VIRULENCE OF ALTERNARIA ALTERNATA INFECTING NONI ASSOCIATED WITH PRODUCTION OF CELL WALL DEGRADING ENZYMES

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Abstract: Different methods of inoculations were tried for proving Koch postulates. Among them, the pin prick plus spore suspension spray was found to be the best method used in a glasshouse. Out of the fifteen isolates of Alternaria alternata, isolate AA1 was highly virulent and AA6 was avirulent. The virulent isolate of A. alternata produced more cellulolytic (C1 and Cx) and pectinolytic (pectin methyl esterase and endo-polygalacturonase) enzymes in vitro than the avirulent one. The activity of cellulolytic enzymes increased with the increase in the age of the culture. Whereas, the pectinolytic enzymes were highly active in 10-day-old culture and the activities decreased with the increase in the age of the culture. Then again, the activity of enzymes produced by the avirulent isolate of pathogens did not decrease and these enzyme activities increased with the increase in the age of the culture.

Key words: Noni, leaf blight, virulent, avirulent, cellulolytic and pectinolytic enzymes and mycelial dry weight

INTRODUCTION

Morinda citrifolia, popularly known as noni, has been generating great enthusiasm within the green health industry for the past two decades. Noni has been promoted as a treatment for a vast array of medical conditions ranging from cancer to sexual dysfunction. In India it is distributed in Tamil Nadu, Karnataka, Andhra Pradesh, Orissa, Madhya Pradesh, Gujarat, Rajasthan and Andaman Nicobar islands; which means; over 1,850 acres (Peter 2009). The intensive cultivation of noni has resulted in the outbreak of several foliar diseases caused by fungi, and can reduce leaf growth and fruit development (Nelson 2006). Leaf blight caused by Alternaria alternata, is one from among all the important diseases that threaten noni production.

Virulence is the measure or degree of pathogenicity of an isolate or race in the host (Singh 2002). It is well established that virulence of different pathogens, of the same host, vary. Even the isolates of same pathogen exhibit differences. Virulence of a particular pathogen is mainly governed by the capacity to produce toxins, enzymes and other physiological factors. Although for many plant pathogens, a capacity to breach the cell walls of their hosts is not required for entry. This is because these pathogens rely on wounds, natural openings or vectors. Many fungal pathogens achieve entry by mechanical force or enzyme activity or a combination of both. A considerable amount of literature has accumulated that implicates degradative enzymes in pathogenesis or virulence. Cell wall degrading enzymes released by pathogens are known to be responsible for pathogenesis. The ability of a pathogen to produce cellulolytic and pectinolytic enzymes determines the degree of cell wall degradation during pathogenesis and inhibition of these enzymes, and ultimately affects the disease development.

A numbers of cell wall degrading enzymes have been shown to be produced by plant pathogens (Chenglin et al. 1996) which are known to facilitate cell wall penetration and tissue maceration in host plants. These microorganisms produce metabolites in culture media and plant tissues which were involved in the disease syndrome (Wood et al. 1972). Several species of Alternaria were known to produce different types of metabolites (Bhaskaran and Kandaswamy 1978). The aims of the present study were to select the best method of plant inoculation and to understand the pathogenicity mechanism, and also to assess virulence of the isolates and its relevance to the production of cell wall degrading enzymes and pathogenesis.

MATERIALS AND METHODS

Collection, isolation and establishment of isolates
The infected noni leaves showing typical symptom of leaf blight were collected from Tamil Nadu, Karnataka and Kerala. The pathogens were isolated by the tissue segment method (Rangaswami 1958). The fungus was...
purified by the single spore isolation technique (Ricker and Ricker 1936) and the purified isolates were maintained on PDA slants for further studies.

Methods of Inoculation

Five different methods of inoculation were tried for pathogenicity testing, to select the best method which may be standardized for proving pathogenicity and to study pathogen virulence. The isolate AA1 of *A. alternata* was used for this study. While in a glasshouse, two month old seedlings were inoculated through different methods (mycelial inoculation, pin prick plus mycelial inoculation, spore suspension spray, pin prick plus spore suspension spray, and spore injection). The seedlings were exposed to 80 per cent RH, both 24 h prior and after inoculation, by covering the seedlings with a polythene bag and spraying water inside the bag. The inoculation was done during the cool evening hours. Three replications were maintained in each method. Fifteen days after inoculation, the disease intensity was recorded in all methods.

Assessment of PDI

Plants showing the symptoms of leaf blight were assessed as per the severity grade from 0–9 and the per cent disease index, were calculated. The disease intensity of leaf blight was assessed as proposed by TNAU (1980), according to a grade chart: 0 – No. infection, 1 – 0 to 10.00, 3 – 10.1 to 15.00, 5 – 15.1 to 25.00, 7 – 25.1 to 50.00, 9 – more than 50.00 per cent of leaf area infected.

The per cent disease index (PDI) was calculated using Mc Kinney’s (1923) formula:

\[
PDI = \frac{\text{Sum of all numerical grades}}{\text{Total number of leaves counted}} \times \text{Maximum category value}
\]

Virulence of isolates

Various isolates of *A. alternata* were inoculated onto two month old seedlings of noni maintained in a glasshouse, using the pin prick plus spore suspension spray method (Lakshmanan et al. 1990). Three replications were maintained for each isolate with five seedlings for each replication. The symptoms were observed 15 days after inoculation. Ten leaves per plant were randomly selected for each isolate and the PDI was calculated according to Mc Kinney’s formula (1923).

Production of cell wall degrading enzymes

To study the *in vitro* production of pectinolytic and cellulolytic enzymes, the isolates; namely AA1 and AA6, were grown on Czapek dox broth (pH 7–7.5) wherein the carbon source was substituted with one per cent pectin (for pectic enzymes) or one per cent carboxy methyl cellulose (for cellulolytic enzymes). The media were inoculated with a 9 mm diameter of a nine day old actively growing culture disc, of each isolate. The culture filtrates were obtained after inoculation at room temperature (27±1°C) for 5, 10, 15 and 20 days and centrifuged at 3000 g, for 20 min. For the assay of pectinolytic enzymes, the culture filtrates were dialyzed for 18 h against distilled water at 40°C. The dialysate served as enzyme source. As dialysis was found to reduce the activity of cellulolytic enzymes (Bateman 1964), the culture filtrates as such were used for the assay of cellulases.

Assay of cellulolytic enzymes

**Cellulase (C<sub>i</sub>) activity**

Cellulase (C<sub>i</sub>) activity was assayed by the method of Norkrans (1950). The assay mixture contained 1 ml of cellulose solution (the concentration of which was adjusted to give approximately 0.85 absorbance at 610 nm), four ml of 0.1 M phosphate buffer (pH 7.0), and 5 ml of enzyme source. The absorbance of the assay mixture was determined at 610 nm in a Spectronic – 20 immediately upon the addition of the enzyme source, and again after an incubation period of 24 h at 27°C. The enzyme activity was expressed in units (1 unit = change in absorbance of 0.01).

**Cellulase (C<sub>c</sub>) activity**

Cellulase (C<sub>c</sub>) activity was assayed by the viscosimetric method of Hancock et al. (1964) using Carboxy methyl cellulose (CMC) as the substrate. Two ml of enzyme extract was added to four ml of 1.2 per cent carboxy methyl cellulose solution buffered at pH 5.0 with sodium citrate buffer. The loss of viscosity of the CMC solution was determined by means of an Ostwald-Fenske viscosimeter size 150 at five min. intervals, for up to 15 min. The enzyme source which boiled for 10 min. at 100°C, served as the check. The results were expressed as the per cent loss in viscosity in 15 min.

\[
V = \frac{T_0 - T_1}{T_0 - T_w} \times 100
\]

where:

- **V** – per cent loss of viscosity,
- **T<sub>0</sub>** – flow time in seconds at zero time,
- **T<sub>1</sub>** – flow time of the reaction mixture at time **T<sub>1</sub>**,
- **T<sub>w</sub>** – flow time of distilled water.

Assay of pectinolytic enzymes

**Macerating enzymes**

Macerating enzyme activity was assessed by the method described by Mahadevan (1965). Potato discs of nine mm diameter and 30 µm thickness were obtained by using a hand microtome. Ten potato discs were placed in a sterile Petri plate and 10 ml of the culture filtrate were added. The uninoculated medium served as the control. The coherence of the potato discs was tested at different time intervals using a sterile glass rod. The enzyme activity was expressed as the time taken in hours for maceration of potato discs.

**Pectin methyl esterase (PME)**

Pectin methyl esterase activity was estimated following the procedure described by Gupta (1970). Pipetted out 20 ml of pectin solution and its pH were adjusted to 7.0 using one N sodium hydroxide. To this, 10 ml of enzyme solution was added and its pH was adjusted immediately
to 7.0 in the pH meter by adding one N sodium hydroxide. The enzyme substrate mixture was incubated for 24 h. The pH was readjusted to seven with 0.02 N sodium hydroxide, which was equal to the enzyme activity and the enzymatic activity, and was expressed in terms of units (one unit is 0.1 ml of 0.02 N sodium hydroxide used).

Endo-polygalacturonase (endo PG)

Endo-PG activity was estimated by the standard viscosimetric method (Hancock et al. 1964) using three ml of enzyme source, one ml of 1.2 per cent sodium polypectate and one ml of 0.5 M ammonium acetate buffer. The loss in viscosity of the pectate solution was determined by means of the Ostwald-Fenske viscosimeter size 150, at five min intervals for up to 15 min. The enzyme source which boiled for 10 min. at 100°C served as the check. The results were expressed as the per cent loss in viscosity in 15 min.

Statistical analysis

All the experiments were repeated once with similar results. The data were statistically analyzed (Gomez and Gomez 1984). The treatment means were compared by Duncan’s multiple range test (DMRT). The package used for analysis was IRRISTAT version 92-a developed by International Rice Research Institute Biometrics Units, The Philippines.

RESULTS

Collection and isolation of pathogens

Leaves showing typical symptoms of leaf blight were collected from fifteen noni growing areas of Tamil Nadu, Karnataka and Kerala. From these leaf samples, a total of fifteen isolates were isolated and purified.

Methods of inoculation

The results indicated that the intensity of disease symptoms varied significantly with that of different methods of inoculation. Among which, the pin prick with spore suspension spray method was the best, which recorded 58.60 PDI, followed by the pin prick with mycelial inoculation method (52.23 PDI). The lowest PDI was recorded in the mycelial inoculation method (16.37 PDI) (Fig. 1). Hence, for further studies, the pin prick plus spore suspension spray was used.

Virulence of the isolates

The data from the virulence study clearly indicated that different isolates varied with respect to virulence. The maximum PDI was recorded in AA1 (61.00 PDI) followed by AA10 (60.33 PDI), AA11 (56.34 PDI) and AA2 (54.07 PDI). The minimum disease intensity was recorded in AA6 which showed 10.33 PDI. The results clearly indicated that isolate AA1 was highly virulent and isolate AA6 was least virulent. Hence, these two isolates were used for production of cell wall degrading enzymes (Fig. 2).

Fig. 1. Effect of different inoculation methods on severity of leaf blight disease under glasshouse conditions

Fig. 2. Virulence of A. alternata isolates on noni seedlings under glasshouse conditions
Production of cell wall degrading enzymes

*In vitro production of cellulolytic enzymes*

The enzyme production increased with an increase in the incubation period. The virulent isolate of *A. alternata* produced more cellulolytic enzymes (C₃ and C₄) than the avirulent ones. The C₃ activity was found to be more (12.00 units) in 20-day-old culture filtrate of virulent isolate of *A. alternata* than the avirulent ones (0.90 units). Similarly, the C₄ enzyme activity was found to be more (76.00 per cent loss of viscosity), in 20 day old culture filtrate of virulent isolate than the avirulent ones (0.90 units). Mycelial dry weight also increased with an increase in the incubation period. The dry weight of mycelium was higher in the virulent isolate (480 mg) as compared to the avirulent ones (95 mg) (Table 1).

Table 1. Production of cellulolytic enzymes by *A. alternata*

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Incubation time [days]</th>
<th>C₃</th>
<th>C₄</th>
<th>Mycelial dry weight [mg]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Virulent (AA1)</td>
<td>5</td>
<td>6.50 d</td>
<td>42.50 d (40.68)</td>
<td>240.00 d</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>7.30 c</td>
<td>51.80 c (46.03)</td>
<td>375.00 c</td>
</tr>
<tr>
<td></td>
<td>15</td>
<td>10.50 b</td>
<td>71.00 b (57.41)</td>
<td>475.00 b</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>12.00 a</td>
<td>76.00 a (60.66)</td>
<td>480.00 a</td>
</tr>
<tr>
<td>Avirulent (AA6)</td>
<td>5</td>
<td>0.20 c</td>
<td>00.50 c (4.05)</td>
<td>70.00 d</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>0.20 c</td>
<td>00.70 b (4.79)</td>
<td>75.00 c</td>
</tr>
<tr>
<td></td>
<td>15</td>
<td>0.70 b</td>
<td>00.90 a (5.44)</td>
<td>85.00 b</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>0.90 a</td>
<td>00.90 a (5.44)</td>
<td>95.00 a</td>
</tr>
</tbody>
</table>

1 enzyme activity in units; 2 per cent loss of viscosity; 3 mean of three replications

Values in parentheses are arcsine transformed values

*In vitro production of pectinolytic enzymes*

In the virulent isolate, the production of the pectinolytic enzymes viz., macerating enzymes, Pectin methyl esterase (PME) and Endo polygalacturonase (Endo-PG) increased for up to 10 days and thereafter the activities of the enzymes decreased. Maximum enzyme activity was observed in the culture filtrate 10 days after incubation. In the avirulent isolate, the enzyme production increased with an increase in the incubation period (throughout the experimental period) but it was much less than the virulent isolates. The mycelial weight increased with an increase in the incubation period in both virulent and avirulent isolates of *A. alternata* (Table 2).

Table 2. Production of pectinolytic enzymes by *A. alternata*

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Incubation time [days]</th>
<th>Macerating enzymes⁴</th>
<th>Pectin methyl esterase⁴</th>
<th>Endopolygalacturonase⁴</th>
<th>Mycelial dry weight [g]⁴</th>
</tr>
</thead>
<tbody>
<tr>
<td>Virulent (AA1)</td>
<td>5</td>
<td>15.00 c</td>
<td>04.50 c</td>
<td>23.50 c (28.99)</td>
<td>250.00 d</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>15.00 c</td>
<td>10.75 a</td>
<td>59.70 a (30.99)</td>
<td>385.00 c</td>
</tr>
<tr>
<td></td>
<td>15</td>
<td>18.00 b</td>
<td>08.75 b</td>
<td>40.00 b (39.23)</td>
<td>590.00 b</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>24.00 b</td>
<td>08.00 b</td>
<td>24.00 b (29.33)</td>
<td>695.00 a</td>
</tr>
<tr>
<td>Avirulent (AA6)</td>
<td>5</td>
<td>no maceration</td>
<td>01.00 c</td>
<td>04.30 d (11.96)</td>
<td>060.00 d</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>no maceration</td>
<td>02.75 b</td>
<td>10.10 c (18.53)</td>
<td>075.00 c</td>
</tr>
<tr>
<td></td>
<td>15</td>
<td>no maceration</td>
<td>05.75 a</td>
<td>20.36 b (26.82)</td>
<td>089.00 b</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>no maceration</td>
<td>05.75 a</td>
<td>22.24 a (28.13)</td>
<td>110.00 a</td>
</tr>
</tbody>
</table>

1 time taken for maceration (h); 2 enzyme activity in units; 3 per cent loss of viscosity

⁴ mean of three replications; values in parentheses are the arcsine transformed value

**DISCUSSION**

In the present study different methods of inoculation of pathogen for proving Koch postulates were tried. Results indicated that pin prick plus spore suspension spray method of inoculation significantly recorded the maximum leaf blight symptom. The maximum infection caused in the pin prick plus spore suspension spray method might be attributed to the ideal conditions like injury, ready to entry of the pathogen into the plant, ready access to the food available to the pathogen, and possibly the absence of competition and antagonism by microorganisms (inside the plant). Similar inoculation methods were reported by Anand (2002) in chili fruits against *A. alternata*.

Plant pathogens usually express several virulence functions that increase their ability to colonize and damage host plants. Variation in the virulence of *A. brassicaceae* (Awasthi and Kolte 1989), *A. alternata* (Anand 2002) and *A. tenuis* (Muthulakshmi 1990) was reported earlier. In our study among fifteen isolates of *A. alternata*, the maximum disease intensity was recorded with the isolate AA1 followed by AA11, and the least disease incidence with AA6. Variation in the virulence may be attributed to the genetic make up of the isolate. In addition the cultivar susceptibility, environmental factor, and the toxins and enzyme production capacity of pathogenic isolates, determines the virulence.
Fungal pathogens are able to produce a variety of enzymes degrading the plant cell wall, and these enzymes help the pathogens in penetration and colonization of their host plants (Yang et al. 2005). In the present investigation, the pathogen *A. alternata* produced both cellulolytic enzymes (C₁ and C₂) and pectinolytic enzymes (macerating enzymes, PME and endo PG) \textit{in vitro}.

Cellulose is a major structural constituent of the cell wall of host plants. Many phytopathogenic fungi produce cellulases in culture adaptively which hydrolyse cellulose and its derivatives (Marimuthu et al. 1974; Muthulakshmi 1990). The results obtained in the present study, indicated that the pathogens *A. alternata* produced C₁ and C₂ \textit{in vitro} and the activity of these enzymes increased with the increase in the age of the culture. The virulent isolates of *A. alternata* produced more cellulolytic (C₁ and C₂) enzymes than the avirulent ones. Similarly, Anand et al. (2008) reported high cellulase activity in the culture filtrate of virulent *A. alternata* and *C. capsici*. Anand et al. (2008) also reported that *A. alternata* and *C. capsici* produced extra cellular enzymes which degraded CMC and cellulose. Many phytopathogenic fungi produce cellulases in culture adaptively which hydrolyse cellulose and its derivatives (Marimuthu et al. 1974; Muthulakshmi 1990).

In the present investigation, the virulent isolates of *A. alternata* produced more macerating enzymes, PME and Endo-PG \textit{in vitro}, than the avirulent ones. All the pectinolytic enzymes were highly active in the culture filtrate up to 10 days of age and thereafter, the activity slowly decreased. The maceration of potato discs increased with the increase in age of the culture \textit{in vitro}, up to 10 days. Maximum pectinolytic enzymes in the culture filtrate occurred up to 10 days, indicating the role of these enzymes in pathogenicity. This corroborated with the observation of Muthulakshmi (1990) and Anand et al. (2008) in the case of *A. alternata*, causing fruit rot of chilli. The enzyme PG hydrolytically cleaves pectin in such a manner that the β-1, 4-glycosidic bonds of the chain are split (Bateman and Miller 1966). PME removes the esterified methyl group from the pectin chain hydrolytically (Goodman et al. 1967).

All these cell wall splitting enzymes are mostly adaptively secreted, secreted by the pathogen in the presence of appropriate substrates. Pectinolytic enzymes were produced only in the presence of pectin in the medium. Cellulolytic enzymes were produced only in cellulose containing medium. Jha and Gupta (1988) reported, that the combination of glucose and pectin (or) polypectate induced secretion of endo-PG and endo-PMG in *A. triticina* infecting wheat.

The production and activity of pectinolytic and cellulolytic enzymes detected \textit{in vitro} suggest their active role in disease development by the pathogen in noni leaves. Since the fungi *A. alternata* is intercellular in the host, the production of these enzymes appears to facilitate the dissolution of host cell wall and middle lamella and help the entry and establishment of the pathogen in the host, and is possibly responsible for playing a vital role in pathogenesis through cell wall degradation and disintegration of tissues.

**ACKNOWLEDGEMENTS**

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**REFERENCES**


**POLISH SUMMARY**

**WIRULENCJA GRZYBA **

**ALTERNARIA ALTERNATA**

**PORAZAJĘCEGO DRZEWO MORINDA CITRIFOLIA ORAZ WYTWARZANIE PRZEZ PATOGENA ENZYMÓW DEGRADYJĄCYCH ŚCIANY KOMÓRKOWE**