Seed-borne pathogenic fungi on some soybean varieties

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Abstract. Soesanto L, Hartono ARR, Mugiastuti E, Widarta H. 2020. Seed-borne pathogenic fungi on some soybean varieties. Biodiversitas 21: 4010-4015. The present study was conducted to detect and identify seed-borne pathogenic fungi in some soybean varieties and their effect on seed germination. Experiment was performed in a completely random design with eight treatments and four replicates. Eight soybean varieties i.e., Malabar, Kaba, Dering, Detam I, Sinabung, Dena, Gepak Kuning, and Slamat were selected for the investigation. Seed borne fungi were isolated using blotter test and agar plate techniques. The variables observed were morphological/cultural characteristics, microscopic features of fungi, and percentage of seed germination. A total of eight fungi namely Aspergillus flavus Link, Aspergillus niger van Tieghem, Cladosporium oxysporum Berk. & M.A. Curtis, Colletotrichum dematium (Pers. et Fr.) Grove f.sp. truncate (Schw.) Arx, Curvularia pallescens Boedijn, Fusarium solani (Mart.) Succ., Melanosphora zamiæ Corda, and Nigrospora sp. Mason were isolated from different varieties of soybean seeds. The highest seed germination was found to be 80.5% in Daring and Detam I varieties, respectively, and the lowest 53% was recorded in Dena variety.

Keywords: Identification, seed-borne pathogenic fungi, seed germination, soybean

INTRODUCTION

Soybean (Glycine max L. Merril) is the third most important food commodity after rice and corn (Bowo et al. 2016). Soybean is one of the government’s priorities because it produces vegetable proteins that contain significant nutritional content, is safe for consumption, and its price is relatively cheap compared to other protein sources, such as meat, milk, and fish (Sharma et al. 2014). However, soybean production during the last 6 years (2011-2017) in Indonesia, especially in East Java, decreased by 7.16% and 45.25% respectively (BPS 2018). This makes Indonesia a soybean importer (Ningrum et al. 2018).

. Attack of plant pathogens is one of the reasons for low soybean productivity and 55 fungal species has been isolated from soybean seeds. The most frequent genera were Alternaria, Diaportha, and Fusarium (Escamilla et al. 2019). Certified seeds do not guarantee to be free from seed-borne pathogens (Bishaw et al. 2013). Infected seeds can directly affect plant growth and become a source of infection in the field (Pérez-Pizá et al. 2019). Healthy seeds are one reason for increasing crop production. Inspection of seed quality, physiological and pathological health is absolutely necessary as seeds can be carriers of plant diseases (Dutta et al. 2014; Sharma et al. 2015). Disease-transmitted seeds can affect seed germination, mortality in the nursery stage, and the development of disease at the younger stage. The impact on farmers is economic loss, because maintenance and labor costs are not comparable with declining production yields (Dutta et al. 2014).

Pathogens can be found in seeds before or after germination (Garuba et al. 2014). Seed-borne pathogens are agents that are found internally or externally in seeds and have the potential to cause diseases in plants (Gupta et al. 2017; Pedraza et al. 2018). Several pathogenic fungi like Rhizopus, Alternaria, Curvularia, Diaporthe, Mucor, Corynespora, Cercospora, Colletotrichum, Phoma, Pythium, Fusarium, Aspergillus, and Cladosporium have been isolated from soybean seeds (Saylendra and Fatmawaty 2010; Kinnikar et al. 2015; Escamilla et al. 2019).

Seed is the most easily planted part mainly for trade. Seed-borne diseases can be spread through insects, water, wind, agricultural equipment, and transportation. These activities need to be monitored to avoid transmission of disease from seeds to plants (Dutta et al. 2014). Therefore, the aim of the study is to identify pathogenic fungi in several soybean varieties and to investigate their effect on seed germination.

MATERIALS AND METHODS

Sample collection

Eight different varieties of soybean i.e. Malabar, Kaba, King, Detam 1, Sinabung, Dena, Yellow Gepak, and Slamat were collected from the Laboratory of Plant Breeding, Jenderal Soedirman University. The seed samples were brought to the laboratory and stored at room temperature for subsequent studies.

Isolation of fungi on blotter test

Four-hundreds (400) soybean seeds per variety were selected for the blotter test. Surface of soybean seeds was sterilized by immersing in sodium hypochlorite solution (1%) for 3 min. and rinsed with distilled water. Soybean seeds are put into a Petri dish which contains 3 sheets of
sterile filter paper that have been moistened with sterile aquadest. In this method, three pieces of filter paper were soaked in sterilized distilled water and placed on the bottom of the petri dish. Soybean seeds were taken randomly from each variety and then placed on moist filter paper. The Petri dishes were then incubated at room temperature (± 26°C) for 7 days. The growth of fungus was observed on 8th-day using stereomicroscope and compound microscopy. In addition, the percentage of seed germination was also observed (de Camargo et al. 2017).

Isolation of fungi on agar plate
40 soybean seeds per variety were used for the agar plate. Soybean seeds were sterilized with 70% alcohol for 30 seconds, followed by 1% NaOCl for 3 minutes and then washed with sterile distilled water. The seeds were placed into a petri dish containing PotatoDextrose Agar (200 g of potato extract, 18 g of agar, 20 g of dextrose, and 1000 mL of distilled water), 10 seeds per petri dish and then incubated for 7 days at room temperature. A pure culture of each fungus was maintained on PDA for identification purposes (Wain-Tassi et al. 2012).

Fungal identification
Isolated fungal species were identified on the basis of their gross colony morphology (shape, size, and color of colony) and microscopic feature (characteristics of mycelium, shape, size and color of conidia and conidiophores, setae, etc.) as described by (Benoit and Mathur 1970; Ellis 1971; Domsch et al. 1980; Watanabe 2002; Gaddeyya et al. 2012; Alsohaili and Bani-Hasan 2018; Amteme and Tefa 2018).

Seed germination (%)
The percentage of seed germination was calculated by applying the formula of ISTA (1996):

\[
\text{Seed germination} \% = \frac{\text{(Number of seed germinated)}}{\text{(Number of total seeds)}} \times 100
\]

Data analysis
Seed germination was analyzed using the F test with a confidence level of 95% and further tested with a HSD level of 5%.

### RESULTS AND DISCUSSION

#### Isolation and identification of seed-borne pathogenic fungi

Results of (Table 1) blotter test method showed that all the eight varieties of soybean were infected with seed born pathogenic fungi. Fungi were found to colonize in the tested seeds. A total of eight fungal species namely *Aspergillus flavus* Link, *Aspergillus niger* van Tieghem, *Cladosporium oxysporum* Berk. & M.A. Curtis, *Colletotrichum dematium* (Pers. Et Fr.) Grove f.sp. *truncate* (Schw.) Arx, *Curvularia pallescens* Boedijn, *Fusarium solani* (Mart.) Sacc., *Melanospora zamiae* Corda, and *Nigrospora* sp. Mason were isolated from soybean seeds.

Based on Tables 1 and 2, it can be seen that more fungi were obtained in the blotter test than PDA. The blotter test method can detect pathogenic fungi found on the surface of seed. A total of eight fungi were isolated by the blotter test and only four fungi were isolated by PDA. Rao et al. (2015), reported that blotter test detected more fungi than agar plate. Pathak and Zaidi (2013) stated that blotter method was found to be the best method for the isolation of mycoflora whether borne externally or internally. In general, on both blotter methods and PDA significantly difference among soybean varieties were found for fungal isolates.

### Table 1. Fungi isolated from different varieties of soybean seeds by blotter test

| Fungi                 | MI   | Kb   | Dr   | Dt   | Sn   | Dn   | Gk   | Sl   |
|-----------------------|------|------|------|------|------|------|------|------|
| *Aspergillus flavus*  | +    | +    | -    | -    | +    | +    | -    | +    |
| *Aspergillus niger*   | -    | -    | -    | -    | +    | -    | -    | -    |
| *Cladosporium oxysporum* | +    | +    | +    | +    | +    | +    | +    | +    |
| *Colletotrichum dematium* | -    | -    | +    | -    | -    | +    | -    | -    |
| *Curvularia pallescens* | +    | -    | +    | -    | -    | -    | -    | +    |
| *Fusarium solani*     | +    | +    | +    | +    | +    | +    | +    | +    |
| *Melanospora zamiae*  | -    | -    | -    | -    | -    | -    | -    | -    |
| *Nigrospora* sp.      | -    | +    | -    | -    | -    | -    | -    | -    |

Note: +: found, -: not found, MI: Malabar, Kb: Kaba, Dr: Dering, Dt: Detam I, Sn: Sinabung, Dn: Dena, Gk: Gepak Kuning, Sl: Slamet

### Table 2. Fungi isolated from different varieties of soybean seeds by PDA

| Fungi                 | MI   | Kb   | Dr   | Dt   | Sn   | Dn   | Gk   | Sl   |
|-----------------------|------|------|------|------|------|------|------|------|
| *Aspergillus niger*   | -    | -    | -    | -    | -    | -    | -    | +    |
| *Cladosporium oxysporum* | -    | +    | -    | -    | -    | +    | -    | -    |
| *Colletotrichum dematium* | -    | -    | +    | -    | -    | -    | -    | -    |
| *Fusarium solani*     | +    | -    | +    | -    | -    | +    | -    | +    |

Note: +: found, -: not found, MI: Malabar, Kb: Kaba, Dr: Dering, Dt: Detam I, Sn: Sinabung, Dn: Dena, Gk: Gepak Kuning, Sl: Slamet
Tables 1 and 2 showed that the most common seed-borne fungi were found in the Slamet variety. It is assumed that Slamet variety has vertical resistance (qualitative) that is organized by one gene and only to several types of pathogens. Varieties with vertical resistance are easily infected with new pathogens. Based on Nelson (2003), vertical resistance is usually inherited by a single gene or a number of genes. Local varieties have horizontal resistance known as quantitative resistance as a type of resistance that is resistant to all types of pathogens. Horizontal resistance varieties show little sensitivity to pathogens, but have the ability to slow the rate of disease progression. *Cladosporium oxysporum* and *Fusarium solani* were recorded as the dominant fungal species on blotted test and PDA (Tables 1 and 2). Kinniker et al. (2015) reported that the pathogenic genus *Cladosporium* and *Fusarium* often infected the soybean seeds. The morphological characteristics of the seed-borne pathogenic fungus are as follows.

**Aspergillus flavus** Link

*Aspergillus flavus* was found in Malabar, Kaba, Sinabung, Dena, and Slamet varieties. Macroscopically, *A. flavus* colonies were yellowish-green, rough conidium structure, and granular in shape (Figure 1.A). Praja and Yudhana (2017) observed that colonies of *A. flavus* were first white and then finally turn into yellowish-green color. The reverse side showed yellowish to brown color. *A. flavus* has a rounded shape with a flat edge (Mathur and Kongsdal 2003). Mycelia was in the form of fine threads (Hameed et al. 2012; Krijgsheld et al. 2013).

Macroscopically, *A. flavus* has a hypha that was insulated, fialid, metula, erectile conidiophores, colorless, and has foot cells (Figure 1.B). According to Procop et al. (2016), *A. flavus* has hyphae insulated, erectile conidiophores, unbranched, colorless, the edges form round vesicles that carry conidia. Conidiophores were clear, rough, not pigmented, and less than 1 mm long (Gautam and Bhadauria 2012).

**Aspergillus niger** van Tieghem

*Aspergillus niger* found in Sinabung and Slamet varieties. Macroscopically, *A. niger* colonies were black with white edges and spread (Figure 2.A). According to Koneman (1979) and Praja and Yudhana (2017), *A. niger* has black colony with white edges, the lower surface of the colonies was yellowish to brown.

Conidia were spherical and black in color. Vesicles were spherical, and conidiophores were colorless (Figure 2.B). The *A. niger* fungus has a visibly cotton-like filamentous called hyphae, has a large, rounded carrier head, black conidia, with conidiophores that contain many pigments (Chang et al. 2020), Walsh et al. (2018), Mathur and Kongsdal (2003), Gautam et al. (2011), and Wangge et al. (2012) resulted in long and colorless vesicles, round to semi-round vesicles, conidia round to semi-round and brown to black.

**Cladosporium oxysporum** Berk. & M.A. Curtis

*Cladosporium oxysporum* was found in all soybean varieties. The colonies of *C. oxysporum* were white to grayish brown to blackish-green color and shaped like cotton-like to velvet shape (Figure 3.A). Briceño and Latorre (2007) and Guan et al. (2016) stated that *C. oxysporum* has light green brownish to blackish-green colony, and growth is relatively slow.

Microscopic observation of the *C. oxysporum* has an elliptical conidium, one of the ends is tapered and there are black spots, sometimes there is a divider (Figure 3.B). Based on Lamboy and Dillard (2007), the microscopic features of *C. oxysporum* are hyphae that are insulated, conidiophores are lateral or terminal in hyphae, conidia are chain-shaped and smooth-walled. An elliptical conidium is found in a branched-chain. The *C. oxysporum* has a small colony that is blackish green and pollinated (Figure 3b). Spores develop at the ends of complex conidiophores that grow from one insulated mycelium. Usually brownish.

**Colletotrichum dematium** (Pers. et Fr.) Grove f.ssp. *truncata* (Schw.) Arx.

The fungus *C. dematium* is found in the Ringer, Detam I, Sinabung and Gepak Kuning varieties. Macroscopically a single aserulovus colony and / or group, sometimes fused, aservulus blackish-brown was observed (Figure 4.A). Based on Soesanto (2015), *C. dematium* fungus has a dark aservulus and develops on patches in the host tissue.

Acerulovus of *C. Dematium* has 1-4 bulkhead, blackish-brown color and, its size was longer than conidia. Conidia were hyaline, curved with pointed ends (Figure 4.B.). According to Haggag and Singer (2013), *C. dematium* colonies form concentric rings, curved conidia with tapered ends, and setae. Hyphae were brown in color, 3-4 septate. Setae were dark brown in color. *Curvularia pallescens* Boedijn

**Curvularia pallescens** Boedijn

*Curvularia pallescens* found in Malabar, Dering, Detam I, and Slamet varieties. Macroscopically the *C. pallescens* spread and form like feathers (Figure 5.A). According to Hoog et al. (2001), *C. pallescens* has a scattered, hairy colony in the middle, often developing concentric zones.

Conidia of *C. pallescens* was light brown to brown, mostly straight and slightly curved, has 3 baffles and 4 cells, the third cell was larger than other cells (Figure 5.B). Hoog et al. (2001), stated that conidia can look slightly bent but mostly straight. The conidia has 3 septae giving rise to 4 cells. The third cell from the base looks swollen compared to the cells around it. Its texture looks smooth and pale brown to brown (Dey et al. 2016).

**Fusarium solani** (Mart.) Sacc.

*Fusarium solani* found in all varieties, namely Malabar, Kaba, Dering, Detam I, Sinabung, Dena, Yellow Gepak, and Slamet. *F. solani* has white to creamy mycelium, clumping like a thick cloud, and droplets were found on the surface of the colony (Figure 6.A). Similar observations were reported by Hafizi et al. (2013).
Figure 1. Macroscopic and microscopic *Aspergillus flavus*. A. *A. flavus* colony at 50x magnification; B. Vesicle and conidia of *A. flavus* at 400x magnification

Figure 2. Macroscopic and microscopic *Aspergillus niger*. A. *A. niger* colony at magnification 30x; B. Vesicle and conidia of *A. niger* magnification at 400x

Figure 3. Macroscopic and microscopic *Cladosporium oxysporum*. A. *C. oxysporum* colony at 30x magnification; B. Conidia at 400x magnification

Figure 4. Macroscopic and microscopic *Cladosporium dematium*. A. *C. dematium* colony at 30x magnification; B. Conidia at 1000x magnification

Figure 5. Macroscopic *Curvularia pallescens*. A. *C. pallescens* colony at 30x magnification; B. *C. pallescens* conidia at 400x magnification

Figure 6. Macroscopic and microscopic *Fusarium solani*. A. *F. solani* colony magnification at 63x magnification; B. *F. solani* conidia at 400x magnification

Figure 7. Macroscopic and microscopic *Melanospora zamiae*. A. Colony of *M. zamiae* at 30x magnification; B. Conidia at 400x magnification

Figure 8. Macroscopic and microscopic *Nigrospora* sp. A. *Nigrospora* sp. at 50x magnification; B. *Nigrospora* sp. Conidia at 400x magnitude.
In *F. solani* microconidia were 1-2 celled, colorless, and oval shape. Macroconidia had 3-4 septa and colorless (Figure 6B). Based on research by Hafizi et al. (2013) reported that in single-cell microconidia were oval to kidney-shaped while, macroconidia were thick-walled, generally cylindrical with dorsal ends and ventral ends parallel.

**Melanospora zamiae Corda**

*Melanospora zamiae* found only in Slamet variety, the fungal colony is yellow and has a neck, black color (Figure 7A). According to Mathur and Kongsdal (2003), *M. zamiae* has a superficial peritheium fungus, light white to golden yellow, translucent, round with short to long necks, dark mass, containing ascospores usually seen flowing from the neck.

Microscopically the conidia were lemon-shaped, brown to almost black color (Figure 7B). According to Mathur and Kongsdal (2003), *M. zamiae* has ascospores of lemon-shaped (citriform) and dark brown to almost black color.

**Nigrospora sp. Mason.**

*Nigrospora* sp. was found only in the Kaba variety. Blackish gray mycelium (Figure 8A). Hao et al. (2020) reported that *Nigrospora* sp. has brownish-gray colonies with white edges, solid surface texture and rounded colonies. It has a hyphae insulated and colorless, single conidia, round in shape, and deep black or not transparent (Figure 8B). According to Hao et al. (2020), *Nigrospora* sp. has colorless conidiophores and insulated hyphae, conidia were black and unicellular.

**Seed germination ability**

From Table 3, germination rate of soybean seeds was found to be different in each variety. The highest seed germination was found in the Daring and Detam 1 soybean varieties as 80.5%, respectively, and the lowest one was found in Dena variety as 53%. The highest (80.5%) seed germination was found in the Daring and Detam 1 varieties and the lowest (53%) was found in Dena variety.

Based on Table 3, Dering and Detam 1 varieties was classified as high seed germination varieties. Whereas, Sinabung, Dena, Gepak Kuning, and Slamet varieties were classified as low seed germination varieties. The statement in accordance with the germination standard which is classified as high for almost all seeds if the results of the seed germination test showed a value of ≥ 80% (Rahayu and Suharsi 2015).

**Table 3.** Percentage of seed germination in various varieties

| Varieties  | Seed germination (%) |
|------------|----------------------|
| Malabar    | 79.0 d               |
| Kaba       | 78.0 d               |
| Dering     | 80.5 d               |
| Detam      | 80.5 d               |
| Sinabung   | 68.5 c               |
| Dena       | 53.0 a               |
| Gepak Kuning| 71.5 c              |
| Slamet     | 59.5 b               |

Note: Number followed by different letters in the same column show significantly different from HSD at the 5% level.

Although 4 pathogenic fungi infected Daring and Detam 1 varieties (Tables 1 and 2), still these varieties showed the highest percentage of seed germination (Table 3). It can be suspected that these two varieties have horizontal resistance so that they were able to withstand the attack of pathogenic fungi. Penfield and MacGregor (2017) state that the percentage of seed germination has been affected by many factors including pathogen attack, genetic factors, and the environment.

Dena variety has the lowest germination rate (Table 3), but associated with 3 fungi (Tables 1 and 2). It was suspected that the Dena variety was frequently planted as 2-3 times resulted in decreasing germination compared to the first planting. Another factor that can trigger low seed germination capacity is uneven imbibition of the seed so that the growth of the seed becomes non-synchronized normal sprouts (Rahayu and Suharsi 2015).

The emergence of many pathogenic fungi may be caused by unsterile conditions of the germination substrate, seed germination tool, storage, and water used. Ghangaoar and Kshirsagar (2013) suggested that seed-borne microbes mostly found in those seeds that do not receive seed treatment. Due to microbial infestation, plant vigor and seed germination capacity decreases.

In conclusion eight pathogenic fungi namely, *Aspergillus flavus, A. niger, Cladosporium oxysporum, Curvularia pallescens, Fusarium solani, Melanospora zamiae*, and *Nigrospora* sp. were found to be associated with soybean seeds. The Daring and Detam 1 varieties showed highest (80.5% each) seed germination while lowest (53%) was found in Dena variety.

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