Nematode extraction from banana roots by the centrifugal-flotation technique.

Abstract — Introduction. The centrifugal-flotation technique allows all nematode species and life stages to be separated from root debris and residual soil particles to facilitate their observation. The principle of the method applied, key advantages, starting plant material and time required are presented. Materials and methods. Necessary laboratory materials, and details of the 19 steps required for extracting nematodes from plant tissue for counting are described. Possible troubleshooting is explained.

France (Guadeloupe) / Musa sp. / pest control / methods / extraction / laboratory equipment / plant nematodes / cell counting

1. Introduction

Application

Nematodes must be extracted from plant tissue for counting and identification of species. The centrifugal-flotation technique allows all nematode species and life stages to be separated from root debris and residual soil particles to facilitate their observation.

Principle

This method has been adapted to bananas from the centrifugal-flotation technique described by Coolen and d’Herde [1]. Root tissues are macerated in a kitchen blender. The suspension of nematodes, plant debris and residual soil particles is poured on a sieve column and then abundantly washed. The biggest debris is retained by the upper sieve and discarded. The smallest debris is eliminated through the mesh of the last sieve. Residues are gently washed in a 500-mL centrifuge tube. After a first centrifugation, the supernatant (water and the lightest debris) is discarded. The centrifuge tube is then filled with a solution of MgSO₄ at a density similar to that of nematodes. After the second centrifugation, the supernatant (water and the lightest debris) is discarded. The centrifuge tube is then filled with a solution of MgSO₄ at a density similar to that of nematodes. After the second centrifugation, the supernatant containing only nematodes is collected on a 5-µm aperture sieve. Finally, nematodes are collected in a beaker or a test tube for counting.
Key advantages
This method:
– facilitates observation and counting since nematodes are collected in a suspension free of root debris,
– allows the collection of all nematodes and life stages (including eggs) whatever their mobility,
– saves time, since nematodes can be counted less than 1 h after the beginning of the extraction process.

Starting material
The method requires root samples collected in the field or from pots (greenhouse experiments).

Time required
About 40–50 min are necessary for four root samples ready to count. One person can extract and count about 16–20 samples a day.

2. Materials and methods

Laboratory materials
The method requires: a centrifuge with four 500-mL tubes (at least 250 mL), a kitchen blender, a vibro-mixer, a set of sieves (at least three) with decreasing mesh aperture (e.g., 250 µm, 50 µm, and 32 µm or 25 µm), four sieves (as many as centrifuge tubes) of 5 µm-mesh aperture, a weighing balance, four funnels (as many as centrifuge tubes) adapted to centrifuge tube size, beakers, graduated tubes (200 mL) or any graduated glass vessel (e.g., beaker), a solution of MgSO₄ at 1.16–1.17 g·L⁻¹, kaolin and counting cells (1 mL).

Protocol
• Step 1
Wash crude root samples thoroughly to eliminate any soil residue.

• Step 2
Chop roots into 0.5–1-cm sections.

• Step 3
Wash roots again to eliminate any residual soil particles.

• Step 4
Mix root sections thoroughly to prepare the aliquot sampling.

• Step 5
Sample an aliquot part of 50 g of root sections.

• Step 6
Pour the root aliquot into a kitchen blender with 200 mL of tap water. Macerate twice for 10 s with a 5-s pause.

• Step 7
Pour the macerated suspension into the column of sieves. Rinse the blender vessel thoroughly to collect all nematodes.

• Step 8
Wash the sieve column abundantly for 1 min (avoiding any splashing).

• Step 9
Discard residues on the top sieve (250 µm).

• Step 10
Gently wash the intermediate sieve (50 µm), giving it a 45° slope and orientating the water flow from below to avoid as much as possible any nematode loss through the meshes. Make all residues flow down along the slope and accumulate along the lower part of the sieve edge. Note: proceed above the 32 µm (or 25 µm) sieve as a security against any casual nematode loss.

• Step 11
Pour residues through a funnel into the centrifuge tube. Wash gently and thoroughly with a washing bottle to collect all residues.

• Step 12
Repeat steps 10 and 11 with the lowest sieve (32 µm or 25 µm).

• Step 13
Add 15 g of kaolin to all centrifuge tubes. Mix the suspension with a vibro-mixer and balance the centrifuge tubes.
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Note: kaolin allows the residue to cement during the centrifugation for easier elimination of supernatant.

- Step 14
Proceed to a first centrifugation (1500 g for 5 min) and discard the supernatant (water + the lightest root debris).

- Step 15
Fill the centrifuge tube with the MgSO₄ solution. Put the residues and the kaolin in suspension again with the vibro-mixer and balance the tubes (using the MgSO₄ solution). Note: this step is crucial. Make sure that the cemented kaolin is well resuspended, since nematodes could be trapped in kaolin curds.

- Step 16
Proceed to a second centrifugation (1500 g for 5 min).

- Step 17
Collect nematodes:
  - collect the supernatant containing nematodes on a sieve of 5-µm aperture,
  - wash gently to eliminate the saline solution residue,
  - wash out nematodes into a graduated tube and adjust the volume of the suspension (generally 100 mL) for counting.
Caution: carry out steps 15 to 17 very quickly since the osmotic pressure may destroy nematodes if they stay too long in the solution.

- Step 18
Put 1 mL of nematode suspension on a counting cell. Counting is done under a microscope or a binocular lens. For one root sample, counting must be repeated at least three times with 1 mL.
Caution: nematodes decant progressively in the tube; don’t forget to shake the tube thoroughly before taking the 1-mL sample.

- Step 19
Results are expressed as density of nematodes per g of root.

Troubleshooting

Four main problems can occur.

(a) No or few observed nematodes out of an apparently infested root sample. This can result from:
  - A density of MgSO₄ solution under 1.16 g·L⁻¹.
  Solution: check the solution density before proceeding.

  - Remaining kaolin curds after the vibro-mixing phase of the second centrifugation (step 16).
  Solution: make sure the cemented kaolin is finely divided and well suspended in the solution.

  - Nematodes decanted in the tube before counting.
  Solution: thoroughly shake the tube before taking the 1-ml sample (step 18).

(b) Nematodes are distorted and not identifiable. This can result from:
  - A density of MgSO₄ solution over 1.17 g·L⁻¹.
  Solution: check the solution density before proceeding.

  - A correct density of MgSO₄ solution but nematodes stayed too long in it.
  Solution: proceed very rapidly during steps 15 to 17 when nematodes are suspended in the solution.

(c) Many root fragments are in the counting suspension: this may be due to a density of MgSO₄ solution which was over 1.17 g·L⁻¹.
Solution: check the solution density before proceeding.

(d) Many chopped nematodes are in the counting suspension: this may be due to too high a speed or too long a running-time of maceration (step 6).
Solution: adjust these parameters to the model of the blender used.

References