



Control of *Ralstonia solanacearum* on Selected Solanaceous Crops in Greenhouse by Selected Soil Amendments

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Authors' contributions

*This work was carried out in collaboration among all authors. Author EKK did *Ralstonia solanacearum* isolates collection, carried out green house experiment, analyzed data and prepared the first draft. Author ZMK provided the working experimental design for treatment and guided (Supervised) the activity. Authors JMM and POO provided general guidance on the experiment and edited the manuscript. All authors read and approved the final manuscript.*

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ABSTRACT

Aims: The aim of this study was to establish the effect of selected soil amendments on *Ralstonia solanacearum* isolates in greenhouse on selected solanaceous crops.

Study Design: The study was laid out as randomized complete block design (RCBD) in split pot arrangement for two seasons in the greenhouse.

Place and Duration of Study: The experiment was carried out in Kenyatta University situated in Kiambu County about 20 km from Nairobi city along Nairobi-Thika road between July, 2017-September, 2017 and between November, 2017- January, 2018.

Methodology: The three host crops of interest (potatoes, tomatoes and capsicum) were inoculated with prepared pure bacterial isolates; 18 (2T-Kiambu-Low Land), 71(2A-Nyeri-Low Land), 67 (2A-Nyeri-High Land), 83 (2T-Kirinyaga-Highland) and MX (18/71/67/83). Potatoes, tomatoes and capsicum were planted in pots each with a radius of 0.07 m (area 0.015 m²).The experiment had a

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total of 450 pots having a total area of 6.93 m². The treatments were Chalim™, Super-hydro-grow polymer + Metham sodium, Metham sodium, Metham sodium & Orange peel, Super-hydro-grow polymer, Brassica tissues, Chalim™ + Super-hydro-grow polymer, Brassica tissue + Orange peel, Metham sodium + Super-hydro-grow polymer and Control (no amendments).

Results: There were significant differences ($P \leq 0.05$) in the bacterial wilt incidences in selected solanaceous crops between control and all the soil amendments used in season 1 and 2. Brassica tissue + Super-hydro-grow polymer was superior in reducing bacterial wilt incidences in selected solanaceous crops in all the *R. solanacearum* isolates from Kenyan highlands and lowlands both in season 1 and 2.

Conclusion: Organic and inorganic soil amendments could serve as a viable control of bacterial wilt in solanaceous crops caused by *R. solanacearum* in the greenhouse.

Keywords: Bacterial wilt; incidences; *Ralstonia solanacearum*; solanaceous crops.

1. INTRODUCTION

The increasing global population coupled with the challenges of environmental degradation and an increasingly variable climate have created a world-wide need for improved food security [1,2]. Bacterial wilt disease caused by *Ralstonia solanacearum* is one of the most important constraints in production of vegetables in the tropical and sub-tropical regions [3]. The pathogen *R. solanacearum* is widespread in tropical, sub-tropical and warm temperate regions and infects more than 450 plant species in 54 families [4,5]. *R. solanacearum*, the causal agent of bacterial wilt disease, is considered one of the most destructive bacterial pathogens due to its lethality, unusually wide host range, persistence and broad geographical distribution [6]. *R. solanacearum* is a diverse species that differs in host range, geographical distribution, pathogenicity and biochemical and physiological properties [7]. This pathogenic species has been divided into five races based on host range [8] and six biovars based on metabolic profiles, related to the ability to metabolize three sugar alcohols and three disaccharides. These globally dispersed and heterogeneous strains cause loss of productivity of many crops, which have major socio-economic impacts [9].

Tim et al. [10] and Joshi et al. [11] reported that bacterial wilt disease is affected by environmental conditions like soil temperature, soil moisture, and soil type (which influences soil moisture and microbial populations). The bacterium enters host plant roots from the soil and colonizes the xylem vessels in the vascular system [12]. Infected plants suffer yellowing, stunting and wilting, and often die rapidly [13]. Symptoms of *R. solanacearum* on tomato include wilting and necrosis as well as vascular browning [14]. Typically, stem and tuber cross-sections

ooze whitish bacterial exudates [15]. Ramesh et al. [16] further reported that *R. solanacearum* infection could spread through contaminated water and weeds in the Solanaceae family. *Ralstonia solanacearum* can be disseminated by farm implements, pollinator insects in banana, irrigation water, infested soil, plant debris, latently infected vegetative propagation materials (such as *Pelargonium* cuttings, potato tubers and banana corms) and through roots damaged by nematodes [17,18]. Latent infection is widespread and has been identified in several asymptomatic host plants including tomato, geranium, squash, and potato [19,20]. An incidence of 55% and 25% has been recorded on chili and potato crops respectively from the major chili and potato producing regions of Ethiopia [21]. Assefa et al. [22] (2015) reported that the percentage wilt incidence of bacterial wilt in Ethiopia was as high as 63% on potato, 55% on tomato and 100% on pepper. Singh et al. [23] observed in a study in India that crop losses of up to 90% were reported in the greenhouse compared to losses of 25-60 % reported for open field tomato.

Bacterial wilt control in various pathosystems has been possible through use of a combination of diverse methods such as host resistance, biofumigation, fertiliser application, soil solarisation, biological control, chemical control, and other cultural practices and integrated disease management schemes [24,25]. However, traditional control measures are not always effective in the control of bacterial wilt, such as the application of bactericides, disease-resistant cultivars and crop rotation [26], thus making it compelling to find a potential soil-borne disease control method to reduce economic loss. Synthetic chemicals have been used for many years to control agricultural biological agents, however, considerable problems have arisen

from the continued application of these chemicals, including development of resistance by the pathogen, high cost, residual effect on soil, pollution of the environment and hazard from handling toxic compounds [27]. The use of soil amendments (SAs) is a widespread means to control soil-borne disease. It has been reported that an SA, composed of urea and calcium oxide (CaO), is effective to control the bacterial wilt of tomato by affecting the pH and nitrite accumulation in the field [28,29]. Calcium carbonate (CaCO₃) could not only serve as a soil amendment to change soil pH but also increase soil Ca²⁺ content. Polymers are widely used for many applications in agriculture: to combat viruses and other crop pathogens, and functionalized polymers are employed to increase the efficiency of pesticides and herbicides, allowing the application of lower doses and thus indirectly protecting the environment [30]. Amendments provide energy and nutrients to soil, drastically changing the environment for the growth and survival of crops and microorganisms [31]. In the current study, the aim of this study was to establish the effect of selected soil amendments on *R. solanacearum* isolates in greenhouse on selected solanaceous crops. This current study will help to reduce bacteria wilt in very important selected solanaceous crops by using organic and inorganic amendments.

2. MATERIALS AND METHODS

2.1 Study Area

The experiment was carried out in Kenyatta University situated in Kiambu County about 20 km from Nairobi city along Nairobi-Thika road. The county enjoys a warm climate with temperatures ranging between 12°C and 18.7°C. The rainfall aggregate for the county is 1000 mm each year. Its geographical coordinates are 1° 10'0" South, 36°50'0" East. Low fertility soils are mainly found in the middle zone and the eastern part of the county which form part of the semi-arid areas. The soils in the midland zone are dissected and are easily eroded [32]. The soils are sandy or clay and can support drought resistant crops such as soya beans and sunflower as well as ranching. The elevation of the main campus is 1720 meters above sea level (ASL) [33].

2.2 Experimental Design and Treatments

The experiment was carried out between July, 2017- September, 2017 and between November,

2017 - January, 2018 and was replicated three times for the two seasons. The experiment was laid out in randomized complete block design (RCBD) in split plot arrangement in the greenhouse. Potatoes, tomatoes and capsicum were planted in pots each with a radius of 0.07 m (area 0.015 m²). The experiment had a total of 450 pots having a total area of 6.93 m². The treatments were Chalim™, Super-hydro-grow polymer and Metham sodium, Metham sodium + Orange peel, Super-hydro-grow polymer, Control, Brassica tissue, Chalim™ + Super-hydro-grow polymer, Brassica tissue + Orange peel and Metham sodium + Super-hydro-grow polymer. The three host crops of interest (potatoes, tomatoes and capsicum) were inoculated with prepared pure bacterial isolates; 18 (2T-Kiambu-Low Land), 71(2A-Nyeri-Low Land), 67 (2A-Nyeri-High Land), 83 (2T-Kirinyaga-Highland) and MX (18/71/67/83). All agronomic practices including, watering, fertilization, weeds, pests and disease control were well managed.

2.3 Preparation of Pot Soil Amendments

Fresh leaves of cabbage plant residues were finely chopped and incorporated into the soil at the rate of 0.01 kg per pot of an area of 0.015 m². The inoculated soil was thoroughly mixed with the finely chopped cabbage plant residue, ensuring that all the residues were well incorporated in the soil pot. Metham sodium, a chemical fumigant was applied in pot of an area of 0.015 m² at the rate of 3.08 ml per pot. This was the positive control. Chalim™ effect was assessed in the inoculated pots after application at the rate of 0.3 g per pot of an area of 0.015 m². Super-hydro-grow polymer was applied in plot of 0.015 M² at the rate of 0.0003 ml per pot using knap-sack sprayer. Combination of Chalim™ + Super-hydro-grow polymer was applied at the rate of 0.3 g per pot and 0.0003 ml per the same pot of an area of 0.015 m² respectively. Metham sodium + Super-hydro-grow polymer was applied in a pot of an area of 0.015 m² at the rate of 3.08 ml per pot and 0.0003 ml per the same pot respectively.

Metham sodium + Orange peel treatment was applied in a pot of an area of 0.015 m² at the rate of 3.08 ml per pot and Orange peel rate of 0.01 kg per the same pot respectively. Brassica tissue + Orange peel treatment were applied at a rate 0.01 kg per pot of an area of 0.015 m² and Orange peel at a rate of 0.01 kg per the same pot respectively.

2.4 Greenhouse Inoculation

The positively identified potato tubers and stem tubers of capsicum and tomato were used to isolate *R. solanacearum*. The five pure bacterial isolates were 18 (2T-Kiambu-Low Land), 71(2A-Nyeri-Low Land), 67 (2A-Nyeri-High Land), 83 (2T-Kirinyaga-Highland) and MX (2T-L/2A-L/2A-H/2T-H (18/71/67/83). Potato tuber a Stems and infected tomato and capsicum plants were cut above the soil level and the cut surfaces were suspended in test tube containing clean water. Bacterial strains were routinely cultured in CPG agar (CPG broth with 15 g of agar/litre) media. These strains were easily distinguished on the basis of colony morphology and colour by using the South Africa semi-selective medium (SMSA-E) at KARI-NARL bacteriology laboratory. Pure bacterial was harvested (30 plates per plant sample) into a 10 L of sterile distilled water to make composite bacterial inoculate to be sprayed in 450 pots. The experimental plant pots was performed in glasshouse where the 450 pots were inoculated each with 10 mL of 3.05×10^9 cfu/mL of *R. solanacearum* isolates 18, 71, 67, 83 and MX. Metham sodium, a known fumigant was used as a positive control. Randomized complete split plot design was used in the pot layout.

2.5 Data Collection and Analysis

Three sample crops (tomato, capsicum and potato) were used. The plants were rated weekly, each Wednesday for bacterial wilt disease incidence from the 18th day after planting where wilted plants were uprooted upon total foliage wilt and recorded though only the incidence at 4th, 7th and 10th weeks after planting (WAP) was considered for evaluation. Plants with visible symptoms (wilted leaves) were recorded as diseased plants. The disease index was calculated as $DI (\%) = 100 \times (\text{number of disease plants} / \text{total number of inoculated plants})$ using the formulas as adapted from Mwaniki et al. [34].

$$Di = (\text{No. of infected plants} / \text{Total number of inoculated plants}) \times 100$$

For proper key diagnostic identification of *R. solanacearum* in the greenhouse and to distinguish bacterial wilt from vascular wilts caused by fungal pathogens, bacterial wilt symptoms was identified by visual observation of typical bacterial wilt disease symptoms such as wilting, vascular discoloration, bacterial streaming in glass of water and browning of the

vascular bundles of the tuber. Milky white strands containing bacteria and extracellular polysaccharide was oozed out from the cut ends of the xylem. The diseased samples were brought to the laboratory and subjected aseptically for detection and confirmation of *Ralstonia solanacearum*.

2.6 Data Analysis

Data that was obtained from soil amendments effect of incidences of *R. solanacearum* on selected solanaceous plants was statistically analyzed by statistical package for social sciences (SPSS) software for Windows, ver. 23 (SPSS, IBM, USA). Chi-square was done to measure the strength of associations between variables. A p-value of <0.05 was considered to be statistically significant.

3. RESULTS AND DISCUSSION

3.1 Influence of Organic and Inorganic Soil Amendments on Disease Incidence on Potatoes

The results of incidences of bacterial wilt on potatoes grown under greenhouse for season 1 and 2 are shown by Figs. 1 and 2 respectively.

Significant differences ($P \leq 0.05$) were revealed in the bacterial wilt incidences in potatoes between control and all the soil amendments used in season 1 and 2 in the five *R. solanacearum* isolate from Kenyan highlands and lowlands. The mean disease index for control and soil amendments; MS+SHG, BT, MS, CM+OP, BT+OP, BT+SHG, CM+SHG, CM and MS+OP for season 1 and 2 were as follows; 2.4, 1, 0.4667, 0.4667, 0.3333, 0.2667, 0.2667, 0.2, 0, 0 and 2.6, 2.067, 0.533, 1.2, 1.667, 0.067, 0, 0.2, 1.333, 2 respectively. These results indicate the suppressive effect of organic and inorganic treatments used in this study. The Brassica tissue + Super-hydro-grow polymer was superior in reducing bacterial wilt incidences in potatoes in all the *R. solanacearum* isolates from Kenyan highlands and lowlands. Brassica species produce glucosinolates which are nematocidal and biocidal. The biocidal action of isothiocyanates produced by Brassica tissue and their potential to manage and suppress phytopathogens has been reported by Brown & Morra [35] and Matthiessen & Kirkegaard [36]. Significant differences ($P \leq 0.05$) were found in the bacterial wilt incidences in potatoes between Brassica tissue alone and Brassica tissue +

Orange peel and Brassica tissue + Super-hydro-grow polymer soil amendments used in both season 1 and 2. Brassica tissue + orange peels, Brassica tissue + Super-hydro-grow polymer and Chalim™ + Super-hydro-grow polymer had a synergetic effect in eliminating *R. solanacearum* in potatoes as opposed to Brassica tissue and Chalim™ used individually as seen in Fig. 1 and 2. Chalim (CaCO₃) in soil reduced bacterial wilt incidence, which was accord with the results of Heyman et al. [37]. The population of *R. solanacearum* was reduced significantly in the

soil with CaCO₃ treatment, which suggested that the effect of CaCO₃ on *R. solanacearum* was mainly related to the role of Ca²⁺ in the greenhouse experiment which increased activity of peroxidase (POD) and polyphenoloxidase (PPO) thus reducing incidences of *R. solanacearum*. Isolate 71 was resistant to Brassica tissue + Orange peel soil amendment causing bacterial wilt while the other isolates were susceptible hence no incidence of bacterial wilt in potatoes to the same treatment in season 2. Variations in the incidence of

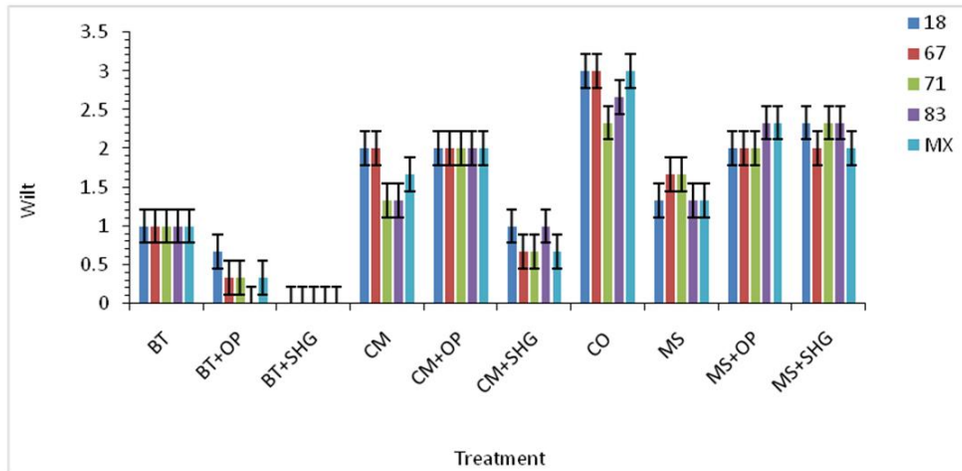


Fig. 1. Greenhouse bacterial wilt incidence in potatoes season 1

BT-Brassicae Tissue, BT+OP -Brassica tissue+Orange peel, BT+SHG- Brassicae Tissue+ Super-hydro-grow polymer, CM- Chalim™, CM+OP- Chalim™+ Orange peel, CM+SHG- Chalim™+ Super-hydro-grow polymer, MS- Metham sodium, Ms+OP- Metham sodium+ Orange peel, MS+SHG- Metham sodium+ Super-hydro-grow polymer : 18 (2T-Kiambu-Low Land), 71(2A-Nyeri-Low Land), 67 (2A-Nyeri-High Land), 83 (2T-Kirinyaga-Highland) and MX (2T-L/2A-L/2A-H/2T-H(18/71/67/83))

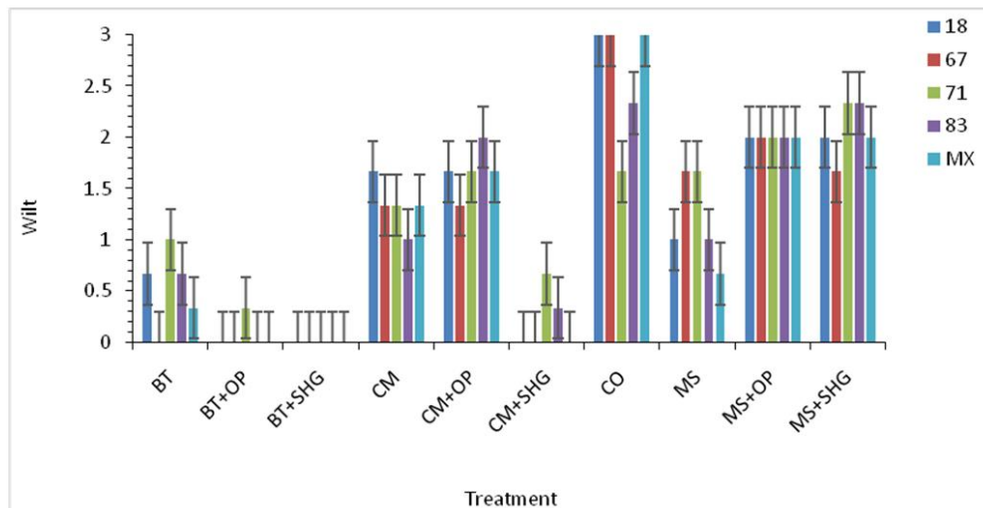


Fig. 2. Greenhouse bacterial wilt incidence in potatoes season 2

bacterial wilt are attributable to the diversity of *R. solanacearum* strains, variations in soil types in different agro ecological zones. There were no significant differences ($P \leq 0.05$) revealed in the bacterial wilt incidences among the isolates in the greenhouse control experiment in potatoes. There was decline in the incidences of bacterial wilt in the second season with subsequent treatment with the same soil amendment except for greenhouse control experiment. Subsequent treatment with inorganic and organic soil treatment tend to drastically reduce *R.*

solanacearum due to suppressive effect of selected soil amendment as opposed to control that was never treated with any soil amendment.

3.2 Influence of Organic and Inorganic Soil Amendments on Disease Incidence on Capsicum

The results of incidences of bacterial wilt on capsicum grown in the greenhouse for season 1 and 2 are shown by Figs. 3 and 4 respectively.

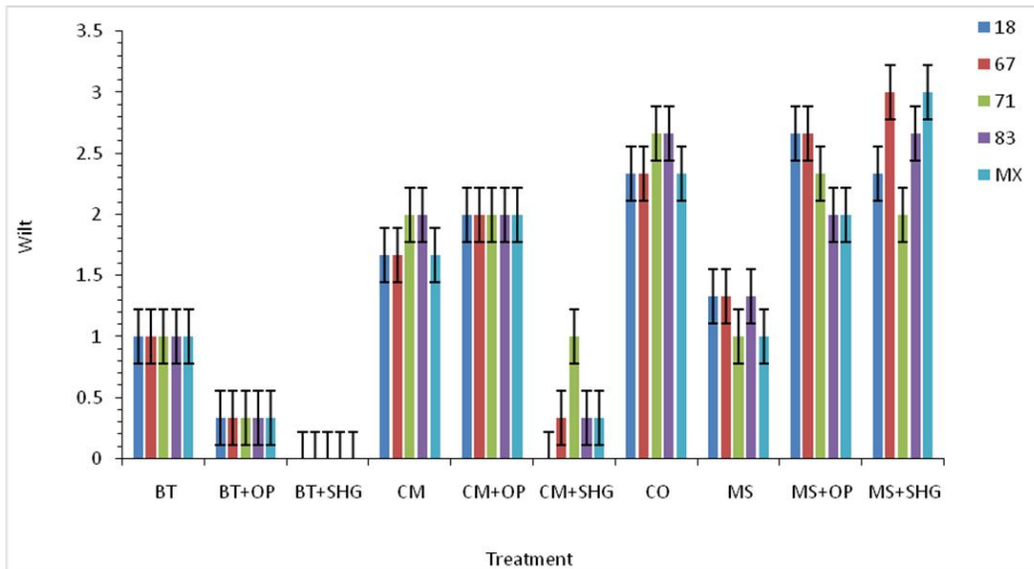


Fig. 3. Greenhouse bacterial wilt incidence in capsicum season 1

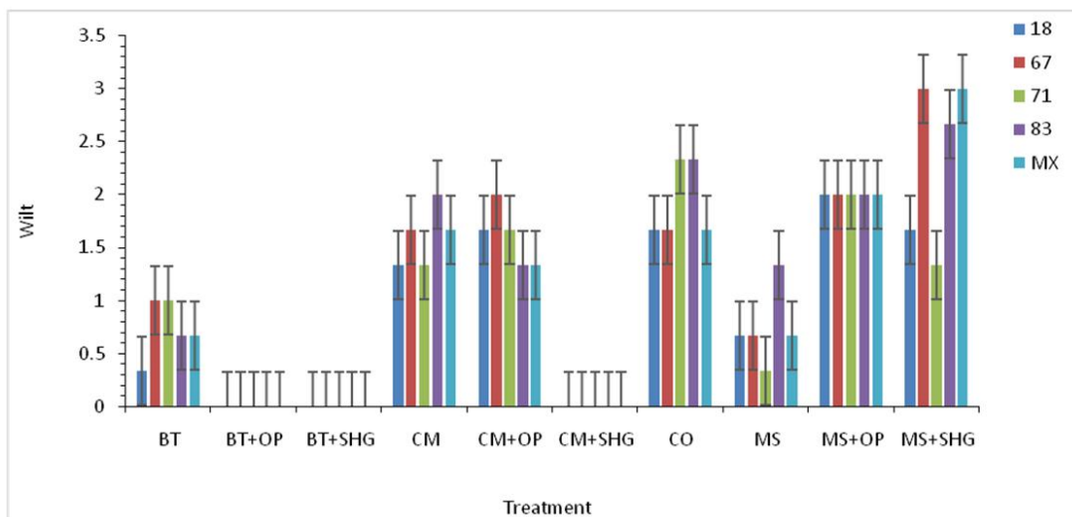


Fig. 4. Greenhouse bacterial wilt incidence in capsicum season 2

There was no significant differences ($P \leq 0.05$) revealed in the bacterial wilt incidences between control and Metham sodium + Super-hydro-grow polymer and Metham sodium + Orange peel in the greenhouse control experiment in capsicum in both season 1 and 2. The Brassica tissue + Super-hydro-grow polymer was superior in reducing bacterial wilt incidences in capsicum in all the *R. solanacearum* isolates from Kenyan highlands and lowlands both in season 1 and 2. Soil amendment in season 2 with Brassica tissue + Orange peel, Brassica tissue + Super-hydro-grow polymer and Chalim™ + Super-hydro-grow polymer recorded zero incidence of bacterial wilt in capsicum in all the isolates. The use of Brassica tissue plant residue in the control of bacterial wilt has been conducted and reported to be effective in a study by [37]. The use of plant tissue in controlling of bacterial wilt has been reported to be eco-friendly relative to the use of inorganic chemicals as well as being readily available to the resource-poor farmers. Combination of Chalim™ + Super-hydro-grow polymer significantly ($P \leq 0.05$) reduced bacterial wilt incidences in capsicum as opposed to use of Chalim™ alone in the two seasons. Metham sodium soil amendment significantly ($P \leq 0.05$) reduced bacterial incidences as opposed to use a combination of Metham sodium + Super-hydro-grow polymer and Metham sodium + Orange peel in the two seasons. The use of Metham sodium + Super-hydro-grow polymer and Metham sodium + Orange peel showed antagonistic effect in control of *R. solanacearum*.

3.3 Influence of Organic and Inorganic Soil Amendments on Disease Incidence on Tomatoes

The results of incidences of bacterial wilt on tomatoes grown under greenhouse for season 1 and 2 are shown by Fig. 5 and 6 respectively.

Significant differences ($P \leq 0.05$) were revealed in the bacterial wilt incidences in Tomatoes between control and all the soil amendments used in season 1 and 2 except in season 1 where Metham sodium + Orange peel and control did not have significant ($P \leq 0.05$) difference as shown on Figs. 5 and 6. The Brassica tissue + Super-hydro-grow polymer was superior in reducing bacterial wilt incidences in tomatoes in all the *R. solanacearum* isolates from Kenyan highlands and lowlands both in season 1 and 2. Brassicaceous materials have been reported to have allelopathic effects as well

as biofumigation effects to soil biota that includes plant parasitic nematodes [38,39,40,41]. The production of biofumigation products including isothiocyanates (ITCs) that has an active ingredient related to that of Metham sodium and dazomet have been reported to be highly toxic to pests and pathogens. Kim et al. [42] reported that, Bacterial wilt of tomatoes caused by *R. solanacearum* is a devastating disease that limits the production of tomato in Korea. Tomato plants are grown worldwide in the field or greenhouse [43]. Isolate 71 was resistant to Brassica tissue + Orange peel soil amendment causing bacterial wilt while the other isolates were susceptible hence no incidence of bacterial wilt in tomatoes to the same treatment in season 2. There was significant difference ($P \leq 0.05$) in bacterial wilt incidence in tomatoes for soil amendment between Brassica tissue soil amendment alone compared to a combination of Brassica tissue + Orange peel and Brassica tissue + Super-hydro-grow polymer in both season 1 and 2. Previous studies have shown that essential oils in citrus, protopine and corydaline alkaloids, lactons, polyacetylene, acyclic sesquiterpenes, hypericin and pseudohypericin compounds are effective toward various bacteria including *R. solanacearum* [44].

In this study, all representative isolates from Kenyan highland and lowland used were pathogenic to tomato seedlings in the greenhouse to varying degrees. Singh et al. [23] showed that the micro-climate inside the greenhouse (Temperatures of 28°C-30°C, 80%-90% RH and wet soil) favors rapid pathogen multiplication and disease. Various findings have reported Ralstonia strains as differing in their virulence [45,46]. Morais et al. [47] reported that information on the pathogenicity and molecular variability of Ralstonia strains will improve our knowledge on the epidemiology and ecology of these pathogens. This is particularly true with respect to latency, survival and aggressiveness of each strain. Bacterial wilt caused by *R. solanacearum* is one of the major diseases of tomato and the disease causes concern for tomato production because it can drastically reduce tomato up to 90% [48,49].

Chalim soil amendment in tomatoes minimally reduced bacterial wilt between the two seasons. Meanwhile, higher Ca^{2+} content in tobacco was associated with less disease, which was agreement with the results of Sugimoto *et al.* [50]; the mechanism may be related to the increased activity of peroxidase (POD) and

polyphenoloxidase (PPO) as reported by Jiang et al. [51] that the severity of tomato wilt can be reduced by increasing activity of POD and PPO in tomato with the increased calcium concentration in tomato tissues which concurs

with our current findings. Mondal et al. [52] also found that Disease incidence in tomato crop was higher compared to other solanaceous crops like brinjal, chilli, capsicum and potato.

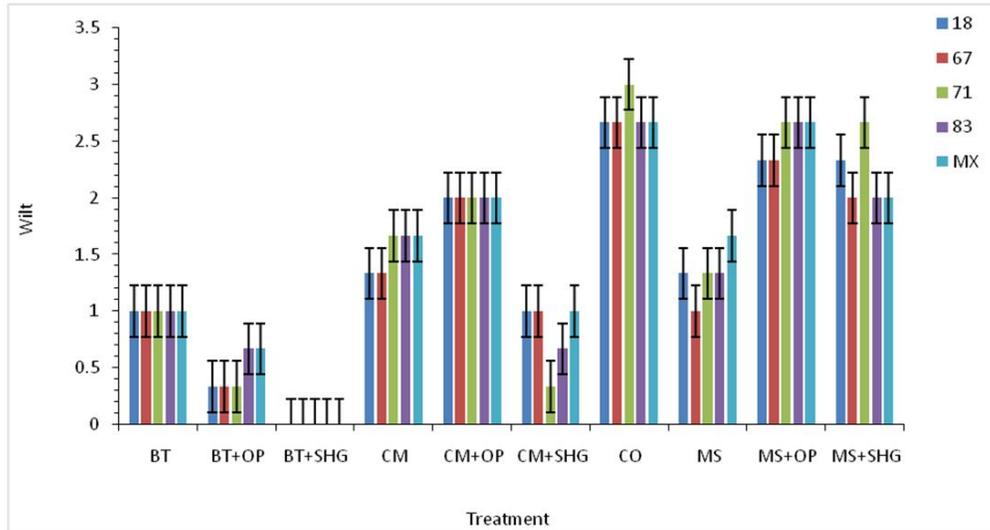


Fig. 5. Greenhouse bacterial wilt incidence in tomatoes season 1

BT- Brassicae Tissue, BT+OP - Brassica tissue+Orange peel, BT+SHG - Brassicae Tissue + Super-hydro-grow polymer, CM - ChalimTM, CM+OP- ChalimTM + Orange peel, CM+SHG - ChalimTM+ Super-hydro-grow polymer, MS - Metham sodium, Ms+OP - Metham sodium+ Orange peel, MS+SHG - Metham sodium + Super-hydro-grow polymer: 18 (2T-Kiambu-Low Land), 71(2A-Nyeri-Low Land), 67 (2A-Nyeri-High Land),83 (2T-Kirinyaga-Highland) and MX (2T-L/2A-L/2A-H/2T-H(18/71/67/83))

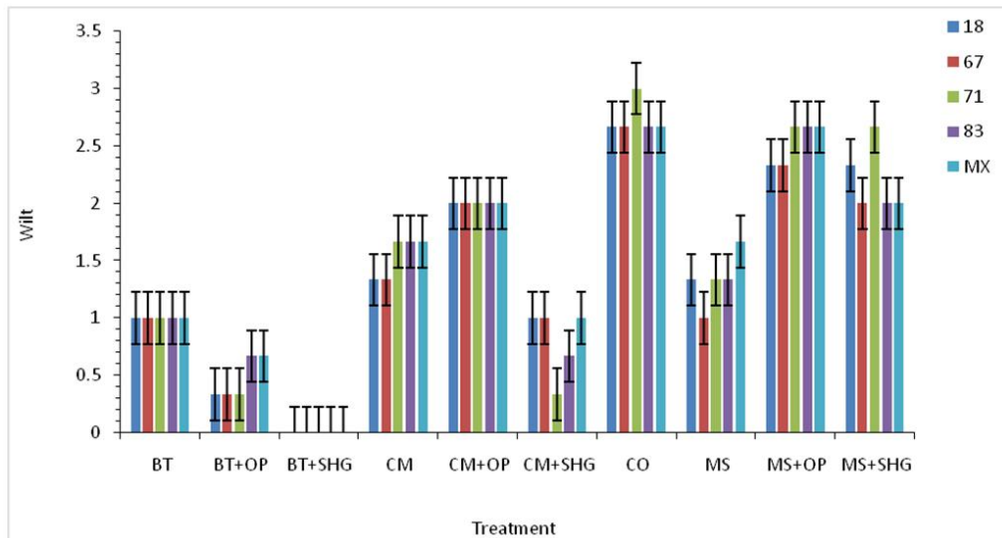


Fig. 6. Greenhouse bacterial wilt incidence in Tomatoes season 2

BT- Brassicae Tissue, BT+OP - Brassica tissue+Orange peel, BT+SHG - Brassicae Tissue + Super-hydro-grow polymer, CM - ChalimTM, CM+OP- ChalimTM + Orange peel, CM+SHG - ChalimTM+ Super-hydro-grow polymer, MS - Metham sodium, Ms+OP - Metham sodium+ Orange peel, MS+SHG - Metham sodium + Super-hydro-grow polymer : 18 (2T-Kiambu-Low Land), 71(2A-Nyeri-Low Land), 67 (2A-Nyeri-High Land), 83 (2T-Kirinyaga-Highland) and MX (2T-L/2A-L/2A-H/2T-H(18/71/67/83))

4. CONCLUSION

In conclusion, our findings showed that organic and inorganic soil amendments could serve as a viable control of bacterial wilt in solanaceous crops caused by *R. solanacearum* in the greenhouse. Brassica tissue + Super-hydro-grow polymer was superior in reducing bacterial wilt incidences in selected solanaceous crops in all the *R. solanacearum* isolates from Kenyan highlands and lowlands both in season 1 and 2. The data presented in this study substantiate the findings that, various *R. solanacearum* isolates from both the Kenyan highland and lowland are causing bacterial wilt disease in various important solanaceous crops grown in the country.

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COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES

1. Beddington J, Asaduzzaman M, Fernandez A, Clark M, Guillou M, Jahn M, Erda L, Mamo T, Van Bo N, Nobre CA, Scholes R, Sharma R, Wakhungu J. Achieving food security in the face of climate change: Summary for policy makers from the Commission on Sustainable Agriculture and Climate Change. CGIAR Research Program on Climate Change, Agriculture and Food Security (CCAFS). Copenhagen, Denmark; 2011. Available: www.ccafs.cgiar.org/commission.
2. Godfray HCJ, Pretty J, Thomas SM, Warham EJ, Beddington JR. Linking policy on climate and food. *Science*. 2011; 331:1013–1014. (DOI:10.1126/science.1202899)
3. Devendra KC, Aundy K, Sajad N. In-vitro Evaluation of *Arabidopsis thaliana* Ecotypes against *Ralstonia solanacearum* Race4, *Int. J. Curr. Microbiol. App. Sci.* 2017;6(5):575-579.
4. Guidot A, Coupat B, Fall S, Prior P and Bertolla F. Horizontal gene transfer between *Ralstonia solanacearum* strains detected by comparative genomic hybridization on microarrays. *ISME J.* 2009;3:549–5. DOI: 10.1038/ismej.2009.14
5. Prior P, Ailloud F, Dalsing BL, Remenant B, Sanchez B, Allen C. Genomic and proteomic evidence supporting the division of the plant pathogen *Ralstonia solanacearum* into three species. *BMC Genom.* 2016;17(90).
6. Wei Y, Moreno CC, Gongora TJ, Wang K, Sang Y, Duran R, Macho AP. The *Ralstonia solanacearum* csp22 peptide, but not flagellin-derived peptides, is perceived by plants from the Solanaceae family. *J Plant Biotech.* 2018;1–14.
7. Sikirou R, Zocli B, Paret ML, Deberdt P, Coranson-Beaudu R, Huat J. First report of bacterial wilt of Gboma (*Solanum macrocarpon*) caused by *Ralstonia solanacearum* in Benin. *Plant Dis.* 2015; 11:1640–1640. DOI: 10.1094/PDIS-02-15-0213-PDN
8. Singh D, Sinha S, Yadav DK, Chaudhary G. Detection of *Ralstonia solanacearum* from asymptomatic tomato plants, irrigation water, and soil through non-selective enrichment medium with hrp gene-based bio-PCR. *Curr Microbiol.* 2014;69:127-134.
9. Jiang Y, Li B, Liu P, Liao F, Weng Q, Chen Q. First report of bacterial wilt caused by *Ralstonia solanacearum* on fig trees in China. *For. Pathol.* 2016;46:256–258. DOI:10.1111/efp.12267
10. Tim M, Prakash P, Carlos L. North Florida Research And Education Center (Nfrec), Quincy; Prakash Pradhanang Post-Doctoral associate, Nfrec, Quincy, FL 32351; Carlos A. Lopes, Researcher, Embrapa Hortaliças, Brasília 70359-970, Brasil; 2008.
11. Joshi M, Srivastava R, Sharma AK and Prakash A. Screening of Resistant Varieties and Antagonistic *Fusarium oxysporum* for Biocontrol of *Fusarium Wilt* of Chilli. *J Plant Pathol Microb.* 2012; 3:134. DOI:10.4172/2157-7471.1000134
12. Muthoni J, Hussein S, Melis R. Management of bacterial wilt *Ralstonia solanacearum* Yabuuchi et al. of Potatoes:

- Opportunity for Host Resistance in Kenya Journal of Agricultural Science. 1995;4(9). [ISSN 1916-9752] E-ISSN. 2012;1916-9760.
13. Meng F. The virulence factors of the bacterial wilt pathogen *Ralstonia solanacearum*. J Plant Pathol Microbiol. 2013;4(3).
 14. Swanson JK, Yao J, Tans-Kersten J, Allen C. Behavior of *Ralstonia solanacearum* race 3 biovar 2 during latent and active infection of geranium. Phytopathology. 2005;95:136–43.
 15. Genin S, Boucher C. *Ralstonia solanacearum*: secrets of a major pathogen unveiled by analysis of its genome. Mol Plant Pathol. 2002;3:111-118.
 16. Ramesh R, Achari GA, Gaitonde S. Genetic diversity of *Ralstonia solanacearum* infecting solanaceous vegetables from India reveals the existence of unknown or newer sequevars of Phylotype I strains. Eur J Plant Pathol. 2014;140:543-562.
 17. Williamson L, Nakaho K, Hudelson B, Allen C. *Ralstonia solanacearum* race 3, biovar 2 strains isolated from geranium are pathogenic on potato. Plant Dis. 2002; 86:987-991.
 18. Kim SH, Olson TN, *Ralstonia solanacearum* race 3, biovar 2, the causal agent of brown rot of potato, identified in geraniums in Pennsylvania, Delaware, and Connecticut (Abstract). Plant Dis. 2003; 87:450.
 19. Cruz APZ, Ferreira V, Pianzola MJ, Siri MI, Coll N, Valls M. A novel, sensitive method to evaluate potato germplasm for bacterial wilt resistance using a luminescent *Ralstonia solanacearum* reporter strain. Mol Plant Microbe Interact. 2014;27:277–285. DOI:10.1094/MPMI
 20. Vrisman CM, Deblais L, Rajashekara G., Miller SA. Differential colonization dynamics of cucurbit hosts by *Erwinia tracheiphila*. Phytopathology. 2016; 106:684-692. Available:https://doi.org/10.1094/PHYTO-11-15-0289-R Link, ISI, Google Scholar.
 21. Bekele B, Abate E, Asefa A. Dickinson M. Incidence of potato viruses and bacterial wilt disease in the west Amhara sub-region of Ethiopia. J Plant Pathol. 2011.93(1):149-157. Available:http://sipav.org/main/jpp/index.php/jpp/article/view/285/151
 22. Assefa M, Dawit W, Lencho A and Hunduma T. Assessment of wilt intensity and identification of causal fungal and bacterial pathogens on hot pepper (*Capsicum annuum* L.) in Bako Tibbe and Nonno districts of west Shewa zone, Ethiopia. Int J Phytopathol. 2015;4:21-28.
 23. Singh, DR, Kumar K, Birah A. Eco-friendly management modules for bacterial wilt (*Ralstonia solanacearum*) of tomato for protected cultivation in a tropical island ecosystem. Biological Agriculture and Horticulture: An International Journal for Sustainable Production Systems; 2014a.
 24. Getachew A, Chemed F, Seid A, Wydra K. Effects of soil amendment on bacterial wilt caused by *Ralstonia solanacearum* and tomato yields in Ethiopia. J Plant Prot Res. 2011;51(1):72–76.
 25. Singh AK, Singh RK, Singh AK, Singh VK, Rawat SS, Mehta KS, Kumar A, Gupta MK, Thakur S. Bio-mulching for ginger crop management: traditional ecological knowledge led adaptation under rainfed agroecosystems. Indian J Tradit Know. 2013;13(1):111–122.
 26. Liu Y, Shi J, Feng Y, Yang X, Li X, Shen Q. Tobacco bacterial wilt can be biologically controlled by the application of antagonistic strains in combination with organic fertilizer. Biol. Fertil. Soils. 2013; 49:447–464.
 27. Tijjani A, Bashir KA, Mohammed I, Muhammad A, Gambo A and Habu M. Biopesticides for pests control: A review. J Biopest Agric. 2016;3(1): 6–13.
 28. Michel VV, Mew TW. Effect of soil amendment on the survival of *Ralstonia solanacearum* in different soils. Phytopathology. 1998;88:300–305.
 29. Michel VV, Wang JF, Midmore DJ, Hartman GL. Effects of intercropping and soil amendment with urea and calcium oxide on the incidence of bacterial wilt of tomato and survival of soil-borne *Pseudomonas solanacearum* in Taiwan. Plant Pathol. 1997;46: 600–610.
 30. Manjunatha SB, Biradar DP, Aladakatti YR. Nanotechnology and its applications in agriculture: A review. J Farm Sc. 2016; 29(1):1-13.

31. Lazarovits G, Tenuta M, Conn KL. Organic amendments as a disease control strategy for soilborne diseases of high-value agricultural crops. *Austr Plant Pathol.* 2001;30(2):111-117.
32. Kago EK, Kinyua ZM, Maingi JM, Okemo PO. Influence of organic and inorganic soil amendments on soil pH and macronutrients. *Journal of Agriculture and Ecology Research International.* 2019;1-10.
33. Jaetzold R, Schmidt H, Hornetz B, Shisanya C. Central Kenya. Agroecological zones and subzones. Ministry of Agriculture, Farm Management Hand book of Kenya. Natural conditions and Farm Management information 2nd Edition Part B, Central Province: 2006;II:434-438.
34. Mwaniki PK, Birech R, Wagara IN, Kinyua ZM and Freyer B. Distribution, Prevalence and Incidence of Potato Bacterial Wilt in Nakuru County, KENYA. *Inter J of Innov Res and Dev.* 2016;5(1).
35. Brown PD, Morra MJ. Control of soil-borne plant pests using glucosinolate containing plants. *Adv Agron.* 1997;61:167-231.
36. Matthiessen JN, Kirkegaard JA. Biofumigation for managing soil-borne pests - progress, pitfalls and prospects. In: Zalucki M, Drew R, White G. eds. *Proceedings of the 6th Australasian Applied Entomology Research Conference.* University of Queensland. 1998;1:364- 372.
37. Heyman F, Lindahl B, Persson L, Wikstrom M and Stenlid J. Calcium concentrations of soil affect suppressiveness against *Aphanomyces* root rot of pea. *Soil Biol. Biochem.* 2007;39:2222-2229.
38. Gruver LS, Weil RR, Zasada IA, Sardanelli S and Momena B. Brassicaceous and rye cover crops altered free-living soil nematode community composition. *Appl Soil Ecol.* 2010;45:1-12.
39. Hartman GL, Hong WF, Hayward AC. Potential of biological and chemical control of bacterial wilt. In: Hartman GL, Hayward AC. eds. *Bacterial wilt.* Proceedings of ACIAR conference, 1992, Kaohsiung, Taiwan. ACIAR Proceedings No. 45. Brisbane, Australia, Watson Ferguson and Company. 1993;322-326.
40. Bailey KL, Lazarovits G. Suppressing soil borne diseases with residue management and organic amendments. *Soil and Tillage.* 2003;72(2):169-180.
41. Schonfeld J, Gelsomin A, van Overbeek LS, Goris-sen A, Smalla K, van Elsas JD. Effects of compost addition and stimulated solarisation on the fate of *Ralstonia solanacearum* biovar 2 and indigenous bacteria in soil. *FMES Microbiology Ecology.* 2003;43:63-74.
42. Kim SG, Hur OS, Ro NY, Ko HC, Rhee JH, Sung JS, Ryu KY, Lee SY, Baek HJ. Evaluation of resistance to *Ralstonia solanacearum* in tomato genetic resources at seedling stage. *Plant Pathol J.* 2016; 32(1):58-64.
43. Peralta IE, Spooner DM. History, origin and early cultivation of tomato (*Solanaceae*). In: Razdan MK, Mattoo AK, eds. *Genetic improvement of solanaceous crops. Tomato.* Enfield, NH:Science Publishers. 2007;2:1–27.
44. Maruti J, Dhanavade CB, Jalkute KD, Sonawane K, Jai SG. Study antimicrobial activity of lemon (*Citrus lemon* L) peel extract. *Br J Pharmacol Toxicol.* 2011;2 (3):119-122.
45. Li Y, Feng J, Liu H, Wang L, Hsiang T, Li X, Huang J. Genetic diversity and pathogenicity of *Ralstonia solanacearum* causing tobacco bacterial wilt in China. *Plant Dis.* 2016;100:1288–1296.
46. Rodrigues L, Destefano S, da Silva M, Costa G, Maringoni AC. Characterization of *Ralstonia solanacearum* strains from Brazil using molecular methods and pathogenicity tests. *J Plant Pathol.* 2012; 94:505–516.
47. Morais TP, Lopes CA, Tebaldi ND, Luz JMQ. Occurrence and diversity of *Ralstonia solanacearum* populations in Brazil. *Biosci. J.* 2015;31:1722–1737.
48. Fujiwara K, Aoyama C, Takano M, Shinohara M. Suppression of *Ralstonia solanacearum* bacterial wilt disease by an organic hydroponic system. *J Gen Plant Pathol.* 2012;78:217–220.
49. Aslam MN, Mukhtar T, Hussain MA, Raheel M. Assessment of resistance to bacterial wilt incited by *Ralstonia solanacearum* in tomato germplasm. *J Plant Dis Protect.* 2017;124:585–590.
50. Sugimoto T, Watanabe K, Yoshida S, Aino M, Furiki M, Shiono M. Field application of calcium to reduce phytophthora stem rot of

- soybean, and calcium distribution in plants. Plant Dis. 2010;94:812–819.
DOI: 10.1094/PDIS-94-7-0812
51. Jiang JF, Li JG, Dong YH. Effect of calcium nutrition on resistance of tomato against bacterial wilt induced by *Ralstonia solanacearum*. Eur J Plant Pathol. 2013; 136:547–555.
52. Mondal B, Bhattacharya I, Khatua DC. Crop and weed host of *Ralstonia solanacearum* in West Bengal. J Crop Weed. 2011;7(2):195-199.

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