IDENTIFICATION OF THE BACTERIUM TOMATO STEM CANKER

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ABSTRACT

Diseased tomato samples were collected from greenhouse was evaluated for isolation, pathogenicity and biochemical tests. The symptoms of the infected tomato plants were as sudden wilting after curled on leaves and necrotic streak regions developed at the crown and base of the stem and the cavities deepen and expand up and down, brown discoloration and necrosis occurring on xylem and phloem vascular. All of ages of tomato plant were susceptible to bacteria when the weather condition favorable and immediately, seen collapse symptom on tomato plant at once fail and die. The bacterium was isolated from diseased plant in all regions on nutrient Agar; a yellow bacterium was isolated from infected tomato plant in greenhouses and fields in Abu-Ghraib, Rashiedia and Qanat Al-Geiaysh nurseries in Baghdad provinces of Iraq. The bacterium was found gram positive, rod-shaped, non-motile and capable an aerobic growth and based on the morphological and biochemical characteristics revealed that this bacterium belongs to: *Clavibacter michiganensis* subsp. *michiganensis*. (smith) pathogenicity and hypersensitivity of the bacterium Cmm showed the disease index were 18.33, 6.66, 16.66, 5, 0% for tomato seedlings were inoculated treatments as the wounding roots, without wounding roots, crown of the stem, petiole and control respectively.

Keywords: Clavibacter Michiganensis Subsp, Michiganensis, Tomato, Canker

1. INTRODUCTION

Bacterial canker of tomato (*Lycopersicon esculentum* Mill.), caused by *Clavibacter michiganensis* subsp. *michiganensis* is one of the most destructive diseases of tomato and has caused major economic losses in commercial tomato production worldwide the disease can occur on other members of the Solanaceae plants as alternative hosts (Agrawal *et al*., 2012; Fatmi and Schaad, 2002; Gleason *et al*., 1993; Leandro, 2011). The severity of disease is very high, the presence of less than 1% infected seeds can cause 60-70% crop loss, the epidemics in the fields and green house can be induced by few plants (Burokiene *et al*., 2005; Strider, 1969). The disease was first reported at the beginning of the twentieth century in Michigan (USA) and currently it is present worldwide (Leandro, 2011), it caused losses in Canada, most European countries, morocco, Kenya, South Africa, Australia and New Zealand (Agrawal *et al*., 2012). Symptoms of systemic infection, particularly wilting, usually appear first and localized symptoms such as marginal necrosis and leaflet spotting may appear first. To further complicate matters, localized infections also can progress into the vascular bundles and lead to systemic symptoms under certain circumstances. Marginal necrosis of leaflets frequently is an early symptom of localized infection. Sometimes referred to as the “firing stage” this appears first as distinct brown, dried margins on lower leaflets (Gleason *et al*., 1993).

The first of tomato bacterial canker symptoms were found in the fields and green houses in Iraq in Abu-Ghraib, Rashiedia and Qanat Al-Geiaysh nurseries in Baghdad. The study aims to describe tomato plant bacterial canker
pathogens and to test pathogenicity, determine morphological and biochemical of the bacterial pathogen.

2. MATERIALS AND METHODS

2.1. Sampling and Isolation

The disease ratio was determined by counting infected tomato plants per 100 plants per greenhouse. Tomato stem canker was identified in the fields and greenhouses in Abu-Ghraib, Rashiedia and Qanat Al-Geaysh nurseries in Baghdad provinces of Iraq. The symptoms appear on the leaves, petioles and fruits, pale yellow to brown streaks develop along the stems and under of petioles, these darken streaks sometimes open, resulting dark brown cankers and pith necrosis symptoms (Fig. 1). The vascular parenchyma in particular has a mealy appearance resulting from bacterial degradation and ooze production (Fig. 2). The samples were collected and kept in refrigerator at 4°C in the laboratory, the infected leaves and stems samples were washed and cut 0.5-1 cm length from the margin of infected parts sterilized in 1.5% sodium hypochlorite for 2 min then rinsed by sterile distilled water, dried by sterilized filter paper, cultured on nutrient agar NA as 4-5 pieces/plate and incubated at 28°C for 2-3 days. The bacterial cultures were purified by a single cell isolation by streaking on NA medium as described by (Kelman, 1954) with a sterile circumstance to getting pure bacterial colonies.

2.2. Hypersensitivity Reaction Test

The Hypersensitivity Reaction (HR) is a rapid indicator test for pathogenicity. Inoculated a Cnm bacterial suspension from 24-48 h culture grown on NA in tobacco leaves by injection into the intraveinal areas of tobacco leaves (Burokiene et al., 2005; Klement and Goodman, 1967), HR determined after 24 h.

2.3. Pathogenicity Test

Tomato seedling two weeks old were planted in sterilized soil by autoclave under 121°C and pressure 1.5 Kg.2Cm. for 20 min. with three plant were inoculated for each treatment and covered with polyethylene bags to kept the humidity, by bacterial inoculums suspension was prepared by using two-day-old bacterial cultures by sterilize needle dipped in bacterial suspension and the test was carried out as follow:

- Seedling inoculated after wounding and cutting the roots by pulling the seedling from peat moss discs and transplanted in the pots after dipping the roots in bacterial suspension
- Seedling inoculated without wounding the roots
- Seedling inoculated after making wounds on the crown of the stem, then injected with bacterial suspension
- Seedling inoculated in the petiole
- Seedling treated without inoculation as control

![Fig. 1. Stem canker and pith necrosis symptoms](image-url)
Each treatment replicated three times and left in
green house to evaluation disease severity depending on
the scale of (Foster and Echand, 1973) as follow degrees:

- No infection
- Willing more than 1/3 of the leaves
- Willing more than 2/3 of the leaves
- Willing more than 2/3 of the leaves but the top
leaves didn’t wilt
- All the leaves wilting and plant dying

The disease index was calculated on Tomato seedling
by Equation (Praveena and Naseema, 2004) by follow:

\[
\text{Disease index (DI) } \% = \frac{\text{No. plant degree } 0 \times 0 \ldots \text{No. plant in degree } 4 \times 4}{\text{No. of plant all degree } \times \text{max. degree of infection}}
\]

2.4. The Biochemical Tests

Biochemical tests had been used after isolation and
purified the culture to support the identification of isolates to the genus and species level depending on morphological and biochemical properties of colonies, Cell shape, Motility, Colony Characteristics, Gram-
staining, Anaerobic growth, Catalase reaction, Production
\(H_2S\), Indole, Red methyl reaction, Acid production, Gelatin
analysis, Nitrate reduction (Bergey et al., 1923; Fahy and
Persely, 1983; Schaad, 1980).

3. RESULTS

3.1. Isolation and Diagnosis

The results showed that isolate which infected tomato
plants in the field and green house in Baghdad provinces,
belonged to *Clavibacter michiganensis* subsp.
michiganensis*. Based on the above tests, which were
conducted in the laboratory on samples that include
isolation, purifying the cultures, HR and pathogenicity
test and morphological biochemical tests.

3.2. Hypersensitivity Reaction Test

The results showed after 24 h as necrosis spotting on
the leaf as compared to control treatment no necrosis
spot was showing on the leaf and no wilting.

3.3. Pathogenicity Test

The disease symptoms were observed in all
treatments tomato seedlings and The disease index
were 18.33, 6.66, 16.66, 5.0% for treatments of tomato
seedlings were inoculated by the wounding roots,
without wounding roots, crown of the stem, petiole and
control, respectively as showed in (Table 1) and Re-
isolation of tomato stem canker pathogen from tomato
plants were positive in all cases. There were significant
differences between the treatments at \(P = 0.05\). Symptoms
appeared on all treatments and at different intervals of
incubation periods, the faster of the visible symptoms
were observed in the treatment of stem inoculation,
symptoms that appeared after 10 days and was delayed in
the treatment of the wounded roots to three weeks and
more and also indicated by (Gleason et al., 1993).

Pathogenicity test indicated that the seedlings showed
wilting symptoms earlier dying and also showed that
incubation periods were longer and disease severity
decreased in some treatments. The earliest symptoms
occurred after 7-10 days on the stems of tomato
seedlings treatment either seedling with roots wounded
treated wilted after 3-4 weeks (Behrendt et al., 2002).
Many studies described the mechanisms involved in

![Fig. 2. Infected vascular stems tissues with mealy appearance of vascular parenchyma](image-url)
bacterial wilt carry one of the factors causing wilting plant is the production of exopolysaccharides (Kiraly et al. 1997) which are responsible for the wilting symptoms caused by Cmm either by plugging xylem vessels or through a phytotoxic effect (Jahr et al., 1999) as well as some extracellular enzymes such as endocellulase (Meletzus et al., 1993), pectinmethylsterase (Strider, 1969), xylanase (Beimen et al., 1992), hydralase (Benhamou, 1991). The wilt symptoms caused by C. michiganensis subsp. michiganensis may be confused with other systemic tomato diseases caused by Rhizoctonia solani, Fusarium spp. and Verticillium spp. especially at the first stages of the disease which increases the complexity of the disease and came to the conclusion that the onset of symptoms on the seedlings were different among all treatments due to the different ways to penetrate the causes of bacterial perhaps through wounds in the roots or stems, which explains Several studies suggest that the factor that is most influential in the epidemiology of diseases lesion is the period between infection and the onset of symptoms, where the duration of the incubation ranging from 7-84 day and affected by temperature, plant age, inoculums concentration of pathogen and the longer period for incubation was founded, in older plants, cool temperature or warmer than 25°C and lower inoculums concentration (Burokiene et al., 2005; Gleason et al., 1993).

Table 1. Showing disease index percentage Clavibacter michiganensis subsp. michiganensis of tomato canker on the inoculated tomato seedling

| No. treatment | Disease index (%) |
|---------------|------------------|
| 1-Inoculated the wounding roots | 18.33 |
| 2-Inoculated without wounding roots | 06.66 |
| 3-Inoculated crown of the stem | 16.66 |
| 4-Inoculated petiole | 05.00 |
| 5-Control | 00.00 |

Table 2. Morphological and biochemical characterizes of Clavibacter michiganensis subsp. michiganensis

| Characteristics | Pathogen clavibacter michiganensis subsp. michiganensis |
|-----------------|--------------------------------------------------------|
| Cell shell      | Coryneform curved rod                                  |
| Motility        | Non-motile                                             |
| Colony          | smooth, shining, round                                 |
| Characteristics | yellow colonies, entire margins.                        |
| Gram- staining  | Positive +                                             |
| Growth          | Optimum temp 27-28°C                                    |
| Anaerobic growth| aerobic                                                |
| Catalase reaction| +                                                      |
| H₂S Production  | -                                                       |
| Indole production| -                                                     |
| Red methyl reaction| -                                                      |
| Acide production| +                                                      |
| Gelatin analysis| w+                                                      |
| Nitrate reduction| -                                                      |

+= positive reaction, -= negative reaction, w = weak

Fig. 3. Clavibacter michiganensis subsp. michiganensis cells
3.4. The Biochemical Tests

Bacterium was diagnosing after purified depending upon morphological biochemical properties of colonies, Cells of *C. michiganensis* subsp. *michiganensis* are Gram-positive, Non-motile, non-spore-forming, short coryneform rods (Fig. 3) (Bradbury, 1986; Gartemann et al., 2003; Hayward and Waterston, 1964), which may be straight to Slightly curved shaped had been identified depending on many biochemical tests carried out in the laboratory as showed that bacteria hydrolyzed weakly for gelatin (Berger et al., 1923) studying are representative selection of coryneform bacteria found that *Clavibacter michiganensis* subsp. *michiganensis* hydrolyzed the gelatin very weakly after four weeks of incubation and this similar to what we founded (Table 2).

4. CONCLUSION

The recording of the study clearly indicated that stem canker of tomato (*Lycopersicon esculentum* Mill.) is caused by *Clavibacter michiganensis* subsp. *michiganensis* which was observed in all nurseries in Abu-Ghraib, Rasheeda and Qanat Al-Jaysh in Baghdad province of Iraq. The disease showed a great economic losses in tomato crop, All of ages of tomato plant were susceptible to bacteria when the weather conditions favorable and immediately, seen weakness symptom on tomato plant suddenly collapses and die. And management strategies have not been realized yet due to the lack of identification of the causal organism with molecular analysis technique and lack of appropriate strategy for the integrated management programs for the protection and the eradication of disease. This is helping to spread bacterial infection for interference with fungal infection.

5. REFERENCES

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