

# Disease on Dragon Fruits (*Hylocereus* sp.)

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## First Report on *Fusarium solani*, a Pathogenic Fungus Causing Stem Rot Disease on Dragon Fruits (*Hylocereus* sp.) in Bali

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### Abstract

In recent years, dragon fruit crop (*Hylocereus* spp.) has become increasingly important in Bali Indonesia due to its high nutrient content and healing properties. However, the dragon fruit was reported to be seriously infected with several complex diseases caused by fungi and causing serious losses to the farmers. The study on morphological and molecular characterization the fungal pathogen was conducted to confirm the species of the fungi. Koch Postulate was applied to confirm the causal agent of the disease. There were two isolates of fungi isolated from the stems of diseased plants, namely isolate w1 (from stem of *H. undatus*) and isolate w2 (from stem of *H. polyrhizus*). Based on macroscopic and microscopic characteristics, and analysis of 18S rDNA, both of them were identified as *Fusarium solani*. This is the first report on the *F. solani* the cause of stem rot disease on dragon fruits in Bali.

**Keywords** : stem rot disease, dragon fruit, *Fusarium solani*

### 1. Introduction

Dragon Fruits (*Hylocereus* spp.), which are also known as pitaya, are the fruits of cactus species, especially of the genus *Hylocereus*. There are three species which have high commercial valuable fruits are the species of *Hylocereus undatus*, (red rind, white flesh), *Hylocereus polyrhizus* (red rind, red flesh), and *Hylocereus costaricensis* (red rind, super red flesh). *Hylocereus* spp. is grown commercially in Vietnam, Spain, Malaysia, Japan, Mexico and other tropical and subtropical areas because of its high nutrient content and healing properties. In recent years, this fruit has become increasingly important in Bali Indonesia. The dragon fruit is rich in vitamin, it helps the digestive process due to its fiber, prevent colon cancer and diabetes, neutralize toxic substances such as heavy metals, and helps to reduce cholesterol levels and high blood pressure (He *et al.*, 2012; Zainoldin and Baba, 2009). Recently, dragon fruit has been reported to be seriously infected with several complex diseases caused by fungi and causing serious losses to farmers. Several dragon fruit plants grown in Sobangan Village, Bali Indonesia showed severe symptom of stem rot disease. The disease caused significant yield losses. Isolation of the fungi associated with the diseased-plants showed that *Fusarium* sp. was the most frequent found on the stems showing brown rot symptom.

Various diseases caused by fungi have been reported on dragon fruit in tropical and subtropical countries, such as fruit rot (*Bipolaris cactivora*) (Tarnowski *et al.*, 2010; He *et al.* 2012), stem rot (*Fusarium semitectum*, *Fusarium oxysporium*, *Fusarium moniliforme*) (Hawa *et al.*, 2010), anthracnose (*Colletotrichum gloeosporioides*) (Masayahit *et al.*, 2009), brown spot (*Botryodiplodia* sp.), basal rot (*Phytophthora* sp.) (Lin *et al.*, 2006), wilt (*Fusarium oxysporum*), stem blight (*Diplodia* sp., *Ascochyta* sp., and *Phytophthora* sp.), black spot (*Alternaria* sp.), speck blight (*Nectriella* sp.), (Wang *et al.*, 2007), and stem lesion (*Septogloeum* sp.) (Zheng *et al.*, 2009).

In order to control the disease, it is necessary to identify the causal agent of the disease. The fungal pathogen can be identified based on cultural and morphological characters. However it could be highly variable depending on the media and cultural conditions that could be the problems in fungal identification. In recent years, the increasing use of molecular methods in fungal identification has emerged as a possible answer to the problems associated with the existing phenotypic identification systems (Mishra *et al.*, 2003). One of the molecular approaches to fungal identification is based on Polymerase Chain Reaction (PCR).

The main area for the development of fungal identification is ribosomal genes, present in all organisms and at high copy numbers aiding detection and the sensitivity of the PCR reaction. The fungal nuclear ribosomal DNA (rDNA) consist of three genes, the large subunit gene (25S), the small subunit gene (18S), and the 5.8S gene, separated by internal transcribed spacer (ITS) regions, in a unit repeated many times. The ITS region is an area of particular importance to fungal identification. It has areas of high conservation and areas of high variability and is an ideal starter for the development of specific PCR primers for identification of fungal species (Atkins and Clark, 2004). Sequences of the ITS regions ITS1 and ITS2 have been used widely in molecular phylogenetic studies because of their relatively high variability and facility of amplification. For phylogenetic

applications, most researchers use sequence alignments that are based on nucleotide similarity (Tippary and Les, 2008).

Suga *et al.* (2000) has investigated phylogenetic relationships of *Fusarium solani* using sequences from the rDNA-ITS region. Mishra *et al.* (2003) has developed a fluorescent-based polymerase chain reaction in ITS region to identify five toxigenic and pathogenic *Fusarium* species. Abd-Elsalam *et al.* (2003) have developed two taxon-selective primers for quick identification of the *Fusarium* genus. These primers, ITS-Fu-f and ITS-Fu-r were designed by comparing the aligned sequences of internal transcribed spacer regions (ITS) of a range of *Fusarium* species. Zhang *et al.* (2006) and O'Donnell *et al.* (2008) have studied phylogeny of *Fusarium solani* Species Complex (FSSC) that cause infection in both humans and plants based on three genes of the ribosomal DNA. The ITS region including 5.8S rDNA sequence of 58 isolates *Candida parapsilosis* in Brazil and Japan was analyzed by Iida *et al.* (2005).

This paper reports on identification of fungal pathogen causing stem rot disease from dragon fruits planted in Bali based on morphological and molecular methods using sequences from rDNA-ITS region.

## 2. Materials and Methods

### 2.1. Fungal Pathogen Isolation and Virulence Test

Fungi were isolated from the diseased stem of *H. undatus* and *H. polyrhizus* from dragon fruit's orchard in Sobangan village, Mengwi Bali. Surface sterilization was carried out by cleaning the symptom margins with 70% ethanol and cut into small blocks (ca 1.5 x 1.5 x 1.5 cm), soaked in 1% sodium hypochlorite (NaOCl) for 3 min, and rinsed in several changes of sterile distilled water (each 1 min). All sterilized samples were placed onto Potato Dextrose Agar (PDA) and incubated at 25 ± 2°C for 7 days. Mycelium growing after 3 days of incubation was transferred to a new PDA to obtain pure fungal cultures. The pure culture was inoculated on healthy dragon fruit stems to confirm the similarity of symptoms in the field. For this purpose the Koch's postulates test was carried out. Dragon fruit stems (30 days in age) were injured with a sterile needle and inoculated by spraying with a suspension of spores of 1.5x10<sup>6</sup> spores/mL. Three plants were inoculated with a fungal isolate. For control, dragon fruit stems were pierced with a sterile needle and sprayed with sterile water. Symptoms were observed for a week after inoculation. The symptoms were compared with the symptoms of the disease in the field. After that, the isolation of pathogenic fungi on infected stem was performed again. Pure cultures of pathogenic fungi were inoculated again on dragon fruit plants and the same inoculation procedure was done to obtain the similarity of symptoms. Fungal isolates obtained can be regarded as the cause of the disease and were used for further identification.

### 2.2. Morphological Characterization

Characterization of the main fungal pathogens was carried out macroscopically, by observing the fungal colony color, colony reverse color, no lines or concentric radii, issued exudates or not, media pigmentation, colony surface, and how the growth of fungi (fast or slow). Microscopic identification was also carried out by observing the shape of hyphae or spores under a microscope, and then the results were confirmed using fungal identification book (Pit and Hocking, 1997).

### 2.3. DNA Extraction

*Fusarium* sp. isolates w1 and w2 were grown on potato dextrose agar (PDA) medium for 3 days at room temperature. The mycelium grown was harvested and grown to a fine powder in a sterile mortar with liquid nitrogen. DNA was extracted by using Phythopure™ DNA Extraction Kit (GE Healthcare, UK) according to manufacturer's instructions.

### 2.4. Molecular identification and phylogenetic relationships of fungal pathogen

Identification of fungal isolates was performed based on molecular genetic analysis using the internal transcribed spacer region (ITS). PCR amplification used ITS 5 F: 5' - GGAAGTAAAAGTCGTAACAAGG - 3' and ITS 4 R: 5' - TCCTCCGCTTATTGATATGC - 3' (White *et al.* 1990). Amplification was performed on a volume of 25

μL with the reaction mixture: 10 μL nuclease free water, 12.5 μL Go taq green master mix™, ITS5 and ITS4 each 0.5 μL, 0.5 μL DMSO, and 1 μL of DNA template. PCR amplification for regional ITS consists of: pre denaturation 95 °C for 90 seconds, followed by 95 °C for 30 seconds with 35 cycles, annealing 55 °C in 30 seconds, extension 72 °C in 90 seconds, and final extension 72 °C for 5 minutes. The product was purified and then sequenced. The nitrogen base sequence was analyzed using automated DNA sequencer (ABI PRISM 3130 Genetic Analyzer) (Applied Biosystems).

Sequencing raw data were trimmed and assembled using ChromasPro program version 1.5. The assembled data were BLASTED with genomic data that has been registered in NCBI / National Center for Biotechnology Information (<http://www.ncbi.nlm.nih.gov/BLAST/>). Some data sequence which is a result of blast nearest species and is a type strains of each species were taken from the data in the NCBI gene bank. Then the data were analyzed again with the sequence alignment using MEGA version 5.0 program (Tamura *et al.* 2011) and bootstrap used is 1000 replications (Felsenstein, 1985).

### 3. Results and Discussion

#### 3.1. Fungal Isolates

There are 10 fungal isolates were obtained from isolation of fungi associated with the stem disease of dragon fruits grown in Sobangan Village, Bali Indonesia. Based on the Koch's postulates test, there were two isolates of *Fusarium* sp. namely isolate w1 (from *H. undatus*) and isolate w2 (from *H. polyrhizus*) caused the stem rot disease with the symptom similar to the symptom occurred in the field. The symptom appeared on stem as dark brown stem rot as shown in Figure 1.



**Figure 1.** Symptoms of the stem rot diseases on *Hylocereus* sp. under field conditions.

#### 3.2. Macroscopic Characteristic

*Fusarium* sp. isolate w1 grown on PDA produced abundant mycelium (sometimes in aerial, depends on cultural condition), white colony appearance, cream colony reverse, yellow to brown pigmentations, and fast growing (3.75-4.45 cm in 3 days) whereas *Fusarium* sp. isolate w2 produced abundant-powdery mycelium (sometimes in aerial), white colony appearance, yellow colony reverse, yellow pigmentations, and fast growing (4.35-5.25 cm in 3 days).

In a similar study, Chandran and Kumar (2012) studied for cultural, morphological variability in 13 isolates of *Fusarium solani* (Mart.) Sacc., incitant of dry root-rot of citrus, they reported that *F. solani* isolates were characterized as fast growing, moderately growing, and slow growing. The five isolates produced pale pink to dusky red color pigmentation in PDA medium and potato dextrose broth culture. The remaining all isolates produced pale yellow to dark yellow pigmentation. Most of the isolates produced profuse sporulation, whereas others produced moderate sporulation. Madhukeshwara (2000) has studied cultural variability among six isolates of *F. udum* causing wilt of pigeon pea. All the isolates varied with each other in terms of growth, mycelium, pigmentation, and sporulation. Most of the isolates produced cottony white raised mycelium, pale yellow to dusky red color pigmentation and moderate to profuse sporulation on PDA medium.

#### 3.3. Microscopic Characteristics

Microscopic characters such as size, shape, septation, and color of conidia were studied using PDA medium. *Fusarium* sp. isolate w1 had longer macroconidia (1-4 septates; 18.4-31.7  $\mu\text{m}$  in length), curved in shaped with hyaline in color, whereas *Fusarium* sp. isolate w2 had shorter macroconidia (1-3 septates; 15.3-24.8  $\mu\text{m}$  in length), curved in shaped with hyaline in color. *Fusarium* sp. isolate w1 produced less microconidia while *Fusarium* sp. isolate w2 produces abundant microconidia (1 septate; 5.7- 8.5  $\mu\text{m}$  in length), round to oval in shaped. Furthermore both the fungi had the septate hyphae and chlamydospores in pairs with hyaline in color and located in the middle of hyphae as presented in Figure 2.

Pitt and Hocking (1997) reported that the main characters used to distinguish species of *Fusarium* are the size and shape of the macroconidia; the presence or absence of microconidia; and the presence of chlamydospores. Kawuri *et al.* (2012) reported that *Fusarium oxysporum* had macroconidia curved in shaped with four septates and foot cell, 31 $\mu\text{m}$  length and smooth surface, whereas microconidia has rough surface at 4.6  $\mu\text{m}$  length. Chandran and Kumar (2012) reported that the number of septa in macro conidia and micro conidia of *Fusarium solani* (Mart.) Sacc. are 3-5 and 0-1 respectively and the color is hyaline. The shape of macro conidia is sickle shaped with blunt ends and micro conidia is round to oval shaped. The chlamydospores located in middle of hyphae (intercalary), on tip of the hyphae (terminal) and some chlamydospores were seen in middle of macro conidia.









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