | No amp. |
| CMV | Prime diagnostics | Cucumber mosaic virus | Unknown | 2021 | Leaf | Capsicum annuum | No amp. | No amp. | No amp. |
| TSWV | Prime diagnostics | Tomato spotted wilt virus | Unknown | 2021 | Leaf | Solanum lycopersicum | No amp./37.83 | No amp. | No amp. |
| MX5 | USDA | Tomato mottle mosaic virus | Mexico | 2009 | Leaf | Solanum lycopersicum | 37.78/No amp. | No amp. | No amp. |
| PV-0196 | DSMZ | Alfalfa mosaic virus | Unknown | 1988 | Leaf | Solanum lycopersicum | No amp. | No amp. | No amp. |
| BaCV | Field isolate | Bacopa chlorosis virus | Netherlands | 2016 | Leaf | Sutera cordata | No amp. | No amp. | 28.15/28.44 |
| DLVd | Field isolate | Dahlia latent viroid | Netherlands | 2020 | Leaf | Dahlia sp. | No amp. | No amp. | No amp. |
| 0088-17 | Field isolate | Tomato mosaic virus | Netherlands | 2017 | Leaf | Solanum lycopersicum | No amp. | No amp. | No amp. |
| TCDVd | NVWA | Tomato chlorotic dwarf viroid | Unknown | 2014 | Leaf | Solanum lycopersicum | No amp. | No amp. | No amp. |
| CEVd | NVWA | Citrus exocortis viroid | Unknown | 2014 | Leaf | Solanum lycopersicum | No amp. | No amp. | No amp. |
| CLVd | NVWA | Columnea latent viroid | Unknown | 2014 | Leaf | Solanum lycopersicum | No amp. | No amp. | No amp. |
| TPMVd | NVWA | Tomato planta macho viroid | Unknown | 2014 | Leaf | Solanum lycopersicum | No amp./38.73 | No amp. | No amp. |
| Tomato | Rijk Zwaan | Solanum lycopersicum | Netherlands | 2021 | Leaf | Solanum lycopersicum | No amp. | No amp. | No amp. |
| Tomato | Rijk Zwaan | Solanum lycopersicum | Chile | 2021 | Seed | Solanum lycopersicum | No amp. | No amp. | No amp. |
| Tomato | Rijk Zwaan | Solanum lycopersicum | Mexico | 2020 | Seed | Solanum lycopersicum | No amp. | No amp. | No amp. |
| Eggplant | Rijk Zwaan | Solanum melongena | Netherlands | 2021 | Leaf | Solanum melongena | No amp. | No amp. | No amp. |
| Water | IDT | no virus | N/A | 2021 | N/A | N/A | No amp. | No amp. | No amp. |

## Annex E. Supplementary data validation report


3.2 Analytical

Sensitivity Limit of $D$

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3.2 Analytical

Sensitivity Single Se

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3.3 Selectivity

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3.4 Repeatability

Data.xlsx

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3.5 Reproducibility Data.xlsx

## 8

3.6 Diagnostic
performance Data.xl

## Annex F. Protocol for detecting Pepino mosaic virus (PepMV) in tomato seed by ELISA

## Materials

- Seed extraction buffer (Table F.1)
- Antiserum PepMV (Prime Diagnostics)
- Phosphate substrate
- ELISA buffers (Table F.2, F. 3 and F.4)
- Controls (Table F.5)

Table F.1. Seed extraction buffer.

| Compound | Amount/L |
| :--- | :---: |
| Sodium chloride $(\mathrm{NaCl})$ | 8.0 g |
| Sodium phosphate dibasic heptahydrate $\left(\mathrm{Na}_{2} \mathrm{HPO}_{4} .7 \mathrm{H}_{2} \mathrm{O}\right)$ | 2.17 g |
| Potassium dihydrogen phosphate $\left(\mathrm{KH}_{2} \mathrm{PO}_{4}\right)$ | 0.2 g |
| Potassium chloride $(\mathrm{KCl})$ | 0.2 g |
| Sodium sulphite $\left(\mathrm{Na}_{2} \mathrm{SO}_{3}\right)^{a}$ | 1.0 g |

${ }^{a}$ Add after autoclaving, use within 24 hours after preparation.
Note: If a different seed extraction buffer is used, it must be verified in a comparison using uniform positive control material that it does not lead to a different positive / negative outcome.

Table F.2. Coating buffer - pH 9.6.

| Compound | Amount/L |
| :--- | :---: |
| Sodium carbonate $\left(\mathrm{Na}_{2} \mathrm{CO}_{3}\right)$ | 1.59 g |
| Sodium bicarbonate $\left(\mathrm{NaHCO}_{3}\right)$ |  |
| Add de-ionized water up to 1 L , adjust pH and autoclave buffer at $121{ }^{\circ} \mathrm{C}$ for 15 min.$$ |  |

Table F.3. Conjugate buffer - pH 7.4.

| Compound | Amount/L |
| :--- | :---: |
| Sodium chloride (NaCl) | 8.0 g |
| Sodium phosphate dibasic heptahydrate $\left(\mathrm{Na}_{2} \mathrm{HPO}_{4 .} .7 \mathrm{H}_{2} \mathrm{O}\right)$ | 10.85 g |
| Potassium dihydrogen phosphate $\left(\mathrm{KH}_{2} \mathrm{PO}_{4}\right)$ | 1.0 g |
| Tween 20 | 0.5 mL |
| Add de-ionized water up to 1 L , adjust pH and autoclave buffer at $121^{\circ} \mathrm{C}$ for 15 min. |  |
| Bovine Serum Albumin ${ }^{\text {a }}$ | 8.0 g |

${ }^{\text {a }}$ Add after autoclaving, use within 24 hours after preparation.
Table F.4. Substrate buffer - pH 9.6.

| Compound | Amount/L |
| :--- | :---: |
| Diethanolamine | 97 mL |

Table F.5. Types of controls used.

| Control type | Description |
| :--- | :--- |
| Buffer control (BC) | The buffers and reagents used in the ELISA, with no seed/tissue <br> matrix or target pathogen |
| Negative process control (NPC) | Tomato seed-free of Pepino mosaic virus |
| Positive process control (PPC) | Positive matrix that contains the target pathogen and is tested <br> at the same time, using the same assay as the corresponding <br> samples |

## 1. General requirements

- Seed extracts and controls must be prepared at the same time, under the same laboratory conditions and stored under the same conditions.
- Seed extracts and all controls must be stored at 4 to $7{ }^{\circ} \mathrm{C}$ until the assay begins. It is recommended to perform the bioassays within 20 hours following seed extraction.


## 2. Preparation ELISA test

2.1. Coat the ELISA plates with coating antibodies (e.g., PepMV Prime diagnostics) in coating buffer (Table F.2).

Note: If different antisera and buffers are used, or even different lot numbers, it is necessary to verify their performance.
2.2. Store the coated plate $\sim 7^{\circ} \mathrm{C}$ overnight to maximum 24 hours before use.

## 3. Extraction of the virus from the seed

3.1. Soak the 250 seeds of each subsample, the positive process control (PPC), and the negative process control (NPC) in 10 mL seed extraction buffer (Table F.1) for 30 to 60 min .
3.2. Macerate the samples for 90 sec using an Interscience BagMixer or equivalent equipment.
3.3. Store seed extracts at $4-7^{\circ} \mathrm{C}$.

## 4. ELISA test

4.1. Clear out the coating buffer from the ELISA plate.
4.2. Put the seed extract of each grinded sample in duplicate in the plate.
4.3. Store the ELISA plate with samples overnight at $\sim 7^{\circ} \mathrm{C}$.
4.4. Wash out the samples from the ELISA plate.
4.5. Conjugate the ELISA plate with the conjugate with the conjugate antibodies (e.g., PepMV Prime diagnostics) in conjugate buffer (Table F.3.).
4.6. Store the conjugate plate for 4 hours at $28^{\circ} \mathrm{C}$.
4.7. Dissolve 10 mg of phosphatase substrate per 10 mL substrate buffer (Table F.4.).
4.8. Wash out the conjugate buffer from the ELISA plate.
4.9. Add the substrate buffer with phosphatase substrate to the ELISA plate.
4.10. Measure after 2 hours the ELISA plate extinction using a spectrophotometer at 405 nm and 620 nm .

## 5. Evaluation of the test results

5.1. To calculate the signal-to-noise $(S / N)$ ratio for an ELISA, divide the average PPC optical density (OD) value by the average NPC OD value.
5.2. Cut-off determination: use the vendor's recommended cut-off (e.g., an S/N ratio (TC:NPC) of 2). Alternatively, use the average NPC OD value plus three times the standard deviation of the NPC OD values. All samples with average OD values at or above the cut-off are considered positive.
6. Validity of test results
6.1. It is the responsibility of the user that an ELISA have a $S / N$ ratio of at least $10: 1$ to ensure sufficient separation between positive and negative results, see ELISA Development Guide (https://resources.rndsystems.com/pdfs/datasheets/edbapril02.pdf)

Note: Test results are only valid when all included controls presented in Table F. 5 give the expected results.

## Annex G. Protocol for detecting Pepino mosaic virus (PepMV) in tomato seed by bioassay

## Materials

- Seed extraction buffer (Table G.1)
- Controls (Table G.2)
- Grinder
- Nicotiana benthamiana plants
- Carborundum
- Protective mask
- Gloves
- Tap water
- ELISA necessities
- Lab disposals


## General notes

- Nicotiana benthamiana is a systemic host for all PepMV strains tested. N. benthamiana is preferred over tomato as an assay plant because the systemic movement of the virus in tomato can be erratic. Furthermore, leaves of $N$. benthamiana are more easily inoculated.
- Although PepMV infection of $N$. benthamiana usually results in conspicuous symptoms, this is not always the case, and symptoms can be caused by other factors than PepMV. Therefore, ELISA of assay plants is required.

Table G.1. Seed extraction buffer.

| Compound | Amount/L |
| :--- | :---: |
| Sodium chloride $(\mathrm{NaCl})$ | 8.0 g |
| Sodium phosphate dibasic dodecahydrate $\left(\mathrm{Na}_{2} \mathrm{HPO}_{4} .12 \mathrm{H}_{2} \mathrm{O}\right)$ | 2.9 g |
| Potassium dihydrogen phosphate $\left(\mathrm{KH}_{2} \mathrm{PO}_{4}\right)$ | 0.2 g |
| Potassium chloride $(\mathrm{KCl})$ | 0.2 g |
| Sodium sulphite $\left(\mathrm{Na}_{2} \mathrm{SO}_{3}\right)^{a}$ | 1.0 g |

${ }^{\text {a }}$ Add after autoclaving. use within 24 hours after preparation.
Note: If a different seed extraction buffer is used, it must be verified in a comparison using uniform positive control material that it does not lead to change in the outcome.

Table G.2. Types of controls used.

| Control type | Description |
| :--- | :--- |
| Buffer control (BC) | The buffers and reagents used in the ELISA, with no seed/tissue <br> matrix or target pathogen |
| Negative process control (NPC) | Tomato seed-free of Pepino mosaic virus |
| Positive process control (PPC) | Positive matrix that contains the target pathogen and is tested <br> at the same time, using the same assay as the corresponding <br> samples |

## 1. General requirements

- Seed extracts and controls must be prepared at the same time, under the same laboratory conditions and stored under the same conditions.
- $\quad$ Seed extracts and all controls must be stored at 4 to $7{ }^{\circ} \mathrm{C}$ until the assay begins. It is recommended to perform the bioassays within 20 hours following seed extraction.


## 2. Seed extraction

2.1. Soak the 250 seeds of each subsample, the positive process control (PPC), and the negative process control (NPC) in 10 mL seed extraction buffer (Table G.1.) for 30 to 60 min .
2.2. Macerate the samples for 90 sec using an Interscience BagMixer or equivalent equipment.
2.3. Store seed extracts at $4-7^{\circ} \mathrm{C}$.

## 3. Inoculation of Nicotiana benthamiana plants

Assay plants should have 4-7 (nearly) fully expanded leaves and should been raised under sufficient light intensity at an average temperature of $20-25{ }^{\circ} \mathrm{C}$. Avoid older assay plants that start to flower. Assay plants should have good turgor at the time of inoculation (Figure G.1).
3.1. Inoculate each seed extract, including the PPC, and NPC, on the two youngest (nearly) fully expanded leaves of two plants, going across the whole surface. Do not use the primary leaf (oldest true leaf).

Inoculate by first dusting the leaves moderately with carborundum (320 mesh grit powder, Fisher scientific or equivalent), while wearing a protective mask, followed by placing a drop of inoculum ( $100-200 \mu \mathrm{~L}$ ) onto the leaf. Smear the drop with fingers (or swab) without applying pressure.

Note: Work with gloves and change them between samples or clean hands thoroughly between samples by using alkaline soap.
3.2. Rinse the plants with tap water a few minutes after inoculation.
3.3. In order to allow the assay plants to become systemically infected, incubate them for at least 14 days under controlled conditions at $25 \pm 5^{\circ} \mathrm{C}$ and with at least 12 hours of light per day.

## 4. Sampling of Nicotiana benthamiana assay plants and leaf extraction

4.1. For each subsample, the PPC, and the NPC, sample and pool leaf material from both assay plants, making sure that the pooled leaves weigh $\sim 10 \mathrm{mg}$. Select younger leaves that have expanded during the preceding weeks and not the inoculated leaves.
4.2. Process samples immediately, store at $4^{\circ} \mathrm{C}$ for at most 48 hours or freeze until use. If the samples were frozen, process them as soon as they are thawed.
4.3. Grind each pooled leaf sample in 0.5 mL seed extraction buffer. Process extracts immediately after grinding, store at $4^{\circ} \mathrm{C}$ for a maximum of 24 hours or freeze until use.


Figure G.1. Nicotiana benthamiana plant front view (A) and top view (B).

## 5. ELISA

5.1. Run a double-antibody-sandwich (DAS-)ELISA (Albrechtsen, 2006; Clark and Adams, 1977) on the extracts obtained at step 4.3. and controls mentioned in Table G.2.
5.2. A subsample is regarded as positive (infectious PepMV present) when the signal in ELISA is above the decision threshold.

Note: The source of antiserum is critical. In the comparative test study, the antiserum supplied by Wageningen University \& Research Prime Diagnostics was used (https://www.wur.nl/en/show/Prime-Diagnostics2.htm). If different antisera and buffers are used, or even different lot numbers, it is necessary to verify their performance.

## 6. Validity of test results

Test results are only valid when all included controls presented in Table G .2 give the expected result.

## REFERENCES

Albrechtsen, S.E. (2006). Testing methods for seed-transmitted viruses: principles and protocols. Wallingford, UK: CABI Publishing.

Clark, M.F. and Adams, A.N. (1977). Characteristics of the microplate method of enzyme-linked immunosorbent assay for the detection of plant viruses. Journal of General Virology, 34, 475-483.


[^0]:    ${ }^{1}$ Available at: https://www.worldseed.org/our-work/seed-health/ishi-method-development-and-validation/
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[^1]:    ${ }^{\text {a }}$ Five mg PepMV infected leaf tissue in buffer, used to spike seed extract.
    ${ }^{\mathrm{b}}$ Includes six or more reactions without amplification.
    ${ }^{c}$ LOD ELISA: Twice the average of the healthy controls ( $\mathrm{S} / \mathrm{N}$ extinction ratio of 2.0).
    ${ }^{d}$ LOD SE-qPCR: The lowest virus dilution in which PepMV can still be detected in all nine replicates.

