First Report on Infection of *Eucalyptus pellita* Seeds by *Ralstonia solanacearum* †

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Abstract: Bacterial wilt is one of major threats to eucalyptus plantations which may cause significant losses. Until now, study about bacterial wilt on *Eucalyptus pellita* in Indonesia has been very limited, especially about the presence of the pathogen on or in the seeds. This study aims to provide evidence of the existence of the *R. solanacearum* bacterium on or in *E. pellita* seeds. Detection of seed-borne bacteria is determined by several approaches such as (i) direct detection using universal and selective media in the laboratory, (ii) the nursery test, and (iii) species-specific molecular detection. The results of our study indicate that *R. solanacearum* can be detected from eucalyptus seeds using universal and selective media in the laboratory, nursery test, and molecular-based detection using the Enrichment PCR method. The bacterial inoculum is also proven to exist both on the surface of and inside the eucalyptus seeds. This is the first report that *R. solanacearum* is a seed-borne pathogen in *E. pellita* seeds. Previous studies in different agricultural systems show that the effective method used to control the pathogen is through seed treatments using biological, physical, and chemical approaches.

Keywords: bacterial wilt disease; detection; pathogen; plantation; seedborne

1. Introduction

Forest plantation in Indonesia has been rapidly increasing in the last few decades and reached 11 million hectares last year [1]. Ecologically, forest plantations are able to reduce pressure on natural forests, quickly fix carbon and stimulate the restoration of natural vegetation. However, monoculture plantations are challenged to sustainably manage pest and disease risks [2]. Traditionally, the use of healthy (pathogen-free) seeds is one approach that can be taken to reduce the risk of disease. So far, most of the reports on seed-borne pathogens on forest trees have dealt with fungi only. Information on seed transmission of other pathogens, especially bacteria, was very limited [3]. Take for example the bacterial wilt pathogen of eucalyptus, *Ralstonia solanacearum*. The pathogen has previously been reported to be a seed-borne pathogen in many agricultural plants including eggplant, tomato, chili, potato, and ginger [4–6]. However, it was yet to be proven to be seed-transmitted in eucalyptus. Bacterial wilt is one of major threats in eucalyptus plantations which can cause significant losses [7]. Until now, study about bacterial wilt on *E. pellita* in Indonesia has been very limited, especially regarding the presence of the pathogen on or in the seeds. This study aims to provide evidence of the existence of the
**2. Materials and Methods**

Seed testing was performed in the laboratory using universal and selective media. *E. pellita* seeds (1 g) from different seed lot numbers were crushed using pestle and mortar and the sap liquid (0.1 mL) was taken for plating into (2,3,5-triphenyl tetrazolium chloride (TZC) for universal medium and modified TZC for the selective medium [8]. The agar plate was incubated at 28 °C for 48–72 h. A single colony of irregularly shaped, fluid and round pink bacterium is positive for the colony of virulent *Ralstonia*.

A nursery test was performed through germinating seeds of four different *E. pellita* seed lot numbers. The seedlings were maintained in the nursery until they reached 60 days old. The seedings were symptomless. A total of 60 healthy-looking seedlings were collected for pathogen isolation using the procedure described elsewhere [9]. The stems were cut transversely and placed in a soaked container and sterilized with 5.25% NaOCl twice before being rinsed with sterile water. The stem pieces were then inserted into a test tube containing 10 mL of sterile distilled water and incubated in an incubator shaker for 30 min. The water stem immersion (0.1 mL) was taken and grown on TZC media and incubated at 28 °C for 72 h.

Molecular detection was carried out through Enrichment PCR (En-PCR) using universal *Ralstonia* primer. The seeds from seven seed lot numbers were divided into two groups, with or without surface sterilization. Surface sterilization was conducted using NaOCl (5.25%) before the seeds were soaked in sterilized water. The sample (0.5 g) was crushed using pestle and mortar and the sap liquid (1 mL) was taken and 3 mL of TZC Enrichment medium added and the mixture incubated for 24–48 h. Genomic DNA was extracted using the Bacteria Genomic DNA Kit (Geneaid) according to the manufacturer’s instructions. To confirm that all tested strains belong to the complex *R. solanacearum*, PCR amplification was performed using species-specific primers 759 (5′-GTCGCCGTCAACTTCC-3′) and 760 (5′-GTCGCCGTCAGCAATGCGGAATCG-3′) to amplify a 280-bp fragment [10]. Each reaction mixture (25 µL) contained 1× reaction buffer, 1 U Taq DNA polymerase (Green GoTaq), 25 mM MgCl2, 2.5 mM dNTP mix, 10 µM of each primer and 20 ng of DNA template. Amplification was carried out in a 2720 Thermal Cycler (Applied Biosystem). The cycling conditions used were: denaturation at 94 °C for 2 min followed by 30 cycles of denaturation at 94 °C for 30 s, primer binding at 55 °C for 30 s, elongation at 72 °C for 1 min, and a final elongation step at 72 °C for 5 min. Water was included in every PCR assay as a negative control.

**3. Results and Discussion**

Bacterial wilt infection in eucalyptus seedlings is usually without apparent symptoms or known as latent infection [8]. The nursery test and laboratory detection need to be done to quantify the infection rate in seedlings. Table 1 shows the presence of the *Ralstonia* bacterium in symptomless seedlings with the infection rate varying between 6 and 42% depending on the seed lot number. These findings indicate the potential of bacterial wilt disease to be transmitted through seeds. Planting seedlings with latent infection into the field may cause an outbreak of the disease [11]. Seed health testing is therefore necessary to obtain pathological quality assurance that the seeds are clean and not contaminated, to minimize pathogen dispersal through seed transmission [12].
Table 1. Infection rate (%) of *Ralstonia solanacearum* in symptomless *Eucalyptus pellita* seedlings.

| Seed Lot Number | Infection Rate (%) |
|-----------------|--------------------|
| 12073           | 42.4               |
| 12074           | 6.7–36.7           |
| 12075           | 8.6–19.7           |
| 12076           | 8.3                |

The seed samples without surface sterilization treatment produced positive bands, implying that they had the *Ralstonia* bacterium. A positive result was determined from the position of the band that matched the expected product length. The amplified 281-bp-DNA fragment of each sample was obtained. Two of the seven seed lot samples with surface-sterilization treatment were positive, an indication that the bacterial inoculum was present both on the surface of and inside the eucalyptus seeds (endosperm) (Table 2).

Table 2. Detection of *Ralstonia solanacearum* in *Eucalyptus pellita* seeds using Enrichment (En)-PCR.

| Seed Lot Number | Surface | Endosperm |
|-----------------|---------|-----------|
| EP15216AA5      | +       | +         |
| EP15215AA5      | -       | -         |
| EP15219AA5      | +       | -         |
| EP15214AA5      | +       | -         |
| EP15218AA5      | +       | +         |
| EP15217AA5      | -       | -         |
| EP15211AA5      | +       | -         |

+: *Ralstonia solanacearum* was positively detected using En-PCR.

PCR assay amplifies the DNA of target organisms, targeting the species-specific sequences in their genome. In the present study, an efficient DNA isolation protocol and PCR-based detection of the bacterial wilt pathogen in soil and infected plant materials were carried out using primers of 759F/760R. The primers were originally developed for amplification of different *R. solanacearum* isolates [10]. Our results confirmed the success of using these specific primers to locate *R. solanacearum*. Although the disease can be managed using a consortium of antagonistic bacteria [13], detection of *R. solanacearum* at low concentration in seeds and plant tissues, which may be undetected using conventional methods, is very critical. Studies elsewhere show that seed treatment is also effective in controlling the pathogen [14].

4. Conclusions

*R. solanacearum* can be detected from eucalyptus seeds using universal and selective media in the laboratory, nursery test, and molecular-based detection using the Enrichment-PCR method. The bacterial inoculum is present both on the surface of and inside the eucalyptus seeds. This is the first report that *R. solanacearum* is a seed-borne pathogen in *E. pellita* seeds.

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