Detection of Seed Borne Fungi of Blonde Psyllium by Standard Blotter Methods

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ABSTRACT

The experiments were conducted at Department of Plant Pathology, S.K.N. College of Agriculture, Jobner (Rajasthan). Five samples of blonde psyllium seeds were collected from Barmer, Nagaur, Pali, Jodhpur and Jalore districts of Rajasthan and tested for seed borne fungi. Both Blotter and Agar Plate methods were used to detect the maximum number of control and external seed borne fungi. A total of 5 species of fungi were identified using the two selective methods. The pathogenicity of A. alternata was studied on psyllium seeds and seedlings.

Keywords: Alternaria alternata, Standard blotter technique, Potato dextrose method and Plantago ovate, Isabgol.

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Introduction

Blond psyllium (Plantago ovata Forsk.) commonly known as isabgol, is an annual herb with narrow linear rosette like leaves belonging to the family Plantaginaceae. Isabgol is an important cash crop cultivated for its export and being of important medicinal value is reported to have larger demands and is traded in major medicinal markets of the world. Isabgol has pharmaceutical importance to treat dysentery, chronic constipation and chronic diarrhoea and as laxative demulcients, emollients and diuretics. India commands nearly a monopoly in the production and export of the seed and husk to the world market. India is earning about Rs. 1600 million as foreign exchange from the export of blond psyllium products to countries like USA, Germany, France, England, Spain and Belgium [1].

In India, the isabgol crop is mainly grown as commercial crop in Gujarat, Rajasthan and Madhya Pradesh. However, the crop is spreading to other non-traditional parts of the country such as Haryana, Uttar Pradesh and Karnata. In Rajasthan, it is being cultivated in 190081 hectares area with a total production of 99950 tonnes of seeds with an average productivity of 525 kg/ha [2]. In Rajasthan, Isabgol mainly cultivated in Barmer, Jalore, Nagaur, Jodhpur and Jaisalmer districts. Presently, Rajasthan is on
the top in productivity in India [3] reported a number of pathogens viz., Fusarium wilt (Fusarium oxysporum), damping off (Pythium ultimum trow), leaf blight (Alternaria alternata (Fr.) Keissler), downy mildews (Peronospora plantaginis) and powdery mildew (Erysiphe cichoracearum D.C.) affecting this crop. Alternaria blight has become a serious problem in recent years. It has been found that downy mildew affected crop is more prone to be attacked by A. alternata.

It causes considerable damage every year and sometimes become very severe which results in total loss of yield [4]. Hence, present investigations were carried out to test the efficacy of plant extracts and fungicides against leaf blight of isabgol incited by A. alternata.

Materials and Methods

Collection of seed samples

To study the seed borne fungi, seed samples were collected from Barmer, Nagaur, Pali, Jodhpur and Jalore districts of Rajasthan. These samples were collected from farmers, where isabgol is generally cultivated.

From each district, 5 samples were collected from farmer’s houses belonging to different villages situated 5-10 km away from district headquarter in different geographical directions where isabgol is generally cultivated and stored by using old traditional practices, under variable environmental conditions. At each district headquarter; all collected seed samples were mixed to represent a composite sample of that particular district. Samples were collected 4-6 weeks before sowing in cloth bags, brought to the laboratory and then stored at 10 ± 1 °C temperature for further studies.

In all, 5 working samples were analyzed by different procedures suggested by [5, 6, 7].

Isolation of mycoflora associated with isabgol seeds

Five seed samples i.e. IG-1, IG-2, IG-3, IG-4 and IG-5 were used separately for the isolation of mycoflora from the isabgol seeds. Two incubation methods viz., (a) Standard Blotter Method and (b) Agar Plate Method [5] were employed for this purpose.

Standard blotter method

Four hundred seeds from each sample were analyzed after surface sterilization and unsterilization. Blotter papers were cut into 9 cm diameter circle and sterilized at 1.045 kg sq cm for 20 minutes.

Three circles of blotter paper were placed at the bottom of each sterilized Petri plate aseptically and moistened by sterile distilled water. Twenty five seeds were placed at equal distance in each Petri plate.

For sterilized condition, seeds were treated with 0.1% mercuric chloride (HgCl$_2$) solution for one minute and rotated continuously to avoid clumping followed by three washing with sterile distilled water, were used.

These Petri plates were incubated at 25 ± 1 °C for 12 h of light alternating period. The seeds were examined on 8th day of incubation for the presence of seed borne fungi.

Standard agar plate method

Two hundred seeds of each of the 5 samples were analyzed after surface sterilization with 0.1 per cent mercuric chloride solution for one minute, followed by 3 washings with sterile distilled water.
Twenty five seeds were equispaced aseptically per Petri plate (9 cm diameter) containing 20 ml of PDA (250 ppm streptomycin was added to PDA just before pouring into the Petri plates to check bacterial contamination).

The plates were incubated at 25 ± 1 °C for 12 h of light alternating with 12 h of dark period. The fungal colonies emanating from seeds were examined from 4th to 7th days of incubation.

Isolation of mycoflora from isabgol seeds were carried out and maintained on 2 per cent PDA medium. Observations on per cent incidence of seed mycoflora were recorded in both Blotter and Agar Plate Methods.

Per cent incidence of fungus (PIF) was calculated as follows:

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\text{PIF} = \frac{\text{Number of seeds sowing fungal growth}}{\text{Total number of seeds incubated}} \times 100
\]

Results and Discussion

Isolation of mycoflora associated with isabgol seeds

Standard blotter method

Five fungal species belong to four genera in addition to un-identified mycoflora were isolated from sterilized and un-sterilized isabgol seeds (Table 4.1 and Fig. 1). Fungi and their respective per cent incidence from sterilized and un-sterilized seeds were Alternaria alternata (1.00-11.25), Aspergillus niger (1.50-8.25), Curvularia clavata (0.25-3.75), C. fallax (0.00-2.50), Fusarium semitectum (1.50-7.25) and un-identified mycoflora (0.00-6.25). Per cent incidence of Alternaria alternata, out of total per cent mycoflora was recorded maximum in un-sterilized seeds of Jalore sample (29.41) followed by Barmer (27.02), Jodhpur (25.80) and Pali (22.54) while it was minimum in Nagaur sample (21.50).

Per cent incidence of Alternaria alternata including other fungi was less in sterilized seeds as compared to un-sterilized seeds.

The sterilized and un-sterilized seeds were plated on moist blotter paper in sterilized Petri plates for detection of fungal flora.

The total frequency of mycoflora was observed maximum in unsterilized seeds of sample of Jalore (38.50%) followed by Pali (25.70%), Barmer (18.50%) and Jodhpur (15.50%) and it was minimum in Nagaur (12.75%), whereas, in surface sterilized seeds, the incidence of mycoflora was maximum in the sample of Jalore (11.25%) followed by Pali (9.25%) and Barmer (7.25%) and it was minimum in samples of Jodhpur (6.00%) and Nagaur (6.00%).

Standard agar plate method

Five fungal species belong to four genera in addition to un-identified mycoflora were isolated from isabgol seeds (Table 2 and Fig. 2). Fungi and their per cent occurrence recorded were Alternaria alternata (1.50-5.00), Aspergillus niger (2.00-4.50), Curvularia clavata (0.50-2.50), C. fallax (0.00-2.50) and Fusarium semitectum (2.00-3.50).

Total per cent mycoflora was maximum in Jalore sample (19.00%) followed by Pali (13.00%), Nagaur (12.00%), Barmer (12.00%) and Jodhpur (9.00%).

Per cent incidence of Alternaria alternata, out of total per cent mycoflora was highest in Jalore (27.77%) followed by Barmer (25.00%), Pali (21.15%), Nagaur (20.83%) and Jodhpur (16.86%).
Fig. 1 Incidence of mycoflora associated with isabgol seeds analysed by Standard Blotter Method

Fig. 2 Incidence of mycoflora associated with isabgol seeds analysed by Standard Agar Plate Method
Table 1 Incidence of mycoflora associated with *isabgol* seeds analysed by Standard Blotter Method

| Mycoflora                  | Barmer US | S | Nagaur US | S | Pali US | S | Jalore US | S | Jodhpur US | S |
|----------------------------|-----------|---|-----------|---|---------|---|-----------|---|------------|---|
| Alternaria alternata       | 5.00      | 1.25 | 2.75       | 1.25 | 5.75     | 2.00 | 11.25     | 2.00 | 4.00       | 1.00 |
| Aspergillus niger          | 4.00      | 1.75 | 2.25       | 1.50 | 5.50     | 2.25 | 8.25      | 2.25 | 3.25     | 1.50 |
| Curvularia clavata         | 1.25      | 0.75 | 0.50       | 0.25 | 3.75     | 1.00 | 2.75      | 1.25 | 1.25     | 0.50 |
| Curvularia fallax          | 2.25      | 1.50 | 2.00       | 1.00 | 0.75     | 0.00 | 2.50      | 1.50 | 1.50     | 1.00 |
| Fusarium semitectum        | 4.25      | 1.50 | 3.25       | 2.00 | 6.00     | 2.75 | 7.25      | 2.00 | 2.50     | 1.50 |
| Unidentified mycoflora     | 1.75      | 0.50 | 2.00       | 0.00 | 3.75     | 1.25 | 6.25      | 2.25 | 3.00     | 1.50 |
| **Total mycoflora**        | **18.50** | **7.25** | **12.75** | **6.00** | **25.70** | **9.25** | **38.25** | **11.25** | **15.50** | **6.00** |
| *Alternaria alternata*     | **27.02** | **17.24** | **21.56** | **20.83** | **22.54** | **21.62** | **29.41** | **17.77** | **25.80** | **16.66** |

* Seed tested 400/ sample, US- Un-sterilized, S- Sterilized

During present investigation five seed samples of *isabgol* were collected from Barmer, Jalore, Jodhpur, Nagaur and Pali districts of Rajasthan during 2013-14. Examination of dry seed samples revealed the presence of deformed (shriveled), discoloured (black and brown coloured) and damaged (mechanically and insect) seeds, in addition to impurities like plant debries (pieces of leaves and stem), seeds of other crops, inert material (stones and sand) and apparently healthy seeds.

It is likely that different types of seed mycoflora during development of seeds and there storage might have caused such deformation and discoloration of seeds. Occurrence of such deformation and discoloration along with impurities have also been reported during examination of dry seeds of pearl millet, barley and *isabgol* by Singh and Singh, 1983; Randhava and Aulakh, 1984 and Meena and Maharshi, 2013, respectively. For the isolation of mycoflora from *isabgol* seeds, two standard methods viz. Blotter and Agar Plate Methods were used to know the incidence of mycoflora. The Blotter Method displayed 5 fungi viz. *Alternaria alternata, Aspergillus niger, Curvularia clavata, Curvularia fallax* and *Fusarium semitectum* along with un-identified mycoflora. It also revealed that the seed sample from Jalore was infected and infested with maximum number of fungi and the sample from Jodhpur showed minimum incidence. The incidence of *Alternaria alternata* has been noted following standard Blotter and Agar Plate Methods. In the present study, maximum counts of *A. alternata* and other fungi were observed by blotter (un-sterilized and sterilized seeds).
method as compared to agar plate (sterilized seeds) method. Other fungi detected were Aspergillus niger, Curvularia clavata, C. fallax, Fusarium semitectum and un-identified fungi. This study is in conformity to earlier findings of [8, 9, 10, 11, 12] who isolated Alternaria sp., Aspergillus sp., Curvularia sp., Fusarium sp. and other fungi from seeds of isabgol.

In general, little variation was observed in blotter and agar plate methods for the presence of Alternaria alternata and other fungi recorded on seeds. Detection of A. alternata was higher in blotter method in comparison to agar plate method. This variation might be due to the reasons that some of the weak and slow growing fungi could not grow in agar culture in comparison to fast growing saprophytic fungi. Pre-surface sterilization of seeds and use of substratum in the method employed may be another reason [13, 14]. To have a complete spectrum of the mycoflora, it seems essential to deploy both the methods. [15] Also observed that blotter and agar plate methods are equally valuable and supplementary to each other.

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