USE OF RAPD, ENZYME ACTIVITY STAINING, AND COLONY SIZE TO DIFFERENTIATE PHYTOPATHOGENIC FUZARIUM OXYSPORUM ISOLATES FROM IRAN

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ABSTRACT

Fusarium oxysporum is a common soilborne plant pathogen with a worldwide distribution. Fusarium yellows disease of chickpea (Cicer arietinum) caused by F. oxysporum is one of the most destructive soilborne disease which is a major production constraint in chickpea-growing regions of Iran. Three laboratory methods “amplification of genomic DNA using random primers, enzyme activity staining, and colony size determination” have been used to discriminate between highly virulent (HV) and weakly virulent (WV) isolates of F. oxysporum. On the basis of colony size (a traditional morphological method) and the ability of isolates to produce pectic enzymes, five HV isolates were differentiated from three WV isolates. The HV isolates formed large colony (ranging from 10.1 to 12.6 mm in diameter) and showed the same enzyme pattern, while the WV isolates produced small colony (ranging from 5.8 to 7.8 mm in diameter) and had not detectable enzyme activity in the stained overlaying gel. Twelve arbitrary 10-mer primers were tested on these 8 isolates of F. oxysporum by Polymerase Chain Reaction (PCR). Cluster analysis of the data from the DNA amplification by Random Amplified Polymorphic DNA (RAPD), differentiated HV from WV isolates. The results obtained from RAPD test confirmed the classification of these eight isolates based on pathogenicity test, colony size, and enzyme activity staining into two groups (HV and WV).

Key words: Fusarium oxysporum- RAPD- colony size- enzyme activity staining

INTRODUCTION

Fusarium yellows disease of chickpea (Cicer arietinum) caused by the vascular wilt pathogen Fusarium oxysporum is a major production constraint in chickpea-growing regions of Iran (1). Isolates of F. oxysporum form a major component of the fungal flora of most cultivated soils and most of these isolates are not perfectly identifiable using phenotypic characters (6). Molecular tools have been used to characterize the diversity among pathogenic isolates of F. oxysporum (2,4,9,12). Molecular markers, such as Random Amplified Polymorphic DNA (RAPD) have been used extensively as genetic markers in different populations (2,4). RAPD markers have some advantages in that they are easy to generate, require only very small amounts of genomic DNA and do not require the use of radioisotopes. RAPD markers can overcome the limitation of other markers because they have the potential to identify a large number of polymorphisms with good coverage of entire genome.

This paper describes three laboratory tests and techniques that confirm the classification of F. oxysporum isolates based on pathogenicity test, as highly virulent (HV) and weakly virulent (WV) isolates. These tests and techniques include colony size, enzyme activity staining and RAPD.

MATERIALS AND METHODS

Fungal isolates and growth conditions

Eight isolates (five highly virulent “HV” and 3 weakly virulent “WV”) of F. oxysporum from aerial sections of chickpea from different geographical locations in Iran were collected (17) (Table 1). Isolates were maintained in wheat seed medium (50 g autoclaved soaked wheat seed in 500 ml flask) at 4°C. Isolates
were grown in shake culture in pectic zymogram (PZ) medium containing 2.64 g, (NH₄)₂ SO₄, 0.34 g KH₂PO₄, 0.14 g MgSO₄.7H₂O, 10 g Citrus pectin (Sigma), one litter distilled water, pH adjusted to 4.5 (14). After 6 days growth at 26ºC a liquid culture filtrate was obtained (as crude enzyme) by Whatman filter paper No. 1 and stored at -20ºC until using for detecting the enzyme activity. For colony size measurement, the isolates were grown on Modified D-medium contained 0.14 g MgSO₄.7H₂O, 2.6g (NH₄)₂SO₄, 0.34 g KH₂PO₄, 0.14 g NaCl, and 20g agar supplemented with 20g L-sorbose in one litter distilled water. To reduce carmelization, the L-sorbose was autoclaved separately from the other ingredients.

Enzyme activity staining

For rapid characterization of pectic enzyme activity, 10 µl of culture filtrate was mixed with equal volume of sample buffer containing, a solution of 50 mM Tris-HCl pH 6.8, 2% (w/v) Sodium Dodecyl Sulphate (SDS), 5% (v/v) of 2-β mercaptoethanol, 10% (v/v) glycerol, and 0.05% w/v) bromophenol blue. The proteins were analyzed by Sodium Dodecyl Sulphate Polyacrylamide Gel Electrophoresis (SDS-PAGE). The separated proteins were assayed for polygalacturonase activity with an ultrathin pectate agarose gel by staining with 0.05% ruthenium red for 30 min as described by Ried and Collmer (13).

(a) Preparation of pectate agarose overlay gel

Thin pectate overlay gels (0.75mm) for detection of enzyme activity were cast by using a gel support film for agarose gels (FMC Bio products) on one of the glass plates of vertical apparatus. The overlay contained, in addition of 1% agarose (Sigma), 0.1% polygalacturonic acid in 100 mM potassium acetate buffer, pH 4.5 and 10 mM EDTANa2 (11). The agarose solution was boiled to dissolve the agarose and cooled to 75ºC. The gel mold was heated to 60ºC before casting.

(b) Detection of polygalacturonase activity in SDS-PAGE

After electrophoresis, the gel was washed by three 200 ml changes of 10 mM potassium acetate buffer, pH 4.5 (5). The pectate agarose overlay was laid directly on top of the gel and incubated at 37ºC for 120 min in a moisture lunch box. The overlay was removed and stained with 0.05% (w/v) ruthenium red for 30 min. To enhance recovery of renature enzyme activity, bovine serum albumin (BSA) at 10 µg/ml was incorporated into separating gels as described by Lacks and Springhorn (10).

Colony size

Isolates were grown on PDA plates for 7 days. With a thin platinum needle, 20 stab transfers were made from the PDA to a plate of D-medium. Plates were incubated for 72 hours at 25ºC. The diameters of 20 colonies of each strain were then measured from the underside of the plate using a binocular dissecting microscope equipped with an ocular micrometer. Colony diameters were measured to the nearest 0.1 mm (6). Dendrogram was produced by cluster analysis, using the Unweighted Pair-Group Method Analysis (UPGMA).

DNA extraction

Lyophilized mycelium of different isolates was homogenized and genomic DNA was extracted according to Zamani et al. (16).

Amplification of genomic DNA using RAPD primers

Amplification of DNA fragments was carried out by Polymerase Chain Reaction (PCR) using 10-mer arbitrary primers. Amplification reactions were performed in 50 µl reaction volumes containing one unit of Taq DNA polymerase, 2 µmol/ml each of dATP, dCTP, dGTP and dTTP, 1.5 µmol/ml primer, and 60 ng of genomic DNA.

The reaction mixture was overlaid with sterile mineral oil (50 µl) to prevent evaporation during PCR cycling. The programe comprised 34 cycles of denaturation at 94ºC for 2 min, primer annealing at 32ºC for 2 min, and extension of primer at 72ºC for 2 min. After the cycling steps were completed, the mixture was held at 72ºC for 4 min to allow complete extension of amplified products. A total of 12 Primers were used (Table 2). Amplified DNA fragments were analyzed by electrophoresis in 2% agarose gel in TBE buffer.

RAPD product scoring and Data analysis

Data were compiled as a binary 0/1 matrix by the presence (1) or absence (0) of a band at particular position. Only major RAPD bands were considered for statistical analysis. Dendrogram were produced by cluster analysis using UPGMA.

RESULTS

Colony size

Fifteen isolates of  F. oxysporum derived from different geographical regions have been previously studied based on pathogenicity test. From these fifteen isolates five highly virulent isolates (F15, F18, F23, F47 and F59) and three weakly virulent isolates (F2, F21 and F58) were distinguished. (17). These eight isolates were grown on D-medium containing L-sorbose.

The growth of different isolates tested on medium, and each isolate had a characteristic mean colony diameter (Fig. 1). Isolates F02, F21, and F58 produced small-size colonies, with mean diameters 7.8, 6.4, and 5.8 mm, respectively (Table 1). Isolates F15, F18, F23, F47, and F59 formed large colonies with mean diameters: 11.5, 10.1, 12.6, 11.1, and 10.1 mm, respectively (Table 1). A dendrogram constructed by UPGMA cluster analysis showed a grouping of these isolates into two clusters. One cluster with five isolates (large colony) were highly virulent and second cluster containing three isolates (small colony) were weakly virulent (Fig. 2).
Methods to differentiate phytopathogenic *F. oxysporum*

**RAPD analysis**

In order to compare the overall similarity between these isolates at DNA level, 12 decamer oligonucleotide primers were used which generated polymorphic bands. Example of polymorphic bands detected in amplified DNA from different isolates is shown in Fig. 3.

The data from polymorphic bands were analyzed by UPGMA method. Comparison of the results from pathogenicity test of these isolates with the results of cluster analysis of RAPD polymorphic bands demonstrated that among 12 primers, only the dendrogram obtained from R2 primer (Table 2) was able to differentiate the weakly virulent isolates F2, F21, and F58 as a separate cluster (Fig. 4).

**Table 2.** Sequence of the primers used in this study.

<table>
<thead>
<tr>
<th>Primer Identification</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>81</td>
<td>5'-ACGGTCTTGG-3'</td>
</tr>
<tr>
<td>82</td>
<td>5'-GGCGCTAGCA-3'</td>
</tr>
<tr>
<td>171</td>
<td>5'-GAAACAGCGG-3'</td>
</tr>
<tr>
<td>172</td>
<td>5'-GGAGCCCAC-3'</td>
</tr>
<tr>
<td>173</td>
<td>5'-GGAGGGTGTT-3'</td>
</tr>
<tr>
<td>174</td>
<td>5'-ACGATCGCGG-3'</td>
</tr>
<tr>
<td>R1</td>
<td>5'-CGGCCACCCCT-3'</td>
</tr>
<tr>
<td>R2</td>
<td>5'-CGCGTGCAG-3'</td>
</tr>
<tr>
<td>R3</td>
<td>5'-ACGATCGGG-3'</td>
</tr>
<tr>
<td>PU1</td>
<td>5'-AGATGCAGCC-3'</td>
</tr>
<tr>
<td>PU2</td>
<td>5'-ACGGATCCTG-3'</td>
</tr>
<tr>
<td>PU3</td>
<td>5'-ACTGGGACTC-3'</td>
</tr>
</tbody>
</table>

**Figure 1.** Comparison of large colonies (F15) and small colonies (F58) of *F. oxysporum* isolates on sorbose medium incubated for 72 hr at 25°C.

**Table 1.** Source, virulence and colony size of different isolates of *F. oxysporum* used in this study.

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Origin</th>
<th>Virulence</th>
<th>Colony size (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F15</td>
<td>Orumieh, Iran</td>
<td>H</td>
<td>11.59 ± 0.77 L</td>
</tr>
<tr>
<td>F18</td>
<td>Orumieh, Iran</td>
<td>H</td>
<td>10.07 ± 0.42 L</td>
</tr>
<tr>
<td>F23</td>
<td>Unknown</td>
<td>H</td>
<td>12.61 ± 0.96 L</td>
</tr>
<tr>
<td>F47</td>
<td>Lorestan, Iran</td>
<td>H</td>
<td>11.14 ± 1.15 L</td>
</tr>
<tr>
<td>F59</td>
<td>Tabriz, Iran</td>
<td>H</td>
<td>10.09 ± 0.45 L</td>
</tr>
<tr>
<td>F58</td>
<td>Tabriz, Iran</td>
<td>W</td>
<td>5.82 ± 0.49 S</td>
</tr>
<tr>
<td>F21</td>
<td>Orumieh, Iran</td>
<td>W</td>
<td>6.46 ± 0.54 S</td>
</tr>
<tr>
<td>F02</td>
<td>Orumieh, Iran</td>
<td>W</td>
<td>7.85 ± 0.41 S</td>
</tr>
</tbody>
</table>

H = Highly virulent; W = Weakly virulent; L = Large colony; S = small colony.

**Figure 2.** UPGMA clustering analysis of colony size data of HV and WV isolates of *F. oxysporum*.

**Figure 3.** Amplification of genomic DNA using R2 primer.
Enzyme activity staining

In order to study whether pectic enzymes are present in the culture supernatant, an attempt was made to stain the gel for pectic enzyme activity, using pectate agarose overlay. An equivalent amount of enzyme was loaded onto the gel for all isolates (HV and WV). The obtaining results indicated that all highly virulent isolates show the same enzyme activity pattern (Fig. 5). Comparison of the banding patterns or enzyme activity for the highly virulent and weakly virulent isolates demonstrated that, a high intensity band is present in the highly virulent isolates which is absent in weakly virulent isolates (Fig. 5).

DISCUSSION

In this study, we have evaluated three laboratory methods for their ability to discriminate between highly virulent and weakly virulent isolates of *F. oxysporum*: amplification of genomic DNA using random primers, enzyme activity staining, and colony size determination.

In a previous study (17) five highly virulent and three weakly virulent isolates have been identified. In the present investigation, these eight isolates were characterized by other laboratory methods. In the first test, colonies of the different isolates of *F. oxysporum* were grown on a sorbose-containing medium and their diameters were measured. The second test assessed the ability of the isolates to produce pectic enzymes, as tested by enzyme activity staining on the gel. By cluster analysis of the results from these two tests, these eight isolates have been divided into two distinct clusters. Comparing the arrangement of isolates in Fig. 2 (based on colony size) and Fig. 5 (based on enzyme activity staining) with the classification in pathogenic groups (Table 1), one can see that the classification of the isolates by pathogenicity test is in accordance with the results of these two tests. Further analysis of the results revealed that isolates F15, F18, F23, F47, and F59, which were highly virulent, formed large colonies (ranging from 10.1 to 12.6 mm in diameter) and produced detectable pectic enzyme activity in overlaying gel, while isolates F02, F21, and F58, which were weakly virulent, formed small colonies (ranging from 5.8 to 7.8 mm in diameter) and showed not detectable pectic enzyme activity in overlaying gel. Correll *et al.* (6) reported that virulent isolates of *F. oxysporum* have a characteristically large colony size which is in agreement with our finding.

When comparing the RAPD analysis to classification into pathogenicity groups, the results using 12 primers did not present good correlation between amplification patterns and pathotype classification.

However, primer R2 alone allowed differentiation of these isolates, which was strongly supported by pathotype, colony size and overlaying classification. As evidenced in our study and in others as well, RAPD markers were effective for detecting polymorphism in *F. oxysporum* (3,7,8,15).

Finally, from these results it can be concluded that these three rapid methods (RAPD, colony size, and enzyme activity staining) could be employed to distinguish highly virulent from weakly virulent isolates of *F. oxysporum*. A fast diagnosis of the highly virulent in an early stage of infection may provide advantages for phytosanitary procedures. Nevertheless, standardization of the detection system will be required.

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RESUMO

Uso de RAPD, coloração de atividade enzimática e tamanho de colônia para diferenciação de isolados fitopatogénicos de *Fusarium oxysporum* obtidos no Irã

*Fusarium oxysporum* é um fitopatógeno amplamente difundido nos solos. A doença amarela do grão-de-bico (*Cicer arietinum*), causada por ele é destrutiva e um grande problema nas regiões produtoras do Irã. Três métodos de laboratório: amplificação aleatória de DNA genômico (RAPD), coloração de atividade enzimática e determinação das dimensões de colônia, foram usados para diferenciar isolados de *Fusarium oxysporum* altamente virulentos (HV) e fracamente virulentos (WV). Baseado no tamanho da colônia (método morfológico tradicional) e na capacidade dos isolados produzirem enzimas pépticas, cinco isolados HV foram diferenciados de três WV. Enquanto os isolados HV formaram colônias grandes (entre 10,1 e 12,6 mm de diâmetro) e apresentaram o mesmo padrão enzimático, os isolados WV produziram colônias pequenas (entre 5,8 e 7,8 mm de diâmetro) e não tiveram atividade enzimática detectável no gel de cobertura corado. Doze primers arbitrários foram testados nos 8 isolados de *Fusarium oxysporum* por reação de polimerase em cadeia (PCR). As análises dos dados da amplificação de DNA pelo método de amplificação aleatória do DNA polimórfico (RAPD) diferenciaram os isolados HV dos WV. Os resultados obtidos pelo teste RAPD confirmaram a classificação dos 8 isolados pelos testes de patogenicidade, tamanho da colônia e atividade enzimática de coloração nos dois grupos, HV e WV.

Palavras-chave: *Fusarium oxysporum*, RAPD, tamanho de colônia, coloração de atividade enzimática.

REFERENCES