Culturable endophytic fungal diversity in cassava tubers of Indonesia

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ABSTRACT
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INTRODUCTION
Endophytic fungi inhabit plant tissue and do not cause disease symptoms in their host plants. These fungi are present in tissues of seeds, flowers, fruits, stems, roots, and tubers, and protect host plants from various environmental stresses. A symbiotic mutualism relationship occurs between endophytic fungi and their associated host plants (Chadha et al. 2014). Indonesia is one of the largest developing country consuming cassava tubers. This tuber has great starch content, low price, and easy to grow even in low fertile soils. Cassava is expected to substitute starch from rice with a very high crop failure rate due to a large number of pests, diseases, and climate changes that can cause drought (Sucipto et al. 2015). However, the community of endophytic fungi inhabiting cassava tubers is not well studied in Indonesia.

One endophyte Guignardia endophyllicola was isolated from cassava stalks originating from agricultural land and supporting forest in Parakan Salak Village, Sukabumi (Suciamih and Supriyati 2016). On the other hand, endophytic fungi belonging to Penicillium, Fusarium, and Aspergillus were isolated from cassava growing in the Nature Reserve Mount, East Nusa Tenggara (Ilyas 2006).

In this study, we focused on the isolation and molecular identification of endophytic fungi isolated from cassava tubers collected from Bogor, Lembang, Makasar, Serang, Singaraja (Buleleng), Subang, Wonogiri, and Wonosobo of Indonesia. Moreover, information on the diversity of endophytic fungi associated with cassava tubers is cultivated in Indonesia and it is still lacking especially the molecular identification of the endophytic fungi.

In general, identification of endophytic fungi has been mostly carried out by morphological observations. However, it has challenges to identify at the species level. At first, endophytic fungi may be Mycelia sterilia (Lacap et al. 2003; Hidayat et al. 2016), where reproductive structures are not formed for morphological characterization. Therefore, in this study, we report the diversity of endophytic fungi in cassava tubers originating from several regions in Indonesia by identifying genetic differences using sequences of ITS rDNA with ITS4 and ITS5 primers to complement morphological characters.

MATERIALS AND METHODS
Materials
Cassava (Manihot esculenta) samples were collected from October 13, 2016 to December 19, 2017 in traditional markets in Bogor 106°43′30″-106°51′00″ Longitude East (LE) and 03°30′ – 6°41′00″ Latitude South (LS); Serang 105°7′-105°22′ LE and 5°50′-6°21′ LS; Singaraja 08°03′40″-08°23′00″ LS and 114°25′-115°15′27′-28′ 28′ LE; Subang 6°34′15.96″ LS and 107°05′-6′21′ LS; Wonogiri 7°32′-8°15′ LS and 110°41′-111°18′ LE; Wonosobo 7°8′8″ LS and 109°55′54″ LE.

Endophytic fungi isolation
Fresh cassava tuber samples were peeled and washed under running water for 10 min. The tubers were cut into 2 × 2 cm. The tuber pieces were then sterilized with 70%
alcohol for 1 min and followed by 1% sodium hypochlorite solution for 2 min. The tuber samples were soaked and shaken with sterile distilled water three times for each 5 min. The sterilized samples were dried with sterile tissue paper for 3 h. The surface of the cassava samples was sliced off and the inside part of cassava tubers was taken aseptically and placed on a petri dish containing potato dextrose agar (PDA) medium and chloramphenicol. Each petri dish contained three pieces of 1.5 × 1.5 cm cassava tubers from one sampling location. The success of surface sterilization was carried out using 10 µL of the last rinse distilled water of the cassava tuber, washed, and smeared on a sterile PDA medium using a micropipette. The test was carried out to find out if there were contamination on the surface-sterilized cassava tubers. The cultures were incubated at 28°C for 2 weeks. The fungal growth was monitored daily.

DNA isolation

For DNA isolation, the fungus was cultured on PDA slants for four days at 30°C. The fungal mycelium was transferred into a 1500 µL Eppendorf tube containing 500 µL nuclease free water (NFW). Centrifugation was carried out at 9,600 × g for 10 min. The supernatant was discarded and pellet was crushed by using a sterile pestle until the entire mycelium was smooth. DNA isolation was performed using the DNA protocol Phytopure™ kit extraction (GE Healthcare, UK) with modifications. Reagent 1 was added as much as 300 µL into the pellet and resuspended with a micropipette for cell lysis. RNase (20 µM) was added as much as 3 µL, resuspended and incubated at 37°C for 30 min. A total of 200 µL of reagent 2 was added and shaken until blended. The samples were incubated at 25°C for 10 min and the ice temperature for 20 min. The mycelium suspension was added with 500 µL of cold solvent (phenol:chloroform: isopropanol = 1:1:1), then shaken with sterile distilled water three times for each 5 min. The sterilized samples were dried with sterile tissue paper for 3 h. The surface of the cassava samples was sliced off and the inside part of cassava tubers was taken aseptically and placed on a petri dish containing potato dextrose agar (PDA) medium and chloramphenicol. Each petri dish contained three pieces of 1.5 × 1.5 cm cassava tubers from one sampling location. The success of surface sterilization was carried out using 10 µL of the last rinse distilled water of the cassava tuber, washed, and smeared on a sterile PDA medium using a micropipette. The test was carried out to find out if there were contamination on the surface-sterilized cassava tubers. The cultures were incubated at 28°C for 2 weeks. The fungal growth was monitored daily.

Molecular identification of endophytic fungi

The hyphae of the fungal isolate maintained in a petri dish were taken inoculated in the potato dextrose agar (PDA) and incubated for 5 days at 25°C. DNA isolation was performed using the DNA protocol Phytopure™ kit extraction (GE Healthcare, UK) with modifications.

PCR amplification of ITS rDNA sites was carried out with ITS 5 (forward: 5’–TCTCCCGCTTATTGATATGC–3’) and ITS 4 (reverse: 5’–TCCGTAGGTGAACCTGCGC–3’) primers (White et al. 1990) by using Gene Amp® PCR System 2400. The PCR reaction for each sample was carried out in 50 µL of the reaction mixture that contained 2 µL of DNA genome with a concentration of approximately 100 ng/µL. Each primer (ITS4 and ITS5) was added at 1.25 µL with a concentration of 10 µM (Innis et al. 2012), in amount of 25 µL of Go Taq Green Master Mix PCR 2x, and 20.5 µL of NFW. The PCR condition that was used as follows: pre-denaturation was at 94°C for 2 min, followed by 35 cycles at 94°C for 15 sec in denaturation process, annealing was at 55°C for 30 min, elongation was at 72°C for 1 min, and final elongation was at 72°C for 5 min. The PCR results were visualized with agarose gel 1% 60 volts for 90 min, then it was immersed in a solution of ethidium bromide for 15 min, and it was rinsed with distilled water. The observation was carried out under UV light.

Molecular analysis was performed at the First Base Company in Malaysia. The results of the DNA sequence were edited by using ChromasPro and nucleotide base results blasted by using the basic local alignment search tool nucleotides (BLASTN) platform. After the blast results were obtained, the out-group and reference data can be downloaded from GenBank (www.ncbi.nih.gov/Genbank). All data were aligned by using the molecular evolution and genetic analysis 7 (MEGA 7) platform. The phylogenetic tree was constructed by using the Maximum Likelihood (ML) Tamura-Nei model or Neighbor-Joining (NJ) by using MEGA 7.

Morphological identification of endophytic fungi

Morphological identification was carried out by macroscopic and microscopic morphological observations (Barnett and Hunter 1998). Macroscopic observations were carried out by observing the fungal diameter size, colony color on PDA media, colony surface structure, and colony edges. Microscopic observations were carried out by observing the reproductive structure using Riddell’s classic slide culture method (Riddell 1950).

RESULTS AND DISCUSSION

A total of 14 isolates of endophytic fungi from cassava tubers were isolated and identified based on molecular characterization. These endophytes were found in cassava tubers from eight regions of Indonesia. Based on molecular characterization the endophytic fungi were from genus Aspergillus (4 isolates), Fusarium (6 isolates), and other four isolates included Lasiodiplodia sp., Nectria pseudotrichia, Penicillium citrinum, and Schizophyllum commune (Table 1).

Previous study reported that endophytic fungi were isolated from other parts of cassava such as petiole (Suciatmih and Supriyati 2016) and root (Ilyas 2006). This study provided important information regarding the diversity of endophytic fungi that was found in fresh cassava tubers. Endophytic fungi also can be found in other plants besides cassava, such as quina (Hidayat et al. 2016a), maize (Suriani and Muis 2017), and mangosteen (Akmalasari et al. 2013). The phylogenetic construction tree of Fusarium spp. was used as reference sequence retrieved GenBank.
**Aspergillus spp.**

A total of four isolates of *Aspergillus* were successfully isolated from cassava tubers, originating from areas of Bogor (FS7), Wonogiri (FS8), Serang (FS9), and Singaraja (FS10). Based on molecular analysis of phylogenetic trees, it identified as three strains of *Aspergillus* sp. (FS7, FS8, and FS9 strains) and one *A. fumigatus* strain (FS10 strain). *Diaporthe angelicae* CBS 123215 was used as out-group. Based on the ITS rDNA sequence analysis, the FS7, FS8, and FS9 strains belonged to the same clade as *A. flavus* 26, 10, 14, 24, C1931, MB42, and 12.25. Meanwhile, *A. oryzae* FS1 125, and *A. parasiticus* DAOM 225948 with a bootstrap value was 97%, led the species name cannot be determined. FS10 strain was *A. fumigatus* with a bootstrap value is 100% based on phylogenetic tree analysis. Based on the phylogenetic analysis, there were three species of fungi in the same clade with the FS7, FS8 and FS9 strains, they were *A. flavus*, *A. oryzae*, and *A. parasiticus*. These isolates belonged to the genus *Aspergillus*. Based on ITS rDNA sequences analysis, strain FS7, FS8, and FS9 identified at genus level (Figure 1.A).

Colony of *Aspergillus* sp. FS7, FS8, and FS9 spread over the PDA media with a 6 cm diameter after 4 days of incubation time. Macroscopic observation showed that the colony was green and had a velvety surface (Figure 1.B, 1.D, and 1.F). The FS7, FS8, and FS9 strains have good conidiophore, insulated hyphae, round vesicles, contained 1.D, and 1.F). The FS7, FS8, and FS9 strains and have green colony, round conidia, smooth surfaces. *Aspergillus. flavus* and *A. parasiticus* have green colony characteristics, round shape conidia, and smooth surfaces. *Aspergillus. flavus* and *A. parasiticus* have green colony characteristics, round shape conidia, and smooth surfaces. *Aspergillus. flavus* and *A. parasiticus* have green colony characteristics, round shape conidia, and smooth surfaces. *Aspergillus. flavus* and *A. parasiticus* have green colony characteristics, round shape conidia, and smooth surfaces. *Aspergillus. flavus* and *A. parasiticus* have green colony characteristics, round shape conidia, and smooth surfaces. *Aspergillus. flavus* and *A. parasiticus* have green colony characteristics, round shape conidia, and smooth surfaces. *Aspergillus. flavus* and *A. parasiticus* have green colony characteristics, round shape conidia, and smooth surfaces. *Aspergillus. flavus* and *A. parasiticus* have green colony characteristics, round shape conidia, and smooth surfaces.

The growth of *A. fumigatus* FS10 strain spread unevenly on PDA media, 6 cm in diameter after 4 days of incubation. Colony surface was dark green powder. Microscopically, *A. fumigatus* has insulated hyphae, conidiophores that were upright, not contain septa, unbranched, and conidiophore’s tip swollen to form vesicles. Fialid was vesicles bottle-shaped formed directly over vesicula that produced the round conidia (Figure 1.E and 1.G). *Aspergillus fumigatus* was fungi that harm humans. These fungal spores can cause respiratory diseases in humans, such as asthma, cystic fibrosis, and aspergillosis (Dagenais and Keller 2009). However, further research is needed to determine the positive and negative impacts of the strains derived from cassava tubers.

**Fusarium spp.**

Six strains of *Fusarium* spp. were isolated successfully from cassava tubers from areas of Wonosobo (FS1 and FS3), Wonogiri (FS2 and FS4), and Singaraja (FS6 and FS10). Molecular identification showed that isolates FS1- FS6 were *Fusarium oxysporum* (FS1), *F. lichenicola* (FS2 and FS4), *F. falciforme* (FS3 and FS6), and *F. solani* (FS5) as depicted in Figure 2.

The FS1 strain formed monophyletic clade with *F. oxysporum* CBSI 133023T and *F. oxysporum* YNMG-1 with a bootstrap value was 64% (Figure 2.A); thus, strain FS1 was identified as *F. oxysporum*. The fungal colonies of *F. oxysporum* grew to fill a 6 cm diameter petri dish for 2 days of incubation in PDA media. The FS1 strain has a thick white mycelium like cotton. After 2 days, the colony turned purplish (Figure 2.B). The microscopic morphology observation of *F. oxysporum*, showed this fungus has much microconidium, oval shape, without insulation and hyalin. Macroconidia were not found, but had monophylalid conidiophore (Figure 2.C).

The FS2 and FS4 strains formed the same clade as *F. lichenicola* NRRL32434 with high bootstrap values (98%). Thus, the FS2 and FS4 strains were identified as *F. lichenicola* species. The FS2 strain grew of 3 cm in 6 cm diameter size PDA media for 4 days with white mycelium growth on the edge of the colony (Figure 2.D). The FS4 strain grew up to 4 cm diameter on a 6 cm-sized PDA media petri dish with 4 days of incubation time. The mycelium was white and thick like cotton. Based on macroscopic observation, the FS2 and FS4 strains changed PDA media color became somewhat reddish. Microscopic observation showed that microconidia have oval shape and hyalin without bulkhead (Figure 2.E). During the observation, macroconidia were not found in FS2 and FS4 strains.

| Place of origin | Date of isolation | Isolate code | Species             |
|-----------------|------------------|--------------|---------------------|
| Bogor           | October 16, 2016 | FS7          | *Aspergillus* sp.   |
| Wonogiri        | January 23, 2017 | FS7          | *Aspergillus* sp.   |
| Serang          | November 7, 2016 | FS9          | *Aspergillus* sp.   |
| Singaraja       | February 17, 2017| FS10         | *A. fumigatus*      |
| Wonosobo        | January 16, 2017 | FS3          | *Fusarium falciforme*|
| Singaraja       | February 17, 2017| FS10         | *F. falciforme*     |
|                  |                  |              |                     |
| Subang          | November 7, 2016 | FS11         | *Nectria pseudotrichia* |
| Subang          | November 7, 2016 | FS13         | *Penicillium citrinum* |
| Bogor           | October 16, 2016 | FS14         | *Schizophyllum commune* |
The FS3 and FS6 strains formed a monophyletic clade with *F. falciforme* NRRL 32757 and the value of bootstrap was 88%. Thus, it was identified as *F. falciforme*. In the FS3 strain, the colonies grew and covered the PDA media with a 6 cm diameter on the third day (Figure 2.F). Macroscopic observation revealed that this strain was white and cottony. The microconidia were oval, conidiophores without septa, and hyaline (Figure 2.G). Meanwhile, macroconidia were slightly curved with pointed ends, one-bulkhead, and hyalin. The FS6 strain grew to cover the petri dish media with a 6 cm diameter on the third day. The colonies were white and thin than FS3 and had the same microscopic appearance as the FS3 strain.

The FS5 strain was identified as *F. solani* with a bootstrap value of 91% formed the same clade as *F. solani* NRRL 28579T (Figure 2.A), colonies on PDA media, grew
up to 3 cm in diameter after two days of incubation (Figure 2h). Colonies were characterized by a slightly yellowish-white color, thin-shape with cottony hyphae that were aerial and the colonies edge was irregular. Microconidia were oval without a bulge, and hyalin (Figure 2.1). **Fusarium** was also found in the other plants, such as maize (Suriyani and Muis 2017), chili (Nurzannah et al. 2014), quina (Hidayat et al. 2016b), and *Cenntelia asiatica* (Radiastuti et al. 2019). **Fusarium** generally produces a secondary metabolite, such as fumonisin. According to Onana et al. (2015), fumonisin has a negative impact on humans, which can stimulate cancer cell activity in humans. Fumonisin can be found in daily food products, such as cassava chips. Cassava chips that are stored for a long time (over 180 days) could increase fumonisin in cassava chips (Onana et al. 2015). This fungus plays a role as a biocontrol agent or pathogen resistance in several plants (Rubini et al. 2005).

In previous studies, *F. oxysporum* (FS1 strain) was able to increase resistance in tomato plants because it can kill pathogenic fungi, such as *Phytophthora infestans*, which causes tomato blight, and *Cladosporium fulvum*, which causes tomato leaf mold disease (Fuchs et al. 1997) and acts as a biocontrol agent in hydroponic plants (Thongkamngam and Jaenaksorn 2017). In banana plants, this fungus can kill pests and nematode worms that can damage crop yields (Waweru et al. 2014), whereas, in cassava plants, this fungus can cause dry rot disease (Boas et al. 2017). *F. lichenicola* (FS2 and FS4 strains) have similarities with *Cylindrocarpon lichenicola*. This fungus was isolated from cassava tubers that collected from Wonosobo and Singaraja Acremonium falciforme was isolated from cassava tubers from Wonogiri area. This fungus belongs to tropical fungi that grow in high rainfall areas show different types of inhabited fungi strains. This fungus is commonly found in cassava tubers and can cause tuber rot (Boas et al. 2017). Cassava tubers that grow in different areas show different types of inhabited fungi strains. This can be due to the seasonal variations and environmental factors, both biotic and abiotic factors (Irawati et al. 2014). Further research is needed to determine the positive and negative impacts of the **Fusarium** spp.

**Lasiodiplodia sp.**

One isolate of **Lasiodiplodia** sp. FS12 was isolated successfully from cassava collected from Makasar (Figure 3). **Lasiodiplodia** sp., phylogenetic tree construction of *Diplodia mutilata* CBS 136015 and Botryosphaeria obtusa CBS 11255 were used as out-groups. The results showed that the FS12 strain forms a clade that was similar to *L. egyptiaceae* BOT10, *L. euphorbicola* CMM 3609, *L. hortozuganensis* IRAN500C, *L. parva* CBS 45678T, CBS494.78, and Botryosphaeria rhodina CBS16496T, with the value of the bootstrap was 48% (Figure 3.A). The genus of **Lasiodiplodia** was also found in Brazil which inhabited cassava (Boas et al. 2017). This endophytic fungus is found in other plants, such as *Albizia chinensis* (Rakhmawati and Prihatini 2016).

Morphologically, the colonies of FS12 grew covering up the surface of the PDA media, 6 cm diameter size of Petri dishes on the third day of incubation. The FS12 strain has grayish-white mycelium which can turn to blackish with thick and smooth shape like cotton characteristics (Figure 3.B). This strain can turn the color of PDA media into black. Microscopically, these colonies have insulated hyphae and no reproductive organs (Figure 3.C).

This fungus often causes diseases such as rotting fruit and damage on the stems. It is also known as endophytic fungi (Slippers and Wingfield 2007). **Lasiodiplodia** usually grows in areas with tropical and subtropical climates. This fungus often causes diseases in plants (Abdollahzadeh et al. 2010). Boas et al. (2017) reported that **Lasiodiplodia** spp. can cause black rot in cassava. A previous study showed that the presence of fungus **Lasiodiplodia** spp., on the gatotan (fermentation from cassava tubers that is stored for a few days before being steamed) and does not have any bad effect on human health (Purwandari et al. 2014). Of course, this matter needs further research.

**Lasiodiplodia** is an anamorph form of Botryosphaeria. In this study, strain FS12 formed a clade equal to *L. egyptiaceae* BOT10, *L. euphorbicola* CMM3609, *L. hortozuganensis* IRAN1500C, *L. parva* CBS456.78 and CBS 494.78, and Botryosphaeria rhodina CBS16496T. The use of other genes, such as elongation factor α and β-tubulin, is required to separate FS12 strain in separate clades (Jang et al. 2012).

**Nectria pseudotrichia**

The FS11 strain was identified as **Nectria pseudotrichia**. Based on the molecular analysis, bootstrap value was 81% (Figure 4). The FS11 strain formed a monophyletic clade with *N. pseudotrichia* CBS551.84 (Figure 4.A). *Cosmospora coccinea* AR2741 was used as an out-group. This fungus was isolated from cassava tubers collected from Subang. The FS11 strain grew covering up the PDA media, up to 6 cm in diameter after five days of incubation, has a smooth colony surface, and white color (Figure 4.B). Microscopically, these fungi have septate hyphae (Figure 4.C). However, reproductive organ was not found in this isolate. Endophytic fungi in the genus **Nectria** were also found in the other plants, such as *Annona squamosa* (Yunianto et al. 2012). However, *N. pseudotrichia* fungi was not found as endophyte in the other plant species. This fungus lives in tropical and subtropical areas. *N. pseudotrichia* looks like saprophytic. However, some species are endophytes. This species functions as a plant pathogen in the tropics region (Hirooka et al. 2012).
Figure 2. Phylogenetic tree of *Fusarium* spp. based on NJ analysis with 1000 bootstrap replicates. A. *Penicillium citrinum* AX4602 was used as an out-group. Morphological identification of: B. Colony of *Fusarium oxysporum* FS1; C. Microconidia of *Fusarium oxysporum* FS1; D. Colony of *Fusarium lichenicola* FS2; E. Microconidia of *Fusarium lichenicola* FS2; F. Colony of *Fusarium falciforme* FS3; G. Microconidia of *Fusarium falciforme* FS3; H. Colony of *Fusarium solani* FS5; and I. Microconidia of *Fusarium solani* FS5.
Figure 3. Phylogenetic tree of: A. Lasiodiplodia sp. based on ML analysis with 1000 bootstrap replicates, Diplodia mutila CBS136015 and Botryosphaeria obtusa CBS112555 was used as the out-group. Morphological identification of endophytic fungi: B. colony of Lasiodiplodia sp. FS12; C. septate hyphae of Lasiodiplodia sp. FS12.

Penicillium citrinum

The FS13 strain was isolated successfully from cassava collected from Subang area. The molecular identification results showed that the FS13 strain formed a clade that was the same as Penicillium citrinum NRRL1841T, NRRL35449, NRRL35448, NRRL35434, P-1637 with a high bootstrap value of 100% (Figure 5). Thus, it can be ascertained that this strain was P. citrinum. Paecilomyces variotii DTO63F5 used as outgroup (Figure 5.A). Besides as a part of an endophyte collected from cassava, P. citrinum has also been found on other plants such as rice grains (Reddy et al. 2010) and quina (Hidayat et al. 2016a). Another study also reported Penicillium capsulatum found in Centella asiatica (Radiastuti et al. 2019).

Based on macroscopic morphological observations, it was found that colonies started with white and turned bluish-green color after 5 days of incubation. The colony was hard-skinned and the surface looks like powder (Figure 5.B). Microscopically, this fungus has a metula and fialid shaped like a bottle and branched conidiophores. The conidia were round and arranged in a basipetal fashion (Figure 5.C).
Figure 4. Phylogenetic tree of: A. *Nectria pseudotrichia* based on ML analysis with 1000 bootstrap replications. *Cosmospora coccinea* AR2741 was used as an out-group. Morphological identification of endophytic fungi: B. Colony of *Nectria pseudotrichia* FS11 in the PDA media; C. Septate hyphae of *Nectria pseudotrichia* FS11.
Penicillium citrinum is a fungus that usually grows in plantations or forests of tropical areas (Amaria et al. 2013). The characteristics of the *P. citrinum* are bluish-green colonies, powdery surfaces, aseptate hyphae, insulated conidiophores, branched, and it has smooth walls. *Penicillium* can act as a biopesticide and biofertilizer because it produces mycotoxin compounds such as citrinin to inhibit the growth of pathogens. In addition, *Penicillium* is used as a decomposer due to its ability to produces cellulose and endoglucanases, which can degrade dead plants to fertilize the soil (Amaria et al. 2013). Citrinin is a compound that is toxic and harmful if it is consumed by humans or animals and can cause chronic diseases. These compounds can contaminate soil, water, plants, and agricultural products (Khosravi et al. 2012).

*Schizophrillum commune*

The FS11 strain was molecularly identified as *Schizophrillum commune* with a bootstrap value of 76% (Figure 6). This strain was anamorph form collected from cassava in Bogor, which formed monophyletic together with KF679517 *S. commune* and *S. commune* MUCL 29305, formed a sister clade with several *S. radiatum*. *Mortierella elongata* MAFF425591 was used as the out-group (Figure 6.A). Based on macroscopic morphological observations, this fungus covered the PDA media with a 6 cm diameter on the fifth day of incubation. The colonies formed fluffy mycelium, which was white and thin (Figure 6.B). Microscopically, *S. commune* has aseptate hyphae, and formed no reproductive structure (Figure 6.C).
**Figure 6.** A. Phylogenetic tree of *Schizophyllum commune* based on ML analysis with 1000 bootstrap replicates, Mortierella elongate MAFF425591 was used as out-group. Morphological identification of endophytic fungi: B. Colony of *Schizophyllum commune* FS14; C. Septate hyphae of *Schizophyllum commune* FS14.

*S. commune* can also be found as an endophyte in other host plants, such as *Panax ginseng* (Zhai et al. 2017) and *Piper hispidum* (Orlandelli et al. 2012). This fungus has previously been found in Gabon ebony fruit and belongs to an endophytic fungus that can grow in tropical environments (Douanla-MeliLanger 2012). *S. commune* is also a saprophytic fungus that belongs to the Basidiomycota, and it is often found in woody plants in the form of teleomorphs, especially in the pith of plants. *S. commune* can break down cellulose and hemicellulose, but cannot break down lignin, due to cellulose enzymes dominate its extracellular activity. These fungal spores are classified as pathogens when inhaled by humans which can attack the human respiratory system and causing various diseases, such as sinusitis, allergies, and respiratory tract infections. In the form of teleomorph, this fungus is edible and has a compound namely *Schizophyllan*, acts as an anti-tumor, anti-cancer, and can increase endurance. This compound is a homopolysaccharide and water-soluble (Salahuddin 2008).

In conclusion, a total of fourteen endophytic fungal strains in cassava tubers have been isolated from 8 areas, and identified as six Fusarium strains, namely *Fusarium solani* (1 strain), *F. lichenicola* (2 strains), *F. oxysporum* (1 strain), and *F. falciforme* (2 strains). Four Aspergillus strains, namely *Aspergillus* sp. (3 strains) and *A. fumigatus* (1 strain). Followed by four other species such as *Lasiodiplodia* sp, *Nectria pseudotrichia*, *Penicillium citrinum*, and *Schizophyllum commune*. Molecular identification is needed to confirm its proper morphology. The use of other genes, such as elongation factor α and β-tubulin, is required to separate strains into separate clades. Further research is needed to determine the positive and negative impacts of the fungus found in cassava tubers.

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**REFERENCES**

Abdollahzadeh J, Javadi A, Goltapeh EM, Zare R, Phillips A. 2010. Phylogeny and morphology of four new species of *Lasiodiplodia* from Iran. Persoonia: Mol Phylogeny Evol Fungi 25: 1-10.

Akmalasari I, Purwati ES, Dewi RS. 2013. Isolasi dan identifikasi jamur endofit tanaman manggis (*Garcinia mangostana* L.). BIOSFERA: A Sci J 30: 82-89.

Amaria W, Tauifq E, Harni R. 2013. Seleksi dan identifikasi jamur antagonis sebagai agens hayati jamur akar putih *Rigidoporus microporus* pada tanaman karet. J Ind Beverage Crops 4: 55-64.

[Indonesian]
