**Screening for resistance to *Xanthomonas vasicola* pv. *musacearum* (Xvm)**

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**Introduction**

*Xanthomonas vasicola* pv. *musacearum* (Xvm) previously known as Xanthomonas campestris pv. musacearum (Studholme et al., 2020) is a plant pathogen that causes banana wilt also known as banana Xanthomonas wilt (BXW). Among banana diseases, Xanthomonas wilt, is reported to be the most devastating disease in East and Central Africa causing 100% yield loss, since it kills the entire plant and prevents fruit formation (Nakato *et al*., 2018).

Xanthomonas wilt, is commonly transmitted by contaminated garden tools, infected planting materials, and vectors like insects, birds, and browsing animals, (Nakato *et al*., 2014). Xvm survives on plant residues for 3 months (Nakato *et al*., 2018) hence it can be effectively controlled by uprooting, chopping, removing, and burying diseased stems, sterilizing and using clean garden tools as well as removing male buds early enough using a fork stick, which will prevent and control cross transmission to other non-infected plants, (Nakato *et al*., 2014).

The infection results into wilting and yellowing of youngest leaves or the male buds, premature ripening of fruits, which results from internal vascular bundle discoloration mainly in the pseudo stem producing a yellow discharge “ooze”, (Biruma *et al*., 2007).

Among banana cultivars however, a wild form *Musa Balbsiana* has proved resistant to this infection, genetic studies are conducted to determine the source of resistance, which resistance genes can be transferred to other genotypes through conventional breeding, and genetic engineering (Sekiwoko *et al*., 2006)

Besides *Musa Bulbsiana`s* resistance to Xvm, it belongs to BB genome subgroup hence not preferred for breeding since the available edible bananas belong to AA genome subgroup, however with molecular characterization, present studies reveal the presence of resistant genes in A genomes banana subgroups especially *Musa acuminata*, which can be effectively used in breeding of edible bananas and plantain, (Nakato *et al*., 2019).

This screening protocol is developed to provide guidance for the Banana breeding program to determine resistance of banana genotypes to *Xanthomonas vasicola* pv. *musacearum* (Xvm) which causes Xanthomonas wilt of banana.

**Materials and methods**

***Plant materials***

TC generated plant material including:

1. Test genotypes (Parental genotypes and newly developed hybrids)
2. Tolerant check
   1. Monyet
3. Land race controls:
   1. TM-28 OBINO LEWAI
   2. Mbwazirume
   3. Mchare

***Other materials (For screenhouse evaluations)***

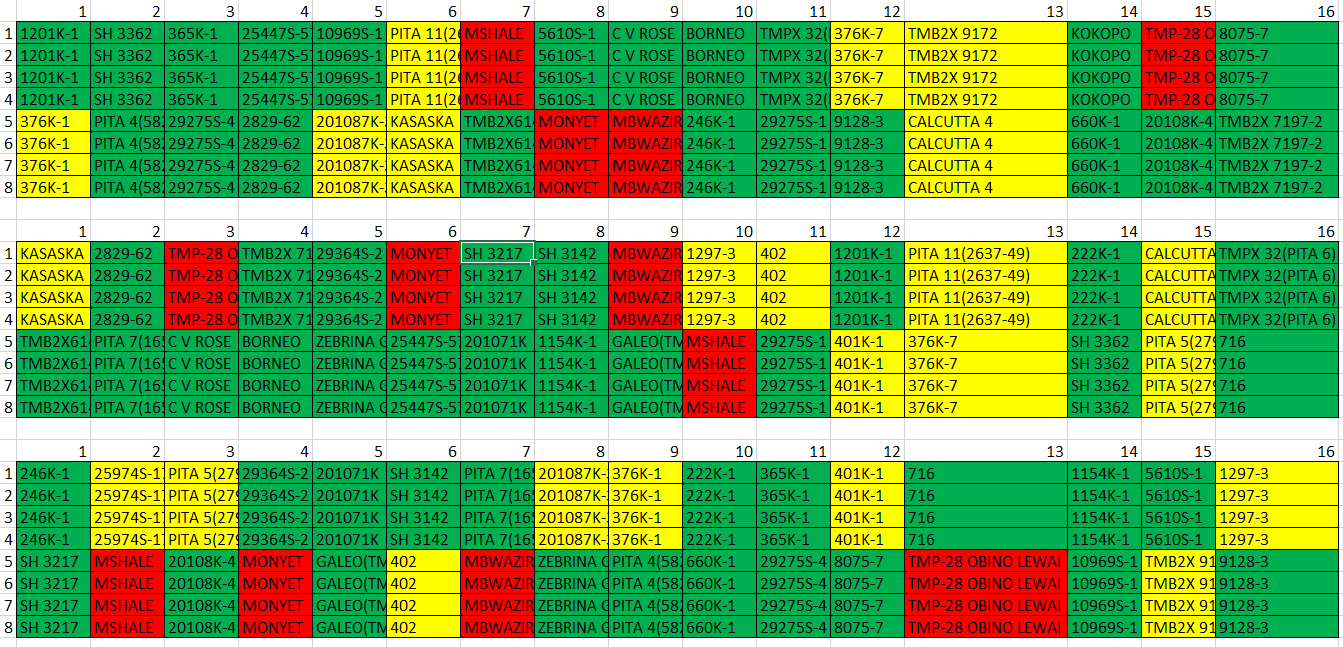
1. Sterile forest soil
2. Sterile sand
3. 13-liter plastic pots
4. Watering cans
5. Xvm inoculum

**Experimental design**

The experiment will be potted and established in an open space with no environmental control but minimal management practices like watering during dry spells.

***Parental genotypes***

Partially replicated experimental design (P-Rep) with three blocks will be adopted with each parental genotype occurring in duplicate and the checks in triplicate for the entire experimental set up. Each plot will constitute three plants per genotype. P-rep designs are useful when plant materials and space are limiting factors. In-fact, P-rep designs allow for repeated trial evaluations at different locations. The experimental design for the available 51 parental genotypes including checks is shown in Figure 1 below. The average efficiency 0.99 indicating that the design is optimal.



**Figure 1.**P-rep lay out for banana Xanthomonas wilt screening for the available 51 parental lines including checks

***Hybrids***

The augmented design resulting into an incomplete block design where standard checks are replicated in each block and test genotypes will be adopted for the hybrids. Such augmented designs are very useful where very large numbers of hybrids are produced with limited planting materials and reduced space.

**Methodology**

***Generation of plants***

The genotypes to be screened are generated from Tissue culture (TC), grown in the nursery to harden for 8 weeks under controlled environmental conditions (humid chamber and the screen house) and later transferred to planting bags containing a sterilized mixture of garden soil, Manure and sawdust mixed in a ratio of 3:1:1 respectively from where they are left to fully grow for 3 months.

***Xvm inoculum preparation***

Total volume of inoculum to be prepared is dependent on the number of genotypes to be inoculated.

1. Xvm cultures are aseptically introduced in YPG broth and incubated at 25oC - 28oC for 48hours with shaking, to optimize cell growth.
2. Record the culture cell concentration (Absorbance/Optical Density) is using a spectrophotometer with a light wavelength of 600 (O.D600).
3. Adjust the culture cell concentration to desired concentration of 108 Colony Forming Units (CFU) per ml which is approximately 0.5 OD600 using C1V1 = C2V2

***Plant inoculation*** (Nakato *et al*., 2018)

1. 1ml of the 108 CFU/ml cell concentration is introduced into the youngest open leaf via the petiole/midrib, for all test genotype.
2. Negative controls are inoculated using sterile distilled water.
3. Record date on which inoculation has been completed.
4. Observe plants weekly for disease symptom development for 4 months.

**Data collection and Analysis (Nakato *et al*., 2019)**

1. Data is collected on a weekly basis, proceeding the date of inoculation for a period of 15 weeks.
2. Parameters recorded include:
3. Number of functional leaves
4. Number of chlorotic leaves (leaf wilting)
5. Death of entire plant (if any) per genotype are recorded

**NB:** Every new observed Chlorotic leaf is tagged with a tape to eliminate errors due to counting other chlorotic leaves caused by normal leaf synesis, (If possible other chlorotic leaves should be eliminated by cutting them off)

Disease incubation, days to symptom development per genotype can be extracted from the weekly data records. Disease Index (DI), Days of Post inoculation (DPI) and Area Under Disease Progression Curve (AUDPC) are computed using the formulae below:

**Disease Index (DI)**

=(((1\*A) + (2\*B) + (3\*C))/Number of Plants) x100

Where

**A** = Number of plants with inoculated leaf showing symptoms

**B** = Number of plants with uninoculated leaves showing symptoms

**C** = Number of wilted Plants

**Days of Post Inoculation (DPI)**

= Time interval (Counting number of Days from inoculation to symptom development and complete wilting).

**Table 1: Disease classification based on DI**

|  |  |
| --- | --- |
| Disease Classification | |
| Resistant(R) | No plants wilted |
| Tolerant (T) | <30% of plants wilted |
| Moderately Susceptible (MS) | >30% and <50% of plants wilted |
| Highly Susceptible(HS) | >50% of plants wilted |

Area Under Disease Progression Curve (AUDPC):

Where:

**t** = time in weeks of each Reading

**Y** = Percentage of affected plants at each Reading

**N** = Number of Readings

**i** = Reading

**Appendix**

* + 1. **Isolation of Xvm**

**Xvm isolation media (Mwangi *et al*., 2007)**

1. Wilbrink media
2. YPGA (5g/l Yeast, 5g/l Peptone, 10g/l Glucose and 15g/l Agar)
3. YDCA (10g/l Yeast, 20g/l CaCO3 20g/l Dextrose and 15g/l Agar)
4. Cellobiose-cepholexin Agar (CCA)

**Materials**

* Well labeled Plant Samples (preferably pseudo stem)
* Laminar flow hood
* Toothpicks
* Cling Film
* Wilbrink Isolation media plates or YPGA media plates (Yeast 5g/l, Peptone 5g/l, Glucose 10g/l and Bacteriological Agar 15g/l)
* Sterile Toothpicks
* 5-fluorouracil for eliminating fluorescent Pseudomonads (Nakato V *et al*., 2018)
* Cycloheximide for Fungal growth inhibition

1. Media is sterilized by autoclaving at 121 Ibs pressure for 15 minutes and dispensed 20mls poured into petri dishes after cooling to 30oC.
2. The sample initials are recorded in the Laboratory sample book.
3. Using a sterile knife, the two outer most sheaths are peeled off and discarded from the 30-cm portion of the pseudo-stem.
4. The remaining exposed outer surface is sterilized with 70% ethanol in a laminar flow hood and placed on absorbent tissue paper to dry.
5. Aseptically cut the sterilized stems into two cross-section portions and keep slightly slanting for 15-10 minutes until Bacterial Ooze is discharged.
6. The Ooze is picked using a sterile toothpick, and directly streaked on the media plate (Wilbrink or YPGA).
7. The plate is sealed using a cling film (to prevent contamination) and incubated at 25oC for 3 days.
8. Yellow mucoid convex shaped colonies are picked from media plate and re-cultured on YPGA media to obtain pure colonies.
9. Confirmation can be done using Xvm specific markers (Nakato *et al*., 2019)
10. Pure colonies are stored on 20% glycerol 50% YPG broth for long term storage at -80oC
11. **Wilbrink media Preparation, (Wang *et al*., 2018)**

|  |  |  |  |
| --- | --- | --- | --- |
| **Wilbrink Media:** | 1000ml | 500ml | 200ml |
| Bacto Peptone | 5g | 2.5g | 1g |
| Sucrose | 10g | 5g | 2g |
| K2HPO4 | 0.5g | 0.25g | 0.1g |
| MgSO4·7H2O | 2.5g | 1.25g | 0.5g |
| Na2SO3 | 0.25g | 0.125g | 0.05g |
| Agar | 15g | 7.5g | 3g |
| **YPGA Media:** | | | |
| Yeast | 5g | 2.5g | 1g |
| Peptone | 5g | 2.5g | 1g |
| Glucose | 10g | 5g | 2g |
| Agar | 15g | 7.5g | 3g |

Table 2: Media for Xvm isolation

1. **Pictorial of Xvm symptoms**

|  |  |  |  |
| --- | --- | --- | --- |
| IMG_256  Image 1: Leaf Bxw Symptoms | IMG_256  Image 2: Yellow discharge (Ooze) in Pseudostem | |  |
| IMG_256  Image 3: uneven Ripening and rotting of fruits | | IMG_256  Image 4: Wilting of the male bud bracts | |

1. **Pure Xvm colonies on YPGA media**

|  |  |
| --- | --- |
| IMG-20210506-WA0017  Image 5: KB5 Xvm culture on YPGA media plate | IMG-20210506-WA0018  Image 6: Bcc 280 Xvm culture on YPGA media plate |

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