Isolation of seed borne myco flora of wheat
(Triticum aestivum L. em. The ll.) Seed samples

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Abstract
Seed borne myco flora of Wheat in two district of southern Rajasthan (Chittorgarh and Udaipur) were surveyed. A total of 80 seed samples, 40 of each district, were collected during 2015-16. Data were recorded for major seed borne myco flora, which were identified and quantified using the blotter and agar plate method as recommended by ISTA (International Seed Testing Association). Eight wheat seed borne myco flora viz., Alternaria alternata, Aspergillus flavus, A. niger, Curvularia lunata, Fusarium moniliforme, Rhizopus stolonifer, Mucor spp. and Trichoderma viride were detected and isolated from eighty seed samples by using standard blotter paper and agar plate methods.

Keywords: wheat, pathogen frequency, seed borne, myco flora, embryo

1. Introduction
Seed borne diseases have been found to affect the growth and productivity of crop plants. A seed borne pathogen present externally or internally or associated with the seed as contaminant may cause seed abortion, seed rot, seed necrosis, reduction or elimination of germination capacity as well as seedling damage resulting in development of disease at later stages of plant growth by systemic or local infection. Seeds are regarded as highly effective means for transporting plant pathogens over long distances. Besides, the mold fungi which grow on the seed substratum produce mycotoxins which are hazardous to humans and animals (Halt, 1994) [9]. Studies were carried out to study the composition of seed borne myco flora occurring in wheat. The significance of sustainable agricultural production is hidden in the use of quality seed. It is the most crucial and vital input for enhancing productivity. Since seed is the custodian of the genetic potential of the cultivars, the quality of the seed determines the limits of productivity to be realized in a given cropping system. Though seeds are of great economic interest and also contribute a major part of diet, they play a vital role in associating microorganism, which prove hazardous for the seed or the new plant created from it, so, any infections agent (bacteria, fungi, nematode, etc.) which is associated with seeds having potential of causing a disease in a seedling or plant, is teemed as seed borne pathogen (Agarwal, 1996) [1].

Hence, the storage fungi are especially insidious because they invade seeds stored at moisture contents that practical grain men consider safe and often cause serious damage before their presence even suspected. Therefore, with few exceptions spoilage of stored fungi, this may be introduced during the post-harvest handling process. It is well known fact that several fungi are known to cause considerable damage to seeds in storage and produce various activities. It is in view of this that the current study aimed at detecting seed borne fungal pathogen wheat seed sample of two districts (Chittorgarh and Udaipur) of southern Rajasthan.

2. Materials and Methods
2.1 Experimental location
The experiment was conducted at Department of Plant Pathology, Rajasthan College of Agriculture (RCA) MPUAT, Udaipur during 2015-16.

2.2 Sources of experimental materials
Eightly seed samples of wheat were collected for the isolation and identification of seed borne fungi from Chittorgarh and Udaipur district of southern Rajasthan. 40 seed sample were collected from each district. All materials except seeds, which used in this experiment, were sterilized using 70% ethyl alcohol. The Petri plates were sterilized at 180 °C for two hours in hot air oven whereas, Media (Potato Dextrose Agar, Richard’s broth), blotter paper and distilled water were sterilized in an autoclave at 1.045 kg/cm² pressures for 20 min. Seeds are surface sterilized by dipping in 0.1 per cent mercuric chloride (HgCl₂) solution for 2-3 minutes.
followed by three washing with sterilized distilled water. Fresh polythene bags were sterilized with 5% formalin solution. Cotton blue and lacto phenol were used for staining of the fungal cytoplasm and for providing a light blue background, against which the walls of hyphae can readily be seen (Aneja, 2004) [3]. Plating of the seed component: Standard blotter method and agar plate method described by the International Seed Testing Association (ISTA 1976) [13], was used for the isolation of the seed borne fungi associated with the wheat seed samples.

2.3 Isolation of seed myco flora by standard techniques

For the isolation of seed myco flora (external and internal) of wheat, standard blotter paper with slight modification and agar plate method were used, standardized by International Seed Testing Association (ISTA, 1976) [13].

2.3.1 Standard blotter method

(Lantos et al., 2002) [14] Four hundred seeds from each sample were analyzed under sterilized and unsterilized conditions. Blotter paper were cut into 9 cm diameter circle and sterilized at 1.045 kg cm-2 for 20 minutes. These circles of blotter paper were placed at the bottom of each sterilized Petri plates aseptically and moistened by sterilized distilled water. Sixteen (1+6+9) seeds were placed at equal distance in each Petri plate. For sterilized conditions, seeds treated with 0.1% mercuric chloride (HgCl2) solution for three minutes followed by three washing with sterile distilled water were used. (Habib et al., 2007) [8]. The Petri plates were incubated at 25±2°C for 12 h of light alternating with 12 h of dark period. The seeds were examined starting from third days to eight days of incubation for the presence of seed borne myco flora.

2.3.2 Agar plate method (APM)

Potato Dextrose Agar (PDA) was used as a basal medium for the isolation of seed myco flora. Four hundred unsterilized and sterilized (with 0.1% mercuric chloride solution) for 2 to 12 minutes followed by three washing with sterile distilled water (Habib et al., 2007) [8]. Seeds of each sample were analyzed for isolation of myco flora. Sixteen (1+6+9) seeds were placed aseptically per Petri plate (9 cm diameter) containing 20 ml PDA. The plates were incubated at 25±2°C for 12 h of light alternating with 12 h of dark period. The fungal colonies go out from seeds were examined from 3 to 10 days of incubation (Henselova and Hudcova, 2001; Gwary et al., 2006) [10, 6].

2.4 Slide preparation and identification

The samples of fungi were taken randomly from each crop. These samples were identified on the basis of colony characteristics and microscopic examinations. Standard books and research papers were consulted during the examination of these fungi (Aneja, 2004; Rifai, 1969; Barnet and Hunter, 1999) [2, 20, 3]. The binocular compound microscope was used to determine the type of fungus in each plate. The seed borne fungi were identified using identification keys and cross-checked for each seed plates to identify the type of fungus growing on each seed. After seven days of incubation, fungal species found growing on the surface of seeds, were identified.

3. Results

(A) Detection of seed myco flora by standard techniques

(a) Standard blotter paper method (SBPM)

Incubation of all the wheat seed samples by standard blotter paper method revealed the presence of seven different fungi viz. Alternaria alternata, Aspergillus flavus, Aspergillus niger, Curvularia lunata, Fusarium moniliforme, Rhizopus stolonifer and Mucor spp.

The unsterilized and sterilized wheat seeds were placed over moist blotter paper in sterilized Petri plates for detection of external seed borne myco flora. In unsterilized wheat seed samples, the total incidence of myco flora recorded were (33.89%) and (36.0%) respectively in Chittorgarh and Udaipur districts unsterilized seed samples while Sterilized seeds showed maximum percentage (12.09%) of myco flora in Chittorgarh district seed samples and minimum (10.68%) in Udaipur district seed samples.

Alternaria alternata was observed maximum (2.63%) in Chittorgarh unsterilized seed samples and minimum in Udaipur (0.44%) sterilized seed samples. However, Aspergillus flavus was found maximum (9.61%) in unsterilized and minimum (3.94%) in sterilized seed samples of Udaipur district. In Udaipur district unsterilized seed samples Curvularia lunata was found maximum (5.15%) and minimum in Udaipur (0.64%) sterilized seed samples, while Fusarium moniliforme was found higher (7.4%) in unsterilized seed samples of Udaipur and lower in (1.54%) sterilized seed samples of Udaipur district. Rhizopus stolonifer was observed maximum in Chittorgarh (3.60%) sterilized seed samples and minimum in Udaipur (0.43%) sterilized seed samples. While Mucor spp. was found maximum (4.0%) and minimum (1.24%) in Chittorgarh unsterilized and Udaipur sterilized seed samples respectively.

(b) Agar plate method

The unsterilized and sterilized seeds were placed over PDA medium in sterilized Petri plates. All the fungi detected with seeds of wheat by using standard blotter method were also detected by the agar plate method. Alternaria alternata was found maximum (1.96%) and minimum (0.44%) in Udaipur unsterilized and sterilized seed samples respectively. Aspergillus flavus was observed maximum (8.90%) and minimum (1.55%) in Chittorgarh unsterilized and sterilized seed samples. In Udaipur district unsterilized seed samples Aspergillus Niger was found maximum (5.2%) and minimum (0.27%) in Chittorgarh district sterilized seed samples. Curvularia lunata was observed maximum (5.3%) and minimum (0.96%) in Udaipur district unsterilized and sterilized seed samples respectively. In Chittorgarh district seed samples Fusarium moniliforme was found maximum (5.06%) in unsterilized and minimum (2.37%) in sterilized seed samples. Rhizopus stolonifer was found maximum (3.7%) and minimum (0.025%) in Chittorgarh unsterilized and sterilized seed samples. Maximum (3.06%) incidence of Mucor spp. was found in Chittorgarh unsterilized seed samples while minimum (0.25%) in Udaipur unsterilized seed samples. Trichoderma viride was observed only in Udaipur district seed samples with frequency of maximum (2.08%) in unsterilized and minimum (0.25%) in sterilized seed samples.

(B) Isolation, purification and identification of the seed borne myco flora

The seed borne myco flora detected by standard incubation technique were isolated by transferring on potato dextrose agar medium in Petri plates. The plates were incubated at 25±2°C for the fungal growth.

The seed borne myco flora was identified on the basis of cultural, morphological characters using selective standard methods and literature (Holliday, 1980; Rifai, 1969; Rape and
as mention below and the wheat seed borne myco flora identity was confirmed Alternaria alternata, Aspergillus flavus, Aspergillus niger, Curvularia lunata, Fusarium moniliforme, Rhizopus stolonifer, Mucor spp. and Trichoderma viride.

4. Discussion
Seeds of many crops are subjected to invasion by pathogen during development and before or after harvest by a great variety of myco flora. Seed carrying a pathogen serves as the primary source of infection and has very important role in the epidemiology of disease. Myco flora present on or in the seed may result in prolonged dormancy, reduce germination and seedling survival (Christensen and Lopez, 1963) [3]. In dates back to 1729, Michelli first demonstrated seed transmission of a pathogen while, Tillet (1755) [22] confirmed and established that Tilletia caries Tull, the fungus responsible for bunt disease of wheat is seed borne. Since then the knowledge of seed borne disease of crop has been greatly increased and now there is hardly any cultivated crop where at least one seed borne fungal parasite is not known. Consequently, it has become clear in Plant Pathology that seed borne myco flora is largely associated with the occurrence of number of diseases. In present study wheat seed sample were collected from two districts of Rajasthan namely Udaipur and Chittorgarh for seed myco flora studies. Forty seed samples were collected from each district. Seeds of these samples were tested for Dry seed inspection of wheat seed samples revealed the presence of deformed, discoloured, damaged seeds and other inert material together with healthy seeds. It is likely that development of different types of seed myco flora during storage may lead to such deformation of seed. Presence of such kind of seeds and other impurities in “Moth” has also been reported by Sharma (1986) [21]. In present investigation, methods viz., blotter paper method, agar plate method were used for detection of seed myco flora of wheat. Alternaria alternata, Aspergillus flavus, A. niger, Curvularia lunata, Fusarium moniliforme, Rhizopus stolonifer, Mucor spp. and Trichoderma viride were isolated from seeds. Similarly, Singh et al. (2011) found many fungal species associated with wheat seeds and their effect on germination. On examination of seed myco flora by agar plate method and blotter method. Total sixteen fungal species were isolated from test cultivars by the standard techniques. Fungi isolated and identified were Alternaria alternata, A. solani, Aspergillus candidus, A. flavus, A. fumigatus, A. niger, A. terreus, Curvularia lunata, Fusarium roseum, F. semitectum, Penicillium citrinum, P. rubrum, Rhizopus stolonifer and Trichoderma harzianum. However, Bashir et al. (2012) [4] also reported five fungal species namely Rhizopus nigricans, Mucor spp., Penillium jensenti, Aspergillus niger, and Fusarium moniliforme. Similarly, Zrani (2013) [23] observed ten seed borne fungi in wheat (Alternaria spp., Aspergillus spp., Aureobasidium spp., Cladosporium spp., Dreschslera spp., Penicillium spp., Rhizoctonia spp., Stemphylium spp., Mucor spp. and Rhizopus spp.). Agar plate method gave higher number of fungi as compared to blotter test. It may be possible that variation may be due to reason that week and slow growing fungi could not grown well on blotter paper as compared to agar plate tests (Neergaard and Saad,1962) [17]. The blotter paper and agar plate test revealed that per cent incidence of all species isolated from sterilized seeds were low as compared to unsterilized ones and in some sterilized seed samples. Pathak and Razia (2013) [18] also found that Fusarium moniliforme, Rhizopus spp., Mucor spp., Alternaria alternata, Aspergillus niger, A. flavus, Curvularia lunata, Dreschslera spp., Alternaria spp. and Penicillium spp., were mostly frequently isolated from the wheat seeds, which is similar to result, observed in present investigation. Similar results with agar plate test also have been reported by Bashir et al. (2012) [4] and Gohari et al., (2007) [16].

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Seed myco flora</th>
<th>Chittorgarh seed sample</th>
<th>Udaipur seed sample</th>
<th>Chittorgarh seed sample</th>
<th>Udaipur seed sample</th>
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<tbody>
<tr>
<td></td>
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<td>Unsterilized</td>
<td>Sterilized</td>
<td>Unsterilized</td>
<td>Sterilized</td>
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<tr>
<td>1</td>
<td>Alternaria alternata</td>
<td>1.93 (7.93)</td>
<td>0.61 (4.43)</td>
<td>1.96 (8.03)</td>
<td>0.44 (3.79)</td>
</tr>
<tr>
<td>2</td>
<td>Aspergillus flavus</td>
<td>8.9 (17.34)</td>
<td>1.55 (7.14)</td>
<td>8.11 (16.59)</td>
<td>2.13 (8.38)</td>
</tr>
<tr>
<td>3</td>
<td>Aspergillus niger</td>
<td>1.36 (6.69)</td>
<td>0.27 (2.97)</td>
<td>5.2 (13.17)</td>
<td>1.96 (8.04)</td>
</tr>
<tr>
<td>4</td>
<td>Curvularia lunata</td>
<td>4.2 (11.82)</td>
<td>1.43 (6.77)</td>
<td>5.5 (13.30)</td>
<td>0.96 (5.61)</td>
</tr>
<tr>
<td>5</td>
<td>Fusarium moniliforme</td>
<td>5.06 (12.99)</td>
<td>2.37 (8.78)</td>
<td>4.9 (12.78)</td>
<td>3.2 (10.30)</td>
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<tr>
<td>6</td>
<td>Rhizopus stolonifer</td>
<td>3.7 (11.08)</td>
<td>0.25 (0.90)</td>
<td>3.63 (10.97)</td>
<td>1.0 (5.73)</td>
</tr>
<tr>
<td>7</td>
<td>Mucor spp.</td>
<td>3.06 (10.07)</td>
<td>2.32 (8.68)</td>
<td>0.25 (8.28)</td>
<td>0.86 (5.31)</td>
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<tr>
<td>8</td>
<td>Trichoderma viride</td>
<td>0.0 (0)</td>
<td>0.0 (0)</td>
<td>2.08 (9.62)</td>
<td>0.25 (8.26)</td>
</tr>
</tbody>
</table>

| Seed without myco flora | 69.19 | 91.17 | 67.8 | 89.2 |
| Seed with myco flora | 30.81 | 8.83 | 32.2 | 10.8 |
| SE± | 0.139 | 0.21 | 0.149 | 0.074 |
| CD at 5% | 0.417 | 0.632 | 0.448 | 0.223 |
| CV | 2.21 | 6.86 | 2.38 | 2.07 |

* The value in parentheses is angular transformed

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<tr>
<td>1</td>
<td>Alternaria alternata</td>
<td>2.63 (9.32)</td>
<td>0.65 (4.62)</td>
<td>2.60 (9.27)</td>
<td>0.44 (3.73)</td>
</tr>
<tr>
<td>2</td>
<td>Aspergillus flavus</td>
<td>9.6 (18.09)</td>
<td>4.0 (11.53)</td>
<td>9.61 (18.05)</td>
<td>3.94 (11.44)</td>
</tr>
<tr>
<td>3</td>
<td>Aspergillus niger</td>
<td>3.1 (10.29)</td>
<td>2.18 (8.48)</td>
<td>4.75 (12.58)</td>
<td>2.45 (9.0)</td>
</tr>
<tr>
<td>4</td>
<td>Curvularia lunata</td>
<td>4.93 (12.82)</td>
<td>1.06 (5.90)</td>
<td>5.15 (13.06)</td>
<td>0.64 (4.59)</td>
</tr>
<tr>
<td>5</td>
<td>Fusarium moniliforme</td>
<td>6.03 (14.58)</td>
<td>1.99 (7.12)</td>
<td>7.4 (15.77)</td>
<td>1.54 (7.12)</td>
</tr>
<tr>
<td>6</td>
<td>Rhizopus stolonifer</td>
<td>3.6 (10.92)</td>
<td>0.61 (4.46)</td>
<td>2.63 (9.32)</td>
<td>0.43 (3.75)</td>
</tr>
<tr>
<td>7</td>
<td>Mucor spp.</td>
<td>4.0 (11.52)</td>
<td>1.60 (7.26)</td>
<td>3.86 (11.32)</td>
<td>1.24 (6.38)</td>
</tr>
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</table>

Table 1: Per cent incidence of seed myco flora associated with seed samples of wheat tested by agar plate method

Table 2: Per cent incidence of seed myco flora associated with seed samples of wheat tested by standard blotter paper method
Seed without myco flora | 66.11 | 87.91 | 64 | 89.32
Seed with myco flora | 33.89 | 12.09 | 36 | 10.68
SEm± | 0.298 | 0.147 | 0.160 | 0.206
CD at 5% | 0.904 | 0.447 | 0.488 | 0.626
CV% | 4.13 | 3.55 | 2.18 | 5.44

* The value in parentheses is angular transformed

Fig: Methods for detection of seed borne myco flora of wheat
5. References