

Detection of *Paracidorvorax citrulli* in Cucurbit Seeds by SE-qPCR

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Crop: Watermelon (*Citrullus lanatus* var. *lanatus*)

Melon (*Cucumis melo*)

Squash (*Cucurbita pepo*)

Root stock squash (*Cucurbita maxima*)

Pathogen: *Paracidovorax citrulli* (former *Acidovorax citrulli*)

Version: 3.1 (January 2026)

PRINCIPLE

Detection of infectious *Paracidovorax citrulli* bacteria on cucurbits seeds is done by growing out seeds under environmental conditions highly conducive to producing disease symptoms followed by a biological assay to confirm the presence of infectious *P. citrulli*.

A seed extract qPCR assay (SE-qPCR) may be used as a pre-screen. The test is complete if no *P. citrulli* is detected and the seed lot is considered healthy. However, as qPCR detects both viable and non-viable bacterial DNA, a positive pre-screen SE-qPCR is followed by the grow-out to determine the presence of infectious *P. citrulli*.

The complete method process workflow is presented in Figure 1.

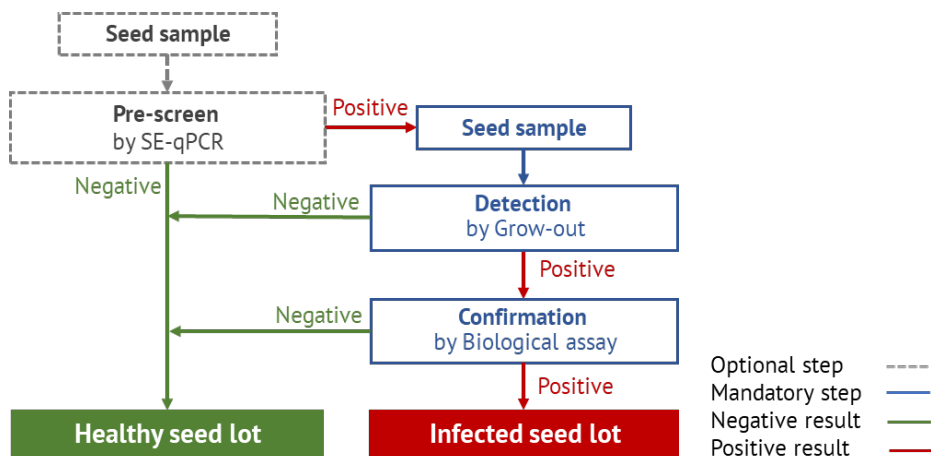


Figure 1. Method process workflow.

METHOD VALIDATION

The seed extract qPCR (SE-qPCR) has been validated by seven independent laboratories using blended naturally infected seed samples through an ISHI comparative test (Kleinhesselink, 2017). Seed from the following cucurbit crops were used during validation: melon (*Cucumis melo*), watermelon (*Citrullus lanatus*), squash (*Cucurbita pepo*) and root stock squash (*Cucurbita maxima*). Recent field observations and additional experimental data are published in an addendum validation report (Willmann *et al.*, 2025), which justifies the removal of the ZUP qPCR assay from the SE-qPCR protocol.

RESTRICTIONS ON USE

Before using this protocol routinely, it is necessary to verify its performance, especially when material and consumables from different suppliers are used. Technical details on the reagents/material used in the validation study (e.g., supplier's information) are provided in the protocol and the validation report. This test method is suitable for untreated seed.

The method may also be suitable for seed treated with some physical and/or chemical processes. It is the responsibility of the testing laboratory to verify that treatment processes do not present an antagonistic effect to method utility.

Guidance for molecular testing methods is provided such that it accommodates modular assay components. DNA isolation, inhibition control selection, qPCR reaction mixture, cycling parameters, multiplex reactions and evaluation specifications (threshold/cut-offs) are variables that may differ between testing laboratories. It is the responsibility of the testing laboratory to verify that the selected combination of assay components meet the following minimum recovery and detection parameter, *viz.* 100% detection of *P. citrulli* across triplicate 5,000 seed subsamples in all qPCR reactions where each 45 mL aliquot contains ~12 CFU/mL, or in a 5,000 seed subsample to which 250 mL buffer is added, 100% detection of *P. citrulli* in all qPCR reactions from triplicate samples of relevant crops, where each sample is spiked with ~3000 cells of *P. citrulli* is required.

METHOD EXECUTION

To ensure process standardization and valid results, it is strongly recommended to follow the [Best Practices for Seed Health Tests](#) developed by ISHI.

In line with the guidance provided in [ISF's view on indirect seed health tests](#), if a sample is determined qPCR positive, the seed lot is considered to be *suspect*. A final decision on the status of the seed lot can be taken only after a test confirms the presence of viable *P. citrulli* in the sample and its pathogenicity.

SAMPLE AND SUBSAMPLE SIZE

The minimum recommended sample size is 10,000 seeds up to 30,000 seeds per lot, with a maximum subsample size of 5,000 seeds. If the SE-qPCR is a positive and a confirmatory test is conducted, a new sample must be drawn for the grow-out.

REVISION HISTORY

Version	Date	Changes (minor editorial changes not indicated)
1.0	July 2017	First version of the protocol.
1.1	August 2018	NSHS grow-out protocol v1.1 combined with the ISHI pre-screening protocol, as together they form a method for the detection of <i>A. citrulli</i> in seed of Cucurbit crops.
2.0	May 2021	Protocol presented in accordance with ISHI's protocol guidelines. Reference to grow-out protocols included in method description. Sample size adjusted to align with grow-out assay.
3.0	October 2025	Removal of ZUP qPCR assay. Pathogen name updated to <i>Paracidovorax citrulli</i> . Control names updated to adhere to the new ISHI Best Practices.
3.1	January 2026	Aac R2 primer sequence corrected (final "T" missing since 2.0 version).

Protocol for detecting *Paracidavorax citrulli* in Cucurbit Seeds

I. PRE-SCREEN BY SEED-EXTRACT qPCR (OPTIONAL STEP)

For PCR methods, in-house method optimization is often necessary by changing certain parameters as described in [Best Practices for PCR Assays in Seed Health Tests](#).

Materials

- Container
- Orbital shaker
- Seed extraction buffer (Table I.1)
- 5× PBS (Table I.2)
- Sorbitol solution (Table I.3)
- T₁₀E_{0.1} (Table I.4)
- Nutrient Broth (Table I.5)
- Liquid KB (Table I.6)
- Controls (Table I.7 to I.10)
- qPCR mix, primers (Table I.11) and equipment
- DNA isolation kit
- Centrifuge
- Lab disposables

Table I.1. Seed Extraction Buffer.

Compound	Amount/L
5× PBS ^a	200.0 mL
Tween TM 20	0.5 mL

^a 5× is given as an example; extraction buffer can be prepared from alternate concentrations.

Notes: Prepare directly before use. Final solution is 1× PBS buffer + 0.05% (w/v) Tween 20.

Table I.2. 5× PBS.

Compound	Amount/L
Sodium chloride (NaCl)	40.0 g
Di-sodium hydrogen phosphate dodecahydrate (Na ₂ HPO ₄ .12H ₂ O) ^a	14.5 g
Potassium dihydrogen phosphate (KH ₂ PO ₄)	1.0 g
Potassium chloride (KCl)	1.0 g

^a 14.5 g Na₂HPO₄.12H₂O can be substituted with 5.76 g of Na₂HPO₄ or 7.20 g Na₂HPO₄.2H₂O

Notes: Adjust pH of the final solution to 7.4 with NaOH or HCl, if necessary. Autoclave before use.

Table I.3. Sorbitol solution.

Compound	Amount/L
D-sorbitol	0.62 g
1 M Tris-HCl, pH 8.0	1.0 mL
0.5 M EDTA, pH 8.0	100.0 μ L
β -mercapto-ethanol (98%)	20.0 μ L

Table I.4. $T_{10}E_{0.1}$ pH 8.0.

Compound	Amount/L
PCR grade water	990 mL
1 M Tris-HCl, pH 8.0	10.0 mL
0.5 M EDTA, pH 8.0	200.0 μ L

Table I.5. Nutrient Broth.

Compound	Amount/L
Nutrient Broth	8 g

Note: Autoclave before use.

Table I.6. Liquid KB.

Compound	Amount/L
Proteose peptone N°3 from Difco	20 g
Glycerol	7.95 mL
MgSO ₄ .7H ₂ O	1.5 g
K ₂ HPO ₄	1.5 g

Note: Autoclave before use.

Table I.7. Types of controls used.

Control type	Description
Inhibition control (IC) ^a	<i>A. cattleyae</i> spike (Table I.8) or <i>X. euvesicatoria</i> spike (Table I.8)
Positive process control (PPC)	Seed extract from known <i>P. citrulli</i> infected seed (Table I.9)
Negative buffer control (NBC)	Extraction buffer
Negative process control (NPC)	Seed extract from seed known to be free of <i>P. citrulli</i> target
Positive amplification control (PAC)	<i>P. citrulli</i> DNA and IC DNA
Non template control (NTC)	PCR grade water or $T_{10}E_{0.1}$ pH 8.0 or DNA isolation kit elution buffer

^a The IC also serves as Internal amplification control (IAC).

Table I.8. IC spike recommendations.

Species	<i>Xanthomonas euvesicatoria</i> (Xe)	<i>Acidovorax cattleyae</i> (Acat)
Isolate	ATCC 11633™ (deposited as <i>X. vesicatoria</i>)	ATCC 33619™ or ZUM3739
Concentration	OD ₆₀₀ = 0.100×10 ⁻³	OD ₆₀₀ = 0.100, 1:50 dilution
Volume	100 µL spike/25 mL extraction buffer	5 µL spike/25 mL extraction buffer

Notes: Bacterial suspensions may be used freshly prepared or following proper preservation and storage.

It is recommended that the Cq values are monitored using a control chart to ensure there is no drift or degradation over time for all controls subject to storage. Examples of freshly prepared and glycerol preserved control preparations are given below.

IC-Xe: Use a 2-3 day-old culture of Xe on YDC agar media. Prepare a cell suspension of this culture corresponding to OD₆₀₀=0.100. Perform three, ten-fold (1:10) serial dilutions to prepare the spike solution Xe: OD₆₀₀=0.100×10⁻³.

IC-Acat: Prepare a 10 mL KB culture suspension in a new 50 mL reaction tube by inoculating *A. cattleyae* from a KB agar plate not older than 10 days. Incubate the culture suspension at 27 °C on a shaker overnight. Determine the optical density of the overnight culture at 600 nm using a photometer set to “absorbance”. Dilute the overnight culture to OD₆₀₀=0.100 using sterile saline. Prepare the *A. cattleyae* glycerol stock according to Table I.9. Mix well the preparation by inverting the tube 10 times. Prepare aliquots of the glycerol stock in 1.5 mL reaction tubes and store at -80 °C. Once frozen, thaw for use only, do not re-freeze.

Table I.9. IC-Acat glycerol stock solution.

Compound	Amount
15% (v/v) glycerol ^a	49 mL
Acat O/N culture adjusted to OD ₆₀₀ = 0.1	1 mL

^a Autoclave before use.

Table I.10. PPC spike recommendation.

Species	<i>Paracidovorax citrulli</i>
Isolate	ATCC 29625™
Concentration	OD ₆₀₀ = 0.100×10 ⁻³
Volume	100 µL spike/25 mL extraction buffer

Notes: Inoculate a tube containing 9 mL nutrient broth with a 1 µL loop of *Paracidovorax citrulli*. Cap tube loosely and place in a tube rack at an approximate 45°-angle. Incubate/shake overnight (~18 hours) at 200 rpm at 37 °C. After incubation period, prepare a cell suspension of this culture corresponding to OD₆₀₀=0.100. Perform three, ten-fold (1:10) serial dilutions to prepare the spike solution of *Paracidovorax citrulli*: OD₆₀₀=0.100×10⁻³.

One mL of *Paracidovorax citrulli* OD₆₀₀=0.100×10⁻³ bacterial suspension is approximately equal to 3000 cells and can be used as a spike to evaluate laboratory performance of the method against the minimum recovery and detection parameter stated in the restrictions on use section of this document.

Table I.11. Primer and probe sequences and references.

PCR assay	Target	Primer	Primer sequence (5' – 3')	Source
Contig 21	Pc	Aac F1	ACC GAA CAG AGA GTA ATT CTC AAA GAC	Sudarshana, 2010
		Aac R1	GAG CGT GAT GGC CAA TGC	
		Aac P1	6FAM – CAT CGC TTG AGC AGC AA – MGBNFQ	
Contig 22	Pc	Aac F2	GAA AGT GGT TGT TCT GGT GAT CAA	
		Aac R2	TTC GGA GGA CTC GGG ATT T	
		Aac P2	6FAM – ATG GTC TGC GAG CCA G – MGBNFQ	
IC-Xe	Xe	Xcv F1	CCT CGA TGG GCA CCT GAT T	Monsanto, 2012
		Xcv R1	CGT CGA TTG CCG GGT ACT	
		Xcv P1	6FAM – ATC GCG GCC AAG AA – MGBNFQ	
IC-Acat	Acat	Acat2-F	TGT AGC GAT CCT TCA CAA G	Koenraadt <i>et al.</i> , 2014
		Acat2-R	TGT CGA TAG ATG CTC ACA AT	
		Acat1-Pr	VIC – CTT GCT CTG CTT CTC TAT CAC G – BHQ1	

1. Extraction of bacteria from the seed

- 1.1. Place seed subsamples in a container appropriate for seed and buffer volume.
- 1.2. Add the extraction buffer to each subsample at a ratio of 2.0 mL buffer per 1.0 g of seed. Due to buffer absorbance by the seeds, subsamples of large seed (e.g. squash) may require a slightly increased buffer to seed ratio (up to 2.5 mL per 1.0 g of seed).
- 1.3. An inhibition control (IC) spike is required. Each subsample being evaluated for *P. citrulli* is spiked with the IC and the spike volume is adjusted to a standardised concentration (i.e. added at a fixed ratio to the required extraction buffer volume). See Table I.8. Suspension may be prepared daily or in bulk (stored at -80 °C in a glycerol solution).
- 1.4. Both a positive process control (PPC) and negative process control (NPC) subsample are required (Tables I.7 and I.10). It is recommended that both the PPC and the IC spikes be added to the NPC subsample (seed known to be free of *P. citrulli* and the IC target).
- 1.5. Incubate all subsamples and controls on an orbital shaker for 1-2 hours at room temperature (~23 °C) at a speed sufficient to agitate the samples (~120 rpm).

2. Collection of Target Bacteria by Differential Centrifugation

- 2.1. Transfer 45 mL of seed extract per 5,000 seed subsample into a 50 mL centrifugation tube.

Note: If multiple containers were used to incubate a subsample, equal volumes of seed extract must be collected from each container to generate a total volume of 45 mL.

- 2.2. Centrifuge subsamples at 1,000 × g for 5 min.
- 2.3. Decant the supernatant into a fresh 50 mL centrifugation tube and discard pellet.
- 2.4. Centrifuge supernatant at 3,200 × g for a minimum of 15 min.

OPTIONAL: If additional PCR inhibitor removal is desired, a sorbitol treatment can be used at this point.

2.4.1. Decant supernatant from step 2.4

2.4.2 Add 1 mL of sorbitol solution (Table I.3) to the pellet and re-suspend.

2.4.3 Incubate the suspension for 20-60 min at room temperature.

2.4.4 Centrifuge the suspension at $1800 \times g$ for 10 min.

2.5. Carefully decant and dispose of the supernatant while preserving as much of the pellet as possible.

2.6. Add a volume of extraction buffer sufficient to resuspend the pellet to achieve the input volume recommended by the DNA isolation kit which will be used, typically 0.5-2.0 mL.

3. DNA isolation

The assay has been validated with the MoBio PowerFood, Machery-Nagel NucleoSpin Plant II and Sbeadex Maxi Plant kits. If a different DNA isolation kit is used, it is necessary to verify its performance.

Recommended Modifications to the DNA kits tested:

i. MoBio PowerFood Kit

- Garnet bead tube is modified from provided 0.5 mL to 2.0 mL screwcap tube.
- EDTA is added to solution PF6 for DNA storage stability at a concentration of 0.1 mM.

ii. Macherey-Nagel NucleoSpin Plant II Kit

I Macherey-Nagel NucleoSpin® 96 Plant II (96-well format, ref #740663). The vacuum manifold option of the protocol is performed with the following modifications:

- Lysis buffer composition: 480 μ L PL1 + 20 μ L of a 10 mg/mL proteinase K stock solution
- Lysis buffer volume used for resuspending pellet 450 μ L
- Lysis incubation time 1 – 20 hours
- Lysate clearing: centrifugation at $3200 \times g$ for 20 min
- PW1 washing step is performed 2×
- Dry membranes for 15 min
- Perform only 1 DNA elution-step
- Option to elute DNA not by vacuum but by centrifugation at $3,200 \times g$ for 2 min

II Macherey-Nagel NucleoSpin® Plant II (single tube format, reference #740770)

- Lysis buffer composition (480 μ L PL1, 20 μ L proteinase K (20 mg/mL stock, no RNase A)
- Lysis buffer volume used for resuspending pellet 450 μ L
- Lysis incubation time 1 – 20 hours
- Lysate clearing: centrifugation at $19,000 \times g$ for 5 min
- Centrifugation steps involving silica column performed at $19,000 \times g$
- PW1 washing step is performed 2 times
- Only 1 elution step using 100 μ L PE, pre-heated at 70 °C

4. qPCR assay

The assay has been validated with the ABI TaqMan® master mix (2×) and Quanta perfecta Multiplex qPCR ToughMix. If different qPCR mixtures and amplification programs are used, it is necessary to verify their performance.

4.1. For Contig 21, Contig 22 and IC – Xe qPCR assays, see the reaction mixture in Table I.12, and for the IC – Acat qPCR assay, see the reaction mixture in Table I.13. The cycling parameters are provided in Table I.14.

Note: Although the qPCR assays were validated as singleplex reactions, the [Best Practices for PCR Assays in Seed Health Tests](#) advises to process the inhibition control in the same reaction as the target assay. When using multiplex qPCR mixtures, it is necessary to verify their performance.

Table I.12. qPCR mix for the Contig 21, Contig 22 and IC-Xe qPCR.

Component	Per reaction (in µL)	Final concentration
PCR grade water	6.988	
Forward primer (100 pmol/µL)	0.225	0.90 µM
Reverse primer (100 pmol/µL)	0.225	0.90 µM
Probe (100 pmol/µL)	0.062	0.250 µM
qPCR master mix (2×)	12.50	1×
Template DNA	5.00	
Total	25.00	

Table I.13. qPCR mix for the IC-Acat qPCR.

Component	Per reaction (in µL)	Final concentration
PCR grade water	7.25	
Acat2-F (100 pmol/µL)	0.10	0.40 µM
Acat2-R (100 pmol/µL)	0.10	0.40 µM
Acat1-Pr (100 pmol/µL)	0.05	0.20 µM
qPCR master mix (2×)	12.50	1×
Template DNA	5.00	
Total	25.00	

Table I.14. qPCR cycling parameters.

Step	Temperature	Duration	Description
Hold	95 °C	10 min	Enzyme activation
40 cycles	95 °C	15 sec	Denaturation
(ramp rate 1.6 °C/sec)	60 °C	60 sec	Annealing/elongation

4.2. A positive amplification control (PAC) and a non-template control (NTC) are required (Table I.7). Load PAC DNA template into separate qPCR reactions at a recommended concentration of 2 pg/µL.

Prepare the PAC-Ac, Xe, Acat as follows:

Inoculate a tube containing 9 mL nutrient broth with a 1 μ L loop of bacteria grown on solid media. Cap tube loosely, place in a rack at an approximate 45°-angle and shake overnight at 200 rpm at 37 °C. Process 1 mL of resulting culture using Qiagen DNeasy Blood and Tissue Kit including an RNase-A digestion. Elute DNA with 200 μ L buffer AE. Quantify eluted DNA. Normalize to 2 ng/ μ L stock concentration and 2 pg/ μ L working concentration. PAC concentrations may vary per laboratory and should produce Cq values less than the LOQ of the method.

5. Interpretation of results

Cut-off values must be established by each laboratory for their positive and inhibition controls prior to the assay being used on routine samples. For recommendations on setting cut-off values, see [Real-time PCR, an 'indirect' test used for pre-screening in seed health methods](#).

It is the responsibility of the testing laboratory to determine threshold/cut-off values, such that they meet the performance-based acceptance criteria stated in the restrictions on use section of this method.

Test results are only valid when all included controls presented in Table I.7 give the expected results. NBC/NPC/NTC must be negative (no detection of *P. citrulli*) and IC/PPC/PAC must fall within their expected ranges for the test to be considered valid. Control ranges are determined per laboratory. Note that IC control range limits must be set per crop species in each testing laboratory.

- 5.1. A sample is considered qPCR positive (suspect) for *P. citrulli* if one or more subsamples show detection of *P. citrulli* DNA by one or more *P. citrulli* primer sets.
- 5.2. If a sample is qPCR positive (*P. citrulli* suspect), the detection, identification and confirmation grow-out assay, described in section II, must be performed to reach a final conclusion about the sample and seed lot (See [ISF's view on indirect seed health tests](#)).

II. DETECTION, IDENTIFICATION AND CONFIRMATION BY GROW-OUT ASSAY

- For melon seed, see the protocol for the [Detection of *Acidovorax citrulli* in melon seed by sweat box grow-out](#) on the ISF website.
- For all cucurbits, see version 1.1 of the [Acidovorax citrulli grow-out](#) protocol on the NSHS website.

REFERENCES

Kleinhesselink, K. (2017). Seed Extract and qPCR Method for the Detection of *Acidovorax citrulli* on Cucurbit Seeds. Validation report, International Seed Federation (ISF), Nyon, Switzerland. <https://worldseed.org/our-work/seed-health/ishi-method-development-and-validation/>

Koenraadt, H., van Vliet, A., Jodlowska, A., Woudt, B., Ebskamp, M. and Bruinsma, M. (2014). Detection of *Acidovorax citrulli* in seed extracts of melon and watermelon with TaqMan PCR. 7th ISTA Seed Health Symposium. Edinburgh, United Kingdom, 12 – 14 June 2014.

Monsanto (2012). Cb 1.4 – *Acidovorax avenae* ssp. *citrulli* – Monsanto PCR. Available at: <https://seedhealth.org/cb1-4/>

Sudarshana, P. (2010). ISHI AAC primers evaluation. ISHI-Veg meeting, July 2010.

Willmann, R., Sabban, M., Thomas, L. and Woudenberg, J.H.C. (2025). Justification for Removal of the ZUP (IS1002) qPCR Assay from the SE-qPCR Protocol for *Acidovorax citrulli* Detection in Cucurbit Seeds. Addendum Validation report, International Seed Federation (ISF), Nyon, Switzerland. <https://worldseed.org/our-work/seed-health/ishi-method-development-and-validation/>