Development of a grapevine detached root assay for assessing pathogenicity of *Cylindrocarpon* spp.

B. Pathrose, E.E. Jones, M.V. Jaspers and H.J. Ridgway

*Faculty of Agriculture and Life Sciences, Lincoln University, Lincoln, New Zealand*

*Corresponding author: Blessy.Pathrose@lincolnuni.ac.nz*

**Abstract** An *in vitro* method was developed to determine the relative pathogenicity of *C. macrodidymum*, *C. liriodendri* and *C. destructans* isolates on grapevine roots. Young feeder roots of similar age and diameter were placed individually into Petri dishes containing water-saturated silica sand, with the basal cut ends of the roots inserted into tubes filled with sterile water. Three isolates each of *C. macrodidymum* and *C. liriodendri* and two of *C. destructans* were inoculated as mycelial plugs onto the wounded roots, three replicates per isolate. After 1 month, all inoculated roots had dark brown lesions (range of lengths 28.7–40.4 mm) that were significantly different between isolates, with an isolate of *C. liriodendri* having the longest lesions and an isolate of *C. destructans* the shortest. This assay provides a rapid *in vitro* method to screen large numbers of *Cylindrocarpon* spp. isolates.

**Keywords** *Cylindrocarpon* species, detached root assay, lesion length, pathogenicity.

**INTRODUCTION**

Black foot disease of grape, caused by *Cylindrocarpon* species, is a problem in vineyard establishment worldwide. Over the last decade, its incidence has increased significantly in all major viticulture regions throughout the world, including South Africa, New Zealand, Italy, Australia and North America (Halleen et al. 2004a; Petit & Gubler 2005; Alaniz et al. 2009). Although the disease occurs in young and mature vines, losses usually occur during the first 5 years, sometimes even during the first year after planting (Halleen et al. 2006).

The symptoms include delayed or absence of budding, slow growth, reduced vigour, shortened internodes, sparse and chlorotic foliage with necrotic margins, wilting and dieback, as well as necrotic root crowns and development of secondary root systems. Grapevine roots affected by black foot disease show sunken necrotic lesions with a reduction in biomass. The below ground rootstock trunk develops black discoloration and necrosis of wood tissues (Halleen et al. 2004b; Alaniz et al. 2009).

In New Zealand *Cylindrocarpon macrodidymum*, *C. destructans* and *C. liriodendri* are pathogenic to grapevines, although their isolates vary in pathogenicity. Research by Alaniz et al. (2009) showed that specific isolate genotypes could be linked to virulence but the link between genotypic diversity and virulence has not been studied for New Zealand isolates. This research aimed to develop a detached root...
assay to provide a rapid in vitro method that could screen a large number of isolates for their relative pathogenicity to grapevine roots.

MATERIALS AND METHODS

Fungal isolates
Three isolates each of *C. macrodidymum* (C06a = ICMP 16788, Hb4a and Nel1b) and *C. liriodendri* (Co5c, Hb5a and Hb2d = ICMP 16790) and two of *C. destructans* (Mar11e and Mar15a), previously identified using morphological and molecular methods, were grown on potato-dextrose agar (PDA; Oxoid) for 7 days at 20°C with 12:12 h dark:light.

Selection of roots
As the physical characteristics of grapevine roots vary significantly, care was taken to collect roots of similar diameter and pigmentation. The younger outer feeder roots were removed from potted 1-year-old *Vitis riparia × V. rupestris* cv. 101-14 rootstock plants and washed in tap water to remove the soil. The roots of rootstock 101-14 were used as this variety had previously been shown to be susceptible to *Cylindrocarpon* infection.

Detached root assay
The three replicate roots per *Cylindrocarpon* isolate were placed individually into Petri dishes (90 × 15 mm) containing 30 g of silica sand wetted with 9 ml of sterile water. The basal cut end of each root was inserted through a Parafilm™ cap into a 1.7 ml tube filled with sterile water to keep the root hydrated during the experiment. The apical end of each root was cut across and a mycelial plug taken from the growing edge of a fungal colony placed against the cut end. The experiment was duplicated using mycelial plugs grown on either half or full strength PDA. Control roots had plugs of half or full strength PDA placed against the cut ends. The Petri dishes were sealed with clingfilm, incubated at room temperature and after 1 month, the lesion lengths were measured using a digital calliper (Mitutoyo, U.K Ltd).

To reisolate the pathogen, the roots were surface sterilised for 3 min in 0.35% sodium hypochlorite and washed in sterile water twice for 2 min. Each root was sequentially cut (1 cm lengths) and the pieces from both the lesions and the apparently healthy root sections were placed onto PDA amended with chloramphenicol (250 mg/litre). The plates were incubated at 20°C in 12:12 h light:dark for 7 days. Colony and conidium morphology was used to identify the *Cylindrocarpon* spp.

The lesion length data were analysed by ANOVA using GenStat version 12.

RESULTS
All inoculated roots (Figure 1b) had dark brown lesions that were clearly distinguishable from the remaining healthy root. Control roots (Figure 1a) had only minor discolourations at the apical cut ends after 1 month incubation.

![Figure 1](http://www.nzpps.org/terms_of_use.html)

**Figure 1** Experimental set-up for the detached root assay, shown 1 month after inoculation. (a) Control root showing healthy root material and (b) root inoculated with *C. liriodendri* isolate Co5c showing development of the black lesion from the apical end.
The lesion length differed (P<0.05) between isolates (Table 1). The largest lesions were produced on ½PDA by *C. liriodendri* isolate C05c (40.4 mm) and on PDA by *C. liriodendri* isolate Hb5a and *C. destructans* isolate Mar11e (38.9 and 39.3 mm, respectively). The smallest lesions were produced by *C. destructans* isolates Mar11e and Mar15a on ½PDA and PDA (28.7 and 30.0 mm, respectively).

There was a significant interaction between agar type and isolate (P<0.001) (Table 1). Several isolates (Co5c, Nel1b and Mar15a) that produced large lesions when inoculated as colonised ½PDA plugs produced significantly smaller lesions when inoculated as colonised PDA plugs. For isolates Hb4a and Mar11e the reverse was true. Notably, *C. destructans* isolate Mar11e produced the smallest lesion when inoculated as ½PDA and the largest lesion when inoculated on PDA.

**DISCUSSION**

The *in vitro* method developed here was able to show varied levels of pathogenicity between the isolates of *C. macrodidymum*, *C. liriodendri* and *C. destructans* isolates on grapevine roots, which was supported by earlier studies using potted rootstock plants (Probst et al. 2007). The inoculated roots showed the typical *Cylindrocarpon* species infection of black discolouration within 1 month. This was faster than disease development in the trunk bases of inoculated vines, which usually takes 3–6 months (Petit & Gubler 2005; Halleen et al. 2006). This may have been due to the young soft roots selected for the bioassay, the absence of plant defence responses, as the roots were detached from plants, or the absence of competing microbial flora. In addition to the advantage provided by the rapid symptom development, this assay can also be done at any time during the year unlike potted vine trials, which are more seasonally restricted. The potential for greater replication and tighter control over light, temperature and humidity also means that this type of assay has less variability than in potted vine, glasshouse experiments or field trials. In such whole-vine experiments, the variability of the growing season greatly affects results, for example, high temperatures during summer play an important role in symptom expression (Halleen et al. 2006).

This study also showed that the pathogenicity of any isolate’s mycelium inoculum was significantly affected by its growing medium. Isolates that were highly pathogenic applied in ½PDA were less pathogenic in PDA and vice versa. While this was surprising, it may reflect differences in the saprophytic ability of the individual isolates. For example, those that were highly pathogenic when applied in ½PDA may have been better able to colonise and access the plant nutrients than other more saprophytic isolates and thus

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### Table 1

Lengths (mm) of lesions produced on detached roots of rootstock 101-14 1 month after inoculation with PDA or half strength (½) PDA plugs colonised with isolates of *C. liriodendri*, *C. macrodydimum* or *C. destructans*.

<table>
<thead>
<tr>
<th>Species</th>
<th>Isolate</th>
<th>½ PDA</th>
<th>PDA</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>C. liriodendri</em></td>
<td>Co5c*</td>
<td>40.4 a</td>
<td>34.6 b</td>
</tr>
<tr>
<td></td>
<td>Hb5a</td>
<td>32.4 cd</td>
<td>39.3 a</td>
</tr>
<tr>
<td></td>
<td>Hb2d</td>
<td>34.9 bc</td>
<td>34.8 b</td>
</tr>
<tr>
<td><em>C. macrodidymum</em></td>
<td>Co6a</td>
<td>35.0 c</td>
<td>34.3 b</td>
</tr>
<tr>
<td></td>
<td>Hb4a*</td>
<td>29.8 de</td>
<td>37.1 ab</td>
</tr>
<tr>
<td></td>
<td>Nel1b</td>
<td>37.4 ab</td>
<td>35.0 b</td>
</tr>
<tr>
<td><em>C. destructans</em></td>
<td>Mar11e*</td>
<td>28.7 e</td>
<td>38.9 a</td>
</tr>
<tr>
<td></td>
<td>Mar15a*</td>
<td>37.5 ab</td>
<td>30.0 c</td>
</tr>
<tr>
<td>Control</td>
<td></td>
<td>12.2 f</td>
<td>10.0 d</td>
</tr>
<tr>
<td>LSD</td>
<td></td>
<td>3.44</td>
<td>3.38</td>
</tr>
</tbody>
</table>

1Means followed by the same letters within each column are not significantly different according to Fishers LSD P<0.05 test. The symbol * denotes isolates whose lesion lengths differ significantly (P<0.05) between agar types.
were less reliant on the agar medium. In full strength PDA, isolates with greater saprophytic ability can produce a larger biomass and thus be more pathogenic by virtue of inoculum density. Overall the ½PDA treatment gave the broader range of pathogenicity (28.7–40.4 mm lesions) compared to the PDA treatment (30.0–39.3 mm lesions). To better reflect the natural processes of infection in soil, the assay could be modified to use chlamydospores, which may be produced in liquid culture (Yoo et al. 1996).

Future work will use 200 selected isolates to investigate the relationships between genotypic characters with the pathogenicity of isolates on potted vines and on detached roots. Although there are few reports of detached root assays, use of detached leaf assays have been reported frequently for investigations into fungicide efficacy, induced resistance assays and isolate variation in pathogenicity. They are generally acknowledged to provide a quick and economic method that correlates well with similar assays on whole plants.

ACKNOWLEDGMENTS
The authors wish to thank New Zealand Winegrowers Inc. for funding this project and Lincoln University for providing a Doctoral Scholarship.

REFERENCES


