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Molecular Characteristic of *Fusarium oxysporum* from Different Altitudes in East Java, IndonesiaHenik Sukorini^{1,*}, Erfan Dani Septia¹, Lili Zalizar² and Netnapis Khewkhom³¹Department of Agrotechnology, Faculty of Agriculture and Animal Science, University of Muhammadiyah Malang, Jl. Raya Tlogomas no 246 Malang 65145, East Java, Indonesia; ²Department of Animal Science, University of Muhammadiyah Malang; ³Department of Plant Pathology, Faculty of Agriculture, Kasetsart University, 50 Thanon Ngamwongwan, Lat Yao, Chatuchak, Bangkok 10900, Thailand

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Abstract

6 Tomato (*Solanum lycopersicum* L.) is one of the most economically important vegetable crops in Indonesia. Tomato diseases caused by fungi are transmitted by seed or transplants. *Fusarium* wilt disease is a cosmopolitan species caused by *Fusarium oxysporum* Schlecht. Emend. Snyder & Hansen. Among this special attention of disease caused by *F. oxysporum* has been given to stem and root rotting. Six selected *Fusarium* samples from previous research were prepared using a single spore method and cultured in the PDB medium. The Research carried out in the Agrotechnology Laboratory of the University of Muhammadiyah Malang. DNA extraction and PCR used ITS 1 and ITS 4, electrophoresis, and data analysis was achieved at the Genetic and Molecular Laboratory of the Biology Department of the Faculty of Science and Technology, Maulana Malik Ibrahim Malang Islamic State University. Isolate code 3313439 originating from Karangploso soil (515 m a.s.l.) and code 3313428 derived from the soil of Blitar (156 m a.s.l.) showed species similarity to *F. oxysporum* f. sp. *lycopersici* strain CBS249.52. Then for sample 3313426, the roots of Pujon have similarities with the strain of *F. oxysporum* S58. Besides, samples of 3313422 Blitar roots, 3313424 Karangploso roots, and 3313432 Pujon soils (956 m a.s.l.) showed proximity to species *F. oxysporum* f.sp. *pisi* HG423346. The samples were in one clade with the nucleotide base sequences of two other *F. oxysporum* species recorded in the NCBI Genbank database. Differences in species will likely affect the pathogenicity, growth rate, spore production, and disease control management.

Keywords: Fungi, *Fusarium* wilt, Molecular Identification, Plant Pathology, Tomato

1. Introduction

25 Globally, one of the most economically important crops is the Tomato (*Solanum lycopersicum* L.) (Aydi-Ben-Abdallah et al., 2020). Several economically essential tomato diseases caused by fungi are transmitted by seed or transplants. *Fusarium oxysporum* f. sp. *lycopersici* Schlecht. Snyder & Hansen (FOL) is the causal agent of fusarium wilt disease on tomatoes. It is a cosmopolitan species that can be found in all types of soil. Ignjatov et al. (2012) reported healthy plants could become infected by *F. oxysporum* if the soil in which they are growing is infested with the pathogen. FOL spread through short distances, mainly through irrigation water and contaminated farm equipment, and it can spread long distances through infected transplants, soils, etc. (Agrios, 2005). Special attention to disease has been given to the rotting of stems and roots caused by *Fusarium* sp. Based on the symptoms of the disease caused by *Fusarium* sp. indistinguishable. Control of this disease is also still problematic. The use of chemicals such as methyl bromide is quite effective for disease management but impacts humans and the environment. The use of resistant varieties has also been carried out. This method is environmentally friendly but requires a lot of money. The ease with which pathogens form new strains and break the resistance of varieties

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causes this disease to be difficult to control (Xie et al., 2015).

16 Biju et al. (2017) reported three known FOL races (Races 1, 2 and 3) pathogens of tomato cultivars are distinguishable by their principle resistance genes. Races 1 and 2 are grown through the tomato-growing regions globally. Race 3 has been reported in countries such as California, Australia, Southwestern Georgia, and Mexico. Most commercial tomato varieties grown through the world are resistant to race 1 and 2, and a few are resistant to race 3. Certainly, once a region becomes contaminated with FOL, the fungus usually remains indefinitely (Animashaun et al., 2017; Prihatna et al., 2018). Pathogenic isolates from three different heights, namely low, medium, high altitude, have different colony colors, sporulation power, and growth rate. The ability to survive at high temperatures and resistance to Mancozeb 64 % + Metalaxyl 8 % and Benomil 50 % fungicide also varies even though the growth inhibition value is below 60 % (Henik et al., 2021). This, of course, will affect the success of controlling this pathogen. Therefore, molecular identification is required.

Identification can be made in two ways consisting of morphological and molecular character identification. Molecular character identification is based on the similarity of DNA (Alsohaili and Bani-Hasan, 2018)

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The genus *Fusarium* has often served as a testing ground for new speciation concepts in fungi (Hsuan *et al.* 2011). The use of molecular approaches to differentiate species has been tried with many strains usually considered problematic, i.e. not fitting within a given species but not distinguishable from it. Studies of the grouping patterns resulting from studies with amplified fragment length polymorphisms (AFLPs) and phylogenetic lineages based on multiple-gene genealogies provide new means of evaluating relatedness.

This research is a follow-up study from previous studies, namely the control of *Fusarium* sp wilt in tomato plants and *Alternaria* sp in potato plants. Henik *et al.* (2021) reported earlier studies, *F. oxysporum* from tomatoes originating from different altitude and parts of plants with spore shape and size characters. The same, but has other growth characters, so too has different pathogenicity and virulence. Therefore, it is necessary to reveal more about the differences in isolates from regions with different altitudes with molecular character identification.

2. Method

2.1. Single spore isolation technique for *F. oxysporum* isolates

Soil and tomato roots samples were taken from tomato growing areas in Blitar (156 m a.s.l.), Karangploso (515 m a.s.l.), and Pujon (956 m a.s.l.). After preparation, isolates were grown on PDA media (Merck). Then the isolates were made suspended and transferred to the water agar media. A suspension of conidia either from a sporodochium or aerial mycelium was prepared in 5 mL sterile water in a sterile vial. The conidial suspension was scraped by an L-shape inoculating needle several times and streaked across the water agar plate. The plate was incubated for 12 h to 24 h at 25 °C; after that, it was examined under a dissecting microscope (Olympus CX43). A lot of germinating conidia appeared on the inoculation, and by following the streaked lines using the low power of the microscope, single germinating conidia could be observed. Finally, identification of *F. oxysporum* cultures was accomplished (Niemeyer and De Andrade, 2016).

Soil and tomato root samples were taken from tomato growing areas in Blitar (156 masl), Karangploso (515 m a.s.l.), and Pujon (956 m.a.s.l.). Isolation was carried out using the method (El-shafey, 2020) with slight modifications in the medium. Isolation was carried out by growing the sample on PDA (Merck) media until pure isolates were obtained. This will be used for further investigations. 0.1uL of *Fusarium* isolates were then grown in water agar media for 24 h, then a single spore from *Fusarium* was cut and grown on new PDA media in Petri dish for 7 d at 25 °C. The growing isolates were then used for DNA testing and stored in the refrigerator for collection and other purposes. Isolates aged 7 d were then harvested for DNA testing.

2.2. Isolate preparation

Isolates grown on PDA media (Merck) were then developed into PDB media (potato dextrose broth - Merck) and incubated in water bath shakers for 7 d to 9 d until the mycelium grew. The mycelium can be harvested, put into

an Eppendorf tube, then added 500 mL Mili Q water and be ready for extraction with centrifuged at 10 000 rpm (1 rpm=1/60 Hz) for 10 min (Hussain *et al.*, 2012).

2.3. Fungi DNA extraction

PCR Preparation Based on isolate preparation, the supernatant was taken with a micropipette and crushed with pastel until it became colloidal. Colloids were added with Reagent 1 as much as 300 mL, then homogenized using tips from the micropipette. Added 3 µL of RNase, homogenized using tips, and incubated for 30 min at 37 °C in a water bath, then added Reagent 2 for 200 µL, strong shaking for ± 10 min set for 10 min at room temperature, put in the freezer for 20 min. After freezing, added 250 µ of chloroform and 250 µ of phenol. It was homogenized for ± 4 min, centrifuged (DLAB High-Speed Refrigerated Micro-Centrifuge D3024R) at 10 000 rpm for 10 min at the temperature of 14 °C. The supernatant was taken, plus isopropanol, in half of the sample volume (250 µL), then reversed slowly to homogeneous. Centrifuged at 10 000 rpm for 10 min, then the supernatant was removed. Added 50 µL ethanol 99 %, then centrifuged with a speed of 10 000 rpm for 10 min; the supernatant was disposed of by pouring. It was dried up by turning the microtube over the tissue by opening the lid for ± 30 min. Added Nuclease free water as much as 100 µL(lots) and 50µL (a little).

2.4. Polymerase Chain Reaction

The DNA obtained was then multiplied by a polymerase chain reaction. The primers used for this PCR process were ITS 1 and ITS primers 4. The primer ITS1 (5'-TCT GTA GGT GAA CCT GCG G3') and ITS4 (5'-TCC TCC GCT TAT TGA TAT GC-3'). The PCR solution mixture was 1 uL to 2 uL primer, 5 uL DNA samples, and 6 uL PCR mix (Half reaction) (Singha *et al.*, 2017). The PCR condition was run in 5 min 94 °C for pre-denaturation, then followed by 35 cycles where each cycle consisted of 30s, 94 °C for denaturation, 30s, 50°C for annealing, and 1 min temperature 72 °C centigrade for extension; then ended with 1 min cycle with a temperature of 72 °C. The particular annealing temperature was also tested at 48 °C as the annealing temperature recommended by primer pairs.

2.5. Agarose Preparation

mixed 1 % agarose (0.3 g) with 30 mL TBE 1× (from Tris base and boric acid 10×) in an Erlenmeyer tube. Agarose was dissolved by heating it to the microwave for ± 2 min until the solution was dissolved entirely then add ETBR 2 µL and pour it into a gel mold into a gel mold that has been fitted with a comb The gel was allowed to hard for ± 30 min.

2.6. Electrophoresis preparation

After the comb was removed from the mold, then the gel was transferred into the electrophoresis tank. The 3 µL DNA solution was mixed with loading dye 1µL and 2µL on parafilm paper, then the mixture and marker were put into the agarose gel well. Adjust the position of the agarose gel in electrophoresis. After combining DNA loading, dye and water were inserted into the well. The electrophoresis device has adjusted the electrophoresis with 100 V and 25 min later running. Electrophoresis was complete. The agarose gel lifted in the agarose gel was brought into the doc gel illuminated with a UV-

Transilluminator and gel photographed with the gel documentation to visualize the results.

2.7. Measurement of concentration and purity of extracted DNA

The concentration and purity of DNA extracted using NanoDrop with two different wavelengths, wavelengths 260 and 280. The purity of DNA can be determined by dividing the results of the NanoDrop measurements of both wavelengths (Srinivas *et al.*, 2019), while the DNA concentration was obtained directly from the NanoDrop.

2.8. Analyzing the results of reading Gel Doc

Analyzing the results of reading Gel Doc was done qualitatively by observing and determining the band's size that comes out with PCR-electrophoresis results compared to the marker. This information was then compared with the literature to ascertain whether the target DNA location obtained was precisely the location of ITS (Adame-García, 2015).

2.9. Sequencing the results of amplification of DNA samples

The results of the DNA sample amplification obtained were then sent to PT. Indonesian Science Genetics for sequencing. The 1st Base company carried out the sequencing process, Axil Scientific Pte Ltd., Based in Singapore. The material used for DNA sequencing is BigDye® Terminator v3.1 Cycle Kit Chemistry.

2.10. Data Analysis of Sequencing Results

Sequencing results based on ITS 1 or ITS 4 primers for each isolate were aligned and then edited using Mega v.10 software by referring to the chromatogram and being converted to fasta. The data in the form of fasta were further analyzed by looking for similarities using the basic local alignment search tool (BLAST) program from the GeneBank Gen database owned by the National Center of Biotechnology Information / NCBI.

BLAST was used to determine which species has the closest homology. Ambiguous areas in parallel sequences were omitted from the analysis. The gap was considered missing data (Pinaría, *et al.*, 2015). The phylogeny tree's evolutionary history was inferred based on the Neighbor-Joining method (Cañizares *et al.*, 2015). The evolutionary distance was calculated using the Maximum Composite Likelihood method (Chala, *et al.*, 2019). And it was in units of base substitution per site. Evolution analysis was carried out with the MEGA X version application (Nirmaladevi, *et al.* 2016). Preparation of the genetic distance matrix *F. oxysporum* Karangploso sample with several species genes recorded in GenBank was also carried out using the MEGA X version application.

3. Result and Discussion

The isolates used in this research had the highest sporulation and growth rates from previous studies, namely those from roots and soil from three high, medium, and low land altitudes. (Henik *et al.*, 2021). Morphologically, *Fusarium* species are identified by several morphological characteristics. One of the notable

features is the development of various shapes and sizes of macro and microconidia. Other structures that they form are called chlamydoconidia spores (El Kichaoui *et al.*, 2017; Raghu *et al.* 2016), and also identified based on the growth rate and their pigmentations on agar media (Leslie *et al.*, 2008).

Moreover, morphological identification can be quite difficult among the *Fusarium* species (Lievens *et al.*, 2008). The sequence information using ITS regions has been immensely used in the phylogeny and taxonomy of *Fusarium* species (Menezes *et al.*, 2010) as ITS regions have successfully identified them (Chen *et al.*, 2004). ITS is differentiated into ITS1 and ITS2 (genes 18S to 5.8S and 5.8S to 28S, respectively) (Hillis and Dixon, 1991). There are more than 172 000 fungal ITS sequences present in Genbank. The result showed the thickness of DNA bands that varied from those showing thin bands to thick bands (Figure 1). The thickness of the tape is related to the concentration of DNA isolation results. The thin ribbon shows that the DNA concentration produced from the extraction process is low, while the thick band indicates the low concentration of DNA from extraction (Menu, *et al.*, 2018).

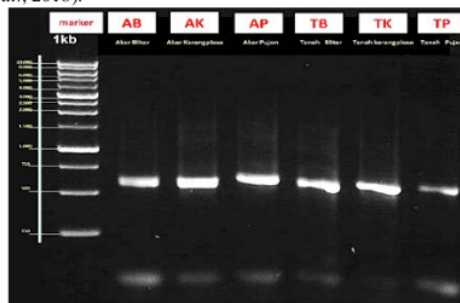


Figure 1. The amplification of *Fusarium* DNA samples using primary pairs ITS 1 and ITS 4 with an annealing temperature of 50 °C at three different altitudes (156 m a.s.l, 515 m a.s.l, and 956 m a.s.l). AB= Blitar root, AK= Karangploso root, AP= Pujon root, TB= Blitar soil, TK= Karangploso soil, TP= Pujon soil.

Electrophoresis results show a band with a smear (Figure 1). Smear is the remainder of the solutions that are still carried during the isolation process and can also be degraded DNA during the isolation process (González-Mendoza, *et al.*, 2015). The process of DNA degradation at the stage of isolation can be caused by mixing the solution using a vortex, which aims to help cell lysis so that some DNA comes out and is fragmented and causes smears when electrophoresed (Campbell *et al.*, 2010).

The results of *Fusarium* sample DNA amplification with a temperature of 48 °C annealing in the Polymerase Chain Reaction (PCR) process showed no amplification in each sample DNA tested. The thin bands below show a size that is much smaller than 250 bp. Allegedly, these bands are a visual form of primers' formation during polymerized Chain Reaction (PCR). Primers-dimers are not the result of the desired target DNA amplification. It can be seen from the size of the resulting tape that it is between 500 bp and 750 bp, but there are still smears.

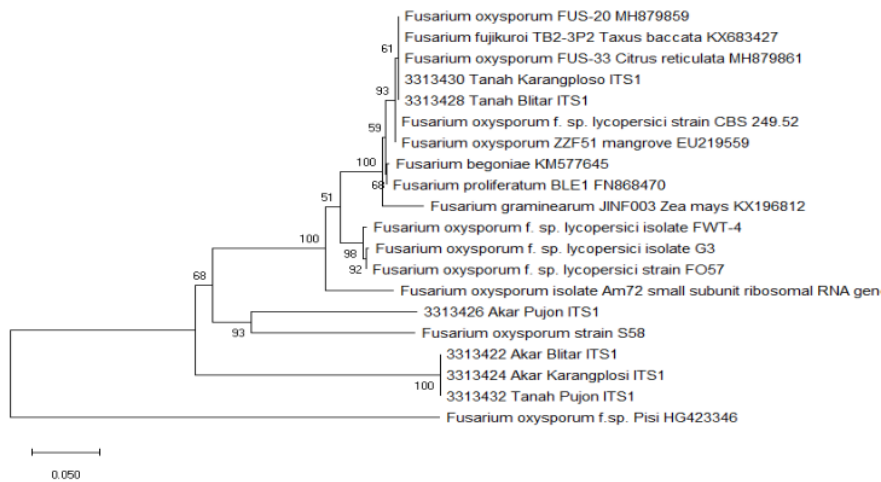


Figure 2. Phylogeny tree of *Fusarium oxysporum* at three different altitudes.

Based on the results of the evolution of tree evolution (phylogeny tree) and the similarity of nucleotide base sequences originating from the Internal Spacer (ITS) region,

F. oxysporum from six samples consisting of soil samples and roots of tomato plants at three different altitudes (156 m a.s.l., 515 m a.s.l. and 956 m a.s.l.), it was found that code 3313439 derived from Karangploso soil and code 3313428 originating from Blitar soil showed similarity of species with *Fusarium oxysporum* f.sp. *lycopersici* strain 249.52. A sample of 3313426 pujan roots has similarities with *Fusarium oxysporum* strain S58. Besides, samples of 3313422 Blitar roots, 3313424 roots of Karangploso, and 3313432 Pujan soil showed closeness to the species *Fusarium oxysporum* f.sp. *pisi* HG423346. That sample is in one clade with a nucleotide base sequence; other *Fusarium oxysporum* species have been recorded in the NCBI Genbank database. This third sample forms a monophyletic group with each other. In other parts of the branching, all groups incorporated in the *F. oxysporum* taxon species are polyphyletic with taxon members of Genus *Fusarium* sp. others. Knowledge of *Fusarium* wilt symptoms as a result of in depth is required. Tomato plants at the same altitudes have different closeness to *Fusarium* species or forma species in one area. It might affect the resistance of heat, fungicides, and different pathogenicity (Henik, *et al.*, 2021) and show variations in pathogenicity, response to management systems, environment, and host differences (Hami *et al.*, 2021).

Campbell *et al.*, (2014) reported that a taxon is monophyletic if its single ancestor produces a whole derivative species in the taxon and not a species in another taxon, polyphyletic. If its members come from several ancestors that are not the same for all members, and paraphyletic if the taxon does not include species with grandmothers, the same ancestor in a member of a species towards another species. Branching like this can occur when two types of molecular characters specific to different homologous DNA species over time change due to various conditions (Nath, *et al.*, 2017). The situation is like a point mutation that removes a nucleotide from the specific DNA sequences and inserts three adjacent nucleotides (Edel-Hermann and Lecomte, 2019). As a result of this situation, DNA sequences that are initially very similar have different lengths and sequences.

At the end of the branching of these three samples, a figure of 100 % shows the high bootstrap value of this branching group. Bootstrap value is one of the measures introduced by (Fredricks, Smith, and Meier, 2005), which offers the stability value of a topology tree. The higher the bootstrap value in a branching, the more fulfilling the topology's validity level requirements. In other words, a high boost value (close to 100 %) does not mean showing the accuracy of a topology tree but rather indicates that each character information of each individual in the branching group "agrees" that a branching member is a group.

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