

PHOTOSYNTHETIC RESPONSES OF *Eucalyptus nitens* AT INITIAL STAGES OF ROOT-ROT INFECTION

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PHOTOSYNTHETIC RESPONSES OF *Eucalyptus nitens* AT INITIAL STAGES OF ROOT-ROT INFECTION. Root-rots are known to be latent diseases that may be present in plants for an extended period without any noticeable expression of symptoms above ground. Photosynthetic responses of *Eucalyptus nitens* saplings artificially inoculated with the root-rot pathogen, *Armillaria luteobubalina* were examined to characterize the initial stages of root-rot infection. This paper studies three photosynthetic parameters, i.e. photosystem II yield (F_v/F_m), chlorophyll content and photosynthetic capacity (A_{max}) for two strains of *A. luteobubalina* over a seven-month period. Root systems were either wounded or left intact before inoculation. A significant difference was observed in the F_v/F_m ratio between the uninoculated control and inoculated saplings. Photosystem II yield was considered the most sensitive parameter for the early detection of root-rot disease. Chlorophyll content and A_{max} decreased for all trees, including controls, during the period of the experiment, and most likely reflected host responses to seasonal change rather than treatment effects. Fungal re-isolations from symptomatic roots of inoculated trees confirmed the presence of *A. luteobubalina*. Findings from this preliminary trial indicated that there were detectable physiological changes associated with early infection of root-rot. However, to detect more widespread physiological changes an experiment of longer duration is needed.

Keywords: *Eucalyptus nitens*, artificial inoculation, chlorophyll content, photosynthetic rate, photosystem II yield, root disease

RESPON FOTOSINTESIS *Eucalyptus nitens* PADA TAHAP AWAL INFEKSI PENYAKIT BUSUK AKAR. Penyakit busuk akar merupakan penyakit yang bersifat laten yang dapat menginfeksi tanaman dalam jangka waktu lama tanpa menimbulkan gejala yang dapat diamati. Oleh karena itu, untuk mengetahui karakter perubahan fisiologis sebelum timbulnya gejala, telah dilakukan percobaan mengenai respon fotosintesis tanaman pada tahap awal infeksi penyakit busuk akar dengan cara menginokulasi anakan pohon *Eucalyptus nitens* dengan patogen *Armillaria luteobubalina*. Inokulasi buatan dilakukan dengan menggunakan dua strain *A. luteobubalina* dan dua variasi perlakuan akar, yaitu : dilukai dan tidak dilukai. Respons fotosintesis diamati dengan cara mengukur tiga parameter fotosintesis, yaitu: efisiensi fotosistem II (F_v/F_m), kadar klorofil dan laju fotosintesis (A_{max}). Pengamatan dilakukan selama tujuh bulan. Perbedaan yang signifikan ditunjukkan oleh data efisiensi fotosistem II (rasio F_v/F_m) antara kontrol dengan perlakuan-perlakuan lainnya. F_v/F_m merupakan parameter yang paling sensitif untuk mengindikasikan serangan awal penyakit busuk akar. Adapun parameter kadar klorofil dan laju fotosintesis (A_{max}) menunjukkan nilai yang menurun baik pada tanaman kontrol maupun perlakuan. Perubahan nilai kedua parameter fotosintesis tersebut lebih ditentukan oleh perbedaan musim. Patogen *A. luteobubalina* berhasil diisolasi kembali dari akar *E. nitens* yang menunjukkan penurunan respons fotosintesis. Hal tersebut menunjukkan bahwa penurunan respon fotosintesis berkaitan dengan adanya infeksi awal penyakit busuk akar. Namun diperlukan percobaan dengan waktu pengamatan yang lebih lama, agar perubahan respon fisiologis lainnya dapat terdeteksi.

Kata kunci: *Eucalyptus nitens*, inokulasi buatan, kadar klorofil, laju fotosintesis, efisiensi fotosistem II, busuk akar

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I. INTRODUCTION

Root-rots significantly reduce the productivity of important crops in tropical countries. For the majority of pathogens causing root-rot diseases, there is no clear indication of early infection (Mohammed, Rimbawanto & Page, 2014). As such, methods for early detection of root-rot disease are important for the pulp, oil palm and rubber industries. Early detection might allow the implementation of effective remedial measures to combat root-rot diseases as long as associated costs are not prohibitive e.g. the removal of infected woody material and/or the targeted application of biocontrol agents in the area infected. However, early detection is difficult because individual trees can appear healthy above ground even when damage to the root system has become severe (Hadfield, Goheen, Filip, Schmitt & Harvey, 1986; Farid, Lee, Maziah, Rosli & Norwati, 2006). For example, in the case of basal stem rot in oil palm, foliar symptoms are only visually observed when the fungus has killed half of the basal stem (Mohammed et al., 2014).

Observation of thin crowns, growth reduction and/or foliage chlorosis have proven to be useful indicators for detecting infected tree for many root-rot diseases (Morrison, Williams & Whitney, 1991; Omdal, Shaw & Jacobi, 2004). At the leaf scale, root-rot diseases can induce many changes to the biochemical, physiological and structural properties of leaves, which may result in a range of visual symptoms including needle chlorosis, production of non-green metabolites, necrosis and desiccation. These changes can potentially be used for monitoring the health and condition of forests (Stone, Coops & Culvenor, 2000; Luyssaert, Raitio, Vervaeke, Mertens & Lust, 2002; Gunthardt-Goerg & Vollenweider, 2007). For example, chlorophyll content has often been advocated as a sensitive indicator of many types of plant stress including drought, nutrient deficiency and diseases (Barry, Stone & Mohammed, 2008). Determining leaf-level physiological responses of the host plants during the initial stage of root-rot infection could provide valuable information that may help with developing

methods for the early detection of root-rot. To date, there has been little research on the effect of root disease on the host's physiology before the appearance of visual symptoms.

Photosynthetic capacity is useful parameter for monitoring these physiological changes. Stressful agents including fungal diseases can reduce the photosynthetic capacity due to their influence on one or more of the partial processes associated with photosynthesis (Dubey, 1997; Pinkard & Mohammed, 2006). This influence may include decreased light-energy utilization, decreased chlorophyll content, the destruction of the chloroplast fine structure, degradation of photosystem (PS) II and alteration of biochemical processes (Sharma & Hall, 1992; Singh & Dubey, 1995; Dubey, 1997; Chou, Bundock, Rolfe & Scholes, 2000; Lopes & Berger, 2001; Meyer, Saccardt, Rizza & Genty, 2001; Berger, Papadopoulos, Schreiber, Kaiser & Roitsch, 2004; Robert, Bancal, Nicolas & Lannou, 2004; Berger, Sinha & Roitsch, 2007).

The degree of inhibition of photosynthesis may be indicative of the aggressiveness of the pathogen (Guest & Brown, 1997). Root pathogens, such as species of *Armillaria*, which occupy and alter the host's vascular tissue (Morrison et al., 1991) may influence the photosynthetic activity indirectly by affecting the pathways of water flow in the xylem. The impact of root-rot on photosynthetic activities may be similar to the disruptions caused by water stress that is associated with decreased stomatal conductance, a lowering of intercellular CO₂, decreased chlorophyll level, changes in ultra-structure of chloroplasts, alteration in electron transport and decreased activity of the enzyme ribulose biphosphate carboxylase (Dubey, 1997).

Physiological aspects of plant-pathogen interactions have been well studied, especially for foliar diseases (Goicoechea, Aguirreola, Cenoz & Garcia-Mina, 2001; Bonfig, Schreiber, Gabler, Roitsch & Berger, 2006; Pinkard & Mohammed, 2006; Rodriguez-Moreno et al., 2007). In contrast, understanding the effects of root-rot diseases on plant physiology, especially in relation to hardwood trees, has received little

attention. This paper observes physiological changes before the appearance of root-rot's visual symptoms.

We used the *Eucalyptus nitens* (H. Deane and Maiden) Maiden and *Armillaria luteobubalina* Watling and Kile model pathosystem to quantify the photosynthetic changes of the host plant in response to root-rot disease. *Armillaria luteobubalina* is a generalist pathogen that has approximately 88 hosts (Shaw & Kile, 1991). This fungal species has been observed to cause root-rot in 3-year- and 6-year-old *E. nitens* plantations in Tasmania, Australia (Tim Wardlaw, personal communication). Findings from this pathosystem model will contribute valuable information about the potential to develop an early detection method of root diseases that are currently threatening the productivity of tropical plantation crops such as hardwoods, rubber and oil palm. In this experiment, the hypothesis that root infection will alter the processes associated with photosynthesis before the visual appearance of the disease symptoms is tested. We undertook a pot trial to characterize early physiological responses, (1) photosynthetic efficiency or photosystem (PS) II yield (F_v/F_m via chlorophyll fluorescence), (2) chlorophyll content, and (3) photosynthetic rate (A_{max}) of *E. nitens* saplings to artificial inoculation with *A. luteobubalina*. Re-isolation of *A. luteobubalina* from symptomatic trees was also conducted in order to confirm whether the measured physiological changes were associated with the fungal pathogen infection.

II. MATERIAL AND METHOD

A. Plants and Isolates

Forty-two two-year-old *E. nitens* saplings were planted in 30-cm-diameter pots containing a potting-mix medium consisting of soil, sand, and pine-bark compost (1:1:1). A previous pilot study showed that this mixed-soil medium was suitable for maintaining the viability of the inoculum. The saplings were fertilised with 15g of a slow-release fertilizer (Osmocote®) and watered daily with drippers until saturated.

The fungal cultures were obtained by isolating

from infected roots of an ornamental olive tree in the Hobart Royal Botanical Gardens, Australia (isolate strain 1) and a *Cupressus* sp. in Cascade Brewery Garden, Australia (isolate strain 2). Molecular analysis identified both isolates as *A. luteobubalina* having 98-100% sequence similarity with the described isolates in GenBank; there was a difference of seven nucleotides between isolates strains 1 and 2 (Agustini, 2010).

B. Fungal Isolations

Pure cultures of *A. luteobubalina* isolate strains 1 and 2 were obtained from sterilized infected root samples grown on 1% malt extract medium with the addition of selected antibiotics (MAT). Specifically, the MAT medium was prepared by autoclaving 1% malt extract agar (MEA) for 30 min at 120°C. Antibiotics (50 ppm penicillin, 50 ppm streptomycin, 25 ppm polymixin and 230 ppm thiabendazole) were added to the autoclaved MEA during cooling (i.e. at < 60°C). Root samples were surface-sterilised through a series of washings in different solutions, i.e. 2 min in tap water, 2-3 min in 20% Chlorox™ (sodium hypochlorite solution), and three times in sterile water. Hyphae that grew from the root samples were sub-cultured onto 2% MEA and incubated in the dark at 21°C for at least one month.

C. Inoculum Preparation and Artificial Inoculation of Plant Material

Branches harvested from young *Eucalyptus globulus* Labill. were prepared as inoculum rods using the method described by Mansilla et al. (2001), with some modification. The colonisation method involved inserting, under sterile conditions, branch segments (5-6 cm in length and 1-2 cm in diameter) that had been autoclaved for 30 min at 120°C into a 200 mL tissue culture vessel (round autoclavable polycarbonate container with lid). This vessel contained 150 mL of sterile MAT medium. Additional MAT medium was added to ensure that the branch segments were completely submerged in agar. These branch segments were inoculated with *A. luteobubalina* by placing seven

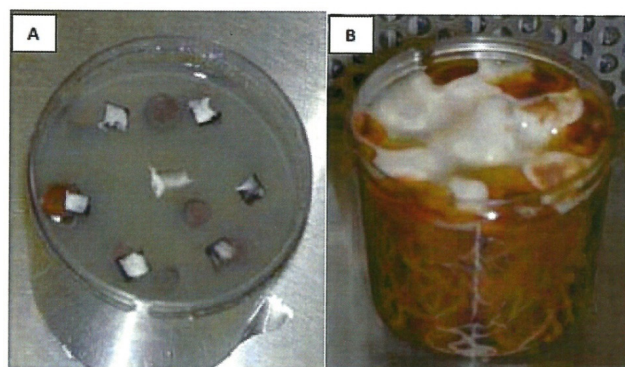


Figure 1. Pieces of agar with mycelia of *A. luteobubalina* on the MAT media (A), Mycelial fans after incubation at 21°C for three months (B)



Figure 2. Inoculum rods (A), Inoculation sites: three holes close to the root collar (B)

colonised mycelial segments (approx. 1 cm²) onto the agar surface (Figure 1A). Each vessel was then closed, sealed with plastic film, and placed in the dark at 21-22°C for approximately three months until the *E. globulus* branches, hereafter referred to as inoculum rods, had been fully colonised with the mycelia and rhizomorphs of *A. luteobubalina* (Figure 1B and 2A). Tissue culture vessels with uninoculated branch segments were also prepared to serve as controls.

Eucalyptus nitens saplings were inoculated by placing fully colonised inoculum rods (Figure 2A) into each pot adjacent to and just touching lateral roots in close proximity to the root collar (Figure 2B). The wounding treatment involved removing a small piece of bark (approx. 0.5-1 cm length) from the lateral roots using a Swiss Army knife.

D. Experimental Layout

A factorial design was applied in this greenhouse-based experiment. Six treatments were tested, including two physical treatments (*i.e.* unwounded and wounded host-root

systems), and two different fungal treatments (*i.e.* *Armillaria* strain 1 and strain 2) and an uninoculated control. The physical treatments were applied in order to examine the ease of pathogen entry into the root tissue. The six treatments were: unwounded-control (UW-P0), wounded-control (W-P0), unwounded-isolate strain 1 (UW-P1), wounded-isolate strain 1 (W-P1), unwounded-isolate strain 2 (UW-P2), and wounded-isolate strain 2 (W-P2). Each treatment consisted of seven replications, resulting in a total of 42 saplings. The *E. nitens* saplings were arranged in a randomised within the block design.

E. Physiological Measurements

Photosynthetic capacity (A_{max}) and maximum quantum yield of photosystem II yield (F_v/F_m) were assessed just prior to inoculation (T_0 , 2 October 2008; Spring) and after the first symptoms were observed (T_2 , 29 April 2009; Autumn). During the seven months between T_0 and T_2 , an intermediate measurement (T_1 , 30 January 2009; Summer) of F_v/F_m was carried out to determine if there was any evidence

of alterations in the plants' physiology prior to the appearance of the visual symptoms. Measurement frequency was decided based on the preliminary trial results which showed no significant differences in the above physiological variables between control and inoculated saplings over a six-month period. It suggested that extensive monitoring during the first six months after inoculation was not warranted.

Physiological assessments of F_v/F_m , A_{max} and relative chlorophyll content were made on three fully-expanded leaves per sapling. The leaves were selected from the third or fourth leaf pair just behind the branch tip. All plants (42 saplings) were assessed. Chlorophyll fluorescence (F_v/F_m) was measured pre-dawn using a chlorophyll fluorometer (OS-30p Opti-Science). Photosynthetic rate (A_{max}) was quantified using a CIRAS infrared gas analyser (PP Systems, Herts, UK) with an artificial light source set to deliver $1500 \mu\text{mol m}^{-2} \text{s}^{-1}$ at the leaf surface and ambient CO_2 concentration (370 – 380 ppm).

A Minolta SPAD-502 chlorophyll meter (Konica-Minolta, Hong Kong, China) was used to obtain relative chlorophyll content. The SPAD was calibrated by directly measuring the chlorophyll concentration of thirty leaves. Fresh leaf discs (dry weight of each disc ~ 0.020 g) were extracted for chlorophyll content with a triple extraction method (Martin et al., 2007). Discs were ground in a mortar with approximately $50 \mu\text{g M}_g\text{CO}_3$, $50 \mu\text{g}$ washed, fine sand and a small volume of liquid nitrogen. Ground leaf material was extracted with three small volumes of 100% cold acetone, and centrifuged for 3 min. Absorbance was read at 470, 645, 663 and 710 nm with a Cary UV-VIS spectrophotometer (Varian Medical Systems, Inc., Palo Alto, CA, USA). Total chlorophyll (Chl *a* and *b*) was calculated using the equations of Lichtenthaler and Buschmann (2001). Using this data, a standard curve was created and the SPAD values converted to chlorophyll concentration ($\mu\text{g/g}$).

F. Fungal Re-isolations

All roots including those from inoculated and control plants were examined at the end of the experiment. Root balls were thoroughly washed and the inoculum rods were removed. Any symptoms and/or signs of infection were recorded and photographs taken. Fungal re-isolations were undertaken from symptomatic roots exhibiting lesions and/or fungal mycelium (Figure 3). This was done to confirm that the causal agent associated with the deterioration of the plants was *A. luteobubalina*. The re-isolations were carried out in the same way as the isolations described above.

Based on the presence or absence of fungal signs and/or root symptoms, four categories were established to describe the infection and root condition:

1. Positive infection by *A. luteobubalina*: as indicated the presence of either mycelial fans and/or lesion with white mycelia (early stage of mycelia fans) on the excavated root; fungal re-isolation was positive for *A. luteobubalina*.
2. Possible infection by *A. luteobubalina*: as indicated by the presence of a small lesion with white mycelia; fungal re-isolation was negative for *A. luteobubalina*.
3. Infection by un-inoculated fungi: Neither mycelia fans nor white mycelia observed; only necrotic tissue or lesion present; fungal re-isolation confirmed fungi other than *A. luteobubalina*.
4. Uninfected: no fungal signs and/or root symptoms and roots appear healthy.

G. Data Analysis

Two-way analysis of variance (ANOVA) performed in XLSTAT2011® was used to analyse the physiological data. Duncan's multiple range tests were used to determine significant differences among treatments.

III. RESULT AND DISCUSSION

A. Changes in Physiological Variables

Physiological variables measured just before inoculation (T_0) were not significantly different

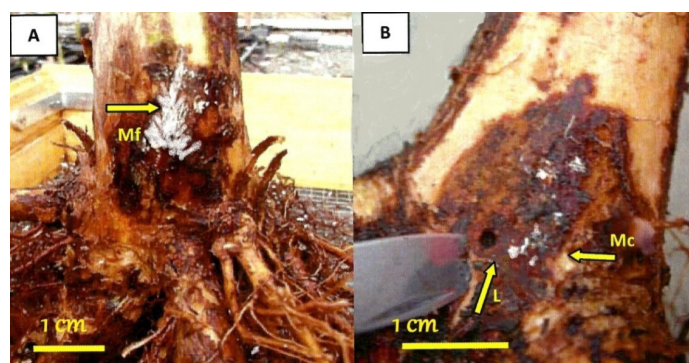


Figure 3. Mycelial fans (Mf) (A), lesion (L) and mycelia (Mc) on the root collar of infected *Eucalyptus nitens* saplings (B)

Table 1. Means \pm (SE) of the efficiency of PS II (F_v/F_m), chlorophyll content and photosynthetic rate (A_{max}) of *E. nitens* saplings inoculated with *A. luteobubalina* isolates over the period of observations at T_0 , T_1 (4 months after T_0) and T_2 (7 months after T_0)

Treatments / Time	Physiological variables		
	Efficiency of PS II F_v/F_m	Chlorophyll content $\mu\text{g/g}$	Photosynthetic rate (A_{max}) $\mu\text{mol/m}^2/\text{s}$
UW-P0 / T_0	0.78 ± 0.01^a	2918.8 ± 231.1^a	12.9 ± 2.2^a
W-P0 / T_0	0.77 ± 0.02^a	2772.1 ± 168.8^a	11.0 ± 0.5^a
UW-P1 / T_0	0.79 ± 0.01^a	2824.3 ± 110.3^a	11.6 ± 1.5^a
W-P1 / T_0	0.78 ± 0.01^a	2750.1 ± 90.0^a	13.5 ± 0.8^a
UW-P2 / T_0	0.79 ± 0.01^a	2552.3 ± 169.5^a	13.0 ± 0.8^a
W-P2 / T_0	0.79 ± 0.00^a	2918.8 ± 157.4^a	13.9 ± 1.1^a
UW-P0 / T_1	0.83 ± 0.01^a	NA	NA
W-P0 / T_1	0.82 ± 0.00^a	NA	NA
UW-P1 / T_1	0.81 ± 0.01^a	NA	NA
W-P1 / T_1	0.82 ± 0.00^a	NA	NA
UW-P2 / T_1	0.82 ± 0.00^a	NA	NA
W-P2 / T_1	0.82 ± 0.00^a	NA	NA
UW-P0 / T_2	0.81 ± 0.01^a	2117.7 ± 119.6^a	10.0 ± 0.4^a
W-P0 / T_2	0.76 ± 0.01^b	1939.0 ± 133.9^{ab}	8.5 ± 0.7^{ab}
UW-P1 / T_2	0.74 ± 0.02^b	1673.7 ± 119.0^{bc}	8.0 ± 1.0^{ab}
W-P1 / T_2	0.75 ± 0.01^b	1490.7 ± 81.3^c	7.7 ± 0.6^b
UW-P2 / T_2	0.73 ± 0.01^b	1629.1 ± 88.9^{bc}	7.2 ± 1.1^b
W-P2 / T_2	0.75 ± 0.01^b	1715.8 ± 70.5^{bc}	8.4 ± 0.5^{ab}

Notes:

- The values followed by different letters in the same column are significant at $\alpha=0.05$, as determined by a Duncan's test-ANOVA for each variable at each time of observation.
- NA = Not attempted

across treatments (Table 1). Three months after inoculation at intermediate assessment (T_1), photosynthetic efficiency of PS II (F_v/F_m) was unaffected by treatments (Table 1). The

first physiological changes were detected seven months after inoculation (T_2) when a significant difference in F_v/F_m between the unwounded controls (UW-P0) and all other treatments

Table 2. Change (\pm SE) calculated as $T_2 - T_0$ of physiological variables (F_v/F_m , chlorophyll content and A_{max}) of *E. nitens* saplings inoculated with *A. luteobubalina* isolates.

Treatments	Photosynthetic variables		
	Efficiency of PS II (F_v/F_m)	Chlorophyll content ($\mu\text{g/g}$)	Photosynthetic rate (A_{max}) $\mu\text{mol/m}^2/\text{s}$
UW-P0	0.03 ± 0.01^a	-801.1 ± 126.1^a	-3.0 ± 1.9^a
W-P0	0.00 ± 0.02^{ab}	-833.1 ± 163.6^a	-2.5 ± 0.8^a
UW-P1	-0.05 ± 0.03^{bc}	-1150.6 ± 107.4^{ab}	-3.7 ± 1.4^a
W-P1	-0.03 ± 0.01^{bc}	-1259.4 ± 99.7^b	-5.8 ± 0.9^a
UW-P2	-0.06 ± 0.02^c	-923.2 ± 188.6^{ab}	-5.6 ± 1.5^a
W-P2	-0.04 ± 0.01^{bc}	-1116.9 ± 140.6^{ab}	-5.5 ± 1.1^a

Note: The values followed by different letter in the same column are significant at $\alpha=0.05$, as determined by separate Duncan's test-ANOVA for each parameter at each time of observation.

was observed (Table 1). In particular, the F_v/F_m saplings in the unwounded-control (UW-P0) treatment increased, while in the other treatments F_v/F_m decreased; reductions in F_v/F_m were significantly greater in the inoculated (for both isolates and wound types) plants than in the UW-P0 treatment (Table 2).

Treatments effects on chlorophyll content (total Chl a and b) and A_{max} were more variable. At T2, there was a significant difference in chlorophyll content between inoculated (for both isolates and wound types) and UW-P0 plants (Table 1). Chlorophyll content decreased over the seven-month period of the experiment but, except for UW-P1 plants, there was no differences between inoculated and control treatments (Table 1). Photosynthetic capacity (A_{max}) decreased in all treatments during this period but there were no significant differences between treatments (Table 2).

Statistical tests show that the response of photosynthetic efficiency of PS II was affected by an interaction between time and treatments (F-ratio = 3.798, P-value = 0.005). For chlorophyll content and photosynthetic rate, the responses were more determined by sampling date (P-value < 0.0001).

Plant physiological changes associated with root-rot disease are not easy to detect. This is because diseases take time to develop before the plant expresses detectable physiological changes to the pathogen infection (Brown et al., 2012). This study found that *E. nitens* saplings grown

under semi-controlled conditions required approximately seven months for developing the first detectable changes in physiological performance following artificial inoculation with *A. luteobubalina* (Table 1).

In this study, F_v/F_m was shown to be the most sensitive physiological variable to detect stress caused by the root-rot pathogen; a significant reduction in F_v/F_m in response to inoculation was observed seven months after treatment. Changes in F_v/F_m have been widely used as a reliable diagnostic indicator of damage in response to various stresses such as extreme temperatures, and water and nutrient stress (Close & Beadle, 2003; Epron, Dreyer & Breda, 1992; Gamon & Pearcy, 1989; Groom & Baker 1992; He, Chee & Goh, 1996; Valladares & Pearcy, 1997). Since root-rot pathogens attack the vascular system of plants, responses to infection may be similar to those observed for drought stress. In drought-stressed plants, thylakoid membranes are the primary site of injury which leads to the decline of PS II activity (Dubey, 1997; Mutava, 2009). However, there was no evidence in this study that the decline in PS II activity was associated with parallel reductions in light-saturated photosynthetic rate A_{max} , but this may be because the reductions in F_v/F_m were not yet of sufficient magnitude. Decreases of PS II activity under stress are associated with photoinhibition where free high energy radicals in the thylakoid membranes cause photo-oxidation of the chlorophyll

(Havaux, 1992; Mutava, 2009). Differences between treatments in chlorophyll content at T_2 in this experiment suggested that this may occur; however between treatment reductions in chlorophyll content between T_0 and T_2 were not significantly different.

While decreases in A_{\max} between T_0 and T_2 were not significant, it is probable that the reduced rate was a response to seasonal changes in light and temperature as the T_0 measurement was done in mid-spring when the daily light exposure and temperature ranged between 3.1 to 32.8 MJ/m² and 10.6 to 30.6°C, respectively, while the T_2 was in late autumn when the daily light exposure and temperature ranged between 2.3 to 19.0 MJ/m² and 10.4 to 26.5°C, respectively. Leaves growing in a high light environment attain greater A_{\max} than leaves growing in a low light environment (DeJong & Doyle, 1985). Reduced A_{\max} can also be caused by reductions in seasonal temperatures (Battaglia, Beadleand & Loughhead, 1996) and overnight frost (Davidson, Battaglia & Close, 2004). Photosynthetic rates have shown to be closely related to chlorophyll content (Boardman, 1977; DeJong & Doyle, 1985), and this is consistent with the reduced chlorophyll content observed in this experiment. Loss of chlorophyll content was also found in *Pinus sylvestris* as seasonal temperatures declined (Ottander, Campbell & Oquis, 1995).

B. Re-isolation from Infected Roots

Root excavation showed that most of the inoculated saplings, both wounded and

unwounded, were infected by *A. luteobubalina* (Table 3). Wounded saplings (W-P1 and W-P2) showed 100% and 85.7% infection of *A. luteobubalina* strain-1 and strain-2, respectively; unwounded ones (UW-P1 and UW-P2), both inoculated with *A. luteobubalina* strain-1 and strain-2 showed only 71.4% infection (Table 3); these results confirm that wounding tends to enhance the possibility of infection. Wounded saplings inoculated with both strains of *A. luteobubalina* showed a greater level of infection than unwounded saplings.

Table 3 also shows that 14.3% and 28.6% of the roots of wounded and unwounded control plants (W-P0 and UW-P0), respectively, were infected by other soil fungi. Identification of these confounding fungal isolates was beyond the scope of this study.

In the field, wounding as well as other factors, such as poor planting, poor drainage and soil compaction can contribute to the increased incidence and severity of *Armillaria* root disease (Hadfield et al., 1986). Intact outer bark may play an important role in protecting roots from invasion by pathogens (Wargo & Harrington, 1991). Root grafts, breakage and associated insect feeding can potentially provide entry points for *Armillaria* and other root pathogens (Harrington, 1986; Rizzo & Harrington, 1988; Whitney, 1961). However, Baumgartner and Rizzo (2006) found that wounding the root collar of grapevine rootstocks did not significantly increase the infection rate of *Armillaria mellea* in a greenhouse trial. This suggests that wounding can trigger host defence reactions

Table 3. Percentages of saplings for which pathogens were re-isolated from roots in each of four categories.

Treatments	Root condition			
	Positively infected	Possibly infected	Infected by other fungi	Uninfected
UW-P ₀	0.0	0.0	28.6	71.7
W-P ₀	0.0	0.0	14.3	85.7
UW-P ₁	71.4	14.3	0.0	14.3
W-P ₁	100.0	0.00	0.0	0.0
UW-P ₂	71.4	28.6	0.0	0.0
W-P ₂	85.7	14.3	0.0	0.0

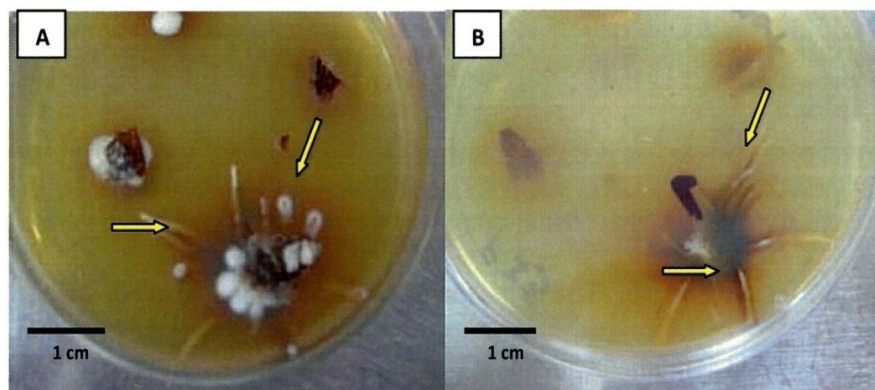


Figure 4. Re-isolated *A. luteobubalina* culture showing typical rhizomorphs (yellow arrows)
Remarks: (A) surface; (B) inverted

(Eyles, Bonello, Ganley & Mohammed, 2010), for example, the production of enzymes that function in lignin synthesis which leads to the reinforcement of the damaged cell wall (Baron & Zambryski, 1995) and/or the release of lytic enzymes or toxic secondary metabolites that may limit hyphal penetration of the inner bark (Wargo & Harrington, 1991). The possibility that wounding may have stimulated a defence reaction of the host in this study was not investigated.

Fungal infection was confirmed by the presence of mycelial fans and lesions on the root or root collar (Figure 3). Fungal cultures that had been isolated from the inoculated and symptomatic plants confirmed the presence of *A. luteobubalina* (Figure 4). The viability of the *A. luteobubalina* isolates on the *E. globulus* inoculum rods after being buried for seven months was low. Positive re-isolations of *A. luteobubalina* from these rods were obtained for three pots only and all were *A. luteobubalina* strain-1; re-isolations from inoculum rods carrying *A. luteobubalina* strain-2 were unsuccessful.

Similarly, there was a low level of successful re-isolations from the inoculum rods that had pseudosclerotial plates. Difficulties in re-isolating from pseudosclerotial plates can be understood since they are an immobile/inactive phase of *Armillaria* and had probably developed in response to the occupation of the rods by decomposing soil fungi (Dowson, Rayner &

Boddy, 1988).

This study confirmed the challenges of detecting the presence of root-rot disease during early infection. Among the physiological variables measured in this study, there was some evidence that at least one photosynthetic variable (*i.e.* F_v/F_m) may provide an early indicator of infection. Root and root collar examination remains the most reliable way to judge whether or not trees are infected. For *Armillaria* root disease, the production of characteristic mycelial fans can be also be used as a diagnostic feature.

IV. CONCLUSION

Physiological responses to artificial infection with the root-rot pathogen, *A. luteobubalina* were examined in *E. nitens* saplings. Among the physiological variables measured, we found that F_v/F_m was the most sensitive parameter for the early detection of root-rot disease. In particular, a significant difference in F_v/F_m between the unwounded control (UW-P0) and other treatments was observed. Chlorophyll content and A_{max} decreased for all trees, including controls during the seven months period and most likely reflected changes in season rather than treatment effects. Successful re-isolation of the root pathogen *A. luteobubalina* from inoculated symptomatic roots confirmed that the physiological changes were associated with the infection by this fungal pathogen. However, the functional changes that led to a reduction

in PS II efficiency in the inoculated saplings require further investigation. The finding from this study demonstrated that several months may be required following infection before any physiological changes can be detected. Root-rot is known to be a latent disease that may be present in plants for an extended period without any noticeable expression of symptoms. Longer periods of observation than were possible in this experiment are recommended for further research with a similar focus of interest.

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